Chemical and isotopic composition of secondary organic aerosol generated by α-pinene ozonolysis

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Abstract. Secondary organic aerosol (SOA) plays a central role in air pollution and climate. However, the description of the sources and mechanisms leading to SOA is elusive despite decades of research. While stable isotope analysis is increasingly used to constrain sources of ambient aerosol, in many cases it is difficult to apply because neither the isotopic composition of aerosol precursors nor the fractionation of aerosol forming processes is well characterised. In this paper, SOA formation from ozonolysis of α-pinene – an important precursor and perhaps the best-known model system used in laboratory studies – was investigated using position-dependent and average determinations of δ¹³C in α-pinene and advanced analysis of reaction products using thermal-desorption proton-transfer-reaction mass spectrometry (PTR-MS). The total carbon (TC) isotopic composition δ¹³C of the initial α-pinene was measured, and the δ¹³C of the specific carbon atom sites was determined using position-specific isotope analysis (PSIA). The PSIA analysis showed variations at individual positions from −6.9 to +10.5‰ relative to the bulk composition. SOA was formed from α-pinene and ozone in a constant-flow chamber under dark, dry, and low-NOx conditions, with OH scavengers and in the absence of seed particles. The excess of ozone and long residence time in the flow chamber ensured that virtually all α-pinene had reacted. Product SOA was collected on two sequential quartz filters. The filters were analysed offline by heating them stepwise from 100 to 400 °C to desorb organic compounds that were (i) detected using PTR-MS for chemical analysis and to determine the O:C ratio, and (ii) converted to CO₂ for δ¹³C analysis. More than 400 ions in the mass range 39–800 Da were detected from the desorbed material and quantified using a PTR-MS. The largest amount desorbed at 150 °C. The O:C ratio of material from the front filter increased from 0.18 to 0.25 as the desorption temperature was raised from 100 to 250 °C. At temperatures above 250 °C, the O:C ratio of thermally desorbed material, presumably from oligomeric precursors, was constant. The observation of a number of components that occurred across the full range of desorption temperatures suggests that they are generated by thermal decomposition of oligomers. The isotopic composition of SOA was more or less independent of desorption temperature above 100 °C. TC analysis showed that SOA was enriched in δ¹³C by 0.6–1.2‰ relative to the initial α-pinene. According to mass balance, gas-phase products will be depleted relative to the initial...
\(\alpha\)-pinene. Accordingly, organic material on the back filters, which contain adsorbed gas-phase compounds, is depleted in \(^{13}\)C in TC by 0.7\(\text{%}\) relative to the initial \(\alpha\)-pinene, and by 1.3\(\text{%}\) compared to SOA collected on the front filter. The observed difference in \(^{13}\)C between the gas and particle phases may arise from isotope-dependent changes in the branching ratios in the \(\alpha\)-pinene + \(O_3\) reaction. Alternatively, some gas-phase products involve carbon atoms from highly enriched and depleted sites, as shown in the PSIA analysis, giving a non-kinetic origin to the observed fractionations. In either case, the present study suggests that the site-specific distribution of \(^{13}\)C in the source material itself governs the abundance of \(^{13}\)C in SOA.

1 Introduction

Secondary organic aerosol (SOA) is formed in the atmosphere by oxidation of volatile organic compounds (VOCs). SOA contributes significantly to atmospheric aerosol and impacts climate, air quality, and health (Stocker et al., 2013; Hänninen et al., 2004; Dockery et al., 1993; Andreae and Crutzen, 1997). Despite their impacts, the mechanisms of SOA formation and processing are poorly understood and significant discrepancies between model predictions and observations are common.

Stable carbon isotopes are increasingly used to investigate aerosols in the atmosphere. Isotopic analysis holds the promise of clearly identifying SOA sources and atmospheric transformations, as long as we know enough details, such as the isotopic composition of the source emissions, the kinetic fractionations occurring in oxidation reactions, and the isotope effects in phase partitioning. The ability to distinguish individual sources, including marine aerosol (Turekian et al., 2003; Ceburnis et al., 2011) and biomass burning (Kirillova et al., 2013), by isotope measurements has been particularly useful when it comes to source apportionment of ambient aerosol (e.g. Sakugawa and Kaplan, 1995; Narukawa et al., 2008; Turekian et al., 2003; Widory et al., 2004; Ho et al., 2006; Huang et al., 2006; Fisseha et al., 2009; Kirillova et al., 2011, 2013; Ceburnis et al., 2011; Fu et al., 2012; Miyazaki et al., 2012; O’Dowd et al., 2014; Masalaite et al., 2015).

In some other cases, measurements of the isotopic composition of SOA have yielded intriguing and/or contradictory results: (Irei et al., 2014) found that SOA formation from oxidation of volatile organics lead to depletion in \(^{13}\)C in the low-volatile fraction of the aerosol. (Fu et al., 2012), however, reported isotopic enrichments in ambient SOA that are distinguishable between biomass burning and primary emissions, possibly hinting at the oxidation of biogenic VOCs.

Recent technical advances have made it possible to quantify isotopic enrichments at specific sites within molecules, in addition to the traditional determination of the enrichment of the sample itself (Bayle et al., 2014b). This development makes possible new analyses of chemical mechanisms and transformations in the environment. Consider a chemical reaction that goes to completion to give a variety of products. The atoms of a reagent are distributed among the products, depending on the branching ratios for the different reaction channels. The branching ratios may themselves be isotope-dependent. For the formation of SOA from a VOC precursor, the isotopic enrichments or depletions that are observed in the SOA arise from the contributions of many oxidation products. Mass balance implies that the complimentary pattern will be observed in gas-phase products. Many interesting research questions arise from the application of site-specific isotopic analysis to atmospheric science. Perhaps the most important question is whether observed average composition of the SOA is due to the isotope dependent changes in product branching ratios within the reaction mechanism or due to the site-specific enrichments of the reagent.

Isotopic substitution can cause reactions to be faster or slower than for the un-substituted case, kinetically fractionating the isotopes and leading to isotopic enrichment or depletion in the products. This is known as the kinetic isotope effect (KIE). If a reaction leads to a single product, the product may initially have a different abundance; but due to the law of mass balance, the product will achieve the same abundance as the reagent as the reaction goes to completion. If a reaction has multiple product channels, enrichment or depletion will occur if there are isotope-dependent changes in the product branching ratios.

The ozonolysis of \(\alpha\)-pinene is often used as a test system for formation of SOA; it is fairly well studied. Figure 1 shows a reaction scheme for \(\alpha\)-pinene ozonolysis, based on the “Master Chemical Mechanism” (MCMv3.1) as described by (Canagaratna et al., 2010). In the first step, ozone adds into the double bond of the molecule, resulting in two branches depending on the usual Criegee mechanism. These two branches proceed by stabilisation, subsequent fragmentation and isomerisation, and subsequent reaction with RO$_2$, HO$_2$, and H$_2$O to yield a wide range of oxidation products from CO, HCHO, and acetone, to many larger oxidised low volatile molecules like pinic acid and pinonic acid and pinonaldehyde. The figure shows only formation of first-generation products. Further reactions including dimer formation (Kristensen et al., 2016) and oligomerisation reactions are not shown.

Generally, reactions involved in atmospheric VOC oxidation fall into three categories: functionalisation, fragmentation, and oligomerisation (Kroll and Seinfeld, 2008; Rudich et al., 2007; Chacon-Madrid and Donahue, 2011). The volatility (Donahue et al., 2006; Jimenez et al., 2009) and oxygen-to-carbon ratio, O:C (Donahue et al., 2011; Kroll et al., 2011), of involved species allow for the characterisation of these processes. The characteristics of the three types of atmospheric VOC reactions and current understanding of how they contribute to isotopic fractionation (Kirillova et al., 2013, 2014) are summarised as follows:
Figure 1. Reaction scheme of α-pinene ozonolysis, based on (Camredon et al., 2010), using the reaction mechanism of MCMv3.1. The boxes show first-generation products, while subsequent dimer formation and oligomerisation are not shown. The O : C ratio typically increases as oxidation proceeds (i.e. oxygen addition via functionalisation). Large, oxygenated product compounds (e.g. pinonic acid, pinonaldehyde) have lower vapour pressures and partition to the particle phase. Small, volatile products including HCHO, acetone, and CO result from fragmentation processes and partition to the gas phase. The coloured symbols highlight how the parent α-pinene’s atoms (denoted in brackets) can form small reaction products including CO, HCHO, and acetone. The symbol colours were chosen based on the heat map in Fig. 7.

