Comparison of isoprene chemical mechanisms at atmospheric night-time conditions in chamber experiments: Evidence of hydroperoxy aldehydes and epoxy products from NO$_3$ oxidation

Philip T. M. Carlsson$^1$, Luc Vereecken$^1$, Anna Novelli$^1$, François Bernard$^2$, Steven S. Brown$^{3,4}$, Bellamy Brownwood$^5$, Changmin Cho$^{1,a}$, John N. Crowley$^6$, Patrick Dewald$^6$, Peter M. Edwards$^7$, Nils Friedrich$^6$, Juliane L. Fry$^{5,b}$, Mattias Hallquist$^8$, Luisa Hantschke$^1$, Thorsten Hohaus$^1$, Sungah Kang$^1$, Jonathan Liebmann$^6$, Alfred W. Mayhew$^7$, Thomas Mentel$^1$, David Reimer$^1$, Franz Rohrer$^1$, Justin Shenolikar$^6$, Ralf Tillmann$^1$, Epameinondas Tsiligiannis$^8$, Rongrong Wu$^1$, Andreas Wahner$^1$, Astrid Kiendler-Scharr$^{1,9}$, and Hendrik Fuchs$^{1,9}$

$^1$Institute of Energy and Climate Research, IEK-8: Troposphere, Forschungszentrum Jülich GmbH, 52428 Jülich, Germany
$^2$Institut de Combustion, Aérothermique, Réactivité et Environnement (ICARE), UPR CNRS, 45071 Orléans, France
$^3$NOAA Chemical Sciences Laboratory, 80309 Boulder, USA
$^4$Department of Chemistry, University of Colorado, 80309 Boulder, USA
$^5$Department of Chemistry, Reed College, 97202 Portland, USA
$^6$Atmospheric Chemistry Department, Max Planck Institut für Chemie, 55128 Mainz, Germany
$^7$Wolfson Atmospheric Chemistry Laboratories, Department of Chemistry, University of York, Heslington, York, UK
$^8$Department of Chemistry and Molecular Biology, University of Gothenburg, 41296 Gothenburg, Sweden
$^9$I. Physikalisches Institut, Universität zu Köln, 50932 Köln, Germany

$^a$now at: School of Environmental Sciences and Environmental Engineering, Gwangju Institute of Science and Technology, Gwangju, South Korea

$^b$now at: Environmental Sciences Group, Wageningen University, 6708 HB Wageningen, the Netherlands

Correspondence: Philip T. M. Carlsson (p.carlsson@fz-juelich.de) and Hendrik Fuchs (h.fuchs@fz-juelich.de)

Abstract.

The gas-phase reaction of isoprene with the nitrate radical (NO$_3$) was investigated in experiments in the outdoor SAPHIR chamber at atmospherically relevant conditions specifically with respect to the chemical lifetime and fate of nitrato-organic peroxy radicals (RO$_2$). Observations of organic products were compared to concentrations expected from different chemical mechanisms: (1) The Master Chemical Mechanism, which simplifies the NO$_3$ isoprene chemistry by only considering one RO$_2$ isomer conformer. (2) The chemical mechanism derived from experiments in the CalTech Caltech chamber, which considers different RO$_2$ isomers conformers. (3) The FZJ-NO3 isoprene mechanism derived from quantum chemical calculations, which in addition to the CalTech Caltech mechanism includes equilibrium reactions of RO$_2$ isomers conformers, unimolecular reactions of nitrate RO$_2$ radicals and epoxidation reactions of nitrate alkoxy radicals. Measurements using mass spectrometer instruments give evidence that the new reactions pathways predicted by quantum chemical calculations play a role in the NO$_3$ oxidation of isoprene. Hydroperoxy aldehydes (HPALD), which are specific for unimolecular reactions of nitrate RO$_2$, were detected even in the presence of an OH scavenger excluding the possibility that concurrent oxidation by hydroxyl radicals (OH) is responsible for their formation. In addition, ion signals at masses that can be attributed to epoxy compounds, which are spe-
cific for the epoxidation reaction of nitrate alkoxy radicals, were detected. Measurements of methyl vinyl ketone (MVK) and methacrolein (MACR) concentrations confirm that the decomposition of nitrate alkoxy radicals implemented in the CalTech Caltech mechanism cannot compete with the ring-closure reactions predicted by quantum-chemical calculations. The validity of the FZJ-NO3 isoprene mechanism is further supported by an accurate simulation of the measured a good agreement between measured and simulated hydroxyl radical (OH) reactivity. Nevertheless, the FZJ-NO3 isoprene mechanism needs further investigations with respect to the absolute importance of unimolecular reactions of nitrate RO$_2$ and epoxidation reactions of nitrate alkoxy radicals. Absolute concentrations of specific organic nitrates such as nitrate hydroperoxides would be required to experimentally determine product yields and branching ratios of reactions but could not be measured in the chamber experiments due to the lack of calibration standards for these compounds. The temporal evolution of mass traces attributed to products species such as nitrate hydroperoxides, nitrate carbonyl, nitrate alcohols as well as hydroperoxy aldehydes observed by the mass spectrometer instruments demonstrates that further oxidation by the nitrate radical and ozone at atmospheric concentrations is not relevant on the typical time scale of one night (12 hours) for typical oxidant concentrations. However, oxidation by hydroxyl radicals present at night and potentially also produced from the decomposition of nitrate alkoxy radicals can contribute to their nocturnal chemical loss.
1 Introduction

Isoprene (C\textsubscript{5}H\textsubscript{8}) is an unsaturated compound and the most abundant hydrocarbon in the atmosphere. Circa 500 Tg per year of isoprene is emitted by plants as co-product of photosynthesis activity (Guenther et al., 2012). The high reactivity of isoprene towards the most important daytime oxidant, the hydroxyl radical (OH), results in a chemical lifetime of a few hours for typical atmospheric conditions, so that the majority of isoprene is oxidized during the day. However, isoprene can also be present in significant quantities after sunset, when the production rate of OH radicals is low, so that oxidation by the nitrate radical (NO\textsubscript{3}) or ozone can gain in importance (Brown et al., 2009; Edwards et al., 2017).

Oxidants add preferentially to the C=\text{C} double bonds in isoprene initiating a cascade of radical reactions. Theoretical studies of the OH-initiated oxidation of isoprene showed that the primary organic peroxy radicals (RO\textsubscript{2}) formed after the OH addition are unstable at atmospheric temperatures. The RO\textsubscript{2} isomers conformers continuously equilibrate through oxygen elimination and re-addition reactions at a time scale that is short relative to the chemical lifetimes of the RO\textsubscript{2} radicals at atmospheric conditions (Peeters et al., 2009, 2014). As a consequence, fast H-shift reactions of minor RO\textsubscript{2} isomers can constitute a large loss process for the entire RO\textsubscript{2} pool. This applies to the 1,6-H-migration reactions of the Z-\ensuremath{\delta}-RO\textsubscript{2} isomers produced from the isoprene + OH reaction (Peeters et al., 2014). These H-migrations lead eventually to the regeneration of OH radicals. Because this type of radical regeneration does not require the presence of nitric oxide (NO), it can significantly enhance radical concentrations in forested environments (Novelli et al., 2020). The OH initiated oxidation of isoprene has been investigated in laboratory (Crounse et al., 2011; Berndt et al., 2019) and simulation chamber studies (Fuchs et al., 2013; Novelli et al., 2020), which contributed to the refinement of the chemical mechanism proposed by the theoretical studies. The results can partly explain high OH radical concentrations observed in field experiments in rainforests (Leleivelde et al., 2008; Whalley et al., 2011).

In contrast to daytime, the loss of RO\textsubscript{2} radicals due to the reaction with NO does not play a role at night in the absence of near emission sources because NO production from the photolysis of NO\textsubscript{2} is stopped and NO is rapidly titrated to NO\textsubscript{2} by the reaction with ozone. In some situations, ozone can also be locally completely consumed in the night if there are high NO emissions for example from traffic or from power plants. In this case, NO can accumulate, but as also. However, for these conditions, the nitrate radical is rapidly lost in the reaction with NO\textsubscript{3}. Therefore, it is unlikely that nitrate RO\textsubscript{2} radicals and NO exist simultaneously. Therefore Thus, nitrate RO\textsubscript{2} from the reaction of NO\textsubscript{3} with organic compounds are expected to react mainly with hydroperoxy radicals (HO\textsubscript{2}), other organic peroxy radicals, the nitrate radical or they may undergo unimolecular reactions.

In previous chamber and laboratory studies investigating the reaction of isoprene with NO\textsubscript{3}, the fate of RO\textsubscript{2} was often assumed to be dominated by RO\textsubscript{2} self- and cross- recombination reactions and RO\textsubscript{2} reactions with NO\textsubscript{3} due to high reactant concentrations (Barnes et al., 1990; Kwok et al., 1996; Perring et al., 2009; Kwan et al., 2012). A chamber study by Schwantes et al. (2015) focussed on the product distribution from the reaction of the impact of the loss of nitrate RO\textsubscript{2} with HO\textsubscript{2} on the product distribution because this reaction pathway is generally the dominant loss path in the atmosphere. Chamber studies by
Rollins et al. (2009) and Ng et al. (2008) were also designed to reproduce atmospheric chemical conditions, for which the nitrate \( \text{RO}_2 \) reacts in various pathways.

Explicit Near-explicit chemical mechanisms such as the Master Chemical Mechanism (Jenkin et al., 2015) and the isoprene mechanism developed by Wennberg et al. (2018) (called CalTech mechanism in this work) were partly built by using results from these studies. In addition, it has been proposed that the nitrate \( \text{RO}_2 \) radicals formed from the reaction of the nitrate radical with isoprene can interconvert at ambient temperature (Wennberg et al., 2018; Vereecken et al., 2021). This can enhance the importance of unimolecular reactions of specific \( \text{RO}_2 \) conformers if the chemical lifetime of the \( \text{RO}_2 \) radicals is long enough that concentrations can re-equilibrate.

Furthermore, the theoretical study by Vereecken et al. (2021) revealed that unimolecular reactions of alkoxy radicals formed in the radical reaction chain subsequent to the addition of \( \text{NO}_3 \) to isoprene lead to the production of epoxide \( \text{RO}_2 \), influencing the distribution of organic products. This newly identified chemistry is only included in the FZJ-NO3 isoprene mechanism published by Vereecken et al. (2021), which focuses on the ability of the updated mechanisms to reproduce \( \text{RO}_2 \) radical concentration measured in experiments in the SAPHIR chamber.

The aim of this study is to compare the \( \text{NO}_3 \) isoprene chemistry of different available explicit mechanisms (MCM, CalTech and FZJ-NO3) with respect to the fate of nitratoo-organic peroxy radicals and the distribution of organic products for a series of chamber experiments for a range of performed at atmospherically relevant night-time conditions.

2 Methods

2.1 Experiments in the SAPHIR chamber

The experiments discussed in this work were performed in the atmospheric simulation chamber SAPHIR chamber (Rohrer et al., 2005) at Forschungszentrum Jülich in 2018. The chamber is a 270 m\(^3\) double-wall reactor. It is operated at a slight overpressure of 35 Pa to prevent ambient air from leaking into the chamber. The space between the 2 films is continuously flushed with pure nitrogen to prevent contamination of the inner chamber. The walls are made of Teflon film (FEP) and are thus chemically inert while the full solar spectrum is transmitted into the chamber (Bohn and Zilken, 2005). Night-time can be simulated by a shutter system that covers the chamber, so that photolysis processes are negligible. Synthetic air used for flushing the chamber and for replenishing losses due to sampling of instruments and leakage is produced from evaporating and mixing high purity liquid nitrogen and oxygen (purity: 99.9999 %, Linde). Inside the chamber, 2 fans are operated to ensure homogeneous mixing of air. The temperature inside the chamber is similar to ambient temperature and ranged between 291 and 308 K with maximum values in the afternoon for the experiments in this work.

Reactive trace gases added to the chamber in the experiments were ozone produced by a silent discharge ozonizer (O3onia), isoprene (\( \text{C}_5\text{H}_8 \), purity: 99 %, Sigma Aldrich), propene (purity: 99.8 %, Linde), CO (purity: 99.997 %, Linde) and \( \text{NO}_2 \) (purity: 99.2 %, 519 ppmv in nitrogen, Linde). Addition of gaseous species were controlled by calibrated mass flow controllers. Isoprene was injected as a liquid with a syringe into a hot volume and the vapour was flushed into the chamber together with the replenishment flow of zero air.
Figure 1. Measurements of radical and trace gas concentrations and NO₃ reactivity in the experiment on 09 August 2018 (Experiment #1) investigating the oxidation of isoprene by NO₃. Between 100 and 200 ppmv propene was present to produce HO₂ radicals by its ozonolysis. OH radicals, which are produced in the ozonolysis reaction, are rapidly converted to HO₂ in the reaction with 70 to 120 ppmv CO that was injected at the start of the experiment. OH reactivity was dominated by the high CO concentration and is not shown. NO₃ reactivity does not include reactivity from organic radicals and NO₂. NO₃ reactivity from isoprene is calculated from measured isoprene concentrations and reaction rate constants recommended in the literature (Mellouki et al., 2021). The difference between measured reactivity and reactivity from isoprene can be attributed to propene in this experiment. Observed RO₂ radicals only include a fraction of the total RO₂ because the LIF instrument cannot detect all RO₂ species formed in the reaction of isoprene with NO₃ (Vereecken et al., 2021).

Four experiments performed on 09, 10, 12, and 13 August 2018 (Experiment #1, #2, #3, #4) are analysed in this work (Table 1, Fig. 1, 2, A1, A2). Before each experiment, the chamber was flushed overnight with a high flow of zero air, so that concentrations of trace gases from previous experiments were below the limit of detection of instruments. The chamber roof was always closed to simulate night-time conditions. Experiments were performed in dry synthetic air. NO₃ was produced by the reaction of NO₂ and O₃. Typical mixing ratios after the injection were 5 ppbv NO₂ and 100 ppbv O₃. NO₃ production rates ranged between 0.9 and 11 ppbv/hour. Highest NO₃ production rates were reached in the experiment on 13 August 2018 (Experiment #4) and lowest rates in the experiment on 10 August 2018 (Experiment #2).

After NO₃ production started, isoprene was added. The injection of all three species was repeated after a few hours, when most of the isoprene had been consumed. Only NO₂ and O₃ were re-injected to enhance NO₃ production in the last part of the experiments, except for the experiment on 10 August 2018 (Experiment #2). Propene was injected in the experiment on 09 August 2018. In the experiment on 09 August 2018 (Experiment #1), propene was injected to enhance HO₂ concentrations by radical production via its ozonolysis. In this experiment, Excess CO was additionally injected to convert OH radicals to HO₂.

In the experiments in this work, no measurable secondary organic aerosol was formed, so that loss of products species on aerosol did not play a role (Brownwood et al., 2021).
Figure 2. Measurements of radical and trace gas concentrations and OH and NO$_3$ reactivity in the experiment on 13 August 2018 (Experiment #4) investigating the oxidation of isoprene by NO$_3$, when the total amount of oxidized isoprene was highest. NO$_3$ reactivity does not include reactivity from organic radicals and NO$_2$. OH and NO$_3$ reactivity from isoprene is calculated from measured isoprene concentrations and reaction rate constants recommended in the literature (Mellouki et al., 2021). NO$_3$ reactivity does not include reactivity from organic radicals and NO$_2$. Observed RO$_2$ radicals only include a fraction of the total RO$_2$ because the LIF instrument cannot detect all RO$_2$ species formed in the reaction of isoprene with NO$_3$ (Vereecken et al., 2021).

The total amount of isoprene that was consumed by NO$_3$ was $(3.2 \pm 0.5)$ ppbv, $(2.5 \pm 0.5)$ ppbv, $(4.8 \pm 0.5)$ ppbv, and $(11.6 \pm 1.2)$ ppbv in the experiments on 09, 10, 12, and 13 August 2018 (Experiment #1, #2, #3, #4), respectively (Brownwood et al., 2021). Approximately 10% of the total isoprene consumed oxidized in the experiment reacted with ozone except for the experiment on 09 August 2018 (Experiment #1), when 25 to 30% of isoprene was lost in the reaction with ozone due to the low NO$_3$ and high ozone concentration. In addition, measurements of OH radicals suggest that up to 10% of isoprene reacted with OH in the experiments without OH scavenger. However, OH concentration measurements were close to the limit of detection of the instrument, so that the fraction of isoprene that reacted with OH is rather uncertain. Overall, the dominant loss for isoprene was due to the reaction with NO$_3$ radicals (80 to 90% of the total loss in most of the experiments).

The chemical conditions in the experiments were chosen such that the chemical loss of nitrated RO$_2$ radicals differed between the experiments (Table 1). Similar as for typical night-time conditions in the nocturnal residual layer in the absence of nearby sources, nitric oxide concentrations were zero, so that RO$_2$ reacted only with either HO$_2$, RO$_2$ or NO$_3$ or re-arranged in unimolecular RO$_2$ reactions (Vereecken et al., 2021).

In the experiment on 09 August 2018 (Experiment #1), the ozonolysis of propene increased the HO$_2$ concentration and therefore increased the relative importance of the peroxy radical loss towards the reaction with HO$_2$. In the experiments on 10, 12 and 13 August 2018 (Experiment #4), the concentrations of NO$_3$ precursor species, O$_3$ and NO$_2$, and of isoprene were
varied. As a consequence, RO₂ concentrations differed between these experiments and therefore also the relative importance of RO₂ + RO₂ loss reactions.

