Atmospheric aqueous phases (cloud and fog droplets, aerosol liquid water) are important reaction media for the processing of water-soluble organic acids (OAs). The photochemistry of these species is known to contribute to the formation of aqueous secondary organic aerosols (aqSOA). OAs arising from the gas-phase oxidation of \( \alpha \)-pinene, such as cis-pinonic acid (CPA) or pinic acid (PA), are considered relevant aqSOA precursors, as they are oxygenated compounds and expected to be sufficiently water-soluble to undergo cloudwater processing. Although PA is a major oxidation product of \( \alpha \)-pinene, the mechanism of its aqueous-phase photo-oxidation has never been studied previously. In this work, we performed the first in-depth investigation for aqueous-phase OH-oxidation of PA. Given that the pH of atmospheric aqueous phases is highly variable, and it has been increasing over the past 40 years, we were particularly interested in the impact of pH on the reaction mechanisms and product yields. Experiments were performed using a combination of offline and online MS techniques to identify PA and monitor the evolution of PA OH-oxidation products. We identified PA as a novel source of important \( \alpha \)-pinene SOA tracers, including 3-methyl-1,2,3-butanetricarboxylic acid (MBTCA), and norpinic acid (NPA). Furthermore, we show a drastic difference in the yield of the oxidation products of PA under acidic and basic pH conditions. Specifically, we observed the yields of NPA and MBTCA at 3.4% and 2.6% under pH 2 but at 10% and 5.0%, respectively, at pH 8. These yield differences are likely due to a charge transfer reaction pathway unique to the aqueous phase. Such impact of pH on the reaction mechanism of OA has never been shown before.

Environmental significance

This work represents the first in-depth mechanistic investigation for the OH-oxidation of pinic acid (PA) in the aqueous phase. PA is a major product of \( \alpha \)-pinene, but its chemistry in the aqueous phase has never been studied. We identified norpinic acid (NPA) and 3-methyl-1,2,3-butanetricarboxylic acid (MBTCA) as major products from PA. We also observed drastically different molar yields of NPA and MBTCA under the acidic and basic conditions, likely due to a charge transfer reaction occurring only to the pinate ion. These results serve as a demonstration that solution pH can alter photochemical mechanisms. The rapidly changing cloud and fog pH in the ambient atmosphere can be affecting photochemistry and the products of organic acids.

1. Introduction

Water-soluble organic acids (OAs) can undergo photochemistry in atmospheric aqueous phases (cloud, fog droplets, and aerosol liquid water). The chemistry occurring in cloudwater, also referred to as cloudwater processing, results in the decay of OAs, and further formation of secondary organic aerosols (SOA). The gas-phase oxidation of biogenic volatile organic compounds (BVOC) are known sources of OAs and other multifunctional compounds found in cloud and fog waters. One of the largest BVOCs emitted annually is \( \alpha \)-pinene; thus, numerous studies focused on its gas-phase kinetics and mechanisms leading to SOA formation. It has been recognized that the aqueous-phase processing of semi-volatile oxidation products of \( \alpha \)-pinene can contribute to SOA formation, but these processes are characterized poorly and limited to a few model compounds, such as cis-pinonic acid (CPA) and limononic acid. Among the multifunctional \( \alpha \)-pinene products, CPA and pinic acid (PA) are first-generation OAs. CPA is the most targeted among the OAs mainly due to its commercial availability, while PA is rarely the target of systematic studies, since it is not readily available, and needs to be synthesized.
PA was identified first by Holloway et al.\textsuperscript{9} in 1955 as the product of liquid-phase ozonolysis of \( \alpha \)-pinene. Forty years later, Christoffersen et al.\textsuperscript{18} initiated a series of smog chamber studies that identified PA as one of the main oxidation products of \( \alpha \)-pinene gas-phase ozonolysis and as an important \( \alpha \)-pinene SOA component. In terms of known physicochemical properties of PA, Bilde et al.\textsuperscript{19} applied tandem differential mobility analysis to measure the evaporation rate and vapor pressure of PA and other semi-volatile oxidation products of \( \alpha \)-pinene. Kołodziejczyk et al.\textsuperscript{20} used a differential scanning microcalorimetry technique to measure PA’s thermal properties, as well as a solid–liquid phase equilibria method for the determination of PA solubility in water. An interesting finding from Kołodziejczyk et al.\textsuperscript{20} is that PA is the most water-soluble among other key \( \alpha \)-pinene SOA oxidation products (PA > \( \text{cis} \)-norpinic acid > CPA > \( \text{cis} \)-norpinonic acid). When cloudwater is present in an air mass, PA and other OAs from \( \alpha \)-pinene will be present exclusively in the aqueous phase.\textsuperscript{21} For chemical reactions of PA, a detailed gas-phase formation mechanism of PA has been proposed by Jenkin et al.\textsuperscript{22} The formation of PA in the gas phase and its particle-phase OH-oxidation has been investigated by many studies,\textsuperscript{5,16,18,23} but there is a lack of information on PA kinetics and thermodynamics in atmospheric aqueous phases, representing a major missing piece in multiphase atmospheric chemistry models.