- Functionalisation describes the addition of oxygenated functional groups to the parent compound. A typical example is the first reaction step in VOC oxidation. In these reactions, products are less volatile and have higher O : C ratios than the parent compound. Functionalisation is typically accompanied by normal kinetic fractionation leading to oxidation products depleted in $^{13}$C (Rudolph and Czuba, 2000; Iannone et al., 2010; Gensch et al., 2011). In the present study, the parent compound α-pinene was fully oxidised, leaving no isotopic imprint on its products (the KIE of α-pinene ozonolysis has not been reported to date).

- Fragmentation describes cleavage of carbon–carbon bonds, possibly followed by addition of oxygen to the fragments. Fragmentation reactions in the condensed phase can result in molecules that are small, like CO$_2$ and CH$_2$O, having higher volatilities than the parent compound and possibly escaping to the gas phase (Kroll et al., 2009). Fragmentation may enrich the aerosol in $^{13}$C through normal kinetic fractionation: when some of the depleted reaction products are lost to the gas phase, the remaining aerosol phase will be enriched (Aggarwal and Kawamura, 2008). The O : C ratio of the products is often higher than for the parent VOC and is typically highest for compounds remaining in the aerosol phase. During ozonolysis of monoterpenes, both functionalisation and fragmentation occur simultaneously (Chacon-Madrid and Donahue, 2011), as also shown in Fig. 1. This increases the O : C ratio but the overall effect on the isotopic balance is not well established.
– Oligomerisation (sometimes also referred to as accretion) describes the building of larger organic structures from monomers, often in the aerosol phase (Kalberer et al., 2004, 2006; Hallquist et al., 2009). One mechanism is that oxygenated organics become linked by dehydration reactions, and this lowers the O:C ratio. The formation of complex organic mixtures in the aerosol phase (Cappa et al., 2008) is expected to show similar characteristics in vapour pressure and O:C ratio. The influence of oligomerisation on the isotopic composition of the aerosol is not clear, but likely small.

Chemical analysis based on proton-transfer-reaction mass spectrometry has proven very useful in ambient and laboratory studies investigating aerosol and gas-phase compounds and their properties (e.g. Holzinger et al., 2010a, b; Shilling et al., 2008; Presto and Donahue, 2006). Advantages of the technique include the soft ionisation, high sensitivity, wide range of detectable compounds, and the possibility of quantifying them. In addition, using a proton-transfer-reaction time-of-flight mass spectrometer (PTR-ToF-MS, shortened to PTR-MS for the remainder of the article) allows chemical characterisation and identification of compounds and estimation of the O:C ratio.

Position-specific isotope analysis (PSIA) of the initial reactant offers a detailed description of bulk isotopic data by mapping the intra-molecular isotope distribution. PSIA by $^{13}$C isotope ratio monitoring by nuclear magnetic resonance (NMR) (irm-$^{13}$C NMR) has proven valuable in interpreting a number of (bio)chemical (Bayle et al., 2014a; Botosoa et al., 2009a; Gilbert et al., 2011, 2012) and physical–chemical processes such as distillation and sorption (Botosoa et al., 2008, 2009b; Höhener et al., 2012), leading to a deeper understanding of the underlying phenomena causing isotope fractionation in nature. It is widely understood that the enrichment or depletion of a product depends on the enrichment of the starting material, the isotopic fractionation occurring in the mechanism of its formation, and the extent of reaction. Using PSIA, we can take this analysis one step further: the enrichment of a product will depend on the position-dependent enrichments of the atoms from which it is formed. For example the ozonolysis mechanism transfers the C9-atom in $\alpha$-pinene into several small, volatile products (see blue squares in Fig. 1). If the C9-position was depleted in $^{13}$C, the gas-phase products containing this atom would be depleted, and the SOA correspondingly enriched, as long as the position-dependent effect was stronger than kinetic isotope effects.

In this study we have designed an experiment to use a combination of position–specific, sample–average, and molecule–average isotopic abundance data to investigate the formation of SOA from the ozonolysis of $\alpha$-pinene. Due to mass balance, the enrichment (or depletion) in SOA must be balanced by depletion (or enrichment) in gas-phase products, since in our experiment $\alpha$-pinene ozonolysis has gone to completion. However, since the reaction mechanism is not known in detail and cannot be determined from our experiment, we are left to interpret an incomplete case. One instructive limiting case is to ask which atoms in $\alpha$-pinene are transformed into known gas-phase products. We argue that the observed isotopic distribution in SOA is likely balanced by the opposite pattern in the gas-phase products, and that this pattern could be produced by site-specific enrichments in the starting material and current incomplete knowledge of the reaction mechanism alone, aside from whatever effects isotopic substitution may have on product branching ratios. Thus, we make the case that position-sensitive isotopic enrichments and depletions, and not just bulk KIEs, are an important element of explaining field observations, especially for natural products which are known to have, as a group, significant site-specific enrichments and depletions relative to fossil and/or synthetic compounds. The goal of this study is to provide detailed isotopic and chemical characterisation of newly formed $\alpha$-pinene SOA and to shed light on the mechanisms that govern isotopic fractionation in the formation of fresh SOA.

2 Material and methods

2.1 Chemical compounds used

Chamber experiments were performed using the following chemicals: (+)-$\alpha$-pinene (Aldrich, > 99 %, batch no. 80796DJV), 1-Butanol (Sigma-Aldrich, > 99.4 %) and cyclohexane (Labscan, 99.5 %). PSIA was performed on several samples of $\alpha$-pinene from Sigma-Aldrich (different lot), Acros Organics, Merck, and Alfa Aesar; see Table 1. The batch of $\alpha$-pinene used in the chamber experiments could not be analysed using PSIA, because the manufacturer does not supply it anymore.

2.2 Chamber design and characteristics

A new aerosol smog chamber was built in Copenhagen based on a steady-state design (King et al., 2009; Shilling et al., 2008; Kleindienst et al., 1999). It consists of a 4.5 m³ teflon bag (the volume / surface ratio is 0.275 m) mounted inside a temperature-controlled insulated room of walk-in size (Viessmann A/S); see Fig. S1 in the Supplement. While details can be found in Sect. S1, chamber operation is briefly summarised here. The chamber was operated in a constant-flow mode: dry air and reactants were constantly flushed into the chamber using mass flow controllers. A syringe pump (NE-300, New Era Pump Systems Inc.) continuously injected a mixture of $\alpha$-pinene and 1-butanol or a mixture of $\alpha$-pinene and cyclohexane into a warmed glass bulb. 1-butanol and cyclohexane were used as OH scavengers and the mixing ratio between $\alpha$-pinene and the OH scavenger was 1 : 600 (v/v). A small flow of clean, dry air (0.1 L min$^{-1}$) directed over a Hg lamp emitting UV light (model 600, Jelkight company inc.) generated ozone which was fed into the chamber separately.