2.2 Instrumentation

A large suite of instruments detected inorganic and organic species during the experiments. Isoprene and its oxidation products were measured by a proton transfer reaction time-of-flight mass spectrometer (VOCUS PTR-MS, Aerodyne, Krechmer et al. (2018)). The instrument was calibrated for isoprene, methyl vinyl ketone and methacrolein. The sensitivity of the instrument for isoprene was higher by a factor of 1.4 in dry air compared to than in humid air in which calibration measurements were performed (Brownwood et al., 2021). Measured concentrations were corrected for this humidity effect. No calibration standards were available for organic nitrate products such as nitrate nitrated alcohols, nitrate carboxyls, and nitrate hydroperoxides and epoxides.

Organic compounds were also detected by 2 other chemical ionization mass spectrometer instruments (CIMS) that used either Br⁻ (Albrecht et al., 2019; Wu et al., 2021) or I⁻ as reagent ions (Tsiligiannis et al., 2022). These instruments detected various oxygenated organic product species, but were not calibrated to provide concentrations. Details of the measurements by the Br⁻ CIMS instrument can be found in Wu et al. (2021) and by the I⁻ CIMS instrument in Tsiligiannis et al. (2022).

The high resolution of the mass spectrometer instruments allowed to attribute the ion mass signals (m/z) to sum formulas of organic compounds (Table A1). In this work, ion signals that were highest among all signals are discussed, most of which can be attributed to products of the isoprene oxidation (Wu et al., 2021; Tsiligiannis et al., 2022). Compared to the CIMS instruments, the precision of measurements by the VOCUS PTR-MS instrument was higher for organic compounds that contain few oxygens. In general, the sensitivity of CIMS instruments can be different for different isomers and functional groups, so that a change in the distribution of isomers isobaric compounds could partly explain observed differences between instruments (Lee et al., 2014a; Xiong et al., 2015, 2016). In addition, changes in the operational conditions of the instrument such as the temperature of the ionization region can lead to a variability of the instrument’s sensitivity (Robinson et al., 2022).

The total organic nitrate concentration was measured by 2 instruments, in which the total NO₂ concentration was detected either by a custom-built (Sobanski et al., 2016) or commercial cavity ring-down instrument (Keehan et al., 2020) after thermal dissociation of nitrate compounds in a heated inlet (TD-CRDS). A common data set from both instruments was created for this campaign. Details of these measurements can be found in Brownwood et al. (2021). These instruments also measured NO₂ in the sampled air in a separate mode or second measurement channel. In addition, NO₂ concentrations were measured by another custom-built cavity ring-down instrument (Liebmann et al., 2018) and a commercial chemiluminescence instrument combined with a blue-light converter (Eco-Physics). NO₂ concentration measurements from all instruments were combined to one common, quality-checked data set (Brownwood et al., 2021). Ozone concentrations were measured by a commercial instrument using UV-absorption (Ansyco).

NO₃ and N₂O₅ concentrations were measured with 2 custom-built instruments applying cavity-ring-down spectroscopy (Wagner et al., 2011; Sobanski et al., 2016). NO₃ was detected at 662 nm and the sum of NO₃ and N₂O₅ in a second channel, in which the inlet and cavity is heated to thermally decompose N₂O₅. Measurements were also combined to one data set taking
also into account that NO$_3$ and N$_2$O$_5$ can be expected to be in a thermal equilibrium for conditions of the experiments in this work.

HO$_2$, OH and RO$_2$ radical concentrations were determined by a laser-induced fluorescence instrument (Fuchs et al., 2011, 2012; Cho et al., 2021). OH radicals are excited at 308 nm in a low-pressure cell and their fluorescence is measured by gated single-photon counting. The fluorescence cell for the detection of only OH radicals was equipped with a chemical modulation reactor (CMR), which allows to account for potential interferences in the measurements (Cho et al., 2021). In another fluorescence cell, HO$_2$ radicals are chemically converted to OH in their reaction with NO. RO$_2$ radicals are converted eventually to OH in a third measurement channel (ROxLIF) that consists of an RO$_2$ converter, in which RO$_2$ and OH radicals are firstly converted to HO$_2$ in the presence of NO and CO, and a fluorescence cell downstream of the converter, in which the sum of all radicals is detected by OH fluorescence after HO$_2$ has reacted with excess NO. Recent studies showed confirmed that not all nitrate RO$_2$ radicals can be detected by the ROxLIF method as they do not form HO$_2$ or OH radicals after reacting with NO (Ashbourn et al., 1998) (Novelli et al., 2021; Vereecken et al., 2021).

OH reactivity ($k_{OH}$, the inverse of the chemical lifetime of the OH radical) was determined by a laser flash photolysis instrument, in which the time resolved decay of artificially produced OH radicals is observed (Fuchs et al., 2017). If, as in this work, the OH-reactivity from inorganic compounds is known, the contribution from organic compounds can be derived and compared to values based on the measurements of single compounds (Tan et al., 2021; Hantschke et al., 2021). In general, differences between measured and calculated OH reactivity can be used to determine if the detection of organic products that are reactive towards OH are complete.

The NO$_3$ reactivity was also measured in this work (Liebmann et al., 2017; Dewald et al., 2020). The concentration of artificially produced NO$_3$ is measured by cavity ring-down spectroscopy after reaction with either ambient or zero air in a flow tube. The NO$_3$ reactivity can be then calculated from the relative change of NO$_3$ concentrations between the two modes. In order to obtain the NO$_3$ reactivity from organic compounds, the contribution of NO$_2$ and NO$_3$ losses in the flow tube were accounted for. NO$_3$ reactivity from HO$_2$ and RO$_2$ radicals is not detected by the instrument due to loss of radicals in the inlet system (Dewald et al., 2020).

### 2.3 Modelling of trace gas concentrations

Trace gas concentrations were calculated using a chemical box model. In this work, three near-explicit chemical models have been applied: (1) The Master Chemical Mechanism version 3.3.1 (MCM) (Jenkin et al., 1997; Saunders et al., 2003; Jenkin et al., 2015), (2) the isoprene oxidation mechanism as introduced in the review article by Wennberg et al. (2018) and available at Bates and Wennberg (2017) (CalTech Caltech), and (3) the NO$_3$ isoprene mechanism based on theoretical calculations by Vereecken et al. (2021) and detailed in the supplement of Vereecken et al. (2021) (FZJ-NO3 mechanism).

The CalTech Caltech mechanism includes reactions of isoprene and isoprene product species but does not include further reactions of organic products that are not specific products from the oxidation of isoprene such as glyoxal or methyl glyoxal. In this work, the CalTech Caltech mechanism is therefore extended with chemistry from the MCM for those species.
Table 1. Chemical conditions in the experiments in this work. Experiments analysed in this work were performed in dry air. Mixing ratios of trace gases give the range of values reached right after their injection.

<table>
<thead>
<tr>
<th></th>
<th>Experiment #1</th>
<th>Experiment #2</th>
<th>Experiment #3</th>
<th>Experiment #4</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>09 August 2018</td>
<td>10 August 2018</td>
<td>12 August 2018</td>
<td>13 August 2018</td>
</tr>
<tr>
<td>(O_3) / ppbv</td>
<td>70–120</td>
<td>40–70</td>
<td>70–110</td>
<td>75–110</td>
</tr>
<tr>
<td>(NO_2) / ppbv</td>
<td>2–6</td>
<td>3–5</td>
<td>4–12</td>
<td>10–25</td>
</tr>
<tr>
<td>isoprene / ppbv</td>
<td>1–2.5</td>
<td>0.5–2</td>
<td>0.3–3</td>
<td>0–8</td>
</tr>
<tr>
<td>propene / ppbv</td>
<td>100–200</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>CO / ppmv</td>
<td>70–120</td>
<td>&lt;0.1</td>
<td>&lt;0.1</td>
<td>&lt;0.1</td>
</tr>
<tr>
<td>(NO_3) / pptv</td>
<td>1–10</td>
<td>5–40</td>
<td>5–60</td>
<td>10–500</td>
</tr>
<tr>
<td>(T) / K</td>
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<td>292–300</td>
<td>288–308</td>
<td>291–298</td>
</tr>
<tr>
<td>data reference</td>
<td>Fuchs et al. (2018a)</td>
<td>Fuchs et al. (2018b)</td>
<td>Fuchs et al. (2018c)</td>
<td>Fuchs et al. (2018d)</td>
</tr>
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</table>

The FZJ-NO3 mechanism only includes the reaction steps subsequent to the initial addition of \(NO_3\) to isoprene, but the chemistry of organic products was not investigated in Vereecken et al. (2021). The chemistry of the trace gases not considered in Vereecken et al. (2021) is taken from the CalTech mechanism. The isoprene OH oxidation scheme is applied as described in the work by Novelli et al. (2020), where the OH oxidation of isoprene was investigated in chamber experiments. Further chemistry of organic products that are not specific for the oxidation of isoprene are taken from the MCM. Chemical loss of first-generation organic products which are not included in either the CalTech or the MCM models is estimated from similarities to other organic products.

In the model runs, the injections of trace gases in the experiments were implemented as source reactions, which are effective during the short period of time during the injection. The rates are adjusted, such that the concentration change of the injected trace gas matches the observed increase in the concentration at the time of the injection. Physical parameters such as temperature and pressure were constrained to measured values. \(NO_3\) was also constrained to measured values, in order to decouple its modelled concentrations from wall reactions of \(NO_3\) and \(N_2O_5\), which are dependent on the chemical conditions of the experiment and hence hard to characterize accurately (Dewald et al., 2020). With \(NO_3\) concentrations constrained to measurements, the measured decay of isoprene, which is dominated by the reaction with \(NO_3\) for most of the time, is well described by the model, confirming that measured \(NO_3\) concentrations are consistent with the chemical loss of isoprene.

3 NO3 oxidation mechanisms of isoprene

The initial reaction steps in the oxidation of isoprene by \(NO_3\) (Vereecken et al., 2021) are similar to the oxidation by OH. H-atom abstraction from isoprene by \(NO_3\) is estimated to be at least 2 orders of magnitude slower than \(NO_3\) addition, based
on the available literature data on aliphatic and allylic H-abstraction reactions (Canosa-Mas et al., 1991; Atkinson et al., 2006) and therefore not further considered in this work.

NO₃ adds to either of the C=C double bonds leading to allyl-resonance stabilized alkyl radicals. Reversible oxygen addition and elimination reactions produce 3 different RO₂ stereoisomers conformers each from the addition of NO₃ on carbon C₁ and C₄ (Fig. 3). The different RO₂ isomers conformers rapidly reach equilibrium concentrations. NO₃ adds preferably on carbon C₁ (yield of 87%). The yield is higher in comparison to the corresponding OH addition (yield of 61%). The additions on the inner carbons (C₂ and C₃) are expected to be of minor importance (Vereecken et al., 2021) and are not further considered in this work.

The isoprene NO₃ mechanisms investigated in this work differ significantly in the treatment of the initially formed RO₂. The FZJ-NO3 mechanism includes 6 RO₂ isomers conformers formed subsequently to the NO₃ addition (Fig. 3). Specifically, the Z- and E-RO₂ isomers conformers of the δ-RO₂ isomers are distinguished. In contrast, the CalTech mechanism only treats δ- and β-RO₂ isomers separately and but does not include the equilibrium reactions between RO₂ isomers conformers.

The MCM simplifies the addition of NO₃ to isoprene even more by only considering the addition of NO₃ on carbon C₁ leading to the δ-RO₂ radical.

It is important to distinguish between Z- and E-RO₂ isomers conformers because isomer-specific unimolecular H-shift reactions need to be considered. Competitive unimolecular H-shift-reactions only occur for the Z-δ-RO₂ (Vereecken et al., 2021) leading to the formation of hydroperoxy aldehydes (HPALD) (Fig. 3). Due to the re-equilibration reactions between RO₂ isomers, these reaction channels can gain in importance if the rate of this RO₂ loss reaction (0.01 to 0.05 s⁻¹) is fast compared to faster than the chemical loss due to bimolecular RO₂ reactions. This will often be the case for night-time conditions, when mainly slow bimolecular RO₂ reactions with NO₃, HO₂ and other RO₂ radicals occur.

The distribution of organic products from the NO₃ oxidation of isoprene depends highly on the competition between the different RO₂ loss reactions. The bimolecular reaction of nitrate RO₂ with HO₂ radicals leads to the formation of nitrate hydroperoxides (NISOPOOH). Whereas one NISOPOOH isomer conformer is the exclusive product of the RO₂+HO₂ reaction in the MCM, the CalTech mechanism and FZJ-NO3 mechanisms include not only different isomers conformers but also the decomposition of the initially formed HO₂-RO₂ reaction complex into an OH radical and a nitrate alkoxy radical with a yield of approximately 50% for nitrate β-RO₂ radicals.

Nitrate alkoxy radicals can also be the product of RO₂+RO₂ reactions, but this reaction channel competes with the production of nitrate carbonyls (NC₄CHO) and nitrate alcohols (ISOPCNO₃). Alkoxy radicals are additionally formed from the reaction of nitrate RO₂ with NO₃ accompanied by the production of NO₂. The nitrate alkoxy radicals are expected to rapidly decompose (Novelli et al., 2021; Vereecken et al., 2021). In the MCM, the decomposition leads exclusively to the formation of one isomer conformer of the nitrate carbonyl product (NC₄CHO) together with an HO₂ radical. A similar mechanism is implemented in the CalTech mechanism and FZJ-NO3 mechanisms for most of the various nitrate alkoxy radical species except for those radicals produced from the most abundant β-1,2-RO₂ isomer, from which nitrate carbonyl species cannot be formed.

In the CalTech mechanism, decomposition of these nitrate alkoxy radicals leads instantly to the formation of methyl vinyl ketone (MVK) or methacrolein (MACR) together with formaldehyde and NO₂. This was determined from chamber
Figure 3. Schematic reaction mechanism of the reaction of isoprene with NO$_3$ as described in Vereecken et al. (2021). This includes fast inter-conversion of nitrate RO$_2$ isomers conformers by oxygen addition and elimination reactions. Only RO$_2$ isomerization reactions (Vereecken et al., 2021) which can compete with bimolecular reactions for typical night-time conditions are shown. Percentage values given next to the structure of RO$_2$ radicals are yields when equilibrium concentrations are established for typical night-time conditions like such as in the experiments in this work.
experiments reported in Schwantes et al. (2015), in which a high yield of MVK was found, when nitrate RO₂ mainly reacted with HO₂. The fate of nitrate alkoxy radicals was also investigated by Vereecken et al. (2021). Quantum chemical calculations show that the decomposition reaction is slow compared to slower than the ring-closure reactions leading to epoxide products. In contrast, 4-membered ring closure (barrier $\sim 25$ kcal/mol, Vereecken (2022)) requires breaking the planar double bond to bring the radical O-atom in an appropriate position for bonding. 5- to 6-membered ring closure (barrier $\sim 13-29$ kcal/mol, Vereecken et al. (2021)) are also favorable.

Differences between the chemical mechanisms also exist concerning the type of chemical loss reactions of first-generation stable organic products. Reactions with OH are considered in all mechanisms applying similar reaction rate constants. In addition, the MCM includes loss of isoprene organic nitrates due to ozonolysis reactions.

4 Results

In the experiments in this work, NO₃ was produced by the gas-phase reaction of NO₂ and O₃. NO₃ production rates ranged between 0.9 and 11 ppbv/hour. Highest NO₃ production rates were reached in the experiment on 13 August 2018 (Fig. 2) and lowest rates in the experiment on 10 August 2018 (Fig. A1). NO₃ mixing ratios were lowest right after the injection of isoprene (1.5 to 6 ppbv) with values between 2 and 20 ppbv due to the consumption in the reaction with isoprene (Fig. 2, 1, A1, A2). The rate with which isoprene was consumed in the reaction with NO₃ varied between experiments and consequently also the production rate of nitrato organic peroxy radicals.

The total amount of isoprene that was consumed by NO₃ was $(3.2 \pm 0.5)$ ppbv, $(2.5 \pm 0.5)$ ppbv, $(4.8 \pm 0.5)$ ppbv, and $(11.6 \pm 1.2)$ ppbv in the experiments on 09, 10, 12, and 13 August 2018, respectively (Brownwood et al. 2021). Approximately 10% of the total isoprene oxidized in the experiment reacted with ozone. In addition, measurements of OH radicals suggest that up to 10% of isoprene reacted with OH in the experiments without OH scavenger. However, OH concentration measurements were close to the limit of detection of the instrument, so that the fraction of isoprene that reacted with OH is rather uncertain. Overall, the dominant loss for isoprene was due to the reaction with NO₃ radicals (80 to 90% of the total loss).

The fate of nitratorganic peroxy radicals formed in the initial addition of NO₃ to isoprene determines the yield and type of organic products, which depends on the availability of reaction partners for bimolecular reactions. Like typical night-time conditions in the nocturnal residual layer in the absence of nearby sources, nitric oxide concentrations were zero in the experiments in this work, so that only RO₂ reactions with other radicals, HO₂, RO₂ and NO₃ and potentially unimolecular RO₂ reactions were of importance (Vereecken et al., 2021). Experimental conditions were varied among the experiments to explore the different fates of nitrate RO₂ radicals initially generated.

