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Mechanism of the hydroxyl radical oxidation of methacryloyl peroxynitrate (MPAN) and its pathway toward secondary organic aerosol formation in the atmosphere†

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Methacryloyl peroxynitrate (MPAN), the acyl peroxynitrate of methacrolein, has been suggested to be an important secondary organic aerosol (SOA) precursor from isoprene oxidation. Yet, the mechanism by which MPAN produces SOA through reaction with the hydroxyl radical (OH) is unclear. We systematically evaluate three proposed mechanisms in controlled chamber experiments and provide the first experimental support for the theoretically-predicted lactone formation pathway from the MPAN + OH reaction, producing hydroxymethyl-methyl- α -lactone (HMML). The decomposition of the MPAN-OH adduct yields HMML + NO₃ (\sim 75%) and hydroxyacetone + CO + NO₃ (\sim 25%), out-competing its reaction with atmospheric oxygen. The production of other proposed SOA precursors, e.g., methacrylic acid epoxide (MAE), from MPAN and methacrolein are negligible (<2%). Furthermore, we show that the beta-alkenyl moiety of MPAN is critical for lactone formation. Alkyl radicals formed cold via H-abstraction by OH do not decompose to HMML, even if they are structurally identical to the MPAN-OH adduct. The SOA formation from HMML, from polyaddition of the lactone to organic compounds at the particle interface or in the condensed phase, is close to unity under dry conditions. However, the SOA yield is sensitive to particle liquid water and solvated ions. In hydrated inorganic particles, HMML reacts primarily with H₂O to produce the monomeric 2-methylglyceric acid (2MGA) or with aqueous sulfate and nitrate to produce the associated organosulfate and organonitrate, respectively. 2MGA, a tracer for isoprene SOA, is semivolatile and its accommodation in aerosol water decreases with decreasing pH. Conditions that enhance the production of neutral 2MGA suppress SOA mass from the HMML channel. Considering the liquid water content and pH ranges of ambient particles, 2MGA will exist largely as a gaseous compound in some parts of the atmosphere.

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Introduction

Anthropogenic emissions of nitrogen oxides ($NO_x = NO + NO_2$) strongly influence the oxidative pathways of gaseous hydrocarbons, from both natural and anthropogenic origins. In particular, the

oxidation of isoprene (C_5H_8), arguably the most important nonmethane hydrocarbon emitted to the atmosphere, has a significant sensitivity to the NO_x conditions under which it is oxidized. This NO_x sensitivity translates to regional environmental impacts. For example, the high- NO_x oxidation of isoprene controls the production of tropospheric ozone in regions rich with biogenic hydrocarbons^{1–3} by accelerating the cycling of NO_x and producing NO_x reservoir species (*e.g.*, organic nitrates and peroxynitrates). Isoprene chemistry also affects global climate primarily through the formation of secondary organic aerosol (SOA).^{4–7} However, the lingering uncertainties of SOA formation in various anthropogenically-impacted, but biogenically-influenced, systems preclude accurate simulations of human-induced climate and air quality feedbacks in atmospheric chemical transport models.

A common approach to identify the source of SOA is through tracer compounds, *e.g.*, 2-methylglyceric acid (2MGA)⁹⁻¹¹ in isoprene-derived ambient aerosols. 2MGA and its oligomers

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Paper

Fig. 1 The formation of acyl peroxynitrates (APNs) from the OH-initiated oxidation of aldehydes is favoured under high NO_2 to NO conditions. For methacrolein, the abstraction of the aldehydic H occurs roughly half of the time⁸ in the OH oxidation reaction at room temperature.

are thought to be enhanced under high-NO_x conditions^{12–14} and, furthermore, can be uniquely traced to the oxidation of methacrolein, ^{14,15} a first-generation isoprene oxidation product. Chan *et al.*¹³ showed that high-NO₂ chemistry (as opposed to "high-NO_x" where most of the NO_x can be represented by NO) favours the production of SOA *via* the acyl peroxynitrate channel (APN, Fig. 1). Essentially all of the SOA generated in the laboratory studies of the isoprene high-NO_x route was shown to be derived from methacrolein photochemistry, suggesting a major role of the APN from methacrolein, methacryloyl peroxynitrate (MPAN). The suggestion by Chan *et al.* was confirmed by the synthesis of MPAN and the measurement of its SOA formation through photochemistry (tested under high-NO conditions).¹⁶