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Table 1. Manufacturer information and isotopic composition of α-pinene samples. Position-specific isotopic fractionation is given as isotopic difference, $\Delta_i^{13} C = \delta_i^{13} C - \delta_j^{13} C$, of individual C atoms (i) in α-pinene sample j with bulk isotopic composition $\delta_j^{13} C$. Listed are the means of five measurements. See inserted figure for numbering of C atoms. Sample 1 was used in the chamber experiments, but PSIA could not be performed as the sample was no longer available.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Manufacturer</th>
<th>Purity (%)</th>
<th>Code</th>
<th>Lot</th>
<th>$\delta_i^{13} C$ ($%$)</th>
<th>$\Delta_i^{13} C$ ($%$)</th>
<th>$C_1$</th>
<th>$C_2$</th>
<th>$C_3$</th>
<th>$C_4$</th>
<th>$C_5$</th>
<th>$C_6.7$</th>
<th>$C_8$</th>
<th>$C_9$</th>
<th>$C_{10}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Sigma-Aldrich</td>
<td>&gt; 99</td>
<td>268070</td>
<td>80796DFJ</td>
<td>-30.0</td>
<td>4.8</td>
<td>10.5</td>
<td>1.0</td>
<td>-6.0</td>
<td>0.6</td>
<td>-0.3</td>
<td>-6.7</td>
<td>-4.3</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>2</td>
<td>Sigma-Aldrich</td>
<td>&gt; 99</td>
<td>268070</td>
<td>MKB92131V</td>
<td>-27.7</td>
<td>4.1</td>
<td>10.4</td>
<td>1.3</td>
<td>-3.2</td>
<td>1.9</td>
<td>0.0</td>
<td>-6.9</td>
<td>-5.4</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>3</td>
<td>Acros Organics</td>
<td>98</td>
<td>131261000</td>
<td>A031001B</td>
<td>-27.0</td>
<td>6.1</td>
<td>9.6</td>
<td>1.4</td>
<td>-3.8</td>
<td>-0.8</td>
<td>1.2</td>
<td>-5.2</td>
<td>-3.9</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>4</td>
<td>Merck</td>
<td>&gt; 97</td>
<td>818632</td>
<td>521251I</td>
<td>-28.1</td>
<td>7.8</td>
<td>5.8</td>
<td>0.8</td>
<td>4.2</td>
<td>0.0</td>
<td>-1.8</td>
<td>-3.6</td>
<td>-4.5</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>5</td>
<td>Alfa Aesar</td>
<td>98</td>
<td>104941</td>
<td>1017535</td>
<td>-27.8</td>
<td>7.8</td>
<td>5.8</td>
<td>0.8</td>
<td>4.2</td>
<td>0.0</td>
<td>-1.8</td>
<td>-3.6</td>
<td>-4.5</td>
<td>-</td>
<td>-</td>
</tr>
</tbody>
</table>

Table 2. Overview of α-pinene ($^{13} C_{\alpha}^{13} C = (-29.96 \pm 0.08) \%$) ozonolysis experiments using 1-butanol (B) or cyclohexane (C) as OH-scaravenger. Filter IDs are explained in the text. The sampling time, $t$, and the sampled volume, $V$, are given. Isotopic data at 100 $^\circ$C and 150 $^\circ$C and for total carbon (TC) are given as $\Delta^{13} C = \delta^{13} C - \delta_{IC^{13} C}$. The total aerosol mass loading as detected by PTR-MS, $M_{total^{PTR-MS}}$, is listed next. The last column lists the measured O : C ratio averaged over all desorption temperatures.

<table>
<thead>
<tr>
<th>Experiment</th>
<th>Filter ID</th>
<th>$t$ (h)</th>
<th>$V$ (m$^3$)</th>
<th>$100^\circ$C</th>
<th>$150^\circ$C</th>
<th>TC</th>
<th>$M_{total^{PTR-MS}}$ (µg m$^{-3}$)</th>
<th>O : C</th>
</tr>
</thead>
<tbody>
<tr>
<td>B$^a$ (α-pinene,</td>
<td>B1f</td>
<td>47</td>
<td>28.2</td>
<td>2.8 ± 0.2</td>
<td>1.2 ± 0.1</td>
<td>1.2</td>
<td>6.9</td>
<td>0.23</td>
</tr>
<tr>
<td>1-butanol)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>B1b</td>
<td>47</td>
<td>28.2</td>
<td>2.9 ± 0.4</td>
<td>-0.7 ± 0.2</td>
<td>-</td>
<td>0.4</td>
<td>0.21</td>
</tr>
<tr>
<td></td>
<td>B2f</td>
<td>16.8</td>
<td>10.1</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>6.3</td>
<td>0.22</td>
</tr>
<tr>
<td></td>
<td>B2b</td>
<td>16.8</td>
<td>10.1</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
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<tr>
<td>C (α-pinene,</td>
<td>C1f</td>
<td>24.5</td>
<td>14.7</td>
<td>2.4 ± 0.5</td>
<td>0.5 ± 0.1</td>
<td>0.6</td>
<td>8.9</td>
<td>0.23</td>
</tr>
<tr>
<td>cyclohexane)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>C1b</td>
<td>24.5</td>
<td>14.7</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>0.9</td>
<td>0.22</td>
</tr>
<tr>
<td></td>
<td>C2f</td>
<td>26.7</td>
<td>16.5</td>
<td>1.7 ± 0.3</td>
<td>1.0 ± 0.1</td>
<td>0.7</td>
<td>11.7</td>
<td>0.23</td>
</tr>
<tr>
<td></td>
<td>C2b</td>
<td>26.7</td>
<td>16.5</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Handling blank</td>
<td>HB</td>
<td>-</td>
<td>-</td>
<td>10.0 ± 0.8</td>
<td>8.9 ± 0.1</td>
<td>-</td>
<td>0.23$^b$</td>
<td>-</td>
</tr>
</tbody>
</table>

$^a$ CCN data available; see Sect. S2.

$^b$ Surface loading in units of micrograms per square centimetre (µg cm$^{-2}$).

Generated aerosol was sampled after an ozone scrubber on doubly stacked quartz-fibre filters (4.7 cm diameter, QMA 1851, Whatman) for offline chemical and isotope analysis at 10 L min$^{-1}$. Collection times were around 1–2 days in order to provide sufficient amounts of carbon on the filters for isotope analysis; see Table 2. The ozone scrubber had a de-nuder design and used potassium iodide (Williams and Grosjean, 1990). It protected instruments from high ozone levels but also precluded further reaction of collected samples with ozone on the filters.

Several instruments monitored gaseous and particulate matter inside the bag: a scanning mobility particle sizer (SMPS; TSI 3081 DMA and 3772 CPC, 0.0508 cm impactor) was used to measure particle size distributions (10–500 nm diameter if not stated otherwise) and a cloud condensation nuclei (CCN) counter (Droplet Measurement Technologies) gave information on the CCN properties of generated SOA (King et al., 2012). Temperature and relative humidity were measured continuously inside the bag (HygroFlex HF532, Rotronic) and read by the same software that controlled the pressure inside the bag. NO$_x (= NO + NO_2)$ levels were monitored using a chemiluminescence NO$_x$ analyzer (42i, Thermo). The same line fed a UV photometric O$_3$ analyzer (49i, Thermo) to monitor ozone levels.

2.3 α-pinene ozonolysis

All aerosol was generated from the dark ozonolysis of α-pinene under low-NO$_x$ conditions (< 2 ppb) without any seed particles present. Experiments were performed with two different OH scavengers: Experiment B with 1-butanol and Experiment C with cyclohexane. The amount of α-pinene injected using the syringe pump resulted in a steady-state concentration of ca. 60 ppb inside the bag (without oxidants). Ozone concentrations during the experiment were always above 150 ppb, i.e. ozone was always in excess. The temperature was stable at 22 °C and RH < 1 %. Without seed particles present, aerosol formed via new particle formation.
Table 2 gives an overview of the conditions under which experiments B and C were performed.

The natural (e-folding) lifetime of \( \alpha \)-pinene with respect to ozonolysis is \( \tau_{e} = (k_{[O_3]})^{-1} = 40 \text{ min} \), based on an ozone concentration of 150 ppb and a second-order rate coefficient of \( k = 1.1 \times 10^{-16} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1} \) (Witter et al., 2002). The nominal residence time of an air parcel in the chamber is \( \tau_{\text{nominal}} \approx 3.4 \text{ h} \) (see Sect. S1) which exceeds the natural lifetime with respect to ozone loss by a factor of 5. This means that more than 99 % of \( \alpha \)-pinene will have reacted at steady state, and for further evaluation and discussion it is assumed that \( \alpha \)-pinene ozonolysis was complete. As total mass is conserved for all isotopes, the isotopic composition of the ozonolysis products should be equal to that of the initial reagent.