Results of the model calculations are shown in Fig. 4 for the experiment on 09 August 2018 (Experiment #1), when high HO₂ concentrations were present, and therefore the main loss path for RO₂ was the reaction with HO₂. Figure 5 shows results for the experiment on 13 August 2018 (Experiment #4), when RO₂ loss was distributed among all pathways that are relevant.
Figure 4. Comparison of results from model calculations applying the different isoprene NO\textsubscript{3} chemistry mechanisms for the experiment on 09 August 2018 (Experiment #1), when HO\textsubscript{2} concentrations were enhanced and excess CO was present as OH scavenger. MVK, MACR, NISOPOOH, ISOPCNO\textsubscript{3} and NC\textsubscript{4}CHO are produced from all mechanisms, whereas the other species are only produced from either 1,6-H-shift reactions or ring-closure reactions of nitrate alkoxy radicals, which are only implemented in the FZJ-NO\textsubscript{3} mechanism. Grey and black dots are measured values. Measured organic peroxy radical concentrations only include part of the total RO\textsubscript{2} because the LIF instrument cannot detect a fraction of nitrate RO\textsubscript{2} (Vereecken et al., 2021). Organic products were detected by the VOCUS PTR-MS instrument, which was only calibrated for MVK and MACR. All other traces are scaled to match best the results from the FZJ-NO\textsubscript{3} mechanism.

During night-time (Brownwood et al., 2021) and the amount of oxidized isoprene was highest. Results from the other experiments are shown in the Appendix (Fig. A5, A6).

Highest HO\textsubscript{2} concentrations of up to $17 \times 10^8$ cm\textsuperscript{-3} were measured in the experiment on 09 August 2018 (Experiment #1), when HO\textsubscript{2} was enhanced by production of OH radicals in the ozonolysis of propene, which were rapidly converted to HO\textsubscript{2} in the presence of excess CO (Fig. 1, Panel (d)). In the other experiments, measured HO\textsubscript{2} concentrations were between 1 and $5 \times 10^8$ cm\textsuperscript{-3} with highest values in the experiment on 13 August 2018 (Experiment #4). As discussed in Vereecken et al. (2021), the measured HO\textsubscript{2} concentrations are much higher than predicted by model calculations for experiments in this work (up to a factor of 10) except for the experiment on 09 August 2018 (Experiment #1). Although it is possible that part of the measured HO\textsubscript{2} radicals is due to an interference (Vereecken et al., 2021), the HO\textsubscript{2} radical concentrations predicted by the model are too low to explain observed OH radical concentrations for example during the last part of the experiment on
Figure 5. Comparison of results from model calculations applying the different isoprene NO₃ chemistry mechanisms for the experiment on 13 August 2018 (Experiment #4), when the amount of oxidized isoprene was highest. MVK, MACR, NISOPOOH, ISOPCNO₃ and NC₄CHO are produced from all mechanisms, whereas the other species are only produced from either 1,6-H-shift reactions or ring-closure reactions of nitrate alkoxy radicals, which are only implemented in the FZJ-NO₃ mechanism. Grey and black dots are measured values. Measured organic peroxy radical concentrations only include part of the total RO₂ because the LIF instrument cannot detect a fraction of nitrate RO₂ (Vereecken et al., 2021). Organic products were detected by the VOCUS PTR-MS instrument, which was only calibrated for MVK and MACR. All other traces are scaled to match best the results from the FZJ-NO₃ mechanism.

13 August 2018 (Experiment #4) (Section 5.6). Therefore, the measured HO₂ radical concentrations are used in the further analysis in this work.

A large fraction of nitrate RO₂ radicals cannot be detected by the LIF instrument used in this work (Novelli et al., 2021; Vereecken et al., 2021) because the detection scheme of the instruments requires that HO₂ or OH radicals are formed subsequent to the reaction of RO₂ with NO. However, this is only the case for some of the nitrate RO₂ radicals from the reaction of isoprene with NO₃ (Section 2.1). Therefore, measured RO₂ concentrations, which are maximum around 1 × 10⁹ cm⁻³ (Fig. 1 and 2, Panel (d)), need to be regarded as lower limits.

In all experiments, significant amounts (up to 1 ppv) of methyl vinyl ketone (MVK) and methacrolein (MACR) were detected by the VOCUS PTR-MS instrument. With respect to organic products, the VOCUS PTR-MS instrument was only calibrated to quantify the sum of methyl vinyl ketone (MVK) and methacrolein (MACR).
Figure 6. Loss reactions of the most abundant β-1,2-RO₂ species. Coloured arrows indicate the preferred reaction channel for the nitrate alkoxy radical in the different chemical models (red: MCM; yellow: CalTech; blue: FZJ-NO3). Dashed red arrows indicate corresponding reactions of the δ-RO₂ species which is the only RO₂ represented in the MCM. Coloured boxes indicate species that were observed by the VOCUS PTR-MS instrument. Though nitrate carbonyl products (NC₄CHO) cannot be formed from this specific nitrate β-RO₂ from isoprene, they are formed from other nitrate radicals and thus nitrate carbonyls were also observed by the VOCUS PTR-MS instrument. In addition, nitrate carbonyls are major products in the MCM (dashed red arrows) because only one nitrate δ-RO₂ is considered.

In addition, the mass spectrum shows signals that can be attributed to the sum formulas of a number of other product species including non-nitrate and nitrate organic compounds (Fig. 6). Their formation is further discussed in the next section (Section 5) together with results from model calculations applying the three different chemical mechanisms.

VOCUS PTR-MS, Br⁻-CIMS and I⁻-CIMS instruments also recorded ion signals from oxygenated organic compounds in the experiments that can be attributed to the sum formulas of a number of other product species including non-nitrate (HPALD: C₅H₈O₃) and nitrate organic compounds (NISOPOOH: C₅H₉NO₅, NC₄CHO: C₅H₇NO₄, ISOPCNO₃: C₅H₉NO₄) and epoxide products that are expected to be formed subsequent to the ring-closure reaction of alkoxy radicals (Reaction R9, R17, C₅H₈O₄, C₅H₈O₃, C₅H₉NO₆, C₅H₉NO₅, C₅H₇NO₅, Fig. 6).

Ion signals shown in Fig. 4, 5, A5, A6 were the highest signals observed in the mass spectrometer instruments except for the ion signal corresponding to a C₄H₇NO₅ compound observed by the I⁻- and Br⁻-CIMS instruments. A species with this sum
formula cannot be attributed to a major product species expected from the chemical mechanism. This is discussed in detail in Tsiligiannis et al. (2022).

Signals at the mass corresponding to NISOPOOH were highest among all product signals observed by the VOCUS PTR-MS instrument. The signal can include nitrate epoxides that are produced from the ring-closure reactions of alkoxy radicals (Section 5.3) and the reaction of NISOPOOH with OH, which have the same mass. However, their contribution is expected to be low for most of the time in the experiments in this work, specifically in the experiment on 09 August 2018 (Experiment #1), when HO₂ concentrations favoured RO₂ + HO₂ reactions and an OH scavenger was present (Fig. 4, Panel (a)).

Compared to the CIMS instruments, the sensitivity of the VOCUS PTR-MS instrument was higher for organic compounds that contain few oxygens. The CIMS instruments were not calibrated for the organic nitrate species, so that only relative signals can be compared. Signals from all three mass spectrometry instruments (Appendix Fig. A7, A8, A9, A10) can be compared by scaling them to best match modelled concentrations of organic products applying the FZJ-NO3 chemical mechanism.

The relative behaviour of signals is similar for all instruments with a few exceptions: (1) In the experiment on 09 August 2018 (Experiment #1), the signals of the Br⁻-CIMS instrument appear to be systematically lower after 10:00 UTC for unknown reasons (Fig. A7). (2) In the experiment on 13 August 2018 (Experiment #4), the loss rate of C₅H₉NO₄ compounds appears to be slower in the signal of the Br⁻-CIMS instrument than in the other mass spectrometer instruments (Fig. A10, Panel (c)) and expected from model calculations. This could be explained if other (fragments of) products were detected at that mass by the Br⁻-CIMS instrument, but not by the other instruments. (3) The loss rate of C₅H₁₀O₃ compounds observed by the I⁻-CIMS instrument appears to be faster than observed by the VOCUS PTR-MS instrument and expected from model calculations in the experiment on 13 August 2018 (Experiment #4) (Fig. A10, Panel (f)). The difference in the observed temporal evolution of C₅H₁₀O₃ compounds could be explained if the sensitivity of the instrument was lower for the hydroperoxide species than for the epoxide species, both of which are detected at the same mass (Section 5.6). Differences would become most obvious during this part of the experiment because these compounds have vastly different chemical lifetimes with respect to the reaction with OH, which was likely the dominant loss process for this part of the experiment. In some parts of the experiments, measurements by the I⁻-CIMS instrument exhibited an oscillating behaviour, which is most likely an instrumental artefact (for example Fig. A10, Panel (b)).

Some As some species produced from different loss pathways can be structurally different but have the same sum formula. These isobars cannot be distinguished by the mass spectrometers (Fig. 6): (1) Nitrate hydroperoxides (NISOPOOH) have the same mass as some nitrate epoxide species (Reaction R16). This applies for nitrate epoxides formed from the reaction of OH with NISOPOOH, which does not play a major role for conditions of the experiments, but also for specific nitrate epoxide products formed subsequently to the ring-closure reaction of nitrate alkoxy radicals predicted by the FZJ-NO3 mechanism (Vereecken et al., 2021). (2) Hydroperoxy aldehydes (HPALD) produced from unimolecular 1,6-H-shift reactions of the nitrate Z-δ-RO₂ isomers have the same mass as one epoxide product formed also from the ring-closure reaction of nitrate alkoxy radicals (sum formula C₅H₈O₃). NO₂ is eliminated, so that these products do not contain nitrate functional groups.
The temporal behaviour of products depends on their production and destruction rates. The rate of production is essentially the same for all products because they are formed from the same pool of nitrate RO₂ radicals from the reaction of isoprene with NO₃ which is the rate limiting step for their production. The temporal evaluation of their concentrations at later times of the experiment when isoprene had been consumed is determined by the rate of loss processes. Small delays due to consecutive radical reaction steps that apply for some of the products do not play a role on the time scale of the chamber experiment. Therefore, the temporal evolution mainly depends on the rate of loss processes, which can be chemical loss and dilution in these experiments. Loss to the chamber wall seems to be insignificant, because the temporal evolution appears to be consistent with only dilution for some species. This is further discussed in Section 5.9.

Br⁻ CIMS and I⁻ CIMS instruments also recorded signals from oxygenated organic compounds in the experiments. Compared to the CIMS instruments, the sensitivity of the VOCUS PTR-MS instrument was higher for organic compounds that contain few oxygens. The CIMS instruments were not calibrated for the organic nitrate species, so that only relative signals can be compared. Signals from all three mass spectrometry instruments (Appendix Fig. A7, A8, A9, A10) can be compared by scaling them to best match modelled concentrations of organic products applying the FZJ-NO₃ chemical mechanism.

The relative behaviour of signals is similar for all instruments with a few exceptions: (1) In the experiment on 09 August 2018, the signals of the Br⁻ CIMS instrument appear to be systematically lower after 10:00 UTC for unknown reasons. (2) In the experiment on 13 August 2018, the loss rate of C₅H₁₀O₃ compounds appears to be slower in the signal of the Br⁻ CIMS instrument compared to the other mass spectrometer instruments and expected from model calculations. This could be explained if other (fragments of) products species were detected at that mass by the Br⁻ CIMS instrument, but not by the other instruments. (3) The loss rate of C₅H₁₀O₃ compounds observed by the I⁻ CIMS instrument appears to be faster compared to the VOCUS PTR-MS instrument and expected from model calculations. The difference in the observed temporal evolution of C₅H₁₀O₃ compounds could be explained if the sensitivity of the instrument was lower for the hydroperoxide species compared to the epoxide species, both of which are detected at the same mass (Section sec:NISOPOOH). Differences would become most obvious during this part of the experiment because these compounds have vastly different chemical lifetimes with respect to the reaction with OH, which was likely the dominant loss process for this part of the experiment. In some parts of the experiments, measurements by the I⁻ CIMS instrument exhibited an oscillating behaviour, which is most likely an instrumental artefact.

In general, the sensitivity of CIMS instruments can be different for different isomers and functional groups, so that a change in the distribution of isobaric compounds could partly explain the observed differences between instruments (Lee et al., 2014; Xiong et al., 2015, 2016). In addition, changes in the operational conditions of the instrument such as the temperature of the ionization region can lead to a variability of the instrument’s sensitivity (Robinson et al., 2022).

Mainly measurements by the VOCUS PTR-MS instrument are discussed in the next sections. However, the conclusions do not depend on the choice of the instrument as can be seen by the overall good agreement in time-series of ion mass signals at the same mass by all of instruments (Fig. A7, A8, A9, A10). Results are also independent on the choice of scaling measured ion mass signals of the VOCUS PTR-MS instrument to the model results of the FZJ-NO₃ mechanism (A11).
5 Discussion

5.1 Chemical lifetime of nitrate RO₂ radicals

Using the RO₂ chemistry as implemented in the FZJ-NO3 mechanism and measured HO₂ concentrations results in overall loss rates of nitrate RO₂ of around 0.035, 0.005, 0.008 and 0.014 s⁻¹ in the experiments on 09, 10, 12, and 13 August 2018 (Experiment #1, #2, #3, #4). This implies chemical lifetimes between 30 s and several minutes, which are similar to values at atmospheric night-time conditions. RO₂ loss rates are 20 to 50 % lower if the chemistry implemented in the CalTech mechanism or MCM is applied.

Overall, differences of the RO₂ loss rates derived from the three mechanisms are mainly related to differences of the distribution of nitrate RO₂ isomers, for which chemical lifetimes vary. In addition, implementation of additional unimolecular RO₂ reactions shorten their chemical lifetime in the FZJ-NO3 mechanism. Differences of RO₂ loss rates between the chemical mechanisms are lowest for the experiment on 09 August 2018 (Experiment #1), in which the RO₂ loss is dominated by the reaction with HO₂ (Fig. 7, A3) leading to an overall high loss rate, so that unimolecular RO₂ reactions implemented in the FZJ-NO3 mechanism were less competitive.

If HO₂ concentrations are used as derived from model calculations, the total RO₂ loss rates are lower by 30 to 50 % than shown here due to the lower low predicted HO₂ concentrations (Vereecken et al., 2021). The distribution contribution of the different RO₂ loss channels shifts towards higher contributions from RO₂ reactions with other RO₂ radicals and with NO₃ (Fig. A4). In addition, unimolecular reactions further gain in importance due to the longer chemical lifetime of RO₂ radicals.

In the following sections, loss reactions implemented in the three chemical mechanisms and the formation of major organic products are analysed for the chamber experiments.

5.2 Production of nitrate alkoxy radicals

Alkoxy radicals play an important role in determining the differences in the concentrations of organic products, obtained by model calculations applying the three mechanisms (Fig. 4, 5, A5, A6). These differences are not only due to differences in the fate of alkoxy radicals, but also due to differences in the formation rates of alkoxy radicals which are formed from nitrate RO₂ radicals reacting with NO₃, RO₂, and HO₂ radicals.

In all three mechanisms, the initial product from the reaction between nitrate RO₂ and NO₃ is a nitrate alkoxy radical and NO₂ (Fig. 6, Reaction R3). Dewald et al. (2020) analysed NO₃ reactivity measurements performed in the same experiments and concluded that the reaction rate constant of the reaction of nitrate RO₂ with NO₃ would need to be around $5 \times 10^{-12}$ cm³ s⁻¹, which is nearly a factor of 2 higher than the generic reaction RO₂ + HO₂ → NO₃ rate constant based on the measured rate constant for CH₃O₂ + HO₂ → NO₃ used in the MCM and the CalTech mechanisms. With this rate constant, the loss rate of nitrate RO₂ in the reaction with NO₃ is between 0.001 and 0.003 s⁻¹ in the experiments on 10, 12 and 13 August 2018 (Experiment #2, #3, #4), contributing between 5 and 20 % to of the total nitrate RO₂ loss rate if the FZJ-NO3 mechanism is applied (Fig. 7, A3).
Relative distribution of loss rates of nitrate RO₂ for the experiment on 09 August 2018 (Experiment #1), when HO₂ concentrations were enhanced, and for the experiment on 13 August 2018 (Experiment #4), when the amount of oxidized isoprene was highest. The total RO₂ loss rate was 0.035 and 0.014 s⁻¹ in the experiment on 09 August 2018 (Experiment #1) and 13 August 2018 (Experiment #4), respectively. Calculations of the loss rates of RO₂ radicals in bimolecular reactions make use of measured HO₂ and NO₃ concentrations. Total RO₂ concentrations and concentrations of speciated nitrate RO₂ were derived from model calculations applying either the FZJ-NO₃, CalTech or MCM mechanism. The chemical mechanisms differ with respect to the number of nitrate RO₂ isomers that are considered, the type of RO₂ loss reactions and products of loss reactions (Fig. 3 and 6). Reactions leading to nitrate alkoxy radicals are indicated by a dotted pattern.

Rate constants of RO₂ + RO₂ reactions for nitrate RO₂ in the CalTech mechanism were derived from the measurement of isomer specific product distributions in the experiments of Schwantes et al. (2015). From their findings, a low reaction rate constant of \(3 \times 10^{-16} \text{ cm}^3 \text{s}^{-1}\) for the recombination self- and cross-reaction of the most abundant nitrate β-1,2-RO₂ radical was found, orders of magnitude lower than the generic rate constant used in the MCM of \(1.3 \times 10^{-12} \text{ cm}^3 \text{s}^{-1}\). This rate refers to a tertiary radical instead of a primary, it is significantly slower than the rate constant used in the MCM of \(1.3 \times 10^{-12} \text{ cm}^3 \text{s}^{-1}\). Rate constants for other nitrate RO₂ were estimated in the CalTech mechanism to be in the range of \(10^{-12} \text{ and } 10^{-13} \text{ cm}^3 \text{s}^{-1}\). These values are also implemented in the FZJ-NO₃ mechanism, all the rates for the nitrate RO₂ self- and cross-reactions were calculated from structure activity relationship (Jenkin et al., 2019) resulting in an even lower rate constant for the self- and cross-reaction of the tertiary β-1,2-RO₂ of only \(3 \times 10^{-16} \text{ cm}^3 \text{s}^{-1}\) and for the cross-reactions of this radical with other primary nitrate RO₂ of 2 to \(10 \times 10^{-14} \text{ cm}^3 \text{s}^{-1}\). The rates of the reactions within the pool of the other nitrate RO₂ are on the same order of magnitude as the values in the Caltech mechanism.