Although the potential of MPAN + OH chemistry to produce SOA is relatively well established, the chemical mechanism leading to SOA from MPAN photooxidation has been subject of debate. It is also not clear whether the MPAN photooxidation is sensitive to NO_x , although past studies have all been performed in the presence of NO. Fig. 2 shows three possible pathways leading to SOA production that have been proposed by independent works. 13,17,18

The initial step of the MPAN photooxidation is OH addition to the double bond, primarily generating the energetically hot tertiary alkyl radical of MPAN (A*, Fig. 2). Chan *et al.*, ¹³ in a series of photochemical chamber experiments, proposed that the main fate of A* is collisional stabilization (producing A)

followed by reaction with molecular oxygen to form an alkylperoxyl radical (B), where the beta-peroxyl moiety attacks at the carbonyl carbon to form a di-oxoketone (DOK) and NO_2 as a co-product (Fig. 2, mechanism 2). The suggestion by Chan *et al.* was derived by performing methacrolein high- NO_2 photooxidation experiments at low O_2 content (\sim 2%). The authors did not observe an increase in SOA yield and concluded that either O_2 addition is required for SOA formation or is non-competitive at those levels.

From a combined suite of density functional theory and coupled cluster calculations, Kjaergaard *et al.*¹⁷ proposed a rapid ring closure from an acylperoxyl oxygen of A* to form a 3-member lactone, hydroxymethyl-methyl- α -lactone (HMML), and NO₃ as a co-product (Fig. 2, mechanism 3). Furthermore, the authors hypothesize that a significant fraction of the HMML product (36%) is generated with sufficient energy to further decompose to hydroxyacetone (HAC) and CO. Lin *et al.*¹⁸ also identified the formation of HMML *via* density functional theory calculations; however, they hypothesized that HMML may be too unstable to form in the atmosphere.

Lin *et al.*¹⁸ proposed a rapid H-migration induced ring closure at the hydroxyl oxygen, *via* a 6-member intermediate, to an epoxide product called methacrylic acid epoxide (MAE, Fig. 2, mechanism 1). This suggestion was formulated by comparing the composition of SOA (*e.g.*, 2MGA and other products) from the photooxidation of methacrolein and the reactive uptake of laboratory-synthesized MAE onto highly-acidic particles, as well as through quantum chemistry calculations. The mechanism of the uptake was suggested to be nucleophilic ring opening of the epoxide in an analogous reaction to the isoprene epoxydiols.¹⁹

Notable differences between the proposed pathways include: (1) the fate of A^* – whether cyclization/decomposition occurs more rapidly than thermalization and, thus, bimolecular reaction with O_2 , (2) the stability of the products – MAE is the only product stable enough to be detected by current analytical instrumentation, and (3) the subsequent SOA formation mechanism under low humidity conditions – MAE requires highly-acidic seed aerosols to open the epoxide ring, while the

Fig. 2 The three proposed pathways to the formation of 2-methylglyceric acid (2MGA), a tracer for isoprene-derived ambient SOA. Mechanisms 1–3 and acronyms are discussed in the text. IUPAC names of key organic compounds discussed in this work are shown in Table S1 (ESI†).

PCCP Paper

ring opening of the more unstable compounds, DOK and HMML, would likely require only collisions with a surface.

The uncertainties in the MPAN + OH mechanism persist due to ambiguities from previous work. For example, earlier studies of the MPAN + OH system^{20,21} did not measure SOA formation and were conducted with high mixing ratios of NO_r (blind to the formation of nitrogen products). More-recent laboratory studies of Chan et al. 13 and Lin et al. 18 were performed with methacrolein, such that products from MPAN oxidation and from the OH addition route of methacrolein ($\sim 55\%$ probability) were formed simultaneously, complicating the analysis.8 The theoretical study of Kjaergaard et al. 17 has yet to be experimentally validated. In this work, we elucidate the chemical mechanism governing the MPAN photooxidation (and the photooxidation of its chemical analogues) through a series of targeted chamber experiments using laboratory-synthesized MPAN under low humidity and low-NO conditions. We further study the chemical pathway with which the MPAN produces SOA under simulated ambient conditions with higher relative humidity and seed particle concentrations.