The aqueous-phase OH-oxidation mechanism shares many similarities with that in the gas phase; however, mechanisms unique to the aqueous phase may lead to reaction pathways that are absent in the gas phase. Acid–base chemistry\textsuperscript{24} and rapid formation of carboxylic acids from aldehydes are representative examples.\textsuperscript{25} In addition, a unique charge transfer can occur to carboxylate but not carboxylic acid, giving rise to pH-dependent reaction mechanisms (Fig. 1). Briefly, the OH radical can react with an OA through H-abstraction and addition to the “R” group. H-abstraction from an O–H bond is slow, hence the OH radical does not react with the carboxyl group (COOH) rapidly. On the other hand, the OH radical is reactive to the carboxylate group (ROO\(^-\)) by abstracting an electron from it.\textsuperscript{26} Our recent study\textsuperscript{27} investigated, for the first time, the pH dependence of the OH-reactivity of PA. The major finding was that the kinetics of PA’s aqueous-phase oxidation is not affected significantly by the medium pH, likely because reaction with the “R” group dominates the OH reactivity of PA. Currently, there is no information available related to possible mechanistic differences under acidic and basic conditions. The pH-dependence of OH-oxidation is thought to be less significant compared to the sulfate radical (SO\(_4^{2-}\)) and nitrate radical (NO\(_3^\)), which are more effective one-electron oxidants than the OH radical.\textsuperscript{28} The overall objective of this investigation was to better understand the fundamental mechanism of PA OH-oxidation in the aqueous phase, given that it has never been studied previously. In particular, we aimed to identify and quantify major products arising from PA OH-oxidation. The impact of solution pH on the reaction mechanism and product formations was also investigated, as it remains a poorly explored research area.

2. Experimental

2.1 Materials

The following chemicals were purchased from Sigma Aldrich: CPA (98%), hydrogen peroxide (H\(_2\)O\(_2\), 30% in water), sulfuric acid (50% in water), and potassium hydrogen phthalate (>99.95%). Chemicals were also purchased from Fisher Chemical: tricarballylic acid (TCA, 99%), sodium hydroxide (98%), acetonitrile HPLC grade, and formic acid 99.0+%. All chemicals were used without further purification. PA was custom synthesized and purified as previously described.\textsuperscript{24}

2.2 Aqueous phase OH-oxidation

The aqueous-phase photo-oxidation of PA was performed using a photoreactor emitting UV-light centered at 310 nm (UVB). H\(_2\)O\(_2\) was added to the solution as the source of OH radical. As shown in previously published data,\textsuperscript{24} the photoreactor (Rayonet, PRP200) emitting UV light was able to efficiently photolyze H\(_2\)O\(_2\) into OH radicals at atmospherically relevant concentrations, which have been estimated to be 1.4 \(\times\) 10\(^{-16}\) to 8.0 \(\times\) 10\(^{-12}\) M.\textsuperscript{26} We conducted an H\(_2\)O\(_2\) control experiment to confirm that PA does not undergo direct photolysis in the absence of H\(_2\)O\(_2\), and this result has already been published in Amorim et al.\textsuperscript{24} https://www.zotero.org/google-docs/?DVqqW5. The UVB light also does not interact with products detected in this work. However, we note that the UVB light can directly photolyze reaction products, especially those containing carbonyl and peroxidic functional groups. The reaction progress was monitored via offline and online methods, as shown in the schematic of the experimental setup in Fig. 2. Electrospray ionization (ESI) high-resolution liquid chromatography (LC)-mass spectrometry (MS) was employed for the offline analysis, and online particle-into-liquid sampler (PILS) coupled to ESI-MS was used for the online analysis of PA reaction products. The experimental conditions used are summarized in Table 1. PA concentrations in ambient cloudwater are likely very dilute, van Pinxteren et al.\textsuperscript{27} have reported sub-\(\mu\)M concentrations of PA in cloudwater samples collected at a mountain site in Germany. In this work, the initial concentration of PA was chosen to be 114 or 330 \(\mu\)M for the offline and online experiments, respectively. These concentrations were necessary to provide sufficient product signals for the mechanistic analysis. To ensure the

![Fig. 1 A schematic illustration of the carboxylate charge transfer reaction occurring only when pH of the solution is greater than the pK\(_a\) of the carboxylic acid. This charge transfer reaction does not proceed with the carboxyl group (i.e., pH < pK\(_a\)).](https://www.zotero.org/google-docs/?DVqqW5)
accuracy of the initial concentration of PA, the PA stock solution was standardized using proton nuclear magnetic resonance (1H NMR) spectroscopy (see Fig. S1 for details in the ESI).

For the offline reaction monitoring, the effect of pH on the reaction mechanism of PA was evaluated, and the pH of the solution was adjusted by adding H2SO4 50% wt (to pH 2) or NaOH 0.5 M (to pH 8). From the decay rate of PA, it was estimated that the steady-state concentration of OH ([OH]ss) was at 1.4 x 10^-12 M at pH 2 and 7.4 x 10^-13 M at pH 8. This concentration is within the range of the estimated concentrations in ambient cloudwater.26,28 The contribution of direct photolysis to the overall PA decay has been previously investigated and presented by Amorim et al.24 All offline experiments were conducted in triplicates to ensure reproducibility and to enable error analysis for quantitative information.