Only RO₂ concentrations derived from model calculations can be used to estimate the loss rate of nitrate RO₂ in RO₂ + RO₂ reactions (=alkoxy radical production rate) because the instrument detecting RO₂ could only measure low limit
concentrations (Vereecken et al., 2021). This gives average RO₂ loss rates between 0.0005 s⁻¹ and 0.002 s⁻¹. The contribution to the total loss rate is less than 10% in the experiments on 09, 10, 12 August 2018 (Experiment #1, #2, #3, #4) but increased to up to 20% in the experiment on 13 August 2018 (Experiment #4), when also the production rate of nitrate RO₂ was highest (Fig. 7).

A yield of 60% for the formation of alkoxy radicals (Fig. 6, Reaction R4) is generally applied for RO₂+RO₂ radical reactions for primary and secondary RO₂ (Jenkin et al., 2019). In the case of the most abundant nitrato-organic peroxy radical (β-RO₂-tertiary β-1,2-RO₂) from the reaction of isoprene with NO₃, however, the yield is nearly 100% for its self-reaction and 80% if this nitrate RO₂ reacts with other RO₂ because the formation of a nitrate carbonyl product (NC₄CHO) is not possible (Fig. 6, Reaction R5). Formation of peroxides (ROOR) is considered in the Caltech and FZJ-NO3 mechanisms with a small yield of 3.5%. The MCM does not distinguish between nitrate RO₂ isomers. Therefore, this increase in the yield of alkoxy radicals is only implemented in the CalTech and FZJ-NO3 mechanisms. With respect to the total yield of alkoxy radicals, the high yield for the β-RO₂ is partly compensated by the lower rate constants of RO₂+RO₂ radical reactions in the FZJ-NO3 and CalTech mechanisms compared to the MCM.

As discussed in Schwantes et al. (2015), reactions of nitrate β-RO₂ and HO₂ can also result in the formation of nitrate alkoxy radicals together with an OH radical (Fig. 6, Reaction R2). A yield of 50% is assumed in the CalTech and FZJ-NO3 mechanisms (Section 5.3).

Overall, the total yield of alkoxy radicals produced in the reactions of nitrate RO₂ differ significantly between the 3 mechanisms. In the FZJ-NO3 mechanism, the total yield is around 50%. The value is similar in all experiments analysed in this work, but the type of reactions producing the alkoxy radicals shifts depending on the availability of reaction partners in bimolecular reactions of RO₂ radicals (Fig. 7). Lower yields of alkoxy radicals are between 25 and 40% achieved if lower in the CalTech mechanism than in the FZJ-NO3 mechanism is applied. The lower value compared to the FZJ-NO3 mechanism value is mainly due to the shift in the RO₂ isomer distribution towards δ-RO₂ isomers. Lowest total yields of alkoxy radicals between 7 and 40% are obtained if the MCM is applied because the MCM does not include alkoxy radical production from the reaction of nitrate RO₂ with HO₂.

5.3 Fate of nitrate alkoxy radicals

The fate of the alkoxy radicals is very different between the three mechanisms which impacts on the distribution of organic products. In the MCM, the only pathway for nitrate alkoxy radicals produced from isoprene is their decomposition forming a nitrate carbonyl (NC₄CHO) together with an HO₂ radical (Fig. 6, Reaction R7). This pathway is not possible for the alkoxy radical from the β-RO₂ radicals, which are absent in the MCM mechanism but included in the FZJ-NO3 and CalTech mechanisms. Therefore, the overall yield of nitrate carbonyls (NC₄CHO) from the subsequent chemistry of nitrate alkoxy radicals is highest if the MCM mechanism is applied in comparison to the results from the other 2 mechanisms.

In the CalTech mechanism, alkoxy radicals from β-RO₂ radicals decompose exclusively to MVK or MACR together with a formaldehyde and an NO₂ molecule (Fig. 6, Reaction R8 Wennberg et al. (2018)). Therefore, nitrate carbonyl concentrations predicted by the CalTech model are at least a factor of 4 lower compared to those calculated...
when applying the MCM for conditions of the experiments. Small concentrations of nitrate carbonyls are also produced from reactions of nitrate $\delta$-RO$_2$ radicals.

Vereecken et al. (2021) calculated that the ring-closure reactions leading to the formation of nitrate epoxy alkyl radicals are much faster than the decomposition reaction for the nitrate $\beta$-RO alkoxy isomer conformer (Fig. 6, Reaction R9, R17), so that MVK and MACR production from this reaction is suppressed and this becomes the predominant reaction pathway. Products from this reaction the epoxide pathway are discussed in Section 5.4. Differences between NC$_4$CHO concentrations predicted by the FZJ-NO3 and CalTech Caltech mechanism are due to differences in the initial distribution of nitrate RO$_2$ isomers. The FZJ-NO3 mechanism favours the $\beta$-1,2-RO$_2$ radicals (Section 3) that do not produce NC$_4$CHO and overall react slower with other RO$_2$ compared to than with the other nitrate RO$_2$ radicals.

The VOCUS PTR-MS instrument detected ion signals at the expected mass of NC$_4$CHO with the sum formula C$_5$H$_7$NO$_4$ in all experiments. Due to the lack of calibration, this measurement cannot be used to test the validity of any of the three chemical mechanisms. However, NC$_4$CHO concentrations would be roughly consistent with predictions by the CalTech Caltech and FZJ-NO3 mechanisms if a sensitivity similar to that for ketones without nitrate functional groups (acetone, MVK, pentanone, nopinone) is assumed.

MVK and MACR are formed in all three mechanisms from the oxidation of isoprene by OH and ozone. Yields from the ozonolysis of isoprene are 0.17 and 0.41 for MVK and MACR, respectively (Nguyen et al., 2016). In the absence of NO as for typical night-time conditions, MVK and MACR are produced from the reaction of OH derived RO$_2$ radicals with other RO$_2$ or HO$_2$ radicals. The overall yield of NO$_2$ MVK from the OH oxidation of isoprene in experiments in this work depends on the fate of RO$_2$ radicals, but is expected to be small due to the slow RO$_2$ + RO$_2$ reaction rate and small yields in the range of a few percent from the RO$_2$ + HO$_2$ reaction (Wennberg et al., 2018). In addition to the production from OH and O$_3$ reactions, the CalTech Caltech mechanism includes a strong source for MVK through the decomposition of nitrate $\beta$-1,2-RO$_2$ radicals produced from the NO$_3$ oxidation.

The VOCUS PTR-MS was calibrated for MVK and MACR, so that model calculations can be compared to measurements. In all experiments, analysed in this work, measured MVK and MACR concentrations are consistent with predictions by the MCM and FZJ-NO3 mechanisms (Fig. 4, 5). In contrast, predictions by the CalTech Caltech mechanism are up to a factor of 2 to 4 higher than measured values. Discrepancies are highest in experiments in which a high fraction of the nitrate alkoxy radicals are formed from the reaction of nitrate RO$_2$ with NO$_3$ with an alkoxy radical yield of 1 (13 August 2018, Fig. 5) and are lowest in the experiment in which nitrate RO$_2$ mainly reacted with HO$_2$ with an alkoxy radical yield of only 0.5 (09 August 2018, Fig. 4). The good model-measurement agreement for MVK+MACR concentrations obtained using the FZJ-NO3 and MCM mechanisms confirms demonstrates that the decomposition of the nitrate alkoxy radicals is negligible as predicted by Vereecken et al. (2021) and unlike predicted by the CalTech mechanism. Production of MVK and MACR from the decomposition of nitrate alkoxy radicals from isoprene (as implemented in the CalTech mechanism) does not play a role as calculated by Vereecken et al. (2021b).

The high yield of MVK and MACR from the decomposition of $\beta$-RO radicals in the CalTech mechanism was derived from chamber experiments in Schwantes et al. (2015). In their experiments, 54 to 74 % of the nitrate RO$_2$ reacted with HO$_2$, so that
the majority of alkoxy radicals was formed from this reaction. MVK and MACR concentrations, however, were only measured in 2 experiments in Schwantes et al. (2015), one of which was used to determine the MVK and MACR yields from the reaction of HO$_2$ + RO$_2$. The overall yield of the sum of MVK and MACR was relatively low with a value of approximately 15 %. In order to determine the yield of MVK and MACR from the decomposition of alkoxy radicals from the RO$_2$ + HO$_2$ reactions, production from the isoprene oxidation by OH and O$_3$ and from the potential decomposition of alkoxy radicals produced from other reaction channels needed to be subtracted. In the experiments of Schwantes et al. (2015), OH concentrations were not measured, but measurements of organic compounds such as isoprene hydroperoxides (ISOPOOH) were taken as indicator for OH oxidation. The authors used model calculations to estimate the actual OH concentration and therefore MVK and MACR production from OH chemistry. However, as pointed out in Schwantes et al. (2015), this method has a high uncertainty and measured ISOPOOH concentrations tended to be underestimated by the model calculations. This can imply that OH chemistry could have been more important than implied by the model and that the MVK and MACR yield from the decomposition of alkoxy radicals in the experiment could be more uncertain than estimated in Schwantes et al. (2015).

MVK and MACR concentrations were also measured in an experiment in the SAPHIR chamber reported by Rollins et al. (2009), in which low reactant concentrations were present as in this work (10 ppbv isoprene, 20 to 30 ppbv NO$_2$, 40 to 60 ppbv O$_3$). According to model calculations in Rollins et al. (2009) using the MCM 3.2, the fate of nitrate RO$_2$ radicals from isoprene with NO$_3$ was dominated by their reactions with HO$_2$. Measured MVK and MACR concentrations were consistent with the production of MVK and MACR mainly from the ozonolysis of isoprene. Therefore, this result supports that MVK and MACR are not produced from the decomposition of alkoxy radicals from β-RO$_2$ radicals.

In the chamber experiments of Kwan et al. (2012), the fate of nitrate RO$_2$ was dominated by reactions with other RO$_2$ and with NO$_3$, both of which also produce nitrate alkoxy radicals. The total yield for MVK and MACR determined in these experiments was 6 %. The authors state that all measured MVK and MACR could be explained by production from OH reactions because no ozone was present. The low yield of MVK and MACR appears to be inconsistent with the production of MVK and MACR from the decomposition of alkoxy radicals from β-RO$_2$ radicals as proposed in Schwantes et al. (2015). This is further supported by other experiments investigating the reaction of isoprene with NO$_3$ at high reactant concentrations (Barnes et al., 1990; Kwok et al., 1996; Perring et al., 2009), in all of which the yield of MVK and MACR was maximum in the range of a few percent.

5.4 Epoxide products from ring-closure reactions of nitrate alkoxy radicals observed in the experiments in the SAPHIR chamber

Epoxide formation from ring-closure reactions of nitrate alkoxy radicals leading to epoxy-RO$_2$ radicals is only implemented in the FZJ-NO3 mechanism (Fig. 6, Reaction R9, R17, Vereecken et al. (2021)).

Nitrate epoxides can be directly formed from bimolecular reactions of epoxy-RO$_2$ radicals with RO$_2$ and HO$_2$ (Fig. 6, Reaction R12, R16, R19, R26) and from nitrate epoxy alkoxy radicals, epoxy-RO, produced by the reaction of epoxy-RO$_2$ radicals with NO$_3$ (Fig. 6, Reaction R13, R21). One of the epoxy-RO radicals exclusively undergoes a 1,5-H-shift reaction for conditions of the experiments and decomposes to an epoxide and NO$_2$ (Fig. 6, Reaction R14). Another epoxy-RO radical can
decompose into a C\textsubscript{5} nitrate epoxide releasing HO\textsubscript{2} (Fig. 6, Reaction R22). This reaction competes with a 1,5-H-shift reaction, in which a C\textsubscript{4} nitrate together with an HO\textsubscript{2} radical and formaldehyde (HCHO) are formed (Fig. 6, Reaction R23).

Epoxy-RO\textsubscript{2} can also undergo unimolecular reactions (Vereecken et al., 2021) that compete with bimolecular reactions. The fastest unimolecular reaction is a 1,6-H-shift reaction with a rate constant of $3.7 \times 10^{-3}$ s\textsuperscript{-1} at room temperature leading to a C\textsubscript{5} epoxy product (C\textsubscript{5}H\textsubscript{8}O\textsubscript{4}) together with NO\textsubscript{2} (Fig. 6, Reaction R11). This loss rate is lower than the loss rate due to bimolecular reactions, which are on the order of $10^{-2}$ s\textsuperscript{-1} for conditions of the experiments in this work but high enough that low concentrations of this epoxide product may be formed (Fig. 5, Panel (f)).

The VOCUS PTR-MS instrument detected product species that have the same sum formula as the epoxide products that are expected to be formed subsequent to the ring-closure reaction of alkoxy radicals (Fig. 6). Hydroxy nitrate epoxides formed from the reaction of epoxy-RO\textsubscript{2} radicals with other RO\textsubscript{2} radicals have the same sum formula, C\textsubscript{5}H\textsubscript{9}NO\textsubscript{5}, as nitrate hydroperoxides (NISOPOOH) produced from the reaction of HO\textsubscript{2} with nitrate RO\textsubscript{2} from the initial reaction of isoprene with NO\textsubscript{3}. Because of their isobaric nature, the mass spectrometer instruments cannot distinguish between hydroxy nitrate epoxides formed from the reaction of epoxy-RO\textsubscript{2} radicals with other RO\textsubscript{2} radicals and nitrate hydroperoxides (NISOPOOH) because both have the same sum formula, C\textsubscript{5}H\textsubscript{9}NO\textsubscript{5}. The concentration of epoxide C\textsubscript{5}H\textsubscript{9}NO\textsubscript{5} species is expected to be at most 30 to 40\% of the concentration of NISOPOOH in the experiment on 13 August 2018 (Fig. 5), when RO\textsubscript{2} concentrations were highest. Their concentration is expected to be less than 10\% of NISOPOOH in the experiment on 09 August 2018 (Fig. 4, Panel (a)), when RO\textsubscript{2} reactions with HO\textsubscript{2} dominated the overall RO\textsubscript{2} loss. Therefore, ion mass signals corresponding to C\textsubscript{5}H\textsubscript{9}NO\textsubscript{5} species cannot be used to estimate the importance of the epoxidation reaction pathways.

Bimolecular reactions of epoxy-RO\textsubscript{2} can also lead to the formation of products with sum formulas that are specific for the epoxidation chemistry. Different isomers conformers of nitrate carbonyls with the sum formula C\textsubscript{5}H\textsubscript{7}NO\textsubscript{5} are produced from reactions of epoxy-RO\textsubscript{2} with other RO\textsubscript{2} radicals or with NO\textsubscript{3} (Fig. 6, Reaction R19, R22). In addition, C\textsubscript{5}H\textsubscript{9}NO\textsubscript{6} compounds are formed from reactions of nitrate epoxy-RO\textsubscript{2} with HO\textsubscript{2} (Fig. 6, Reaction R12, R26). Mixing ratios of these epoxides are calculated to be highest with mixing ratios of 1 ppbv in the experiment on 13 August 2018 (Experiment #4), when the total isoprene consumption by NO\textsubscript{3} reactions was highest. Values are similar to mixing ratios of other products obtained in this experiment (Fig. 5, Panel (g)).

The mass spectrum measured by the VOCUS PTR-MS instrument shows clear signals at the masses of these compounds. The count rates are much lower compared to signals of other products, although expected concentrations are in the same range. This could be due to a lower sensitivity of the instrument for nitrate epoxides compared to the sensitivity than for other organic nitrates. However, this could also indicate a lower than assumed production rate of alkoxy radicals for example from the reaction of nitrate RO\textsubscript{2} with HO\textsubscript{2} (Section 5.6).

A C\textsubscript{4}-nitrate with the sum formula C\textsubscript{4}H\textsubscript{5}NO\textsubscript{4} produced subsequent to the 1,5-H reaction of the nitrate alkoxy radical (Fig. 6, Reaction R23–R25) was not detected by the VOCUS PTR-MS instrument in the experiments in this work, though significant mixing ratios of up to 0.6 ppbv are calculated by the FZJ-NO3 mechanism in the experiment on 13 August 2018 (Experiment #4), when the amount of oxidized isoprene was highest (Fig. 5, Panel (i)). There is no obvious reason why the sensitivity of
instrument for this compound would be lower compared to other compounds. Only the I− CIMS instrument detected a very small signal (less than 30 cnts) at the corresponding mass, which is at least a factor of 100 smaller than ion signals of masses at other products shown in Fig. A10).

The formation of this compound competes with the decomposition of the epoxy alkoxy radical leading to an epoxy-C\textsubscript{5} compound with the sum formula C\textsubscript{5}H\textsubscript{7}NO\textsubscript{5} that is observed in the mass spectrum of the VOCUS PTR-MS instrument (Fig. 6, Reaction R22). The fact that the C\textsubscript{4} nitrate is not observed in the mass spectrum could indicate that the 1,5-H reaction is not competitive or that the branching ratio of 2 epoxy-alkyl radicals from the nitrate alkoxy radical disfavours the epoxy-alkyl radical that eventually leads to the formation of the C\textsubscript{4} nitrate (Fig. 6, Reaction R17). Rate constants of the epoxidation chemistry calculated in Vereecken et al. (2021) have an uncertainty of a factor of 2 to 4. Therefore, low rate constants that weaken the formation of the C\textsubscript{4} nitrate are within in the uncertainty of calculations.