2.4 Filter handling protocol

Glass vials with plastic stoppers were used to store the quartz filters for off-line analysis before and after the experiments. The glass vials themselves were cleaned in a ceramic oven at 600 °C for 24 h prior to use. The quartz filters were cleaned in the same oven at 600 °C for more than 20 h prior to use. Each filter was stored in a separate glass vial that was wrapped in aluminium foil and stored in a dark freezer (−30 °C) except when loading or during transport between Copenhagen and Utrecht. Two quartz filters (quartz behind quartz, QBQ) were loaded at a time in a cleaned filter holder to account for possible sampling artefacts, such as adsorption and evaporation of organic vapours on or from the filters (Watson et al., 2009; Turpin et al., 2000). In this study the first filter, facing the sample stream, is called the \textit{front} filter, while the second one (stacked below) is called the \textit{back} filter. Storage time between loading and analysis was up to 6 months. During transport the filters stayed in the wrapped vials but were not actively cooled. Prior to analysis the filters were cut into pieces of uniform size (0.5 and 1 cm diameter). Blank filters were treated identically to loaded filters but were not exposed to chamber air. Gloves were used whenever working directly with the filters, and all tools were rinsed several times using first acetone and then ethanol.

2.5 Filter analysis

The filter samples from the smog chamber experiments were analysed for their chemical and isotopic composition at the Institute for Marine and Atmospheric research Utrecht (IMAU). Propagated uncertainties based on at least three measurements are given as 1-sigma errors. Filter IDs (identifications) are composed of a capital letter denoting the experiment (B using 1-butanol or C using cyclohexane as OH scavenger), a digit counting the experiments using that scavenger, and a small letter indicating the filter position: \( b \) for back filter and \( f \) for front filter.

2.5.1 Thermal-desorption chemical analysis of filters

The chemical analysis follows methods described earlier (Holzinger et al., 2010b; Timkovsky et al., 2015) and will only be outlined briefly here. The chemical analysis setup consisted of a two-stage oven. Filter pieces were heated stepwise in the first oven to temperatures of 100, 150, 200, 250, 300, and 350 °C, while the temperature of the second oven was kept constant at 200 °C. SOA components desorbed from the filters according to their volatility and a flow of nitrogen (50 mL min\(^{-1}\)) carried them to the PTR-MS (PTR-TOF8000, Ionicon Analytik GmbH, Austria) situated directly after the second oven. The PTR-MS inlet was heated to its maximum temperature of 180 °C, while the drift tube was operated at 120 °C. This monotonic gradient from the first oven stage to the drift tube inside the PTR-MS reduced cold spots and minimised repeated sample condensation. The PTR-MS detected the desorbed compounds after protonation (addition of \(^1\)H) as ions with a mass-to-charge ratio \((m/z) + 1\).

The PTR-MS had a mass resolution of \( m/\Delta m \approx 4000 \), allowing detection of ions with differences in \( m/z \) larger than 30 mDa. The algorithm for analysing the PTR-MS data is based on a method reported earlier (Holzinger et al., 2010a; Holzinger, 2015). For each experiment (B or C), the ions detected on the front and back filters were combined in a unified-peak list to minimise statistical uncertainty and improve overall mass accuracy (Holzinger et al., 2010a). Ions with \( m/z < 39 \text{ Da} \) were excluded (except 33.03 Da, methanol, and 31.01 Da, formaldehyde) as the PTR-MS mainly detects primary ions in this mass region, which do not originate from the filters. Water clusters with masses 37.026 and 55.038 Da can form in the PTR-MS and were not considered to be aerosol compounds. A total of 685 (753) ions were detected by the PTR-MS in experiment B (C). In order to take the contribution from the blank filter (filter ID “HB” for handling blank) into account, the blank filter loading was calculated from the PTR-MS signal and subtracted from the front and back filters for each ion on the unified-peak list. All concentrations are then reported relative to smog chamber conditions, i.e. as mass concentration per unit volume of air in the smog chamber. Table 2 lists the blank-corrected total mass concentration, \( M_{\text{total}} \). Many ions had negligible concentrations. The 427 (451) ions from the unified-peak list of experiment B (C), comprising 90 % of the total aerosol mass detected by PTR-MS (counting from the ion with highest concentration downwards), were considered for further analysis in order to streamline data analysis and reduce noise.

The data analysis algorithm identified the molecular formulas of the detected ions (i.e. the total number of most abundant stable C, H, O, and N isotopes) based on their exact masses. If the peak resolution did not allow unambiguous identification, several candidates were suggested (Holzinger et al., 2010a). The suggested formulas for all prominent peaks were checked manually on at least two filters and cor-
rected if necessary, including when the suggested formula contained N or $^{13}$C. The former can be excluded due to the low-NO$_x$ conditions in the experiment, and the latter can be easily verified by the corresponding carbon-12 peak at ($m/z - 1$) + 1. Most of the ions were identified unambiguously and typically only ca. 5% of the total mass from front filter desorption was attributed to ions with no clear molecular formula.

The oxygen-to-carbon ratios of the ions were calculated as in (Holzinger et al., 2013) for each filter at each desorption temperature:

$$O:C = \frac{\sum w_i n_{O,i}}{\sum w_i n_{C,i}}.$$  \hspace{1cm} (1)

Here, the sum counts over all identified ions $i$, $w_i$ is the measured amount of ion $i$ in moles (mol) and $n_{C,i}$ and $n_{O,i}$ are the respective number of carbon and oxygen atoms for ion $i$ as given by its molecular formula. Equation (1) gives the ratio of oxygen to carbon atoms in all identified ions.

### 2.5.2 Thermal-desorption isotope analysis of filters

The system used to measure the carbon isotope composition of the filter samples consisted of a two-stage oven, as described in detail by (Dusek et al., 2013). The filter pieces were heated stepwise in the first oven stage to temperatures of 100, 150, 200, 250, 300, 340, and 390°C, desorbing SOA compounds according to their volatility. In the second oven stage, the gaseous compounds were oxidised to CO$_2$ at 550°C using a platinum wool catalyst. The CO$_2$ was dried and purified using two cold traps and a gas chromatography column before it was analysed in a Delta Plus XL isotope ratio mass spectrometer (IR-MS; Thermo). The catalyst was charged using pure O$_2$ and then a filter piece was placed in the oven which was subsequently flushed with helium. The heating of the filter and subsequent purification of CO$_2$ took place in a helium carrier gas flow. Filter samples were bracketed between blank filter samples. IR-MS detects each isotopologue of CO$_2$ as a distinct peak with an associated peak area. The $\delta^{13}$C value was calculated from these areas (see below). Reported data were corrected by taking the corresponding blank filter measurements into account.

Isotope data are commonly reported in delta notation, using Vienna Pee Dee Belemnite (VPDB), as an element-specific international standard for $^{13}$C:

$$\delta^{13}C = \frac{R_{sa}(^{13}C)}{R_{VPDB}(^{13}C)} - 1.$$  \hspace{1cm} (2)

Here, $R_{sa}(^{13}C)$ and $R_{VPDB}(^{13}C)$ denote the isotope ratios ($^{13}$C / $^{12}$C) in the sample and standard, respectively. For typical ambient samples, the reproducibility lies below 0.3% for oven temperatures below 200°C and below 0.5% for oven temperatures above 200°C (Dusek et al., 2013).

In this study isotopic compositions of filter material are discussed relative to the isotopic composition of the initial $\alpha$-pinene, $\delta^{13}$C$_{TC}$, where TC denotes total carbon analysis (see below). Changes in isotopic composition are then reported as an isotopic difference (Coplen, 2011):

$$\Delta(13C) = \delta^{13}C - \delta^{13}C_{TC}.$$  \hspace{1cm} (3)

where $\Delta(13C) > 0$ indicates enrichment and $\Delta(13C) < 0$ indicates depletion in $^{13}$C with respect to the initial $\alpha$-pinene.