For the online analysis (details in Section 2.3.2), the pH of the solutions was kept uncontrolled, i.e., with no addition of acid or base. These salts can cause the suppression of the analyte signal in ESI; thus, no base or acid was added for the online analysis. To overcome the detection limit of the online MS, a higher initial PA concentration (330 μM) was chosen. To maintain a similar [OH]ss, H2O2 concentration was increased accordingly to 20 mM, which resulted in an estimated [OH]ss of 6.69 x 10^-13 M. A duplicated online analysis was conducted to ensure the reproducibility. Given that this experiment required a higher concentration and a larger volume of PA solution, we could not afford more replicates with a limited amount of PA synthesized.

2.3 Offline and online sampling and analysis

2.3.1 Offline (−)-ESI-LC-MS analysis of reaction products. Aliquots of the reaction solution were collected as a function of experimental time for offline LC-MS analysis. The separation and identification of the elemental composition of PA oxidation products were performed offline with an Agilent 6220 accurate-mass time-of-flight (ToF) LC-MS system (Agilent, CA, USA). The chromatographic separation was performed on a Kinetex C18 reverse-phase (50 mm x 2.1 mm I.D., 3.0 μm particle size) column. The C18 column was equipped with a security guard cartridge with a 2.1 mm ID C18 pre-column and kept at 50 °C. Mobile phase A consisted of 0.1% formic acid solution in water (pH 2.8), and mobile phase B was 0.1% formic acid in acetonitrile (ACN), delivered at a flow rate of 0.5 mL min^-1; the injection volume was 2.0 μL. The following mobile phase gradient was applied: 0.0–0.5 min isocratic 1% B, 0.5–5.5 min linear gradient to 95% B, 5.5–6.5 min isocratic 95% B, 6.5–7.0 min linear gradient to 1% B. The column was re-equilibrated at 1% B, and the analysis completed in 12.0 min. The MS acquisition was performed in scan mode (m/z of 0–500) and the detection of compounds was performed using negative ion ESI ((−)ESI) due to its selectivity towards OAs.29 The following ESI conditions were applied: capillary voltage, 3000 V; fragmentor, 125 V; drying gas temperature, 325 °C; drying gas flow, 9 L min^-1, and nebulizer pressure, 20 psi.

2.3.2 PILE-EI-MES analysis of reaction products. A unique online monitoring system was developed to monitor the evolution of PA OH-oxidation and product formation continuously (Fig. 2). An atomizer (TSI model 3076) was used to draw the sample solution from the reaction vessel by using a fast airflow that creates a negative pressure. Therefore, a small proportion of the sample solution is sprayed out as aerosol,
while the rest of the solution returns to the original container. The sprayed aerosol is introduced to a PILS (Brechtel Manufacturing Inc., Model 4001), which collects and extracts the atomized aerosol particles into water in real-time. The operating mechanism of PILS has been discussed extensively in the literature. Briefly, the sample aerosol is mixed with saturated water vapor to grow the particles into larger droplets. Then, the droplets are impacted onto a vertical impactor surface, and the water-soluble species collected on the impactor are washed off with an aqueous solution. This aqueous solution is sent to an ESI-triple-quadrupole MS (Varian, 320-MS) by a peristaltic pump at a flow rate of 18 µL min⁻¹. We note that the online PILS-ESI-MS is used solely for mechanistic analysis, tracking the formation of multi-generational reaction products forming in the reaction system. Quantitative analysis using PILS-ESI-MS is challenging due to numerous factors, including the atomization efficiency, extraction efficiency in PLS, and carry-over contamination in the MS. The triple-quadrupole MS is unit resolution; thus, products with the same nominal mass cannot be distinguished.

For the online PILS-ESI-MS experiments, signals from the PA + H₂O₂ solution were recorded until a stable signal was obtained using the MS. The OH-oxidation is then initiated by turning the UV lamps on. The MS was operated in the continuous selected- ion monitoring (SIM) acquisition mode to track the evolution of selected species. A list of targeted species was generated by consulting the literature and our own offline measurements. To assist product identification, the MS was occasionally operated in flow injection mode, with the experimental solution injected with a syringe pump. During this time, the MS was operated under the MS/MS mode. A list of parameters for the triple quadrupole MS is summarized in the ESI (Table S1†).

2.4 Determination of the acid dissociation constant (pKₐ) of PA

To better understand the acid dissociation equilibria of PA, PA’s pKₐ was experimentally determined via titration with NaOH. The pKₐ value can be determined fairly accurately from the titration curve using the “half-volume” method. Therefore, 25.0 mL of 5.0 mM PA solution was titrated manually with 5.0 mM NaOH. The NaOH solution was standardized previously by titration with potassium hydrogen phthalate. The pH of the PA solution was measured continuously with a calibrated pH meter (Sartorius).