Two further epoxy compounds which have lost the nitrate functional group by eliminating NO\textsubscript{2} are expected to be formed. One has the sum formula C\textsubscript{5}H\textsubscript{8}O\textsubscript{3} and is a product of a fast 1,5-H-shift reaction of an epoxy alkoxy radical (Fig. 6, Reaction R14). The sum formula is the same as hydroperoxy aldehydes (HPALD) that are formed from 1,6-H-shift reactions of the primary nitrate Z-δ-RO\textsubscript{2} radical (Fig. 3). The contribution of the epoxy species to the sum of epoxy compounds and HPALD is calculated to be small with values of less than 15% in all experiments. The VOCUS PTR-MS instrument was not calibrated for either one of the compounds, so that it is not clear if the epoxy compound was detected.

The other epoxy compound without a nitrate functional group is produced from a 1,6-H-shift reaction of one of the nitrate epoxy-RO\textsubscript{2} (Fig. 6, Reaction R11). Due to the relatively low reaction rate constant, only small mixing ratios of maximum 0.15 ppbv of this compound with the sum formula C\textsubscript{5}H\textsubscript{8}O\textsubscript{4} are modelled in the experiment on 13 August 2018 (Experiment #4) (Fig. 5, Panel (f)). Nevertheless, a corresponding signal is observed in the mass spectrum of the VOCUS PTR-MS instrument that can be attributed to this compound.

### 5.5 Epoxide products from nitrate alkoxy radicals observed in previous chamber experiments

In chamber experiments by Kwan et al. (2012), products of the reaction of isoprene with NO\textsubscript{3} were detected by a chemical ionization spectrometer using CF\textsubscript{3}O− as reagent ions. These experiments were performed under conditions where the majority of nitrate RO\textsubscript{2} reacted with other RO\textsubscript{2} radicals. Chamber experiments reported in Schwantes et al. (2015) were performed at lower concentrations and HO\textsubscript{2} concentrations were enhanced, so that reactions of RO\textsubscript{2} with HO\textsubscript{2} were favoured. Similar to the experiments in this work, products that have the sum formulas of nitrate epoxide products expected to be formed in the FZJ-NO3 mechanism were observed in the experiments in Kwan et al. (2012) and Schwantes et al. (2015): (1) C\textsubscript{5}H\textsubscript{9}NO\textsubscript{6} compounds, which appear at the same mass as NISOPOOH, (2) C\textsubscript{7}H\textsubscript{7}NO\textsubscript{5} compounds from epoxy-RO\textsubscript{2} + RO\textsubscript{2} reactions, (3) C\textsubscript{5}H\textsubscript{9}NO\textsubscript{5} compounds from epoxy-RO\textsubscript{2} + HO\textsubscript{2} reactions.

In Kwan et al. (2012) and Schwantes et al. (2015) it is suggested that the product with the sum formula C\textsubscript{5}H\textsubscript{9}NO\textsubscript{6} is a hydroxy hydroperoxy nitrate and that the product with the sum formula C\textsubscript{7}H\textsubscript{7}NO\textsubscript{5} is a hydroxy carbonyl nitrate. Both compounds are suggested to be products from a 1,5-H shift reaction of the nitrate alkoxy radicals from the bimolecular reactions of δ-RO\textsubscript{2} radicals. The authors could not estimate a reaction rate constant for the 1,5-H shift reaction from their
experiments. Vereecken et al. (2021b) calculated a reaction rate of $2.2 \times 10^6 \text{ s}^{-1}$ ($T = 298$ K), which makes the 1,5-H-shift reaction too low to compete with the ring-closure reaction forming epoxy alkyl radicals ($1.2 \times 10^6 \text{ s}^{-1}$, $T = 298$ K) and subsequent O₂ addition. Therefore, C₅H₇NO₆ and C₅H₉NO₆ compounds observed in Kwan et al. (2012) and Schwantes et al. (2015) could rather be nitrate epoxy product instead of the products suggested by the authors. It is worth noting that compounds suggested by Kwan et al. (2012) and Schwantes et al. (2015) would only be produced from nitrate $\delta$ RO₂ radicals that have small yields, whereas the nitrate epoxy products in the FZJ-NO3 mechanism are also produced from the most abundant nitrate $\beta$ RO₂ radicals. This may also explain why compounds with these sum formulas were clearly detected in experiments from all studies.

In the experiments in Kwan et al. (2012) and Schwantes et al. (2015), the C₅H₇O₅ compound without a nitrate functional group was observed, which is consistent with observations in this work. Because HPALD appears at the same mass and HPALD is also produced from OH oxidation, the authors concluded that C₅H₇O₅ is a product from the reaction of isoprene with OH. Nevertheless, their observations of C₅H₇O₅ compounds could also be partly due to the production of epoxy species from the oxidation of isoprene by NO₃ as described in the FZJ-NO3 mechanism.

The other product without a nitrate group that is produced from the ring-closure pathway of nitrate alkoxy radicals in the FZJ-NO3 mechanism, C₅H₈O₃, was not observed in the experiments in Kwan et al. (2012) and Schwantes et al. (2015). The reason could be that the chemical lifetime of RO₂ radicals was too short in the experiments in Kwan et al. (2012), in which high concentrations of reactants were present, so that the 1,6-H-shift reaction of the epoxy-RO₂ radical producing the C₅H₈O₃ compound could not compete with bimolecular reactions. Similarly, RO₂ reactions with HO₂ were favoured in the experiments in Schwantes et al. (2015), so that the 1,6-H reaction may have not been competitive.

Interestingly, similar to the experiments in this work, no organic nitrate with the sum formula C₄H₅NO₄ that is expected to be formed from the ring-closure reactions of nitrate alkoxy radicals (Fig. 6) was observed in the experiments in Kwan et al. (2012) and Schwantes et al. (2015). This further suggests that there is no significant production of this compound.

Overall, observations in this work and in previous chamber experiments are consistent with the formation of epoxy organic compounds, which are expected to be formed from the ring-closure reaction of nitrate alkoxy radicals as calculated in Vereecken et al. (2021b) and implemented in FZJ-NO3 mechanism.

### 5.6 Reaction of nitrate RO₂ with HO₂ Production of nitrate hydroperoxide (NISOPOOH)

The chemical loss rates of nitrate RO₂ towards reaction with HO₂ was 0.032 s⁻¹ (90 % of the total loss rate) in the experiment with high HO₂ concentrations (09 August 2018, Experiment #1). The contribution to the total loss rate was 40 to 50 % with loss rates between 0.002 and 0.007 s⁻¹ in the other experiments (Fig. 6). In general, this reaction can proceed via several reaction pathways (Rollins et al., 2009; Kwan et al., 2012; Schwantes et al., 2015):

\[
\text{nitrte} - \text{RO₂} + \text{HO₂} \rightarrow \text{ROOH} + \text{O₂} \quad (R1)
\]
\[
\text{nitrte} - \text{RO₂} + \text{HO₂} \rightarrow \text{RO} + \text{OH} + \text{O₂} \quad (R2)
\]
Nitrate hydroperoxide (NISOPOOH) is the only product in the MCM (Reaction R1) and a major product in the CalTech and FZJ-NO3 chemical mechanisms (Fig. 6, Reaction R1). The CalTech and FZJ-NO3 mechanisms assume that the yield of nitrate alkoxy radicals is approximately 0.5 if nitrate \( \beta\)-RO\(_2\) radicals react with HO\(_2\) (Reaction R2). The fate of nitrate alkoxy radicals is discussed above (Section 5.3). Predictions of NISOPOOH concentrations by the three mechanisms differ significantly with highest concentrations if the MCM is applied. NISOPOOH concentrations predicted by the FZJ-NO3 mechanism are approximately half of the concentration calculated by the MCM and concentrations predicted by the CalTech mechanism are between both values. This is mainly due to the different distribution of nitrate \( \beta\)- and \( \delta\)-RO\(_2\) radicals in the FZJ-NO3 and CalTech mechanisms.

Signals at the mass corresponding to NISOPOOH were highest among all product signals observed by the VOCUS PTR-MS instrument. The signal can include nitrate epoxides that are produced from the ring-closure reactions of alkoxy radicals (Section 5.3) and the reaction of NISOPOOH with OH, which have the same mass. However, their contribution is expected to be low for most of the time in the experiments in this work, specifically in the experiment on 09 August 2018, when HO\(_2\)-concentrations favoured RO\(_2\)+HO\(_2\) reactions and an OH scavenger was present (Fig. 4).

The VOCUS PTR-MS instrument was not calibrated for NISOPOOH, so that its concentrations could not be determined. The high count rate and the uncertainty in the branching ratio of Reactions R1 and R2 appear to support a high yield of NISOPOOH from the reaction of HO\(_2\) with nitrate RO\(_2\).

NISOPOOH has been detected by mass-spectrometer instruments in previous chamber studies by Ng et al. (2008), Kwan et al. (2012) and Schwantes et al. (2015). Similar to this work, the instruments were not calibrated for NISOPOOH, but the sensitivity of the instrument was calibrated for nitrate alcohols (ISOPCNO\(_2\)). The sensitivity for other organic nitrates like NISOPOOH was estimated from calculations of the dipole moment and polarisability (Ng et al., 2008; Kwan et al., 2012; Schwantes et al., 2015).

In the experiments of Schwantes et al. (2015), HO\(_2\) concentrations were enhanced. NISOPOOH yields were between 0.32 and 0.41, when 54 and 76\% of the nitrate RO\(_2\) were calculated to react with HO\(_2\). The authors calculated that these yields are consistent with a 50\% branching ratio of the reaction of nitrate RO\(_2\) with HO\(_2\) (Reaction R2) to form alkoxy radicals. An uncertainty of \( \pm 20\% \) of the measured NISOPOOH concentration is stated. The uncertainty of the alkoxy radical yield, however, could be higher because the calculation also requires the knowledge of the fraction of isoprene that reacted with NO\(_3\) and the fraction of RO\(_2\) that reacted with HO\(_2\), both of which are uncertain because NO\(_3\) and HO\(_2\) concentrations were not measured. Therefore, a NISOPOOH yield of the reaction of nitrate RO\(_2\) with HO\(_2\) higher than 50\% may also be consistent with the experimental results in Schwantes et al. (2015).

Ng et al. (2008) quantified NISOPOOH concentrations in their chamber experiment, which was performed at high concentrations of reactants (800 ppbv isoprene, 120 ppbv N\(_2\)O\(_5\)). They determined that 50\% of the reacted isoprene resulted in the formation of NISOPOOH, but the fraction of nitrate RO\(_2\) that reacted with HO\(_2\) could not be determined to calculate yields from specific reactions. Therefore, their experiments cannot be used to derive information about potential alkoxy radical formation from the reaction of RO\(_2\) with HO\(_2\). HO\(_2\) concentrations in experiments in Kwan et al. (2012) were presumably small because high
reactant concentrations were used. This explains the relatively small overall NISOPOOH formation of 10% from the reaction of isoprene with NO3.

### 5.7 Production of OH radicals

Alkoxy radical formation from the reaction of nitrate RO2 with HO2 is accompanied by the formation of OH (Fig. 6, Reaction R2), which can be responsible for the formation of products that are specific for the OH oxidation of isoprene that are as observed in experiments that are designed to investigate the NO3 oxidation mechanism of isoprene.

Kwan et al. (2012) assumed that specific C5 organic compounds (HPALD, NISOPOOH, C5 hydroxy carbonyl C5H9O2) and MVK and MACR, all of which were quantified in their chamber experiments, were exclusively formed from OH radicals that are formed as co-product of alkoxy radicals. In this case, the yield of nitrate alkoxy radical formation competing with the formation of NISOPOOH in the reaction of nitrate RO2 with HO2 is 38 to 58%. Although the experiments were performed in the absence of ozone, so that OH was not produced by ozonolysis reactions, this approach gives only an upper limit of the yield because OH as well as some of the organic products may not have been exclusively produced by this assigned reaction pathway. For example, HPALD can also be produced from the oxidation of isoprene by NO3 from 1,6-H reactions of nitrate RO2 (Vereecken et al. (2021b), Fig. 3, Section 5.8).

OH concentrations were measured in the experiments in this work, but concentrations were around the limit of detection of the instrument (a few 10^5 cm^-3) in most experiments. Model calculations for the experiment on 13 August 2018 (Experiment #4), when reactant concentrations were highest, result in significant OH concentrations between 5 and 8 × 10^5 cm^-3 and also model results indicate that OH concentrations could have been in the range of a few 10^5 cm^-3 (Fig. 5, Panel (j)). A large fraction of OH, however, is produced by the reaction of HO2 with NO3, both of which are constrained to measured values in the model calculations. As discussed in Vereecken et al. (2021), model calculations without constraining HO2 to measured values cannot reproduce measured HO2 concentration suggesting shortcomings of the model to describe HO2 source and/or sink reactions.

This is further analysed by comparing results of model runs, in which either HO2 concentrations are constrained to measurements or HO2 is calculated by the model (Fig. 8). In the unconstrained case, modelled HO2 concentrations are much lower than measurements. This reduces the OH concentration by a factor of 3 due to the lower production of OH from the reaction of HO2 with NO3. During the part of the experiment, when isoprene is oxidized by NO3, differences between measured and modelled OH concentrations tend to be smaller if HO2 is not constrained to measured values. At later times of the experiment after 13:30 UTC, when isoprene had been consumed and NO3 concentrations were enhanced by additional injections of NO2 and O3 (Fig. 2, Panel (b)), measurements show a steeper increase of OH concentrations than model calculations with unconstrained HO2. This further indicates that modelled HO2 concentrations might be too low.

If the yield of alkoxy radicals and therefore also of OH from the reaction of nitrate RO2 with HO2 was lower than 50% as assumed in the Caltech Caltech and FZJ-NO3 mechanisms, modelled OH concentrations would be even lower. Sensitivity model runs show that modelled OH concentrations would only decrease by 1 to 3 × 10^5 cm^-3 directly after the isoprene injections, when nitrate RO2 concentrations are also highest, in this case. However, such differences are in the range of the
Figure 8. Comparison of results from model calculations applying the FZJ-NO3 mechanism for the experiment on 13 August 2018 (Experiment #4) with HO2 concentrations being either constrained or unconstrained to measurements. A large fraction of OH is produced from the reaction of HO2 with NO3, so that lower than measured HO2 concentrations in the unconstrained model run lead to lower OH concentrations compared to model results if HO2 concentrations are constrained to measured values. Because HO2 + RO2 reactions contribute significantly to the total loss of RO2, modelled RO2 concentrations are higher in the unconstrained model run. RO2 measurements by the LIF instrument does not include all RO2 radicals (Vereecken et al., 2021), so that measured concentrations are lower than modelled values.
accuracy of measurements, which was a few $10^5$ cm$^{-3}$ due to the subtraction of an OH background signal that was determined by using a chemical modulation system (Cho et al., 2021).

Overall, considering the uncertainties in the measured OH concentrations and in the modelled OH due to the uncertainty in the OH production from the HO$_2$ + NO$_3$ reaction, differences between model results and measured values are too small to draw conclusions about the yield of alkoxy radicals from model-measurement comparison of OH concentrations.

5.8 Production of hydroperoxy aldehydes (HPALD) from nitrate RO$_2$ isomerization reactions

Only the FZJ-NO3 mechanism includes unimolecular loss reactions of nitrate RO$_2$ (Fig. 3). The reaction rate constants of the 1,6-H shift reactions of the Z-δ-RO$_2$ isomers have a strong temperature dependence (Vereecken et al., 2021). Values range between 0.016 to 0.023 s$^{-1}$ for the Z-δ-1,4-RO$_2$ isomer conformer and 0.045 and 0.06 s$^{-1}$ for the Z-δ-4,1-RO$_2$ isomer conformer for temperatures experienced in the experiments in this work.

Although the fraction of Z-δ-RO$_2$ isomer conformer to the total RO$_2$ is only between 5 and 6% for the Z-δ-1,4-RO$_2$ and between 1 and 2% for the Z-δ-4,1-RO$_2$ isomer conformer, the overall bulk RO$_2$ isomerization rate is around 0.002 s$^{-1}$. This value makes the 1,6-H-shift reaction competitive with bimolecular reactions in all experiments except for the one with high HO$_2$ concentration (09 August 2018, Experiment #1). Its contribution to the overall loss rate is expected to be between 10 and 30% depending on the total RO$_2$ loss rate (Fig. 7). This is similar or even higher compared to analogous, much faster 1,6-H-shift reactions in the OH-initiated isoprene oxidation ($k$(298 K) $\approx$ 0.5 s$^{-1}$, Peeters et al. (2014)) due to significantly longer RO$_2$ lifetimes in the night compared to daytime conditions than during the day.

HPALD concentrations predicted by the model applying the FZJ-NO3 mechanism are between 0.1 and 1.2 ppbv depending on the chemical conditions with different availability of reaction partners for competing bimolecular reactions. Lowest values are obtained in the experiment on 09 August 2018 (Experiment #1), when the total RO$_2$ loss rate due to bimolecular reactions competing with HPALD formation from unimolecular reactions was highest. HPALD mixing ratios are calculated to be highest in the experiment on 13 August 2018 (Experiment #4), when the total concentration of oxidized isoprene was high. Approximately 10 to 15% of the HPALD that is predicted by the FZJ-NO3 mechanism is due to OH oxidation of isoprene also producing HPALD from 1,6-H-shift reactions. This can be seen as an upper limit, however, due to the uncertainty of the modelled OH concentration (Section 5.6). It is worth noting that the fast 1,6-H-shift reactions reactions of Z-δ-RO$_2$ isomers from the OH oxidation of isoprene (bulk loss rate $\approx$ 0.006 s$^{-1}$) makes these reactions very competitive with bimolecular reactions for night-time conditions (loss rate in the experiments in this work: 0.005 to 0.014 s$^{-1}$, Section 5.1).