Experimental

Chamber studies

The atmospheric chamber facility used for this work has been described in detail.²² A portion of the present work, namely, control experiments with α-pinene and exploratory experiments with methacrolein, were performed as part of the Focused Isoprene eXperiment at the California Institute of Technology (FIXCIT) chamber campaign.²² Briefly, experiments were conducted in a large (24 m³) teflon chamber using purified dry air (<5% relative humidity (RH) at 298 K) such that initial NO mixing ratios were lower than 100 pptv (measured by G. S. Tyndall, D. D. Montzka, and A. J. Weinheimer at FIXCIT) and initial particle mass concentrations were much lower than 0.01 µg m⁻³ (measured by

SMPS, see following text). For the experiments that were performed under humid conditions, water vapour was added to the chamber using a Nafion membrane humidifier (Perma Pure LLC) and recirculating ultrapure water (18 M Ω , Millipore Milli-Q). Particles were injected (mean particle diameter ~ 70 nm) by atomizing dilute solutions of ammonium sulfate (0.1 M) through a heated wet-wall denuder to deliquesce the particles prior to entering the chamber held at RH 40% or 85%. Corrections for the wall deposition of particles, using ammonium sulfate seed aerosols, were derived from control experiments performed at several water vapour mixing ratios in the chamber. Volatile organics, excluding MPAN, were injected by microliter syringe into a clean glass bulb and quantitatively transferred with dry nitrogen gas into the chamber through a short section of tubing, optionally with gentle heating for the larger hydrocarbons such as α -pinene (<80 °C). The temperature in the chamber enclosure was adjusted to 15-25 °C for experiments as needed. MPAN experiments were performed at 15 °C to minimize the thermal decomposition of MPAN (thermal lifetime ~ 26 h at 15 °C, compared to ~ 5 h at 25 °C, for an NO₂/NO ratio of 10 that is representative of areas outside of urban centres).23

Table 1 lists the laboratory conditions for the main experiments in this work, e.g., the gas-phase oxidation of MPAN (synthesized standard), methacrolein (MACR, Aldrich, 95%), isobutyraldehyde (ISOBUT, Aldrich, >99%), and 2-methyl-but-3-ene-1-ol (231MBO, Aldrich, 98%). Photochemistry under low-NO conditions (HO₂ > NO) was initiated by the near-UV (300–400 nm, $\lambda_{\rm max} \sim 350$ nm) photolysis of gas-phase hydrogen peroxide that has been evaporated into the chamber (Aldrich, 50 wt% in water):

$$H_2O_2 + h\nu \rightarrow OH + OH$$

$$\mathrm{OH} + \mathrm{H_2O_2} \, \rightarrow \, \mathrm{HO_2} + \mathrm{H_2O}$$

Photochemistry under high NO₂ (and typically lower in NO) conditions were initiated by the photolysis of gas-phase methyl

Table 1 Experimental conditions for the laboratory photooxidation of hydrocarbons (HC) used in this work: MPAN, methacrolein (MACR), isobutyraldehyde (ISOBUT) and 2-methyl-but-3-ene-1-ol (231MBO). The light intensity was 100% ($J_{NO_3} \sim 7 \times 10^{-3} \, s^{-1}$) for hydrogen peroxide (HP) experiments and 10-20% for methyl nitrite (MN) experiments. Steady state OH is estimated from hydrocarbon decay (GC) data. α -pinene (α -Pin) and methacrylic acid epoxide (MAE) were added in select experiments to capture NO₃ and observe SOA formation, respectively. Control experiments of HP + $h\nu$, tridecane + $HP + h\nu$, and wall losses (described in Experimental section) are not included in table. Experiments were performed at various temperature and relative humidity (RH). Select experiments were performed with \sim 80 μg m⁻³ of ammonium sulfate (AS) seed particles