### 2.6 Total carbon isotope analysis of $\alpha$-pinene and selected filters

The $\alpha$-pinene used in the smog chamber experiments and selected filters (see Table 2) were transferred into tin capsules (4 x 6 mm capsules from Lüdi AG, Flawil, Switzerland), weighed, and analysed for total carbon isotope composition, $\delta^{13}$C$_{TC}$ in the ISOLAB of the Max Planck Institute for Biogeochemistry in Jena, Germany. The analytical setup comprised an elemental analyser (EA-1100, Carlo Erba, Milan, Italy) which was coupled to a Delta+ IR-MS (Finnigan MAT, Bremen, Germany) through a ConFlo III interface (Werner et al., 1999). The complete system was described by Brooks et al. (2003). All $\delta^{13}$C$_{TC}$ isotope ratios were referenced against the VPDB scale using an in-house working standard which itself is referenced against NBS-22 (National Bureau of Standards, now NIST), with a prescribed value of $-30.03$% (Coplen et al., 2006). Blank tests within each measurement sequence were used for blank correction. The analytical performance was maintained and monitored according to a measurement protocol that was described by Werner and Brand (2001). The estimated uncertainty of the $\delta^{13}$C$_{TC}$ analysis was $0.11$%, based on long-term performance records. The total carbon isotopic composition is reported here as $\delta^{13}$C$_{TC}$ for the initial $\alpha$-pinene used in the smog chamber experiments and as isotopic difference with sub-TC, $\Delta^{13}$C$_{TC}$ = $\delta^{13}$C$_{TC}$ - $\delta^{13}$C$_{TC}$, for all other samples.

### 2.7 Position-specific isotope analysis of $\alpha$-pinene

Quantitative NMR spectrometry tuned for isotopic measurements was used to quantify the relative abundance of each carbon isotopomer of $\alpha$-pinene. The sample preparation for NMR analysis consisted of the successive addition of 100 µL of a relaxing agent, Cr(Acac)$_3$ (Merck) at 0.1 M in the lock substance, Acetone–d$_6$ (EURISOTOP), to a 4 mL vial. Then 600 µL of pure $\alpha$-pinene was added and the mixture was introduced into a 5 mm NMR tube. Quantitative $^{13}$C NMR spectra were recorded using a Bruker AVANCE III connected to a 5 mm internal diameter BBFO probe tuned to the recording frequency of 100.62 MHz. The temperature of the probe was set to 303 K, without tube rotation. The acquisition conditions were those recommended in previous works.
(Bayle et al., 2014b; Silvestre et al., 2009) and are detailed in Sect. S3. Isotope $^{13}\text{C}/^{12}\text{C}$ ratios were calculated from processed spectra (see Fig. S4) as described previously (Bayle et al., 2014b; Silvestre et al., 2009); see Sect. S5. The measured position-specific isotopic compositions are given in delta notation and denoted as $\delta_i(^{13}\text{C})$ where $i$ denotes the position of the C atom; see Fig. 7. Typical accuracy of $\delta_i(^{13}\text{C})$ is 1 ‰.

Total carbon isotopic abundance, $\delta_i^{\text{TC}}(^{13}\text{C})$, was determined by IR-MS using an Integra2 spectrometer (Sercon Instruments, Crewe, UK) linked to a Sercon elemental analyser. Here, $j$ denotes different $\alpha$-pinene samples as listed in Table 1 (PSIA was only performed for samples $j = 2...5$). A precision balance (Ohaus Discovery DV215CD) has been used to introduce 0.5 mg of pure $\alpha$-pinene into tin capsules ($2 \times 5$ mm, Thermo Fisher scientific), before loading them into the elemental analyser. The instrument was referenced against the VPDB scale using international reference materials NBS-22 ($\delta_{\text{VPDB}}(^{13}\text{C}) = 30.03$ ‰), SUCROSE-C6 ($\delta_{\text{VPDB}}(^{13}\text{C}) = -10.80$ ‰), and IAEA-CH-7 PEF-1 ($\delta_{\text{VPDB}}(^{13}\text{C}) = -32.15$ ‰) (IAEA, Vienna, Austria). Instrumental deviation was followed via a laboratory standard of glutamic acid. The corresponding position-specific isotopic difference is reported as $\Delta_i(^{13}\text{C}) = \delta_i(^{13}\text{C}) - \delta_j^{\text{TC}}(^{13}\text{C})$.

3 Results

3.1 Aerosol characteristics

The evolution of SMPS-derived size distributions and total mass concentrations over time show that the aerosol population inside the bag was nearly constant for several days after the first day; see Fig. S2 and panel (a) in Fig. S3. Sampling on filters was started ca. 24 h after the injection of VOCs into the chamber. The CCN activity of SOA generated in this study resembles literature data for $\alpha$-pinene SOA generated in batch mode chambers (Fig. S3b). The integrated SMPS size distribution provides an estimate of the total SOA mass concentration in the chamber: 22 and 25 µg m$^{-3}$ for experiment B and C, respectively, although particles larger than 500 nm are not accounted for. A detailed characterisation of the chamber aerosol can be found in Sect. S2.

3.2 Chemical composition

3.2.1 Blank filter

Very low surface loadings (0.23 µg cm$^{-2}$) were found on the blank filter (HB); see Table 2. No peaks above 220 Da were detected. C$_2$H$_4$OH$^+$ (corresponding to protonated acetone) was the only compound found on the blank filter in concentrations above 5 ng cm$^{-2}$ at 100 °C, while all other ions showed temperature-independent concentrations below that.

High concentrations of acetone desorbed from all filters at 100 °C and may be an artefact from cleaning, although acetone is produced in the reaction, as shown in Fig. 1. The total surface loadings of the front and back filters always exceeded those on the blank filter except for single ions with concentrations near the detection limit, sometimes found desorbing from back filters at temperatures other than 150 °C. Correction for the blank filter concentration then resulted in negative concentrations for these ions for the back filters. These cases were neglected in the further data analysis.

3.2.2 Concentration thermograms

Figure 2 shows the sum of mass concentrations at each temperature step, as measured by PTR-MS. All front filters (B1f, B2f, C1f, and C2f) show a similar profile with most of the mass desorbing at 150 °C. The back filters (B1b and C1b) are used to characterise the positive sampling artifact, namely gas-phase compounds that adsorb to the quartz-fibre filters. Material collected on QBQ back filters can be assumed to mainly consist of adsorbed gas-phase compounds corresponding to a positive gas-phase artefact (Cheng and He, 2015). This is confirmed by the chemical analysis of back filters in this study, which differs considerably from that of the corresponding front filters, as detailed in Sect. S3. The back filters show small mass loadings – roughly 6 and 13% of the masses of their respective front filters. The large mass difference between front and back filters suggests efficient sampling of a dominant aerosol phase on front filters and a small positive sampling artifact. The front filters were not corrected for the sampling artifact.

The total SOA mass concentration in the chamber derived from PTR-MS measurements was of the order of 10 µg m$^{-3}$; see Table 2. Overall, the SMPS measured total mass concent-
trations that were up to 4 times higher than the concentrations measured by PTR-MS. This difference might be due to some factors related to the individual steps of the chemical analysis, e.g. filter sampling, extraction from the filter, and analysis by PTR-MS. An earlier study using an impactor-based thermal-desorption PTR-MS concluded that the total aerosol mass measured was typically 20% lower than the total aerosol mass measured with an SMPS (Holzinger et al., 2010b). The authors estimated conservatively that their PTR-MS setup detected 55–80% of the total aerosol mass. Filter sampling losses of up to 10% were attributed to negative sampling artefacts, i.e. evaporation from the filter, during sampling times of 24 h or longer in earlier work (Subramanian et al., 2004). The maximum desorption temperature during chemical analysis was only 350°C, and previous studies on β-pinene ozonolysis and photo-oxidation of terpenes also showed significant remaining volume fractions at desorption temperatures exceeding 400°C (Emanuelsson et al., 2013, 2014). Finally, charring and fragmentation in the PTR-MS can additionally lower PTR-MS-derived total mass concentrations. Section S4 describes these processes in more detail, as well as other aspects relevant to PTR-MS data interpretation.

Figure 3 shows mass concentration thermograms of specific compounds desorbing from front filter C1f. Table 4 complements information in Fig. 3, and Sect. S6 gives the full list of ions detected by PTR-MS from filter C1f. In Fig. 3 most ions show the highest concentrations at a desorption temperature of 150°C, in agreement with Fig. 2.