The VOCUS PTR-MS instrument shows a clear signal on the mass of HPALD. One epoxide product from the ring-closure reactions of nitrate alkoxy radicals (Fig. 6) has the same mass, but its contribution to the sum signal is expected to be small (Fig. 5). Unfortunately, the instrument was not calibrated for HPALD to compare absolute concentrations.

Although the absolute importance of HPALD formation from H-shift reactions of nitrate RO$_2$ radicals is uncertain, HPALD is clearly formed from the oxidation of isoprene by NO$_3$. This is demonstrated by the observation of a signal at the mass of HPALD in the experiment on 09 August 2018 (Experiment #1), when an OH scavenger was present, so that HPALD could not be produced by OH reactions. In this experiment, HO$_2$ + RO$_2$ reactions were favoured, so that also formation of the epoxides
with the same mass is expected to be small (Fig. 4, Panel (c)). Therefore, the signal on the mass of HPALD can be attributed to HPALD formation from the oxidation of isoprene by NO$_3$ in this experiment.

The relative importance of HPALD formation is expected to be highest for conditions of the experiment on 10 August 2018 (Experiment #2), when the total loss rate of RO$_2$ due to bimolecular reaction are calculated to be between 0.005 and 0.006 s$^{-1}$. In this case, approximately 25 to 30% of the isoprene consumed by NO$_3$ would form HPALD. Brownwood et al. (2021) calculated the yield of total organic nitrates from measurements for the same experiments analysed in this work and found a yield of (94 ± 20)% for this experiment. Values ranged between (112 ± 13)% and (140 ± 24)% in the other experiments.

The lowest yield of organic nitrates is obtained in the experiment with the longest RO$_2$ lifetime (10 August 2018, Experiment #2) supporting that more non-nitrate organic products like such as HPALD are formed in this experiment compared to than in the other experiments. The signal of the VOCUS PTR-MS instrument, however, does not clearly scale with the expected differences in the HPALD yield in the experiments in this work. This and the overall high yields of organic nitrates indicate that the impact of unimolecular reactions producing HPALD might be overestimated in the FZJ-NO3 mechanism. Uncertainties in the quantum-chemical calculations, from which reaction rates are taken in the FZJ-NO3 mechanism, are a factor of 2 to 3, so that unimolecular RO$_2$ reaction might be less competitive with bimolecular RO$_2$ reactions for atmospheric conditions like in the experiments in this work.

HPALD was also observed in chamber experiments in Kwan et al. (2012) and Schwantes et al. (2015). The authors attributed the observations to the OH oxidation of isoprene, but their observations could also indicate HPALD formation from nitrate RO$_2$. Specifically in the experiments in Schwantes et al. (2015) the total loss rate of nitrate RO$_2$ were calculated to be in the range of 0.03 to 0.13 s$^{-1}$, so that 1,6-H shift reactions with rates 0.02 and 0.05 s$^{-1}$ ($T = 298$ K) calculated in Vereecken et al. (2021b) can compete with bimolecular loss reactions.

Overall, experiments in this work and previous chamber experiments demonstrate that HPALD formation from 1,6-H shift reactions of Z-$\delta$ RO$_2$ isomers play a role for atmospheric night-time conditions.

### 5.9 Night-time loss

#### Loss rate of organic nitrate products and hydroperoxy aldehydes (HPALD)

Chamber experiments in this work were designed to also investigate further oxidation of the organic products. This was achieved by re-injecting O$_3$ and NO$_2$ to enhance NO$_3$ production after most of the isoprene had reacted away (Fig. 2, 1, Panel (a)). Highest product concentrations were achieved in the experiment on 13 August 2018 (Experiment #4), when the amount of isoprene that was oxidized was highest. Therefore, the further discussion concentrates on this experiment (Fig. 5). In addition, information from the experiment on 09 August 2018 (Experiment #1) (Fig. 4), when an OH scavenger was present, is used to remove the effect of the OH oxidation.

Reaction rate constants of nitrate products from the oxidation of isoprene with OH and O$_3$ implemented in the CalTech mechanism are listed in Wennberg et al. (2018). They are based on laboratory experiments with synthetic standards of isoprene hydroxy nitrate isomers (Lee et al., 2014b). Values are assumed to be applicable for other organic nitrates such as nitrate carbonyls and nitrate hydroperoxides. Specific additional reaction channels increasing the reaction rate constants are
considered. Only part of the loss reactions listed in Wennberg et al. (2018) are implemented in the code of the CalTech mechanism (Bates and Wennberg, 2017) that is applied in model calculations in this work.

Rate constants for the reaction of the first generation organic nitrates with ozone (Reaction R28, R31, R34, R37) are in the range of \(10^{-19}\) to \(10^{-18}\) s\(^{-1}\) cm\(^{-3}\) to \(10^{-17}\) to \(10^{-19}\) cm\(^3\) s\(^{-1}\) in Lee et al. (2014b) with rates being relevant for only \(\delta\) nitrate alcohols and hydroperoxides making them not relevant on the time scale of one night for typical oxidant concentrations during the night. As only \(\delta\) species are implemented in the MCM, the overall relevance of these loss reactions are overestimated under atmospheric conditions in the MCM (Table 2). As assumed in that work, reaction rate constants can be expected to be similar for the different first-generation organic nitrates according to structure activity relationship (Jenkin et al., 2020). Therefore, up to factor of 10 higher reaction rate constants as implemented in the MCM, which would make ozone loss reactions relevant for atmospheric conditions, are not applicable (Table 2).

In the FZJ-NO3 mechanism, reaction rate constants of organic nitrates with \(\text{OH}\) radicals (Reaction R27, R30, R33, R36) are taken from the CalTech mechanism, but rate constants with ozone and \(\text{NO}_3\) are optimized to best describe the temporal behaviour of the signals observed by the VOCUS PTR-MS instrument at the respective mass (Table 2). Reaction rate constants of loss reactions that lead to loss rates much lower than the dilution rate of the chamber are not relevant for the timescale of one night, are set to upper limit values that equal the loss rate due to dilution in the experiments (\(k_{\text{dil}} = 1.5 \times 10^{-5}\) s\(^{-1}\)). Reaction rate constants are likely even lower because doubling the loss rate from dilution would already worsen the model-measurement agreement of the temporal behaviour of products.

Chemical loss of NISOPPOOH by reactions with \(\text{NO}_3\) (Reaction R29) and \(\text{O}_3\) (Reaction R28) are expected not to be relevant for atmospheric conditions in all mechanisms. This is consistent with the slow decay of the total signal for \(\text{C}_5\text{H}_9\text{NO}_5\) observed by the VOCUS PTR-MS instrument in the experiment on 09 August 2018 (Experiment #1), when OH oxidation was suppressed by the presence of an OH scavenger (Fig. 4, Panel (a)). In this case, the loss rate is consistent with the dilution rate in the experiment.

Nitrate hydroperoxides, NISOPPOOH, are expected to react with OH with a fast reaction rate constant of \(10^{-10}\) s\(^{-1}\) cm\(^{-3}\) in the MCM. A 3 times lower reaction rate constant is implemented in the CalTech and FZJ-NO3 mechanisms. Differences in the OH reaction rate constants explain the faster decay of NISOPPOOH predicted by the MCM compared to the CalTech and FZJ-NO3 mechanisms for the experiment on 13 August 2018.

In the MCM, the rate of the reaction of \(\text{OH}\) with hydroperoxides, NISOPPOOH, is assumed to be fast with a rate coefficient of \(10^{-10}\) s\(^{-1}\) cm\(^3\). In contrast, the Caltech and FZJ-NO3 mechanisms assume a smaller rate coefficient for this reaction, by a factor 3, which can account for the faster decay of NISOPPOOH in the MCM mechanism than in the Caltech and FZJ-NO3 mechanisms.

In the MCM, products of the NISOPPOOH + OH reaction (Reaction R27) are a nitrate alkoxy radical together with an OH radical leading to a zero net loss of OH. In addition, the alkoxy radical produces a nitrate carbonyl (NC\(_4\)CHO) together with an \(\text{HO}_2\) (Section 5.3). In contrast, in the CalTech mechanism, a large fraction of the predicted products are epoxide products (yield: 0.37 to 1.0 depending on the precursor \(\text{RO}_2\) isomer conformer, Schwantes et al. (2015)) together

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with OH analogous to the formation of epoxides in the OH oxidation of isoprene (Paulot et al., 2009). The implementation in these mechanisms is based on the observation of epoxides in chamber experiments in Schwantes et al. (2015). Nitrate epoxides have the same sum formula as NISOPOOH (C₅H₇NO₅), so that the VOCUS PTR-MS instrument cannot distinguish between both compounds. The reaction of OH radicals with nitrate epoxides can be expected to be much slower compared to their reaction with NISOPOOH due to the lack of C=C double bonds. Therefore, the sum signal time series of the sum of both compounds is affected by their different temporal behaviour in the Caltech and FZJ-NO3 mechanisms.

The code of the Caltech mechanism (Bates and Wennberg, 2017) includes less loss reactions implemented as described in (Wennberg et al., 2018). Chemical loss of nitrate epoxides are not implemented in the chemical mechanisms.

<table>
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<th>Caltech</th>
<th>FZJ</th>
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*not implemented; ^b $2.8 \times 10^{-19} \text{s}^{-1} \text{cm}^3$, Wennberg et al. (2018); ^c $3.0 \times 10^{-14} \text{s}^{-1} \text{cm}^3$, Wennberg et al. (2018); ^d $4.4 \times 10^{-18} \text{s}^{-1} \text{cm}^3$, Wennberg et al. (2018); ^e $1.1 \times 10^{-13} \text{s}^{-1} \text{cm}^3$, Wennberg et al. (2018); ^f $2.8 \times 10^{-19} \text{s}^{-1} \text{cm}^3$, Wennberg et al. (2018); ^g $3 \times 10^{-14} \text{s}^{-1} \text{cm}^3$, Wennberg et al. (2018)
that for example the loss rate of the C_5H_9NO_5 compounds in the MCM is only affected determined by the fast loss of NISOPOOH because no epoxides are formed.

For the experiment on 13 August 2018 (Experiment #4) (Fig. 5, Panel (a)), the temporal behaviour of the total ion signal for corresponding to C_5H_9NO_5 species observed by the VOCUS PTR-MS instrument fits best the modelled trace of the FZJ-NO3 mechanism with the low OH reaction rate of NISOPOOH. In addition, the slow decay low chemical loss rate of epoxides improves contributes to the slow decay of the ion signal at that mass improving the model-measurement agreement. This demonstrates that OH reaction rate constants measured in Lee et al. (2014b) for nitrate alcohols can be applied to NISOPOOH as implemented in the CalTech Caltech and FZJ-NO3 mechanisms. In contrast, the fast OH reaction rate constant for NISOPOOH implemented in the MCM cannot describe the observations.

If the MCM mechanism is used, a significant fraction of nitrate carbonyls, NC_4CHO, that are produced from nitrate RO_2 + RO_2 reactions and from the decomposition of specific nitrate alkoxy radicals is expected to be consumed on the time scale of the experiment for the experiment on 13 August 2018 (Experiment #4) (Fig. 5, Panel (d)). For conditions of this experiment, reactions of NC_4CHO with OH (Reaction R30) but also with NO_3 (Reaction R32) for high NO_3 concentrations can be relevant if reaction rate constants of the MCM are applied (Table 2). The faster loss of NC_4CHO calculated using in the MCM compared to using the CalTech Caltech and FZJ-NO3 mechanisms can be partly explained by up to a factor of 3 lower OH reaction rate constants for some isomers conformers of NC_4CHO. In addition, the MCM overestimates the loss of NC_4CHO by the reaction with ozone as discussed above.

The temporal behaviour of the modelled NC_4CHO concentrations are in good agreement with the corresponding signal observed by the VOCUS PTR-MS instrument for the CalTech Caltech and FZJ-NO3 mechanisms. This confirms that there is no significant chemical loss of NC_4CHO only a small fraction of NC_4CHO is expected to be chemically lost for typical night-time conditions in contrast to results if reaction rate constants implemented in the MCM are used.

In addition, a fast loss rate due to the reaction with NO_3 (Reaction R32) as suggested in Wennberg et al. (2018) would lead to a chemical lifetime of NC_4CHO of less than 30 min in the last phase of the experiment on 13 August 2018 (Experiment #4), when NO_3 mixing ratios increased to several 100 pptv (Fig. 2, Panel (b)), but this is not observed (Fig. 5, Panel (d)). Though not fully applicable, structure activity relationship in Kerdouci et al. (2014) gives reaction rate constants lower than 10^{-16} s^{-1} cm^{-3} due to the carbonyl group in the \( \beta \)-position of the C=C double bond supporting the low loss rate due the addition of NO_3. Overall, further oxidation of nitrate carbonyls from isoprene is of minor importance for typical night-time conditions as experienced in these experiments is not relevant.

Similar differences between model predictions in the temporal behaviour like such as for NC_4CHO are seen for nitrate alcohols (ISOPCNO_3, Reaction R33, 34, 35): the MCM predicts a significant faster chemical loss compared to than the CalTech Caltech and FZJ-NO3 mechanisms. A large part of the discrepancy is explained by the fast loss due to the reaction with ozone implemented in the MCM that is not applicable as discussed above. In addition, the reaction rate constant of the reaction of ISOPCNO_3 with OH (Reaction R33) is up to 3 times faster in the MCM compared to than in the CalTech Caltech and FZJ-NO3 mechanisms (Table 2). The good agreement of the temporal behaviour of the signal of the VOCUS PTR-MS
instrument at the mass of ISOPCNO₃ confirms also the low reaction rate constants with OH determined experimentally in Lee et al. (2014b).

HPALD formation from the reaction of isoprene with NO₃ is only implemented in the FZJ-NO3 mechanism. Wolfe et al. (2012) investigated the photo-oxidation of a closely-related compound of HPALD to constrain photolysis rates and reaction rate constants in the reaction with OH (Reaction R36) and O₃ (Reaction R37). A fast reaction rate constant \(5.1 \times 10^{-11} \text{ s}^{-1} \text{cm}^3\) was found. This value is implemented in the MCM, CalTech Caltech and FZJ-NO3 mechanisms (Table 2).

In the MCM, a fast reaction rate constant of HPALD with ozone (Reaction R37) is implemented, which would lead to a short chemical lifetime of 4.6 h for conditions of the experiment in this work (100 ppbv O₃). In addition, the MCM assumes that HPALD reacts with NO₃ (Reaction R38) with a fast reaction rate constant of \(1.2 \times 10^{-14} \text{ s}^{-1} \text{cm}^3\), which would lead to a significant loss of HPALD in the last part of the experiment on 13 August 2018 (Experiment #4), when NO₃ mixing ratios were very high up to several 100 pptv. These assumptions about the reaction rate constants of HPALD with O₃ and NO₃ are inconsistent with the observed temporal behaviour of the signal at the mass of HPALD observed by the VOCUS PTR-MS instrument, which is explained by the loss of HPALD by only its reaction with OH (Fig. 5, Panel (c)). In the experiments on 09 August 2018 (Experiment #1), when OH reactions were suppressed by the presence of the OH scavenger, the temporal behaviour of the HPALD signal is fully consistent with only the loss due to dilution (Fig. 4, Panel (c)).

The reaction rate constant of HPALD with ozone was experimentally determined in Wolfe et al. (2012) to be \(1.2 \times 10^{-18} \text{ s}^{-1} \text{cm}^3\) making the ozone reaction irrelevant for typical atmospheric conditions. Results in the experiments in this work confirm this low value. The temporal behaviour of HPALD implicates that also the reaction of HPALD with NO₃ does not significantly contribute to its chemical loss for typical night-time conditions. There are no experimental values for the reaction rate constant of HPALD with NO₃. Structure activity relationship (SAR) like described in Kerdouci et al. (2014) cannot be applied because the effect of a COOH substituent in the β-position of the C=C double at which NO₃ adds is not considered. Omitting this substituent results in a reaction rate constant similar to the value in the MCM, indicating that a COOH substituent further lowers the reaction rate constant.

The further oxidation of epoxides produced from ring-closure reactions of nitrate alkoxy radicals calculated in Vereecken et al. (2021) have not been investigated so far. The temporal behaviour of signals measured by the VOCUS PTR-MS instrument suggests that their loss rate can be explained by only the dilution rate in the experiments indicating that chemical loss was not significant even in the presence of several 100 pptv NO₃, 100 ppbv O₃ and presumably several \(10^5\) OH in the last period of the experiment on 13 August 2018 (Experiment #4) (Fig. 5, Panel (j)). An upper limit value of the reaction rate constant of the reaction of epoxides with OH of \(1.2 \times 10^{-11} \text{ s}^{-1} \text{cm}^3\) \((T = 298 \text{ K})\) can be assumed similar to the value found for epoxides produced from the reaction of OH reaction of hydroperoxides derived from isoprene (Bates et al., 2014) making chemical loss a minor loss pathway. This means that further oxidation of these compounds is not relevant for typical conditions during night-time.