3. Results and discussion

3.1 Determination of pKₐ

PA contains two ionizable carboxyl functional groups sharing a chemically similar environment; thus, under acidic conditions, the dissociation of both acidic protons will happen independently. Although PA is a di-acid, having two dissociation constants (Kₐ,1 and Kₐ,2), only one equivalence point was noticeable on the titration curve shown in Fig. 3. This indicates that the difference in PA’s first and second dissociation constants is small, and the equivalence point observed is from the merged neutralization of both acidic protons.

Although it is not possible to determine PA’s first and second pKₐ values experimentally via acid–base titration, one can use the “half-volume” method to determine its “effective” pKₐ (pKₐ(eff)). The pKₐ(eff) is useful to estimate the fraction of neutral PA and pinic anion (PAn) in a solution. By applying the method described in Section 2.4, the PA’s pKₐ(eff) was measured to be 4.9. To the best of our knowledge, the pKₐ value of PA has never been determined experimentally. A computationally determined value of 4.48 was used in the literature. Based on our observation, subsequent experiments in this work were conducted under pH values of 2 and 8. The PA’s pKₐ(eff) indicates that the neutral form of PA dominates at pH 2 (99.9%), while at pH 8, the majority of PA will be dissociated fully and be present as PAn. The pH values of ambient cloudwater and fogs fall between 2 and 7. Therefore, in atmospheric aqueous phases, PA can exist as a mixture of its neutral form and as a PAn ion. The ratio of which varies drastically with changing cloudwater compositions. For this reason, it is important to fundamentally study PA oxidation mechanisms under both acidic and basic conditions, in which PA exists only in one of the two forms.

3.2 Offline (−)-ESI-LC-MS for elemental analysis

The products arising from the aqueous-phase PA OH-oxidation were detected and identified using (−)-ESI-LC-MS. Many peaks were detected at retention times shorter than that of PA, which indicates the formation of more oxygenated compounds. Table 2 shows the major oxidation products detected, with their elemental compositions and potential identities. We included all detected peaks with a signal intensity larger than 5.2 x 10⁴ cps, and the m/z values correspond to their deprotonated forms ([(M–H)⁻]). We have also performed CPA OH-oxidation and detected all the oxidation products previously identified by Witkowski et al. (Table S2†). The results of the CPA OH-oxidation experiment provide both confidence for our experimental approach and a means of comparing mechanistic insights of CPA and PA, two structurally related compounds.

3.3 Online PILS-MS for formation and evolution of oxidation products

The evolution of PA OH-oxidation followed by the formation of its oxidation products can be observed in Fig. 4. The UV
Lamps were turned on to initiate OH-oxidation at time 0, which is indicated by a vertical dashed line. To avoid signal saturation, the signal of PA with a naturally existing $^{13}\text{C}$ isotope ($m/z$ 186 instead of $m/z$ 185) was monitored and plotted. Immediately following the onset of OH-oxidation, the PA signal decayed rapidly, and numerous product signals appeared and subsequently reacted away. The online PILS-MS method was applied to monitor some of the major products detected using offline (−)ESI-LC-MS (Table 2). At the beginning of the experiment, a background signal was obtained by sampling the solution containing PA and H$_2$O$_2$ but without light. The background signal for the PA solution shows that some products were already present before the start of the experiment, which is likely attributed to reaction during the storage time. Control experiments (Section 2.2) showed that PA does not react with H$_2$O$_2$ under dark conditions, therefore, the products observed are not due to rapid oxidation as H$_2$O$_2$ is added to PA solution. Furthermore, these impurities are present in trace amounts, as their signal is smaller than that of the PA $^{13}\text{C}$ isotope (1.1% abundance). The $^1\text{H}$ NMR technique used for PA standardization did not detect any impurities, except for formic acid. We are confident that impurities at these concentrations have minimal impact on our mechanistic analysis.

### 3.4 MS$^2$ for further structural confirmation

The structural information of selected PA OH-oxidation products was confirmed using the fragmentation patterns from the triple-quadrupole MS instrument. Fig. 5 shows the fragmentation results for three major oxidation products with $m/z$ 171, $m/z$ 203, and $m/z$ 217. The fragmentation spectra of a few other compounds monitored with MS$^2$ are available in the ESI Fig. S2†. The MS$^2$ spectra were collected by direct infusion of the reaction solution without separation; therefore, the fragmentation patterns represent those of mixed isobaric compounds.

The compound detected as $m/z$ 171 ion produced fragment ions of $m/z$ 153 (−18 au) and $m/z$ 83 (−88 au), indicating the loss of H$_2$O and the presence of up to two carboxylic acids, respectively. The ion detected at $m/z$ 171 was identified as the major PA oxidation product. Previous studies have shown that both gas-phase ozonolysis and aqueous-phase OH-oxidation of α-pinene SOA leads to the formation of compounds of $m/z$ 171.24,41 These compounds have been assigned to be terpenylic