Exp.	HC precursor		Oxidant								
#	Name	[HC] ₀ (ppb)	Source	[Ox] ₀ (ppb)	[OH]ss (cm ⁻³)	Rxn time (h)	Temp. (°C)	RH (%)	[NO] ₀ (ppb)	$[NO_2]_0$ (ppb)	Additional injections
1	MPAN	30	$H_2O_2 + h\nu$	2000	1.5×10^6	2.5	15	< 5	< 0.1	1	α-Pin (50 ppb)
2	MPAN	13	$H_2O_2 + h\nu$	2000	1.5×10^6	5	15	< 5	< 0.1	1	_
3	MPAN	23	$H_2O_2 + h\nu$	2000	$1.5 imes 10^6$	4.5	15	< 5	< 0.1	1	_
4	MPAN	30	$H_2O_2 + h\nu$	2000	1.5×10^{6}	4.5	15	< 5	< 0.1	1	α-Pin (30 ppb) + MAE (15 ppb)
5	MPAN	17	$H_2O_2 + h\nu$	2000	$1.5 imes 10^6$	3.2	15	< 5	< 0.1	1	α-Pin (30 ppb) + MAE (15 ppb)
6	MACR	100	$H_2O_2 + h\nu$	2000	1.5×10^{6}	5	15	< 5	10	230	_ ` ` ` ` ` ` ` ` ` ` ` ` ` ` ` ` ` ` `
7	MACR	50	$MN + h\nu$	140	2×10^7	2.5	25	< 5	< 0.1	50	_
8	MACR	80	$MN + h\nu$	130	1×10^7	2.5	15	< 5	< 0.1	110	_
9	MACR	100	$MN + h\nu$	200	2×10^7	2.5	25	< 5	10	100	_
10	ISOBUT	80	$MN + h\nu$	220	2×10^7	5.5	25	< 5	10	50	_
11	231MBO	200	O_3 , then	$\sim 600 [O_3],$	2×10^7	24	25	< 5	10	100	_
			$MN + h\nu$	200 [MN]							
12	MACR	200	$MN + h\nu$	200	2×10^7	3.5	25	40	10	100	AS seeds
13	MACR	100	$MN + h\nu$	200	2×10^7	2.5	25	85	10	100	AS seeds

Paper

nitrite (synthesized standard, stored in liquid N2) in the presence of various mixing ratios of additional NO2 (standard

$$CH_3ONO + h\nu + O_2 \rightarrow HO_2 + NO + HCHO$$

$$HO_2 + NO \rightarrow OH + NO_2$$

Methyl nitrite was introduced into the chamber by filling a clean, evacuated, 500 mL glass bulb with the desired pressure of the standard, backfilling with N2, and transferring the contents of the bulb into the chamber with a stream of N2. In high-NO₂ experiments, the added NO₂ and that formed from CH₃ONO photochemistry conspire to maintain the NO₂/NO ratio > 10 throughout the experiment. Importantly, the near-UV broadband radiation used in this work does not efficiently photolyze NO₃ via mechanisms that yield a net destruction of NO₃ ($J_{\mathrm{NO_2}} \sim 5.5 \times$ $10^{-4} \text{ s}^{-1}, J_{\text{NO}_3} \rightarrow {}_{\text{NO+O}_2} \sim 4.5 \times 10^{-6} \text{ s}^{-1}).^{24}$

Analytical measurements

MACR and isobutyraldehyde were quantified with a commercial gas chromatograph with a flame-ionization detector (GC-FID, HP 6890N) and calibrated by volumetric injections of commercial standards. Particle size and number were measured by a scanning mobility particle sizer (SMPS), i.e., a custom-built differential mobility analyser coupled to a commercial condensation particle counter (TSI Inc.). Aerosol speciation was measured using a high resolution time-of-flight aerosol mass spectrometer (AMS, Aerodyne). 25 Bulk aerosol species (organic, sulfate, ammonium, nitrate) were calculated using AMS data analysis modules (Pika 1.14D). The instrument's ionization efficiency was calibrated with 350 nm ammonium nitrate particles.

Volatile acids and select polar organic compounds were quantified with a custom-built triple-quadrupole chemical ionization mass spectrometer (CIMS, Agilent/Caltech).26 The CIMS operated in three modes: scanning negative ion mode using CF_3O^- as the reagent (m/z 50–250), scanning positive ion mode using primarily H_3O^+ as the reagent ion (m/z 50–200), and tandem mass negative ion mode (monitoring select precursorproduct ion pairs). The ion chemistry (e.g., detection as $[M + F]^$ and [M + CF₃O]⁻ ions)²⁷ and tandem mass determinations²⁶ of the CF₃O⁻ CIMS have been described previously.