In the presence of aerosol surface, epoxides could be lost by particle uptake, but this was not relevant in the experiments analysed in this work due to the absence of seed aerosol. Loss to the Teflon surface of the chamber was not significant as demonstrated by the consistency of the loss rate with the dilution rate in the experiments.
5.10 OH and NO$_3$ reactivity from products

Overall, night-time oxidation of products from the reaction of isoprene with NO$_3$ appear to be of minor importance. This is further supported by measurements of total OH and NO$_3$ reactivity in the experiments in this work. In Figure 9, measured OH reactivity from organic compounds (Section 2.1) is compared to values calculated from modelled concentrations of products for the experiment on 13 August 2013 (Experiment #4), when the total consumption of isoprene by NO$_3$ was highest. Reaction rate constants for the reactions of organic compounds with OH are applied from the FZJ-NO3 mechanism.

OH reactivity is dominated by isoprene right after each injection (Fig. 9). After isoprene has reacted away, OH reactivity is only approximately 30% of the initial reactivity demonstrating the much lower reactivity of from products compared to than from isoprene. The major organic nitrate and epoxides produced from the reaction of NO$_3$ with isoprene explain approximately 50% of the total reactivity of organic products. Hydroperoxy aldehydes (HPALD), which are partly also produced from the OH oxidation of isoprene, contribute approximately 15% to the OH reactivity from products. A similar contribution is obtained from compounds that are formed from the oxidation of isoprene by OH and O$_3$, ISOPOOH, HCHO, MVK and MACR. At the end of the experiment, 25% of the total reactivity is due to a high number of organic compounds that are produced from minor reaction pathways or secondary oxidation.

The good agreement in the temporal behaviour of the observed and calculated OH reactivity demonstrates is consistent with the low loss rate of products due to further oxidation reactions. In addition, measured OH reactivity values are consistent with OH reaction rate constants implemented in the FZJ-NO3 mechanism, so that further OH oxidation of products are of minor importance is small for night-time conditions, when OH concentrations are typically maximum a few 10$^5$ cm$^{-3}$ (Stone et al., 2012, 2014; Lu et al., 2014; Tan et al., 2017).

OH oxidation of nitrate hydroperoxides is most important due to their fast reaction rate constant and their high concentrations for typical night-time conditions, when HO$_2$ + RO$_2$ reactions can dominate the loss of RO$_2$. However, part of the reactivity from hydroperoxides is invisible for the OH reactivity instrument, because OH is partly produced in their reactions with OH. Approximately 90% of the reactivity is detected assuming an OH yield of 10% as implemented in the Caltech and FZJ-NO3 mechanisms. In contrast, an OH yield of 100% is assumed for NISOPOOH in the MCM, which is likely too high as formation of epoxide products is expected to be a major reaction pathway.

In addition, OH oxidation of HPALD produced from unimolecular reactions of nitrate RO$_2$ can be significant because of the fast reaction of HPALD with OH.

In contrast the absolute values of OH reactivity as well as its temporal behaviour calculated from model calculations using the MCM with high OH reaction rate constants and high yields of NISOPOOH and NC$_4$CHO leads to results that are inconsistent with the observed OH reactivity (Fig. A12). This confirms that the MCM does not reproduce the product distribution and loss rates of products.

Dewald et al. (2020) discussed the NO$_3$ reactivity measured in the experiments also investigated in this work. Consistent with conclusions above that the chemical loss of products by NO$_3$ was not relevant, the authors found that the NO$_3$ reactivity could be fully explained by the reactivity from isoprene and propene in these experiments. This confirms that further NO$_3$
Comparison of measured OH reactivity from organic compounds and OH reactivity (left panel) calculated from concentrations of organic compounds modelled applying the FZJ-NO3 chemical mechanism for the experiment on 13 August 2018 (Experiment #4). The reactivity from hydroperoxide compounds (NISOPOOH, ISOPOOH) is partly invisible for the LP-LIF instrument, because these species produce OH radicals after reacting with it. The OH yield is rather uncertain, but is expected to be less than 10 % for example in the Caltech mechanism. In addition, the relative distribution of OH reactivity from organic products is shown (right panel). OH reactivity from organic compounds is derived by subtracting the reactivity from NO₂ and O₃ calculated from measured concentrations from the measured total OH reactivity. “Other” compounds include a high number of organic compounds that are produced in the reaction of isoprene with OH, O₃ and NO₃ and for which loss by the reaction with OH is implemented in the FZJ-NO3 mechanism. Dashed vertical lines indicate times, when isoprene, NO₂ and O₃ were re-injected. The last injection only included NO₂ and O₃.

Oxidation loss of organic products from the reaction of isoprene with NO₃ due to further NO₃ oxidation is small compared to the dilution rate in the chamber experiments. For typical night-time conditions as in the experiments in this work, is of minor importance.

6 Comparison to previous experiments

The high yield of MVK and MACR from the decomposition of β-RO radicals in the Caltech mechanism was derived from chamber experiments in Schwantes et al. (2015). In their experiments, 54 to 74 % of the nitrate RO₂ reacted with HO₂, so that the majority of alkoxy radicals was formed from this reaction. MVK and MACR concentrations, however, were only measured in 2 experiments in Schwantes et al. (2015), one of which was used to determine the MVK and MACR yields from the reaction of HO₂ + RO₂. The overall yield of the sum of MVK and MACR was relatively low with a value of approximately 15 %. In order to determine the yield of MVK and MACR from the decomposition of alkoxy radicals from the RO₂ + HO₂ reactions, production from the isoprene oxidation by OH and O₃ and from the potential decomposition of alkoxy radicals produced from...
other reaction channels needed to be subtracted. The authors used model calculations to estimate the actual OH concentration. Uncertainties in this calculations may explain the high MVK and MACR yield in Schwantes et al. (2015).

MVK and MACR concentrations were also measured in an experiment in the SAPHIR chamber reported by Rollins et al. (2009), in which low reactant concentrations were present as in this work (10 ppbv isoprene, 20 to 30 ppbv NO₂, 40 to 60 ppbv O₃). According to model calculations in Rollins et al. (2009) using the MCM 3.2, the fate of nitrate RO₂ radicals from isoprene with NO₃ was dominated by their reactions with HO₂.Measured MVK and MACR concentrations were consistent with the production of MVK and MACR mainly from the ozonolysis of isoprene. Therefore, this result supports that MVK and MACR are not produced from the decomposition of alkoxy radicals from β-RO₂ radicals.

In chamber experiments of Kwan et al. (2012), the fate of nitrate RO₂ was dominated by reactions with other RO₂ and with NO₃, both of which also produce nitrate alkoxy radicals. The total yield for MVK and MACR determined in these experiments was 6%. The authors state that all measured MVK and MACR could be explained by production from OH reactions because no ozone was present. The low yield of MVK and MACR appears to be inconsistent with the production of MVK and MACR from the decomposition of alkoxy radicals from β-RO₂ radicals as proposed in Schwantes et al. (2015). This is further supported by other experiments investigating the reaction of isoprene with NO₃ at high reactant concentrations (Barnes et al., 1990; Kwok et al., 1996; Perring et al., 2009), in all of which the yield of MVK and MACR was maximum in the range of a few percent.

Similar to the experiments in this work, products that have the sum formulas of nitrate epoxide products expected to be formed in the FZJ-NO3 mechanism were observed in the experiments in Kwan et al. (2012) and Schwantes et al. (2015): (1) C₅H₅NO₅ compounds, which appear at the same mass as NISOPOOH, (2) C₅H₇NO₅ compounds from epoxy-RO₂ + RO₂ reactions, (3) C₅H₉NO₆ compounds from epoxy-RO₂ + HO₂ reactions.

In Kwan et al. (2012) and Schwantes et al. (2015) it is suggested that the product with the sum formula C₅H₉NO₆ is a hydroxy hydroperoxy nitrate and that the product with the sum formula C₅H₇NO₅ is a hydroxy carbonyl nitrate. Both compounds are suggested to be products from a 1,5-H-shift reaction of the nitrate alkoxy radicals from the bimolecular reactions of δ-RO₂ radicals. The authors could not estimate a reaction rate constant for the 1,5-H-shift reaction from their experiments. Vereecken et al. (2021) calculated a reaction rate of 2.2 × 10⁶ s⁻¹ (T = 298 K), which makes the 1,5-H-shift reaction too low to compete with the ring-closure reaction forming epoxy alkyl radicals (1.2 × 10⁸ s⁻¹, T = 298 K) and subsequent O₂ addition. It is worth noting that compounds suggested by Kwan et al. (2012) and Schwantes et al. (2015) would only be produced from nitrate δ-RO₂ radicals that have small yields, whereas the nitrate epoxide products in the FZJ-NO3 mechanism are also produced from the most abundant nitrate β-RO₂ radicals. This may also explain why compounds with these sum formulas were clearly detected in experiments from all studies.

In the experiments in Kwan et al. (2012) and Schwantes et al. (2015), the C₅H₈O₃ compound without a nitrate functional group was observed, which is consistent with observations in this work. Because HPALD appears at the same mass and HPALD is also produced from OH oxidation, the authors concluded that C₅H₈O₃ is a product from the reaction of isoprene with OH. Nevertheless, their observations of C₅H₈O₃ compounds could also be partly due to the production of epoxy species from the oxidation of isoprene by NO₃ as described in the FZJ-NO3 mechanism.
The other product without a nitrate group that is produced from the ring-closure pathway of nitrate alkoxy radicals in the FZJ-NO3 mechanism, C₅H₈O₄, was not observed in the experiments in Kwan et al. (2012) and Schwantes et al. (2015). The reason could be that the chemical lifetime of RO₂ radicals was too short in the experiments in Kwan et al. (2012), in which high concentrations of reactants were present, so that the 1,6-H-shift reaction of the epoxy-RO₂ radical producing the C₅H₈O₄ compound could not compete with bimolecular reactions. Similarly, RO₂ reactions with HO₂ were favoured in the experiments in Schwantes et al. (2015), so that the 1,6-H reaction may have not been competitive.

Interestingly, similar to the experiments in this work, no organic nitrate with the sum formula C₄H₅NO₄ that is expected to be formed from the ring-closure reactions of nitrate alkoxy radicals (Fig. 6) was observed in the experiments in Kwan et al. (2012) and Schwantes et al. (2015). This further suggests that there is no significant production of this compound.

NISOPOOH has been detected by mass spectrometer instruments in previous chamber studies by Ng et al. (2008), Kwan et al. (2012) and Schwantes et al. (2015). Similar to this work, the instruments were not calibrated for NISOPOOH, but the sensitivity of the instrument was calibrated for nitrate alcohols (ISOPCNO₃). The sensitivity for other organic nitrates like such as NISOPOOH was estimated from calculations of the dipole moment and polarisability (Ng et al., 2008; Kwan et al., 2012; Schwantes et al., 2015).

In the experiments of Schwantes et al. (2015), HO₂ concentrations were enhanced. NISOPOOH yields were between 0.32 and 0.41, when 54 and 76 % of the nitrate RO₂ were calculated to react with HO₂. The authors calculated that these yields are consistent with a 50 % branching ratio of the reaction of nitrate RO₂ with HO₂ (Reaction R2) to form alkoxy radicals. An uncertainty of ±20 % of the measured NISOPOOH concentration is stated. The uncertainty of the alkoxy radical yield, however, could be higher because the calculation also requires the knowledge of the fraction of isoprene that reacted with NO₃ and the fraction of RO₂ that reacted with HO₂, both of which are uncertain because NO₃ and HO₂ concentrations were not measured. Therefore, a NISOPOOH yield of the reaction of nitrate RO₂ with HO₂ higher than 50 % may also be consistent with the experimental results in Schwantes et al. (2015).

Ng et al. (2008) quantified NISOPOOH concentrations in their chamber experiment, which was performed at high concentrations of reactants (800 ppbv isoprene, 120 ppbv N₂O₅). They determined that 50 % of the reacted isoprene resulted in the formation of NISOPOOH, but the fraction of nitrate RO₂ that reacted with HO₂ could not be determined to calculate yields from specific reactions. Therefore, their experiments cannot be used to derive information about potential alkoxy radical formation from the reaction of RO₂ with HO₂. HO₂ concentrations in experiments in Kwan et al. (2012) were presumably small because high reactant concentrations were used. This explains the relatively small overall NISOPOOH formation of 10 % from the reaction of isoprene with NO₃.

Kwan et al. (2012) assumed that specific C₅ organic compounds (HPALD, ISOPOOTH, C₅ hydroxy carbonyl C₅H₈O₂) and MVK and MACR, all of which were quantified in their chamber experiments, were exclusively formed from OH radicals that are formed as co-product of alkoxy radicals. In this case, the yield of nitrate alkoxy radical formation competing with the formation of NISOPOOH in the reaction of nitrate RO₂ with HO₂ is 38 to 58 %. Although the experiments were performed in the absence of ozone, so that OH was not produced by ozonolysis reactions, this approach gives only an upper limit of the yield because OH as well as some of the organic products may not have been exclusively produced by this assigned reaction
pathway. For example, HPALD can also be produced from the oxidation of isoprene by NO₃ from 1,6-H reactions of nitrate RO₂ (Vereecken et al. (2021), Fig. 3, Section 5.8).

HPALD was also observed in chamber experiments in Kwan et al. (2012) and Schwantes et al. (2015). The authors attributed the observations to the OH oxidation of isoprene, but their observations could also indicate HPALD formation from nitrate RO₂. Specifically in the experiments in Schwantes et al. (2015) the total loss rate of nitrate RO₂ were calculated to be in the range of 0.03 to 0.13 s⁻¹, so that 1,6-H shift reactions with rates 0.02 and 0.05 s⁻¹ (T = 298 K) calculated in Vereecken et al. (2021) can compete with bimolecular loss reactions.

Tsiligiannis et al. (2022) showed that a C₄ nitrate with the sum formula C₄H₇NO₅ was observed by the I⁻ CIMS instrument in the experiments in this work and also in several field campaigns, in which isoprene oxidation by NO₃ was important. This compound was also detected in the chamber experiments by the Br⁻ CIMS instrument (Wu et al., 2021), but signals observed by the VOCUS PTR-MS instrument at the respective mass were below the limit of detection. Yields of C₄H₇NO₅ determined in previous chamber experiments in Schwantes et al. (2015) were below 1%.

Mayhew et al. (2022) applied the three chemical models investigated in this work to field observations in an urban location in Beijing in June 2017. Differences between model results were similar as calculated in this work. The comparison of modelled data with measurements, however, is more complex for field experiments than for chamber experiments, because trace gas concentrations are not only impacted by chemical process, but also by transportation. In the field campaign in Beijing, organic nitrates from isoprene were detected by an I⁻ CIMS instrument. The instrument was not specifically calibrated for those compounds, but the same sensitivity as for isoprene epoxides (IEPOX) was assumed. In general, concentrations of measured isoprene-derived organic nitrates were lower than calculations for all three models in the night (Mayhew et al., 2022). As pointed out by the authors, the potential loss of epoxide nitrates due to particle uptake could not entirely explain the model-measurement discrepancies.

Overall, results in the experiments in this work appear to be consistent with results in previous experiments supporting the validity of the FZJ-NO3 mechanism.

7 Conclusions

The oxidation of isoprene by the nitrate radical, NO₃, was investigated in chamber experiments covering different atmospherically relevant chemical regimes. The chemical lifetimes of RO₂ radicals formed in the initial reaction of isoprene with NO₃ were in the range of atmospheric lifetimes with values between 30 s and several minutes due to atmospheric concentrations of reaction partners (RO₂, HO₂ and NO₃). In one experiment, RO₂ + HO₂ reactions were favoured by producing HO₂ and OH radicals in the ozonolysis of propene in the presence of excess CO for the conversion of OH to HO₂ radicals. Results from calculations of three near-explicit chemical models (MCM, CalTech, FZJ-NO3) were compared to measurements.

The MCM simplifies the oxidation of isoprene by NO₃ by forming only one RO₂ isomer conformer, whereas the other 2 chemical mechanisms differentiate between nitrate RO₂ isomers conformers due to the different positions at which NO₃ and O₂ can add (Wennberg et al., 2018; Vereecken et al., 2021b). In addition, a fast equilibrium between these RO₂ isomers...
is implemented in the FZJ-NO3 mechanism (Vereecken et al., 2021b) analogous to RO₂ isomers formed in the OH radical addition to isoprene (Peeters et al., 2014). The formation of different RO₂ isomers in these mechanisms leads to differences in the expected distribution of organic nitrate products, nitrate hydroperoxide (NISOPOOH), a nitrate carbonyl (NC₃CHO), and a nitrate alcohol (ISOPCNO₂). For example, concentrations of nitrate carbonyl products are expected to be much lower in the CalTech and FZJ-NO3 mechanisms compared to the MCM because they cannot be formed from nitrate /β RO₂ conformers not considered in the MCM.

Another critical difference between the three chemical mechanisms is the fate of nitrate alkoxy radicals formed in the radical reaction chain which mainly undergo ring-closure reactions in the FZJ-NO3 mechanisms, whereas decomposition to MVK and MACR is not competitive. Nitrate carbonyl products are exclusively formed in the MCM, whereas abundant RO₂-conformers are assumed to decompose to MVK or MACR together with HCHO and OH in the CalTech mechanism.

Measured concentrations of MVK and MACR in the experiments in this work, however, are consistent with their production from only O₃ and OH reactions with isoprene consistent. This is also in agreement with results in previous chamber experiments in Rollins et al. (2009) and Kwan et al. (2012). In addition, Vereecken et al. (2021b) calculated that ring-closure reactions of nitrate alkoxy radicals are highly competitive leading to the formation of epoxides rather than MVK and MACR. In some of these reaction pathways, NO₂ is released, so that not all epoxide products contain a nitrate group.