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**Table 2** List of major PA OH-oxidation peaks detected by (−)ESI-LC-MS

| MW (g mol$^{-1}$) | [M−H]$^-$ | Elemental composition | Identity$^a$ | Ref.$^b$ |
|------------------|------------|-----------------------|--------------|---------|
| 144              | 143        | C$_{8}$H$_{16}$O$_{3}$ |              |         |
| 156              | 155        | C$_{8}$H$_{16}$O$_{3}$ |              |         |
| 158              | 157        | C$_{8}$H$_{16}$O$_{3}$ |              |         |
| 172              | 171        | C$_{8}$H$_{16}$O$_{3}$ | Terpenylic acid or norpinic acid (NPA) | 40 |
| 188              | 187        | C$_{8}$H$_{16}$O$_{4}$ | Hydroxynorpinic acid | 41 |
| 190              | 189        | C$_{8}$H$_{16}$O$_{4}$ |              |         |
| 200              | 199        | C$_{8}$H$_{16}$O$_{4}$ |              |         |
| 201              | 201        | C$_{8}$H$_{16}$O$_{4}$ |              |         |
| 204              | 203        | C$_{8}$H$_{16}$O$_{4}$ | 3-Methyl-1,2,3-butanetricarboxylic acid (MBTCA) and/or isomers | 42 |
| 216              | 215        | C$_{8}$H$_{16}$O$_{4}$ |              |         |
| 218              | 217        | C$_{8}$H$_{16}$O$_{4}$ |              |         |

$^a$ If missing, the compound has not been identified previously. $^b$ References.

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**Fig. 4** Temporal evolution of PA (monitored as a $^{13}\text{C}$ isomer) and its major products during OH-oxidation, monitored using the online PILS-ESI-MS technique. The vertical dashed line indicates the onset of OH-oxidation.

**Fig. 5** Fragmentation spectra of (a) NPA ($m/z$ 171), (b) MBTCA ($m/z$ 203), and (c) $m/z$ 217 under uncontrolled pH conditions.
acid (a mono-acid) and/or NPA (a diacid). We propose that the peak at m/z 171 in our study is mainly attributed to NPA, based on the observed loss of 88 au, which is indicative of the presence of two carboxylic groups. The formation mechanisms for NPA are shown in Sections 3.6.1 and 3.6.2.

For the compound with m/z 203, fragment ions include m/z 185 (−18 au; neutral loss of H₂O), m/z 173 (−44 au; neutral loss of CO₂), m/z 141 (−62 au; neutral loss of CO₂ and H₂O), and m/z 97 (−106 au; neutral loss of H₂O and two CO₂). Although small, the presence of a product ion of m/z 71 represents a combined loss of three CO₂, which implies the presence of three carboxyl groups in its structure. The fragmentation pattern of the product detected as m/z 203 was similar to that of MBTCA and its isomers.Offline analysis with (−)ESI-LC-MS detected a dominant isomer of m/z 203 at Rᵣ = 0.67 min and a few minor ones. To confirm that the dominant isomer is attributable to MBTCA, we compare the extracted ion chromatogram (EIC) of m/z 203 to that of TCA, which exhibits a peak at Rᵣ = 0.59 min (Fig. 6). Given that MBTCA and TCA are structurally similar triacids (Fig. S3†), we are confident that the major isomer of m/z 203 at Rᵣ = 0.67 min is MBTCA. Although the formation of MBTCA from PA has never been reported before, its formation from OH-oxidation of CPA has been previously confirmed. For comparison, we have also added the EIC of m/z 203 from CPA OH-oxidation to Fig. 6. Interestingly, as opposed to PA, our results show that m/z 203 arising from CPA is attributed to multiple isomers, and the contribution of MBTCA (peak at Rᵣ = 0.63 min) may not be significant.

The product detected as m/z 217 is also a major product of PA OH-oxidation, and it produced fragmentation ions at m/z 199 (−18 au; neutral loss of H₂O), m/z 183 (−34 au, neutral loss of H₂O₂), m/z 173 (−44 au; neutral loss of CO₂), m/z 155 (neutral loss of CO₂ and H₂O), m/z 129 (−88 au, presence of up to two carboxylic acids). The loss of 34 au is a fragmentation pattern unique to the products detected at m/z 217. Previous studies have attributed this pattern to the presence of a hydroperoxy group (ROOH); as such, m/z 217 is most likely to be a pinonic hydroperoxide, forming as a first-generation product of PA as we propose in Section 3.6.1. This result is in agreement with CPA's oxidation product of m/z 215, which is a 2 au difference from PA's product. Witkowski et al. proposed that m/z 215 is a pinonic hydroperoxide. To confirm that m/z 217 from PA photo-oxidation and m/z 215 from CPA photo-oxidation are indeed organic peroxides, we performed iodometry-assisted LC-MS. This technique takes advantage of the selectivity of iodometry towards organic peroxides and serves as a powerful method for qualitative confirmation of peroxides. We applied the same technique described by Zhao et al. on a PA OH-oxidation sample collected at 5 min of the reaction progress (Section 2.2) to ensure that m/z 217 has not been further oxidized. Fig. 7 shows a comparison of the product detected as m/z 217 and its isomers when treated with and without iodometry. We note that all peaks were depleted by iodometry, which is an indication that the products with m/z 217 appear to be peroxides (Fig. 7(a)). A very similar observation was made for m/z 215 from CPA photo-oxidation (Fig. 7(b)), indicating that some of its isomers are organic peroxides that form via a mechanism equivalent to that of m/z 217 in PA photo-oxidation.