Sensitivities of the triple-quadrupole CF₃O⁻ CIMS used in this work were determined based on a calibrated time-offlight CF₃O⁻ CIMS instrument during the FIXCIT campaign.²² The absolute calibration for commercially-available and synthesized standards in the time-of-flight CIMS, using gravimetric, optical, and thermal-dissociation + laser-induced fluorescence methods are described elsewhere.²⁸ Uncertainty is estimated to be $\pm 30\%$. Synthesized standards of organic nitrates were found to have water-dependent sensitivities similar to HNO₃;²⁹ thus the CIMS sensitivity of the pinene nitrooxyhydroperoxide (PNP), for which there are no authentic standards, is assumed to be similar to HNO₃ (estimated uncertainty $\pm 50\%$). For other compounds discussed in this work, theoretical calculations^{30,31} were used to estimate the sensitivity (estimated uncertainty $\pm 50\%$).

Fig. 3 The experimental scavenging of NO₃ into stable products detectable by CIMS: PNP (one of several isomers shown) and HNO3.

The observed ion of methacrylic acid epoxide (MAE, synthesized standard, detected as [MAE·F]-)18 was isobaric with a water cluster $[(H_2O)_2 \cdot CF_3O]^-$ (m/z 121) in the single-ion analysis. Thus, MAE quantification is performed with collision-induced dissociation into its main precursor \rightarrow product ion $(m/z 121 \rightarrow$ m/z 101). When experiments are performed at higher relative humidity, the water signal $(m/z \ 121 \rightarrow m/z \ 103)$ may have an extensive tail that would lead to an over-estimation of MAE. The tandem signal that we measure must then be considered as an upper limit to the MAE concentration. The ion chemistry in the positive ion mode is similar to proton transfer reaction³² mass spectrometry (quantification mainly with $[M + H]^+$ ions in this work). Calibrations for select species in the positive ion mode (e.g., α -pinene and 231MBO) were performed immediately prior to experiments using commercial standards under dry conditions.

CIMS was also used to indirectly monitor nitrogen oxide chemistry (Fig. 3). For example, the reaction of NO₂ with HO₂ generates peroxynitric acid (HO2NO2, PNA), which was used as a sensitive tracer for NO2. α-pinene was used in a subset of experiments to capture NO3 as the pinene nitrooxy hydroperoxide (PNP), which is detectable by CIMS, under the condition where the HO₂/NO ratio is high (Fig. 3, left). α-pinene was chosen as an NO₃ scavenger because its reaction rate coefficient with OH is the lowest for a monoterpene that reacts rapidly with NO₃ (i.e., $k_{NO_3}/k_{OH} \sim 0.12$), ³³ and its reaction rate with NO₃ is at least 4 orders of magnitude larger than the MPAN reaction with NO₃.34 There are varying reports of SOA yield from the α -pinene + NO₃ reaction (0–16%), ^{35–37} possibly decreasing with higher RH.³⁷ Thus, experiments involving α-pinene were not included in the yield calculations from MPAN due to the possibility of interference from the α-pinene + OH or + NO₃ reaction. HNO₃, produced from H-abstraction chemistry of hydrocarbons (Fig. 3, right) when NO2 is low (and thus NO2 + OH is not a significant source of HNO₃), was also used as tracer for NO₃-initiated chemistry.

MPAN, peroxyacetyl nitrate (PAN), and NO2 are measured with a commercial NO2 and acyl peroxynitrate (NO2/APN) analyser (Fitz Aerometric Technologies). In the NO2/APN instrument, NO2, PAN, and MPAN were chromatographically separated, in that order, with a room-temperature deactivated DB-5 column and detected by monitoring chemiluminescence from their reactions with luminol (Fig. S1, ESI†). NO2 was calibrated with a diluted standard mixture (488 ppmv NO2 in N2, Scott Specialty Gases). The sensitivity of PAN in the instrument was inferred from its relationship to NO2.38 MPAN, in isolation, was calibrated using a commercial NOx analyser (Teledyne model T200) due to its quantitative decomposition to NO2 when

exposed to the heated (310 °C) molybdenum catalyst. A similar calibration was obtained by thermally decomposing MPAN to NO2 in a heated stainless steel tube before the NO2/APN analyser. Sampling MPAN from the chamber (before adding other compounds) through room-temperature teflon tubing leads to a small decomposition yield of NO₂ from MPAN (\sim 4%, Fig. S1, ESI†). The NO₂ signal was corrected in the NO₂/APN data as a fraction of the MPAN integrated peak.