A pure compound is expected to desorb from the filter at temperatures between its melting and boiling temperatures. (Dusek et al., 2013) observed this on the same analytical setup for dicarboxylic acids. In principle, pure compounds will be detected by the PTR-MS as an ion of similar mass and only in this temperature window. There are several possible reasons why the same ion is observed over a range of temperatures. Since fragmentation of chemical compounds can occur during thermal desorption in the oven and ionisation in the PTR-MS (see Sect. S4), a fraction of the detected ions is likely fragments of larger (heavier) compounds. This fragmentation can occur at all desorption temperatures and consequently fragments are detected over a range of temperature.

Table 3. Main references cited in this study for comparison of chemical composition. The identifier (last column) denotes the letter the reference corresponds to in Table 4 and Sect. S6. Abbreviations: QP: quadrupole; TD: thermal desorption; TOF: time-of-flight; LC: liquid chromatography; MS: mass spectrometer; GC: gas chromatography; HP: high performance.

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<th>Sample phase</th>
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<th>Identifier</th>
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<td>(Holzinger et al., 2010a)</td>
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<td>LC-MS</td>
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<td>(Jaoui and Kamens, 2003)</td>
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<td>(Jenkin, 2004)</td>
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<td>Gas + aerosol</td>
<td>Modelling</td>
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<td>(Camredon et al., 2010)</td>
<td>α-pinene ozonolysis</td>
<td>Gas + aerosol</td>
<td>MCM Modelling</td>
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</table>

* MCM names.
The ozonolysis of weight dimer esters contribute significantly to SOA from the single compound. Recent studies show that high molecular is reached, which might differ from the melting point of the desorb significantly, when the melting point of the mixture et al., 2008). A specific compound in this mixture will only an SOA particle usually does not consist of a single com-

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<tr>
<td>141.089</td>
<td>(C$_2$H$_4$O)$_2$H$^+$</td>
<td>39.91 (24)</td>
<td>0.03</td>
<td>2,2-Dimethyl-cyclobutyl-1,3-diethanal</td>
<td>a, d (d)</td>
<td></td>
</tr>
<tr>
<td>171.065</td>
<td>(C$_2$H$_4$O)$_2$H$^+$</td>
<td>42.34 (21)</td>
<td>0.03</td>
<td>Not norpinic acid</td>
<td>a, b, c, d −1 (−)</td>
<td></td>
</tr>
<tr>
<td>171.098</td>
<td>(C$_2$H$_4$O)$_2$H$^+$</td>
<td>18.75 (48)</td>
<td>0.07</td>
<td>Norpinic acid (?)</td>
<td>b, c, d</td>
<td></td>
</tr>
<tr>
<td>183.099</td>
<td>(C$_2$H$_4$O)$_2$H$^+$</td>
<td>26.00 (35)</td>
<td>0.03</td>
<td>C109CO, 4-Oxopinonaldehyde</td>
<td>b, c, d +1 (232), +2 (−)</td>
<td></td>
</tr>
<tr>
<td>185.117</td>
<td>(C$_2$H$_4$O)$_2$H$^+$</td>
<td>5.31 (114)</td>
<td>0.12</td>
<td>Pinonic acid, OH-pinonaldehyde, PINONIC, C107OH, C109OH</td>
<td>(a), b, c, d −1 (51), −2 (132)</td>
<td></td>
</tr>
<tr>
<td>187.093</td>
<td>(C$_2$H$_4$O)$_2$H$^+$</td>
<td>11.37 (71)</td>
<td>0.01</td>
<td>Pinic acid, 10-OH norpinic acid, PINIC</td>
<td>b, b, c, d −1 (1), −2 (22), −3 (200)</td>
<td></td>
</tr>
<tr>
<td>199.093</td>
<td>(C$_2$H$_4$O)$_2$H$^+$</td>
<td>12.87 (66)</td>
<td>0.01</td>
<td>Oxopinonic acid, keto-pinonic acid</td>
<td>b, d −1 (27), −2 (105), +1 (380)</td>
<td></td>
</tr>
</tbody>
</table>

pending on the scavenger used, influencing the volatility distribution of products of α-pinene SOA. Overall, an increase of SOA yield is predicted when using cyclohexane as OH scavenger compared to 2-butanol (Jenkin, 2004). Assuming that 1-butanol behaves like 2-butanol (Shilling et al., 2008), the larger desorbed aerosol mass detected by PTR-MS in the cyclohexane experiments is consistent with these considerations. However, the most abundant reaction products were not affected.

The chemical composition of aerosol found in this study was compared to previously published chemical compositions of ambient aerosol and SOA derived from α-pinene ozonolysis (in dry, dark, and low-NOx conditions). Figure 3b shows concentration thermograms of ions with masses attributed to compounds reported previously; see Table 4. The PTR-MS allowed discrimination of particulate reaction products reported earlier (Holzinger et al., 2005, 2010a; Winterhalter et al., 2003; Jenkin, 2004; Jaoui and Kamens, 2003). In many cases the identification could be positively confirmed, but in some cases an original attribution could be falsified (details can be found in Sect. S4). Some references in Table 4 and in Sect. S5 are bracketed to indicate when the assigned formulae for ion masses differed from those in the literature. Compounds predicted by modelling studies are noted by their capitalised Master Chemical Mechanism name in the description field in Table 4 and Sect. S6, confirming the presence of several predicted species. These compounds include pinic acid (compound no. 72 in Sect. S6) and pinonic acid (no. 116 in Sect. S6) which are also shown in Fig. 1.

Figure 4 shows the mass spectra of compounds that desorbed from filter C1f at 150°C. Most ions have a mass below 250 Da, indicating no direct observation of oligomers. However, several fragmentation patterns were detected in the data, as highlighted by the arrows in Fig. 4: the light green arrow connects peaks with a mass difference of 14.016 Da (corresponding to a CH₂ group), and the dark blue arrow connects peaks with a mass difference of 18.011 Da (water). Details on fragmentation patterns can be found in Sect. S4. Fragmentation patterns in the mass spectra indicate that large compounds like oligomers and/or complex organic mixtures were present on the filters, as also reported in the literature (Docherty et al., 2005; Gao et al., 2004; Cappa et al., 2008; Kristensen et al., 2016), but that these compounds decompose during desorption or in the PTR-MS during ionisation before they can be detected as smaller ions. While the arrows in Fig. 4 indicate the possibility that the peaks, separated by a water or CH₂ mass, are from the same molecule, this constitutes only circumstantial evidence and not proof.

### 3.2.3 O : C ratio

Figure 5 shows the measured O : C ratio versus desorption temperature for selected filters. The O : C ratio of desorbed material increases from 0.18 to 0.25 when the desorption temperature increases from 150 to 250°C. At higher desorption temperatures the O : C ratio levels off and remains constant at ca. 0.25. The O : C ratios of material desorbing from the back filters are similar to those of the front filters with averages of 0.21 (B1b) and 0.22 (C1b); see Sect. 2.

The observed increase in the O : C ratio at increasing desorption temperatures (below 250°C) is to be expected if functionalisation yields oxygenated compounds with lowered volatility (Jimenez et al., 2009; Holzinger et al., 2010b). For material desorbing at temperatures above 250°C, this correlation seems to break down and oligomerisation is likely to be more important. Oligomerisation reactions can be accompanied by the exclusion of water (Tolocka et al., 2004) that goes undetected in the PTR-MS, lowering O : C ratios. The desorption temperature of formed oligomers and complex mixtures is also higher than the desorption temperature of their single constituents (Cappa et al., 2008). Therefore oligomerisation can decrease the volatility of compounds without raising the O : C ratios, resulting in the observed plateau of O : C ratios at high desorption temperatures.