3.5 NPA and MBTCA yield at pH 2 and pH 8

As shown in previous sections, the photo-oxidation of PA leads to the formation of many products; among which NPA and MBTCA have been of interest as known tracers of α-pinene SOA, and involved in the formation of SOA. The EICs of NPA and MBTCA formed via PA OH-oxidation at pH 2 and 8 are shown in Fig. 8(a) and (b), respectively. The most noteworthy observation is a significant difference in the signal intensities for NPA and MBTCA, with a greater amount detected under the basic conditions. Therefore, it is evident that the pH of the aqueous phase affects PA's reaction mechanism.
To provide a quantitative comparison between pH 2 and 8, we determine the aqueous-phase molar yield \( Y_{aq} \) based on the ratio of concentration of the product formed \([\text{Product}]\) to change in PA concentration \( \Delta [\text{PA}] \) over reaction time \( t \), as expressed in eqn (1):

\[
Y_{aq} = \frac{[\text{Product}]}{\Delta [\text{PA}]} = \frac{[\text{Product}_{t}]}{([\text{PA}]_{t} - [\text{PA}_{0}])}
\]

\([\text{PA}_{0}]\) represents the concentration of PA remaining, while \([\text{Product}_{0}]\) represents the concentration of either NPA or MBTCA formed at time \( t \). It is important to note that \( Y_{aq} \) varies as a function of time, as PA is depleted in time, and the products also undergo further OH-oxidation. In this work, we found that the \( Y_{aq} \) values were relatively constant until 60% of PA was consumed; therefore, we have decided to report \( Y_{aq} \) by averaging those from all the samples with at least 40% of PA remaining.

There are a few advantages in this method. First, this amount of decay ensures an accurate determination of \( \Delta [\text{PA}] \). Second, multi-generational OH-oxidation significantly complicates the product distribution, and focusing on the onset of the experiment allows more confident identification and quantification of NPA and MBTCA. Third, NPA and MBTCA have not undergone extensive photo-oxidation themselves by this time; therefore, \( Y_{aq} \) obtained this way likely represents the maximum yield of NPA and MBTCA.

As there are no commercial standards available for NPA and MBTCA, these two products were quantified assuming their ionization efficiencies are similar to those of PA and TCA, respectively. These surrogates were chosen due to their structural similarities to NPA and MBTCA (Fig. S3†). Note that TCA has been previously used as a surrogate to study reaction mechanisms of MBTCA. In this work, a PA and TCA calibration curve was prepared and analysed using (−)ESI-LC-MS as described in Section 2.3.1. The signal of MBTCA was related to that of TCA for quantification, while the signal of NPA was related to that of PA.

The \( Y_{aq} \) of NPA and MBTCA calculated with this method are summarized in Table 3. The mean values were obtained from the triplicates, while the uncertainty range indicates 1σ after propagating uncertainties related to all the parameters in eqn (1). The \( Y_{aq} \) values of both compounds are higher by more than a factor of 2 under pH 8. Such an impact of pH on the photochemical products of OAs has been suspected but never been reported in the literature. In Section 3.6.2, we propose a mechanistic explanation for this observation.

For comparison, we have also calculated MBTCA’s \( Y_{aq} \) arising from CPA OH-oxidation. Although a peak attributable to MBTCA has been observed from CPA OH-oxidation, as shown in Fig. 8(c), this peak was below the limit of detection of our (−)ESI-LC-MS technique. As discussed in Section 3.4, numerous isomers of \( m/z \) 203 are detected, and we consider the peak at \( R_{t} = 0.63 \) min to be MBTCA. We focus on \( Y_{aq} \) at \( t = 5 \) min, at which 40% of CPA remains. For this calculation, the contribution of the direct photolysis of CPA must be considered. CPA contains a carbonyl group that absorbs the UVB light equipped in our photoreactor. Judging from our previous study, in which the same lamp setting was employed, 30% of CPA decay was due to direct photolysis, while the remaining 30% was due to OH-oxidation. In this work, we assumed that MBTCA from CPA is solely from the 30% of CPA reacted via OH-oxidation. Based on the detection limit of MBTCA (0.21 \( \mu M \)), the upper-limit estimate for MBTCA’s \( Y_{aq} \) from CPA is 0.1%. MBTCA’s \( Y_{aq} \) from CPA was found to be significantly smaller than its \( Y_{aq} \) from PA. While \( Y_{aq} \) of MBTCA has never been reported in the literature, the gas-phase yield is described to be <1%. The \( Y_{aq} \) determined by our study is low due to our finding that \( m/z \) 203 from CPA is attributable to numerous isomers, and the relative contribution of MBTCA is minor. Overall, our results show that MBTCA’s \( Y_{aq} \) from PA OH-oxidation is much greater than that of CPA, which is the currently known precursor of MBTCA.

3.6 Proposed reaction mechanisms of PA OH-oxidation

3.6.1 General reaction mechanisms. Similar to the aqueous-phase OH-oxidation of other precursors, PA gives rise to numerous reaction products through multi-generational oxidation steps. In this section, we describe the general formation mechanism of early-generation products, with a focus placed on the three species discussed in Section 3.4, as elucidation of an explicit mechanism of PA is infeasible. Fig. 9 shows the temporal profiles of four relevant products with \( m/z \) of 155, 187, 171, and 203, with their signals normalized to their maximum values to emphasize their temporal evolutions.