MPAN

The synthesis of MPAN from the peroxidation of methacrylic anhydride (Aldrich, 94%) and nitration of the methacrylic peracid (MPAA) was performed as suggested by Bertman and Roberts³⁹ with the following revisions: we used 50 wt% H₂O₂ (Aldrich) instead of 30 wt% H₂O₂ and methanesulfonic acid (Aldrich, 99.5%) instead of conc. H₂SO₄. The crude MPAN mixture in water-tridecane was stored frozen and separated in small fractions with tridecane on a silica gel column deactivated by successive solvent washes of methanol, acetone, ethyl acetate, hexanes, and tridecane. Signs of known impurities were checked by CIMS (i.e., methacrylic acid and MPAA) and the NO₂/APN instrument (PAN). MPAN is thought to be explosive when pure⁴⁰ and no attempt was made to remove the solvent. The separated MPAN fraction in tridecane was stored at 0 °C or below and used promptly.

MPAN was injected by gently bubbling air through the tridecane-solvated mixture (submerged in an ice bath) placed inside the 15 °C chamber enclosure over the course of roughly 30 min to achieve 15-30 ppbv in the chamber. Tridecane was chosen as the storage solvent due to its low volatility, and thus, lower extent of co-evaporation into the chamber. To correct for any SOA originating from photooxidation of the solvent, we performed a control photooxidation experiment with tridecane +OH similarly to Exp. 2-3. The CIMS signals of tridecane (positive mode) and tridecane hydroperoxide (negative mode) were used to normalize the 10-50% correction of tridecanederived SOA in MPAN experiments.

2MGA

2-Methylglyceric acid (2MGA) was synthesized and purified according to An et al.41 via oxidation of the C=C bond of methacrylic acid (Aldrich, 99%) using H₂WO₄ (Aldrich, 99%) and 50 wt% H₂O₂ (Aldrich). Proton NMR (in DMSO-d₆) was used to verify the isolated 2MGA with residual CH₃CN (93% 2MGA, Fig. S2A, ESI†). The semi-pure 2MGA (viscous liquid) was then crystallized upon cooling, filtered with CH₃CN, and washed with Et₂O to afford the pure (99%) 2MGA crystalline solid. The CIMS observes 2MGA at the fluoride transfer ion (C₄H₇O₄·HF⁻) and collision-induced dissociation leads primarily to the deprotonated 2MGA ion (C₄H₇O₄⁻, Fig. S2B, ESI†). The gas-phase signal of a 2MGA aqueous solution was measured at various solution pH values in a custom 10 mL glass vial fitted with 3 mm O.D. PTFE sampling tubes. A 0.075 M solution of 2MGA had similar pH to glyceric acid $(pK_a 3.5)^{42}$ at the same concentration. The 2MGA solution was further acidified incrementally by adding droplets of 1-10 wt% H₂SO₄ and the pH of

the solution was measured with a digital pH meter (VWR, Model 8015) that had been calibrated with commercial buffer solutions. The headspace of the vial (at 24 $^{\circ}$ C) was sampled with CIMS at a flow rate of 147 std cm³ s⁻¹ for each solution pH.

High-resolution mass spectrometry

At the end of photooxidation experiments, SOA samples were collected on hydrophilic PTFE-based membrane filters (Millipore, Omnipore, 0.2 µm diameter pores) by pulling chamber air through an activated charcoal denuder, vacuum-sealed, and frozen for further analysis. The SOA material on the filters was gently extracted by wetting with 100-300 µL of acetonitrile and water mixture (2:5 v/v, HPLC grades) for roughly 5 min. The filters were not exposed to heat or ultra-sonication conditions (which may produce free radicals such as OH through cavitation)⁴³ in order to preserve organic species. The extracts were analysed with high-performance liquid chromatography (HPLC) coupled to electrospray ionization mass spectrometry (ESIMS). The separation was performed with a polar embedded C₁₈ column with TMS endcapping (Phenomenex, Synergi™ 4 μ m Fusion-RP 80 Å, 150 imes 2.0 mm) and an eluent mixture of acetonitrile and water (HPLC grades, Aldrich) with 0.5% of formic acid. Generally, the organic constituents eluted together (e.g., the entire oligomer family of 2-methylglyceric acid in the MPAN + OH samples) but are satisfactorily separated from the inorganics. Additionally, an ammonium sulfate solution was analysed with HPLC-ESIMS as an inorganic blank. HSO₄, HSO₄(H₂SO₄)⁻, and HSO₄(H₂SO₄)₂⁻ were the dominant ions observed from the ammonium sulfate solution. These ion signals were subtracted from the SOA mass spectra to remove the remaining inorganic contribution. The mass analyser was a high-resolution (100 000 $m/\Delta m$ at m/z 300) linear-ion-trap (LTQ) Orbitrap™ mass spectrometer (Thermo Corp.) operated in the negative ion mode with a mass range of 80-2000 m/z. The LTQ-Orbitrap was calibrated with commercial standard (LTQ ESI Negative Ion Calibration Solution, Thermo Scientific, Inc.) prior to mass spectral analyses (mass accuracy up to 0.5 ppm at m/z 500).