Literature values for O : C ratios of SOA from α-pinene ozonolysis are generally somewhat higher than the values reported here: Shilling et al. (2008) investigated SOA from α-pinene ozonolysis and reported O : C ratios around 0.33 for aerosol loadings, which are comparable to this study. Aiken et al. (2008) report O : C ratios around 0.3 for laboratory SOA from α-pinene ozonolysis and observe more oxidised aerosol in ambient samples. These two studies (Shilling et al., 2008; Aiken et al., 2008) used aerosol mass spectrometers to determine the O : C ratio, while Holzinger et al. (2010a) employed a PTR-MS, as in the present study. They report a measured O : C ratio of 0.33–0.48 for remote ambient aerosol in the Austrian Alps.

PTR-MS measurements may underestimate O : C ratios because of several factors, including charring and fragmentation due to ionisation; see Sect. S4. Holzinger et al. (2010a) assessed in detail how oxygen loss – common in PTR-MS measurements – lowers O : C ratios. However, this was not taken into account for data reported here.
3.3 Isotopic composition

3.3.1 Total carbon and thermally desorbed material

Figure 6, panel (a), shows the isotopic composition of total carbon on the filters relative to the isotopic composition of the precursor α-pinene. The aerosol on all front filters is enriched in carbon 13 relative to the initial α-pinene, and the enrichment is larger for filter B1f (1.2 ‰) than for filters C1f (0.6 ‰) and C2f (0.7 ‰). Compounds desorbing from back filter C2b, which very likely represent gas-phase material, are depleted by 0.8 ‰.

Figure 6b shows Δ(13C) of thermally desorbed filter material as a function of desorption temperature. Aerosol on the front filters shows a 13C enrichment of 0.2–2.8 ‰ relative to the initial compound. The most volatile fraction that desorbed at 100 °C consistently shows the highest enrichment. SOA compounds desorbing at 100 °C are enriched in 13C compared to 150 °C by about 0.7 to 1.9 ‰; see Table 2. The Δ(13C) values of SOA do not change significantly with temperature at desorption temperatures above 150 °C. SOA formed in the presence of 1-butanol (filter B1f) is enriched by an additional 0.2–1.3 ‰ compared to SOA formed in the presence of cyclohexane (filters C1f and C2f). The higher enrichment observed for SOA formed in the presence of 1-butanol scavenger relative to cyclohexane scavenger may be because the latter has a higher SOA yield. This is corroborated by the PTRMS data; see Table 2. The gas-phase compounds desorbing at 150 °C from back filter B1b are depleted by 0.7 ‰ compared to the initial α-pinene and depleted by 1.9 ‰ with respect to the particulate SOA on filter B1f. This is expected due to isotopic mass balance. The isotopic enrichment of gas-phase compounds on filter B1b at 100 and 350 °C is similar to that of the corresponding particulate matter on front filter B1f. The low concentrations detected on the back filters at these desorption temperatures preclude any in-depth discussion of the enrichment seen at those temperatures.

Values of ΔTC(13C) are close to Δ(13C) values at 150 °C (Fig. 6 and Table 2). Values of ΔTC(13C) represent a convolution of the volume-normalised peak area and Δ(13C), with a dominant contribution of Δ(13C) at 150 °C. Total carbon analysis, however, misses details like the enrichment of the most volatile mass fraction desorbing at 100 °C.

3.3.2 Position-specific isotope analysis of α-pinene

Table 1 lists Δi(13C) for each C atom of all analysed α-pinene samples. Single sites show variations between −6.9 and 10.5 ‰. The α-pinene samples from manufacturers Sigma-Aldrich (different lot from SOA experiments), Acros Organics, and Merck have similar position-specific isotope profiles. It is likely that these three manufacturers sell α-pinene with the same origin (e.g. natural) and use similar preparation techniques. The last sample (from Alfa Aesar) has a slightly different profile which would seem to indicate a different origin and/or purification method. Samples of α-pinene from Acros Organics (j = 3) and Merck (j = 4) have similar Δi(13C) profiles but differ by 1.1 ‰ in their bulk value.

Unfortunately, PSIA could not be performed on the α-pinene used in chamber experiments (it was used up and is no longer available from the manufacturer). The α-pinene used in chamber experiments has a bulk isotopic composition of δ29v13C(1) = (−29.96 ± 0.08) ‰, which differs by up to 3 ‰ from the other α-pinene samples; see Table 1. Given that the bulk isotopic composition is not correlated with the position-specific isotope profiles, and assuming that the manufacturer did not change product origin or purification method, it is probable that the two α-pinene samples from Sigma-Aldrich share similar position-specific isotope profiles. It is therefore assumed for the remainder of the discussion that the α-pinene used for the SOA experiments has the same position-specific isotope profile as the batch from
Table 5. Calculated $\Delta^{(13}C)$ values, assuming that fragmentation yields certain C atoms to be expelled to the gas phase and others to partition to the aerosol phase (details in Sect. 4.1). Calculations based on position-specific measurement results of the Sigma-Aldrich sample in Table 1. See Fig. 7 for numbering of C atoms.

<table>
<thead>
<tr>
<th>Expelled C atom(s)</th>
<th>$\Delta_{\text{gas}}^{(13}C$) ($%$)</th>
<th>$\Delta_{\text{aerosol}}^{(13}C$) ($%$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>$C_1$</td>
<td>4.9</td>
<td>-0.4</td>
</tr>
<tr>
<td>$C_2$</td>
<td>10.5</td>
<td>-1.1</td>
</tr>
<tr>
<td>$C_7$</td>
<td>0.6</td>
<td>0.0</td>
</tr>
<tr>
<td>$C_9$</td>
<td>-6.7</td>
<td>0.8</td>
</tr>
<tr>
<td>$C_8 + C_1 + C_9$</td>
<td>-0.7</td>
<td>0.4</td>
</tr>
<tr>
<td>$C_{10} + C_1 + C_9$</td>
<td>-2.0</td>
<td>1.0</td>
</tr>
</tbody>
</table>

Sigma-Aldrich on which PSIA was performed. The $\Delta_i^{(13}C)$ distribution of that sample is visualised in Fig. 7.

4 Discussion

In this experiment we have run $\alpha$-pinene ozonolysis to completion and analysed the SOA and some of the gas-phase material. After comparing the IR-MS (Fig. 6) analysis with the PTRMS data (Fig. 5), we found that $\Delta^{(13}C$) does not show a correlation with the O : C ratio. The PTRMS data showed that the isotopic enrichment in the SOA is carried by a broad range of oxidation products rather than one or a few dominant products. We argue that the observed isotopic abundances are largely due to a combination of isotope-dependent changes in product branching ratios and unequal partitioning of carbon atoms from specific sites in $\alpha$-pinene into the aerosol and gas phases. Additional effects could include isotope-dependent variations in partitioning of semi-volatile compounds between the gas and condensed phases.

4.1 Can position-specific isotope analysis of $\alpha$-pinene explain the enriched aerosol phase?

Here, a limiting case is presented for the isotopic difference of a number of plausible oxidation products. The underlying assumptions exclude isotope-dependent changes in product branching ratios, as well as effects of temperature, relative humidity, pressure, ozone concentration, etc. This simplistic approach allows for the estimation of the maximum isotopic enrichment in $\alpha$-pinene fragments using the $\Delta_i^{(13}C$) profiles obtained from PSIA. Table 5 shows predicted maximal enrichments or depletions if a single carbon atom or reasonable combinations of three carbon atoms are split off the parent compound. Based on the chemical reaction pathways presented in Fig. 1, volatile reaction products such as acetone, CO, and formaldehyde can reasonably be assigned to specific sites in the parent $\alpha$-pinene. The minor (potentially gaseous) expelled fragment is predicted to have an overall isotopic difference relative to the initial $\alpha$-pinene, $\Delta_{\text{gas}}^{(13}C$), similar to the measured $\Delta_i^{(13}C$) value for the carbon atom’s former position, as seen in Fig. 7. The larger fragment, which would partition to the aerosol phase, is predicted to have an overall $\Delta_{\text{aerosol}}^{(13}C$) value equal to the average of the $\Delta_i^{(13}C$) values of the remaining C atoms. For example, the pathway leading to formaldehyde in the sixth box in Fig. 1 is predicted to deplete formaldehyde by $\Delta_{\text{gas}}^{(13}C$) = 6.7% relative to the initial compound and leave the corresponding major fragment (denoted as “R” in Fig. 1) enriched by $\Delta_{\text{aerosol}}^{(13}C$) = 0.8% here. Here, $\Delta_{\text{gas}}^{(13}C$) denotes the mean of $\Delta_i^{(13}C$) values of C atoms 1–8 and 10. Expelled C atoms from positions with small $\Delta_i^{(13}C$) values, e.g. C2, will only have a small impact on the isotopic composition of the remaining fragment. For expulsion of C2, a depletion of −1.1% is predicted for the aerosol fragment relative to the initial $\alpha$-pinene.