As already discussed in Section 3.4, the major products of \( m/z \) of 171 and \( m/z \) 203 are NPA and MBTCA, respectively. By observing the formation and decay profile of each compound, we propose a series of mechanisms illustrated in Fig. 10. As the initial step of OH-oxidation, H-abstraction by OH radicals can occur at multiple positions on a PA molecule (1–6). For any of these positions, the resulting peroxy radical \( (\text{RO}_2) \) can react with a hydroperoxy radical \( (\text{HO}_2) \), giving rise to an organic hydroperoxide \( (\text{ROOH}) \) with \( m/z \) 217. Fig. 10 shows the case in which H-abstraction occurs at position C-3. H-abstraction at different positions gives rise to various peroxy radicals, which can undergo hydroperoxy radical (ROOH) addition and H-abstraction again.
positions of PA results in different structural isomers of this compound, likely explaining the presence of multiple peaks on its EIC (Fig. 7(a)). Fig. 10 further illustrates the cases in which the initiation step occurs at the 1 and 4 positions, followed by O2 addition and formation of RO2. Reactions of NO are not expected to be significant in atmospheric aqueous phases due to the low solubility of NO.

The peroxy radicals formed mainly react with another RO2 in solution or with an HO2, forming the corresponding alkoxy radical (RO).

The ROs further undergo β-scission to form a variety of possible products, including an aldehyde and a ketone of m/z 155. From Table 2, the accurate mass indicated the elemental composition of C8H12O3, in agreement with those proposed in Fig. 10. We considered that the ion at m/z 155 is an important intermediate product leading to the formation of NPA (m/z 171), and MBTCA (m/z 203).

The m/z 155 intermediates can undergo multiple reaction pathways, and we herein focus on those leading to NPA and MBTCA. In an aqueous solution, aldehydes can be hydrated, forming their geminal diols. Thus, for the aldehyde product of m/z 155 (right), the OH radicals can abstract a hydrogen atom from both aldehydic and geminal diol forms. The respective alkyl radicals formed are also in hydration equilibrium. This geminal–diol pathway is unique to the atmospheric aqueous phase and it does not occur via gas-phase oxidation. The addition of dissolved O2 to the alkyl radicals formed leads to two types of peroxy radicals. A peroxy radical (not shown here), which dissociates rapidly to form NPA. In parallel, the acylperoxy radical shown in Fig. 10 can either hydrate or react with a HO2 to form NPA. The observation that NPA is a major product of PA explains a previous finding by Amorim et al. The OA with m/z 171 exhibited a completely different decay profile during OH-oxidation of α-pinene SOA, indicating that it was forming from other species. The results obtained here can, therefore, confirm that the OA at m/z 171 was likely NPA and was formed from PA present in the SOA extract.

In the case of the ketone product of m/z 155 (left), the OH radical can abstract an H-atom from different sites of the molecule. The mechanism shown in Fig. 10 focuses on the H-abstraction from C-3, followed by the addition of dissolved O2 to form the corresponding RO2. The reaction with another RO2 produces an alkoxy radical that proceeds via β-scission to break the four-membered cyclic structure. The resulting peroxy radical can react with HO2 to form the corresponding carboxylic acid with m/z 187. The elemental composition and fragmentation pattern of the m/z 187 product was confirmed by the results from other species. The results obtained here can, therefore, confirm that the OA at m/z 171 was likely NPA and was formed from PA present in the SOA extract.

Table 3  Calculated aqueous-phase photo-oxidation molar yield of NPA and MBTCA

| Precursor | Yaq NPA (pH 2) | Yaq NPA (pH 8) | Yaq MBTCA (pH 2) | Yaq MBTCA (pH 8) |
|-----------|---------------|---------------|-----------------|-----------------|
| PA        | (3.4 ± 1.2)%  | (10 ± 1.2)%   | (2.6 ± 0.9)%    | (5.0 ± 0.5)%    |
| CPA       | N/A           | N/A           | <0.1%           | <0.1%           |

* Signal was below the limit of detection. The value represents an upper-limit estimation.

Fig. 9  Temporal evolutions of compounds related to the formation of NPA and MBTCA: PA, m/z 155, NPA, m/z 187 and MBTCA. Signals are normalized to their maximum values.

Fig. 10  Proposed mechanism of NPA, MBTCA, and one of the m/z 217 isomers formed from PA OH-oxidation at pH 2. Note that multiple isomers of m/z 217 can form, and the structure shown here is an example.
presented in Sections 3.3 and 3.4. As shown in Fig. 9, the m/z 187 compound appears earlier than MBTCA, which is consistent with it being an intermediate for MBTCA formation. Additionally, the mechanism shown in Fig. 10 follows Enami and Sakamato’s work on MBTCA formation from CPA oxidation, in which the authors also identified m/z 187 as a byproduct of MBTCA. As the last step, the aldehyde functional group of the m/z 187 can be converted into a carboxyl group following the same mechanism discussed in the previous paragraph. This reaction pathway agrees with the mechanism proposed by Witkowski et al. and Enami and Sakamato for MBTCA formation from the aqueous phase oxidation of CPA.