Results and discussion

Photooxidation of MPAN

Fig. 4A shows the reaction progress for a representative low-NO MPAN experiment under dry conditions. Reagents were equilibrated in the chamber prior to the initiation of photochemistry (yellow shaded region). Fig. 4B shows a representative low-NO MPAN experiment where α-pinene was added at the beginning of the reaction to scavenge NO3, and MAE was added at the end (after the reaction mixture has stabilized with lights off) to observe its effects on SOA formation.

A low yield of NO₂ (7 \pm 3%) is observed by the NO₂/APN instrument and confirmed by the negligible CIMS PNA signal. This NO₂ formation is likely due to thermal decomposition of MPAN ($\tau_{288 \, \text{K}} \sim 35 \, \text{h}$ at the NO_x conditions in Exp. 2-3)^{23,44} because a 10 \pm 5% yield of PAN was also observed. The MPAN decomposition produces NO2 and acylperoxyl radical, which is

Paper PCCP

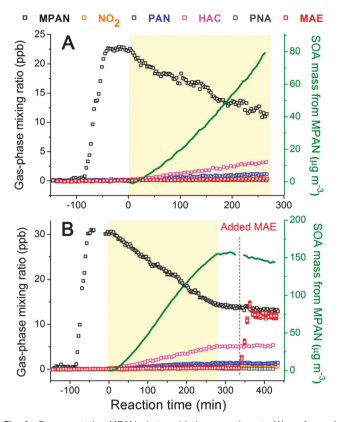


Fig. 4 Representative MPAN photooxidation experiments (A) performed with MPAN and $\rm H_2O_2$ (Exp. 3, base experiment) and (B) the base experiment with the initial addition of α -pinene and subsequent injection of MAE standard (Exp. 4). The yellow shaded areas designate the time period when photochemistry occurs in each experiment. The right mass axis corresponds only to the SOA trace (green line).

subsequently converted to the acyloxyl radical from reaction with either NO or $\mathrm{HO_2}$. The acyloxyl radical decarboxylates to the vinyl radical that ultimately forms PAN and other products through subsequent reactions. The thermal decomposition is calculated to be 10–25% of the total MPAN loss, qualitatively consistent with observed yields. The uncertainty is due to difficulty in determining the $\mathrm{NO_2/NO}$ ratio when $\mathrm{NO_x}$ is low. Neither PAN nor the $\mathrm{NO_2}$ yield was enhanced in the presence of α -pinene (Fig. 4B), suggesting they both originate from MPAN. The lower bound (10%) of the MPAN decomposition is used to correct gaseous molar yields.

In contrast to NO_2 , the experiments performed with α -pinene conclusively demonstrate high yields of an NO_3 co-product, as suggested by mechanisms 1 and 3 (Fig. 2). Fig. 5 shows that upon photooxidation of MPAN in the presence of α -pinene, PNP is formed in $\sim 35 \pm 17\%$ yields with respect to α -pinene loss and $\sim 70 \pm 35\%$ yield with respect MPAN loss. Experiments without α -pinene did not produce interfering species at the mass used for quantification of PNP. Furthermore, experiments performed during the FIXCIT campaign (Exp. # 10, 13 in Table 2 of that work)²² with 30 ppbv of α -pinene demonstrate that PNP is not formed under low-NO photooxidation conditions, similar to those used to oxidize MPAN here (Fig. S3a, ESI†). However, PNP is an abundant product when NO_3 is used as an oxidant

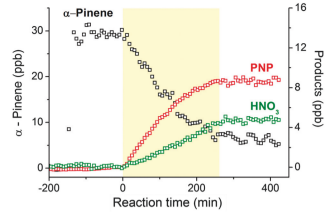


Fig. 5 Representative result from NO_3 scavenging experiment using α -pinene (Exp. 4). The yellow shaded area designates the time period when photochemistry occurs. Note the double-vertical scales.