If three carbon atoms are expelled, as in the case of acetone, the enrichment of the minor fragment relative to the initial $\alpha$-pinene is calculated as the average of the $\Delta_i^{(13}C$) values of the respective expelled positions C8, C1, C9 or C10, C1, and C9; see Fig. 1. The formation of acetone may involve methly migration of either the C8 or C10 atom. The gaseous fragments composed of three carbon atoms are predicted to show $\Delta_{\text{gas}}^{(13}C$) values of −0.7 and −2.0%, and the corresponding $\Delta_{\text{aerosol}}^{(13}C$) values for the larger fragment are 0.4 and 1.0%; see Table 5. These calculations are based on the measured position-specific enrichment for sample 1 in Table 1, but the results and conclusions drawn do not change significantly when performing similar calculations for the other $\alpha$-pinene samples where PSIA data are available (with the exception of the anomalous Alfa Aesar sample).
4.2 Branching ratios and other isotope effects

The observed $^{13}$C enrichment in compounds from the front filters (SOA) and the depletion in compounds from the back filters (gas phase) are generally larger or equal in magnitude to those predicted by position-specific isotope effect analysis (see Table 5). The data shown in Table 5 are limiting values and show that PSIA is capable, in principle, of producing abundances of similar magnitude to what was observed in our experiment. This mechanism does not include concomitant enrichment due to isotope effects on product branching ratios.

The other explanations include isotope-dependent changes in branching ratios in the reaction mechanism (Fig. 1) and incomplete reactions. It has been shown previously in simple systems (e.g. evaporation of solvents and sorption of vanillin) that each carbon position can have its own isotopic fractionation and that different positions can show opposing isotope effects at the same time (Höhener et al., 2012; Julien et al., 2015). In chemical reactions, the substitution of a $^{12}$C atom by $^{13}$C will affect isomerisation and stabilisation dynamics by changing vibrational frequencies with an associated change in zero point energies. Therefore, positions that are not reaction sites can also show isotope effects, which have been termed non-covalent isotope effects (Wade, 1999), as has been observed during the chain-shortening reaction for the bioconversion of ferulic acid to vanillin (Botosoa et al., 2009b). It is generally difficult to predict which position has which isotope effect, but it has been shown that isotopic substitution in ring structures at positions that carry functional groups lead to stronger position-specific isotope effects compared to positions that have no functional groups attached (Höhener et al., 2012; Botosoa et al., 2009b). Similarly, the C atoms in $\alpha$-pinene that are not part of the ring structure might have large position-specific isotope effects. Höhener et al. (2012) note that, for vanillin, such effects can produce large fractionations in the small fragments and small fractions in the large fragments, similar to what we show in Table 5. Accordingly, in $\alpha$-pinene there is an order of magnitude difference between the site-dependent enrichments seen in the C$_{10}$ molecule (6–10‰) and the enrichments seen in the SOA (0.6–1.2‰).

Kinetic isotope effects associated with fragmentation provide a possible explanation for the enrichment of the particle phase with respect to the gas phase. After the first two reaction steps (i.e. formation of the Criegee intermediate and the subsequent decomposition, stabilisation, or isomerisation of it; see Fig. 1) the products will be distributed between the gas and particle phases according to their respective partitioning coefficients. This partitioning is not expected to lead to significant isotopic fractionation within a chemical species. A number of these products are small volatile compounds that partition to the gas phase. The remaining particle phase would be enriched, which is consistent with observations reported previously (Aggarwal and Kawamura, 2008; Kirillova et al., 2013, 2014). Fragmentation followed by functionalisation is expected to be accompanied by elevated O : C ratios in both products, as detailed in the introduction. This is in line with the observed increasing O : C ratios of desorbed material from front filters at increasing desorption temperatures below 250°C (Fig. 5), and with O : C ratios that are generally elevated for back filter material desorbing at all temperatures (Table 2).

Large equilibrium fractionation due to gas-particle partitioning could also explain the observations, given that individual compounds might show isotope effects of 1–2‰ due to partial volatilisation. In an ensemble of compounds, however, most individual compounds are likely to be found predominantly in one phase; only a fraction of products have intermediate volatilities. In the case of $\alpha$-pinene ozonolysis shown in Fig. 1, small first generation products like CO, HCHO, and acetone will be gaseous and the larger entities, including prominent acids, will partition to particles due to their lower vapour pressure. Ongoing reactions (e.g. dimer formation and oligomerisation) will lower their vapour pressure even further. Equilibrium isotope effects of single compounds are therefore diluted and hence less likely to cause the observed isotopic enrichment in SOA.

The present study allows new insight to be gained into observations of isotopically enriched ambient SOA. In some cases isotopic enrichment in ambient aerosol was attributed to photochemical ageing during long-range transport (Kirillova et al., 2013; Pavuluri et al., 2011). However, Fu et al. (2012) note that a normal kinetic isotope effect cannot explain their observation of aerosol that was even more enriched than aerosol from biomass burning. The authors noted that such aerosol occurred predominantly during the day during episodes of high abundance of biogenic SOA. The present study suggests that the site-specific distribution of $^{13}$C in the source material itself governs the abundance of $^{13}$C in SOA.

5 Conclusions

The isotopic and chemical compositions of SOA generated from dark ozonolysis of $\alpha$-pinene were determined using isotope ratio mass spectrometry and thermal-desorption PTR-MS, and PSIA was applied for the first time on $\alpha$-pinene from a series of manufacturers. A key result is the first observations of strong site-dependent enrichments and depletions in $^{13}$C in $\alpha$-pinene, with values between $-6.9$ and $+10.5$‰ relative to the bulk composition. Total carbon from SOA collected on front filters was enriched in $^{13}$C by $0.6$–$1.2$‰ with respect to the initial $\alpha$-pinene precursor. Total carbon adsorbed on the back filters, designed to sample gas-phase compounds, was depleted by $-0.8$‰.

High-resolution data retrieved by a PTR-MS detected more than 400 ions. More than 90% of the total desorbed mass as measured by PTR-MS (from front filters) was un-
ambiguously identified and discussed in the context of the current literature. SOA mainly desorbed from the filters at 150 °C, and larger compounds likely formed in oligomerisation reactions decomposed during extraction or ionisation. Besides the fragments from such oligomers, single constituents of complex organic mixtures formed on the filter were also detected as single ions of lower mass which show significant non-zero concentrations at desorption temperatures higher than 150 °C. The observed constant O : C ratio at desorption temperatures exceeding 250 °C also indicates fragments of larger molecules. At lower temperatures the O : C ratio increases from 0.18 to 0.25, indicating functionalisation reactions during SOA formation. A comparison of the IR-MS results with the PTRMS data shows that Δ(13C) does not correlate with the O : C ratio.

Analysis of the isotopic composition as a function of desorption temperature showed that the isotopic composition of material desorbing at 150 °C was similar to the isotopic composition of total carbon. Functionalisation typically follows fragmentation in monoterpane ozonolysis and was shown to drive the O : C temperature profiles. The partitioning of products between the gas and condensed phases is an active area of research, and both chemical and physical effects must be considered. The partitioning is not simply governed by vapour pressure; some gas-phase products may be incorporated into SOA through oligomerisation. Many variables, including product branching ratios, temperature, pressure, humidity, OH, NOₓ, etc., can in principle have an effect on the isotopic abundance of 13C in SOA and this leaves room for further studies. The present study suggests that the site-specific distribution of 13C in the source material itself governs the abundance of 13C in SOA.


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Competing interests. The authors declare that they have no conflict of interest.

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