Mass signals of most of the organic products expected from the ring-closure reactions of the nitrate alkoxy radicals were detected by the VOCUS PTR-MS instrument demonstrating that the reactions calculated in Vereecken et al. (2021) may indeed be relevant pathways. Signals at the same masses were observed by chemical ionization mass spectrometer in previous chamber experiments (Kwan et al., 2012; Schwantes et al., 2015), but were partly attributed as products from the OH chemistry of isoprene or an 1,5-H-shift reaction of alkoxy radicals, for which Vereecken et al. (2021b) calculated the reaction rate coefficients to be too low to be competitive. One product of the ring-closure reaction of nitrate alkoxy radicals, which has the sum formula C₄H₅NO₄, calculated by Vereecken et al. (2021) to be produced could not be detected by the VOCUS PTR-MS instrument in the experiments in this work and has also not been observed in experiments in Kwan et al. (2012) and Schwantes et al. (2015). Therefore, the reaction pathway leading to this product is likely less important than implemented in the FZJ-NO3 mechanism, but this is within the uncertainty of the calculations in Vereecken et al. (2021).

The formation of hydroperoxy aldehydes (HPALD) from 1,6-H-shift reactions of nitrate Z-δ-RO₂ isomers conformers is only implemented in the FZJ-NO3 mechanism (Vereecken et al., 2021) analogous to RO₂ from the reaction of isoprene with OH (Peeters et al., 2014). The calculated isomerization reaction rate constant would lead to a HPALD yield between 10 and 30 % for conditions of the experiments in this work. High yields of total organic nitrates of around 100 % were determined in Brownwood et al. (2021) for the same experiments, but the lowest total organic nitrate yield of (94 ± 20) % was found in the experiment with the longest RO₂ lifetime due to bimolecular reactions (10 August 2018, Experiment #2) consistent with the production from unimolecular H-shift reactions, which were most competitive in this reaction. Although total organic nitrate yields determined in Brownwood et al. (2021) have a large uncertainty of up to 25 %, the overall high values hint that reaction rate constants of 1,6-H-shift reactions are lower than calculated by Vereecken et al. (2021), which have an uncertainty of at least a factor of 2.
A clear signal at the mass of HPALD was detected by the VOCUS PTR-MS instrument in all experiments in this work. This was also the case in the experiment, when an OH scavenger was present (09 August 2018, Experiment #1), demonstrating that HPALD was formed from the reaction of isoprene with NO₃ and that the HPALD was not only formed from the small fraction of isoprene reacting with OH radicals and ozone in the experiments. This is also consistent with previous chamber experiments by Kwan et al. (2012) and Schwantes et al. (2015), in which HPALD formation was observed but attributed to the production from the reaction of OH with isoprene.

Overall, results from experiments in this work demonstrate that the chemical mechanisms describing the chemistry of NO₃ with isoprene need to consider the different nitrate-RO₂ conformers. Large uncertainties, however, still exist in the exact distribution of the RO₂ isomers and their fate. Specifically, the yield of alkoxy radicals from the reaction of nitrate-RO₂ with HO₂ is uncertain. A yield of 50% is proposed in the CalTech mechanism (Wennberg et al., 2018), but this value is based on measurements of MVK and MACR concentrations that are attributed as products from this reaction pathway in chamber experiments in Schwantes et al. (2015). This attribution, however, is not supported by the theoretical calculations in the study by Vereecken et al. (2021) and the results in this study. The FZJ-NO3 mechanism assumes a 50% yield for alkoxy radical formation, but this number is highly uncertain due to the lack of measurements of absolute concentrations of specific products from the ring-closure reaction of the nitrate alkoxy radicals. Calibration of instruments detecting these compounds is urgently needed in future experiments, in order to determine the absolute importance of these reaction pathways for atmospheric conditions.

In the night, the fate of nitrate-RO₂ includes bimolecular reactions with HO₂ radicals, other RO₂ radicals and NO₃ radicals, all of which are significant for atmospheric conditions. This is in particular true if the reaction rate constants for the RO₂ + RO₂ reactions of the most abundant nitrate-RO₂ isomer from isoprene are rather low compared to other RO₂ radicals as concluded in Schwantes et al. (2015), and if the reaction rate constants for the nitrate-RO₂ + NO₃ reactions are a factor of 2 higher than previously assumed as concluded in Dewald et al. (2020). Nevertheless, the reaction with HO₂ is the dominant loss reaction of nitrate-RO₂ in the experiments in this work with a contribution of approximately 50% to the total loss rate. This is also confirmed by the high signal in the mass spectrum of the VOCUS PTR-MS instrument at the mass of the nitrate hydroperoxides (NISOPOOH) produced from this reaction. For atmospheric concentrations of radicals (Lu et al., 2014; Tan et al., 2017), it can also be expected that the loss of nitrate RO₂ in the reaction with HO₂ has a large contribution to their overall loss.

Tsiligannis et al. (2022) showed that a C₄ nitrate with the sum formula C₄H₇NO₅ was observed by the I−CIMS instrument in the experiments in this work and also in several field campaigns, in which isoprene oxidation by NO₃ was important. This compound was also detected in the chamber experiments by the Br−CIMS instrument Wu et al. (2021), but signals observed by the VOCUS PTR-MS instrument at the respective mass were below the limit of detection. Yields of C₄H₇NO₅ determined in previous chamber experiments in Schwantes et al. (2015) were below 1%.

None of the current chemical models can predict C₄H₇NO₅ yields estimated in Tsiligannis et al. (2022). They could be formed from further oxidation of first-generation C₅ nitrates by OH (Wennberg et al., 2018), but the expected yields in the experiments in this work are small due to the low OH concentrations. In addition, C₄H₇NO₅ compounds were also detected in the experiment, when OH concentrations were suppressed by an OH scavenger demonstrating that they are also formed.
from other reaction pathways. Further investigations are required to quantify the importance of $C_4H_7NO_5$ in the NO$_3$ isoprene oxidation scheme.

In the nocturnal atmosphere, isoprene is not only oxidized by NO$_3$, but also a significant fraction reacts with ozone depending on the availability of nitrogen oxides and ozone (Edwards et al., 2017). It is worth noting that due to the fast reaction rate constant of isoprene with OH, reaction with OH could also contribute to the overall loss of isoprene in the night. Part of the OH radicals can be produced in the subsequent reaction chain of the NO$_3$ oxidation of isoprene (Kwan et al., 2012; Vereecken et al., 2021). In the CalTech and FZJ-NO3 mechanisms, a large fraction is produced from the reaction of nitrate-RO$_2$ with HO$_2$. This again emphasizes the need to quantify the branching ratio of the alkoxy radical formation of this reaction pathway. Fast unimolecular reactions of RO$_2$ from the reaction of isoprene with OH (Peeters et al., 2014) can further gain in importance during the night compared to daytime (Novelli et al., 2020) because of the long chemical lifetime of RO$_2$ radicals in the range of minutes in the absence of NO, which is often the most important reaction partner for RO$_2$ radicals during the day. Therefore, the yield of HPALD produced from the OH reactions with isoprene can be high in the night despite low OH concentrations. HPALD photolysis could then contribute to OH production on the next day (Wolfe et al., 2012).

Mayhew et al. (2022) applied the three chemical models investigated in this work to field observations in an urban location in Beijing in June 2017. Differences between the chemical models with respect to product concentrations were qualitatively like differences discussed in this work but results were additionally impacted by complex chemical and meteorological conditions at the field site. In the field campaign in Beijing, organic nitrates from isoprene were detected by an I$^-$ CIMS instrument. The instrument was not specifically calibrated for those compounds, but the same sensitivity as for isoprene epoxides (IEPOX) was assumed. In general, concentrations of measured isoprene derived organic nitrates were lower than calculations for all three models in the night (Mayhew et al., 2022). As pointed out by the authors, the potential loss of epoxide nitrates due to particle uptake could not entirely explain the model-measurement discrepancies. This again suggests that the chemical mechanism of the isoprene oxidation by NO$_3$ has still large uncertainties.

Low-volatile epoxy and organic nitrate products be taken up by particles and therefore contribute to the secondary organic aerosol formation in the night. Only a small fraction of first-generation organic products are further oxidized for atmospheric night-time conditions. Reaction rate constants of the reactions of nitrate-carbonyl, nitrate-alcohol and epoxides with NO$_3$ and O$_3$ give chemical lifetimes which are longer than a night for typical concentrations of NO$_3$ and O$_3$. Also HPALD does not react efficiently with NO$_3$ and O$_3$. Reaction rate constants of these reactions as implemented in chemical models like such as the MCM, which lead to short chemical lifetimes in the range of hours, need to be revised.

Depending on the availability of OH radicals, first-generation products can partly be oxidized by OH in the night. Because OH concentrations are often very low, however, the majority of these compounds are most likely chemically processed by photolysis and reaction with OH on the next day. Oxidation of isoprene by the nitrate radical is most important in the residual layer, in which anthropogenic emissions including nitrogen oxides and biogenic emissions are present (Brown et al., 2009; Edwards et al., 2017). Oxidation products are therefore expected to be first further oxidized by OH radicals after sunrise within the residual layer, before the residual layer mixes with the convective boundary layer.
Overall, results from experiments in this work demonstrate that the FZJ-NO3 mechanism for isoprene (Vereecken et al., 2021) is valid and previous chemical mechanisms are incomplete to describe the nocturnal oxidation of isoprene. New reaction pathways in Vereecken et al. (2021) can have consequences for the nocturnal loss of reactive nitrogen and formation of secondary organic aerosol. However, large uncertainties still exist in the exact distribution of the different RO₂ isomers formed in the reaction of isoprene with NO₃ and their fate. Specifically, the yield of alkoxy radicals from the reaction of nitrate-RO₂ with HO₂ is uncertain. Calibration of instruments detecting organic nitrate products for specific reaction pathways is urgently needed in future experiments in order to determine the absolute importance of these reaction pathways for atmospheric conditions.

Data availability. Data from the experiments in the SAPHIR chamber used in this work are available on the EUROCHAMP database webpage (https://data.eurochamp.org). Experiment on 09 August 2018 (Experiment #1): Fuchs et al. (2018a), experiment on 10 August 2018 (Experiment #2): Fuchs et al. (2018b), experiment on 12 August 2018 (Experiment #3): Fuchs et al. (2018c), experiment on 13 August 2018 (Experiment #4): Fuchs et al. (2018d).

Author contributions. PC and HF wrote the manuscript, analysed the data and did model calculations of the experiments. SB, MH, JF, AN and HF designed and executed the experiments. LV provided insights into the chemical mechanisms. LH, TH, SK, TM, RT, DR, FR, RW, BB, JF, ET, JC, PD, NS, JF, JS, FB were responsible for measurements used in this work. All authors intensively discussed the manuscript and thereby contributed to the writing.

Competing interests. The authors declare no competing interests.

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Table A1. Organic products expected to be produced from the oxidation of isoprene in this work and the ion mass (m/z) at which they are detected by the mass spectrometry instruments, which measured in the experiments. Evaluation of the ion mass signals of the Br⁻-CIMS instrument includes both major isotopes of Br. Only the chemical structure of one isomer of the same compound is shown.

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<th>molecular weight</th>
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<th>ion mass (m/z) Br⁻-CIMS</th>
<th>ion mass (m/z) I⁻-CIMS</th>
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<td>210 / 212</td>
<td>385</td>
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<td>211 / 213</td>
<td>386</td>
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<td>224 / 226</td>
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<td>226 / 228</td>
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<td>258 / 260</td>
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50
**Figure A1.** Measurements of radical and trace gas concentrations and OH and NO$_3$ reactivity in the experiment on 10 August 2018 (Experiment #2) investigating the oxidation of isoprene by NO$_3$. NO$_3$ reactivity does not include reactivity from organic radicals and NO$_2$. OH and NO$_3$ reactivity from isoprene is calculated from measured isoprene concentrations and reaction rate constants recommended in literature (Mellouki et al., 2021). Observed RO$_2$ radicals only include a fraction of the total RO$_2$ because the LIF instrument cannot detect all RO$_2$ radicals formed in the reaction of isoprene with NO$_3$ (Vereecken et al., 2021).

**Figure A2.** Measurements of radical and trace gas concentrations and OH and NO$_3$ reactivity in the experiment on 12 August 2018 (Experiment #3) investigating the oxidation of isoprene by NO$_3$. NO$_3$ reactivity does not include reactivity from organic radicals and NO$_2$. OH and NO$_3$ reactivity from isoprene is calculated from measured isoprene concentrations and reaction rate constants recommended in literature (Mellouki et al., 2021). Observed RO$_2$ radicals only include a fraction of the total RO$_2$ because the LIF instrument cannot detect all RO$_2$ radicals formed in the reaction of isoprene with NO$_3$ (Vereecken et al., 2021).
Figure A3. Relative distribution of loss rates of nitrate RO$_2$ for the experiment on 10 August 2018 (Experiment #2) and on 12 August 2018 (Experiment #3). The total RO$_2$ loss rate was 0.005 and 0.008 s$^{-1}$ in the experiment on 10 August 2018 (Experiment #2) and 12 August 2018 (Experiment #3), respectively. Calculations of the loss rates of RO$_2$ radicals in bimolecular reactions make use of measured HO$_2$ and NO$_3$ concentrations. Total RO$_2$ concentrations and concentrations of speciated nitrate RO$_2$ were taken from model calculations applying either the FZJ-NO3, Caltech or MCM mechanism. The chemical mechanisms differ with respect to the number of nitrate RO$_2$ isomers that are considered, the type of RO$_2$ loss reactions and products of loss reactions (Fig. 3 and 6). Therefore, the distributions of nitrate RO$_2$ radicals and RO$_2$ concentrations differ between the model runs.

Figure A4. Relative distribution of loss rates of nitrate RO$_2$ for the experiment on 13 August 2018 (Experiment #3) if the FZJ-NO3 mechanism is applied and HO$_2$ is not constrained to measured values. Total RO$_2$ concentrations and concentrations of speciated nitrate RO$_2$ were taken from model calculations.
Figure A5. Comparison of results from model calculations applying the different isoprene NO$_3$ chemistry mechanisms for the experiment on 10 August 2018 (Experiment #2). MVK, MACR, NISOPOOH, ISOPCNO$_3$ and NC$_4$CHO are produced from all mechanisms, whereas the other compounds are only produced from either 1,6-H-shift reactions or ring-closure reactions of nitrate alkoxy radicals, which are only implemented in the FZJ-NO$_3$ mechanism. Grey and black dots are measured values. Measured organic peroxy radical concentrations only include part of the total RO$_2$ because the LIF instrument cannot detect a fraction of nitrate RO$_2$ (Vereecken et al., 2021). Organic products were detected by the VOCUS PTR-MS instrument, which was only calibrated for MVK and MACR. All other traces are scaled to match best the results from the FZJ-NO$_3$ mechanism.
Figure A6. Comparison of results from model calculations applying the different isoprene NO₃ chemistry mechanisms for the experiment on 12 August 2018 (Experiment #3). MVK, MACR, NISOPOOH, ISOPCNO₃ and NC₄CHO are produced from all mechanisms, whereas the other compounds are only produced from either 1,6-H-shift reactions or ring-closure reactions of nitrate alkoxy radicals, which are only implemented in the FZJ-NO3 mechanism. Grey and black dots are measured values. Measured organic peroxy radical concentrations only include part of the total RO₂ because the LIF instrument cannot detect a fraction of nitrate RO₂ (Vereecken et al., 2021). Organic products were detected by the VOCUS PTR-MS instrument, which was only calibrated for MVK and MACR. All other traces are scaled to match best the results from the FZJ-NO3 mechanism.
Figure A7. Comparison of reported signals from three mass spectrometer instruments applying different ionization methods (VOCUS PTR-MS, Br⁻-CIMS, I⁻-CIMS measuring organic products in the experiment on 09 August 2018 (Experiment #1). All signals are scaled to match best the concentrations resulting from model calculations applying the FZJ-NO3 chemical mechanism.

Figure A8. Comparison of reported signals from three mass spectrometer instruments applying different ionization methods (VOCUS PTR-MS, Br⁻-CIMS, I⁻-CIMS measuring organic products in the experiment on 10 August 2018 (Experiment #2). All signals are scaled to match best the concentrations resulting from model calculations applying the FZJ-NO3 chemical mechanism.
Figure A9. Comparison of reported signals from three mass spectrometer instruments applying different ionization methods (VOCUS PTR-MS, Br⁻-CIMS, I⁻-CIMS measuring organic products in the experiment on 12 August 2018 (Experiment #3). All signals are scaled to match best the concentrations resulting from model calculations applying the FZJ-NO3 chemical mechanism.

Figure A10. Comparison of reported signals from three mass spectrometer instruments applying different ionization methods (VOCUS PTR-MS, Br⁻-CIMS, I⁻-CIMS measuring organic products in the experiment on 13 August 2018 (Experiment #4). All signals are scaled to match best the concentrations resulting from model calculations applying the FZJ-NO3 chemical mechanism.
Figure A11. Ion mass signal of the VOCUS PTR-MS instrument scaled to the model results from the MCM, Caltech and FZJ-NO3 models in the experiment on 13 August 2018 (Experiment #4). Only species for which the instrument was not calibrated and which are produced in all models are shown.

Figure A12. Comparison of measured OH reactivity from organic compounds and OH reactivity calculated from concentrations of organic compounds modelled applying the MCM chemical mechanism. Up to 10% of the reactivity from hydroperoxide compounds (NISOPOOH, ISOPOOH) is invisible for the LP-LIF instrument, because these species produce partly OH in their reaction with OH. The exact OH yield is uncertain. 100% yield is assumed in the MCM. OH reactivity from organic compounds is derived by subtracting the reactivity from NO2 and O3 calculated from measured concentrations from the measured total OH reactivity. “Other” compounds include a high number of organic compounds that are produced in the reaction of isoprene with OH, O3 and NO3 and for which loss by the reaction with OH is implemented in the MCM mechanism.