3.6.2 pH-dependent reaction mechanisms. The mechanisms presented thus far are pH-independent, as the carboxylic groups are not involved. In this section, we propose a series of pH-dependent mechanisms that can likely explain the different yields of NPA and MBTCA observed at pH 2 and 8 (Section 3.5). At pH 8, in addition to the H-abstraction reactions occurring over the PA structure, a unique charge-transfer reaction occurs in the aqueous phase, as discussed in Introduction (Fig. 1). At pH 8, PAN undergoes an electron-transfer reaction with OH radicals at two different sites of the molecule, labeled as A and B in Fig. 11. The electron-transfer reaction taking place at site A is followed by a decarboxylation step, which is the key step in this part of the mechanism. A series of radical reactions, previously discussed in Section 3.6.1, take place, resulting in the formation of the deprotonated form of the intermediate compounds of m/z 171 and m/z 187. Here, the compound with m/z 187 is equivalent to the one in Fig. 10. The reactions leading to MBTCA formation have already been discussed in Section 3.6.1. Alternatively, when the charge transfer reaction occurs to site B on a PA molecule, an intermediate with m/z 155 can be formed as a result of the decarboxylation. This compound is equivalent to the aldehydeic of m/z 155 (Fig. 10, right), and the subsequent OH oxidation of this compound gives rise to NPA.

Overall, we propose that the charge transfer reaction, followed by initial decarboxylation of PA and formation of the corresponding alkyl radicals (denoted with * in Fig. 11) are the key steps that serve as additional formation routes for NPA and MBTCA. The mechanisms proposed in Fig. 11 are unique to the aqueous-phase OH-oxidation of PA. In comparison, the molecular structure of CPA differs from PA at site A, in which CPA possesses a ketone functional group, which does not allow the charge transfer reaction. The aqueous-phase formation of MBTCA from PA is, however, possible due to the additional carboxyl group under basic pH conditions. These results agree with the higher yields of MBTCA under pH 8 discussed in Section 3.5. Therefore, we have arrived at the following conclusion. Although reactions occurring at pH 8 lead to similar compounds, as in the case of pH 2, the charge transfer reaction serves as additional formation pathways and thus accelerates the formation of several key oxidation products, including NPA and MBTCA.

4. Conclusions

We have, for the first time, investigated the mechanism of aqueous-phase PA OH-oxidation in detail. Using both offline high mass resolution (−)ESI-LC-MS and a novel online monitoring system (PILS-ESI-MS), we detected a large number of oxidation products. Among them, NPA (m/z 171) and MBTCA (m/z 203) were well-established α-pinene oxidation products, and yet have not been identified as products of PA oxidation. Their identities were confirmed by observing their fragmentation patterns using MS/MS. Organic hydroperoxides with m/z 217 were also detected as major and first-generation products. An iodometry-assisted method was used to confirm their peroxide nature.

Combining observations obtained using (−)ESI-LC-MS and PILS-ESI-MS, we proposed the possible reaction mechanisms for the targeted compounds. The compounds of m/z 155 and m/z 187 serve as intermediates for the products assigned to NPA (m/z 171), and MBTCA (m/z 203). The detection of MBTCA from PA is new, as the current understanding is that MBTCA arises only from CPA. We estimate that the molar yield of MBTCA from PA is at least an order of magnitude larger than that from CPA. We also observed a difference in the yield of NPA and MBTCA under pH 2 and pH 8, with the yield under basic conditions doubling that under acidic conditions. We propose that it is the unique charge transfer reaction occurring only in the aqueous phase that leads to the enhancement of MBTCA and NPA formation. Such an impact of pH on the photochemical products of OAs has never been reported in the literature.

More generally, our results present significant implications to atmospheric chemistry. Measured pH of cloud and fog water fluctuates significantly between 2 and 7, depending on the location and chemical composition of these hydrometeors. Over the past 40 years, the average pH of fog and cloudwater in Europe and the US has increased by two units. This trend...
arises from reduced emissions of acid precursors to the atmosphere. The pH dependence of cloudwater chemistry must be understood to predict the impact of a constantly changing acidity in atmospheric aqueous phases. From Amorim et al. and this work, a complex interplay between acid–base chemistry and photochemistry has become clear. The charge transfer reaction does not significantly change the overall OH reactivity of PA. However, the underlying reaction mechanisms can be altered, changing the yield of reaction products. In the case of PA oxidation, MBTCA is known to contribute significantly to SOA mass due to its low volatility. The high yield of MBTCA from PA, particularly under basic conditions, indicates that aqSOA can increase when the acidity in atmospheric aqueous phases is constantly reduced. Although our study has focused on a single compound, PA, we believe that it serves as a vivid demonstration of how pH can affect the reaction mechanisms of many other atmospherically relevant OAs under cloudwater-relevant conditions. Our results suggest the importance of further understanding the impact of pH on aqueous-phase photochemistry, particularly acids and bases that are sensitive to the highly variable pH of atmospheric aqueous phases.

Conflicts of interest

There is no conflict to declare.

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