while HO_2 is present in the chamber (Fig. S3b, ESI†). Using known rate coefficients of α -pinene with NO_3 and OH, 33 we estimate the steady state NO_3 concentration in Exp. 4 (Table 1) to be 6.5×10^6 mol cm $^{-3}$. The steady state OH concentration ($\sim 1.5 \times 10^6$ mol cm $^{-3}$), which is reproducible within 25% under identical oxidant precursor and light conditions, 22 is derived from separate low-NO photolysis experiments using isoprene as a reference hydrocarbon. Thus, we estimate that the loss of α -pinene is $\sim 40\%$ by NO_3 and $\sim 60\%$ by OH. This is in relatively good agreement with the PNP yield from α -pinene assuming most of the pinene nitrooxy alkylperoxyl radical reacts with HO_2 to form PNP.

HNO₃ formation in MPAN experiments (Fig. 5) occurs likely from the NO₃-initiated H-abstraction of alkanes (e.g., tridecane from the introduction of MPAN) or other saturated compounds, as the reaction of NO₃ with α-pinene and MPAN will result largely in addition rather than abstraction. Using the steadystate radical concentrations derived above and reaction rate coefficients from the literature, 34 we estimate < 0.1% of the MPAN loss was due to reaction with NO3 radicals. HNO3 is observed in all MPAN photooxidation experiments due to the ubiquitous presence of the tridecane solvent, and no enhancement is observed in the presence of α -pinene. HNO₃ provides evidence of another 30 \pm 9% NO₃ yield from MPAN, which taken together with PNP, suggests that MPAN efficiently yields NO₃ and a co-product upon reaction with OH. The high NO₃ yield confirms one shared aspect of mechanisms 1 and 3 (Fig. 2) and, in combination with the low NO₂ observations, allows us to rule out mechanism 2 as a significant contributor to the MPAN photooxidation chemistry.

We observe a large mass yield of SOA from the dry MPAN + OH experiments (Fig. 4, green traces), suggesting that the MPAN photooxidation chemistry is efficient at forming SOA in the absence of NO_x. Approximately 125–145% SOA by mass (with respect to MPAN mass reacted) is formed, and assuming an average molecular weight of 340 g mol⁻¹ (the intensity-weighted average of the MPAN + OH high-resolution mass spectrometry data, assuming similar ESIMS sensitivities for

observed analytes), approximately \sim 60 (+15/–5)% by mole. The uncertainty is derived from corrections (e.g., for tridecane-derived SOA, particle wall loss, and MPAN thermal decomposition) and analysis of the average molecular weight of the SOA. The observation that SOA formation from MPAN + OH is not NO-dependent is consistent with the suggestion in mechanisms 1 and 3 (Fig. 2) that the traditional RO₂ chemistry of the thermalized radical A does not come into play. However, we note the data do not eliminate the possibility that an RO₂ formed after O₂ addition also rapidly decomposes to the same products.

Importantly, the SOA formation from MPAN + OH is prompt and occurs without the injection of inorganic seed aerosols. It is possible the SOA forms via nucleation (e.g., from the chemistry of larger VOC like tridecane) or growth in the presence of pre-existing seed particles that are under the size detection limit of the SMPS and AMS. Even if SOA growth occurs on dry, pre-existing, nanoparticles, most of the SOA formation occurs on primarily-organic particles after the first few condensation cycles. This observation favors the proposed mechanism 3 over mechanism 1 due to the need for MAE to undergo reactive uptake partitioning, which is a slow process for this compound even in the presence of hydrated, acidic sulfate particles that would accelerate ring-opening of the epoxide ($\gamma \sim 5 \times 10^{-4}$ when seed aerosols are 50% H₂SO₄ by mass).

Of the putative SOA precursors, MAE is the only compound that has been detected by mass spectrometry due to its chemical stability. Throughout all experiments, we observed small to negligible yields of MAE in the gas phase (2 \pm 1%, upper limit, see Experimental), whether the precursor was MPAN or MACR. FIXCIT experiments of MACR under low-NO conditions also did not observe any appreciable MAE production. 22 Lin et al. 18 similarly observed a small gas-phase yield of MAE (~1% from MACR, Fig. S1 of that work), but suggested that MAE is so rapidly converted to SOA mass that most of it is observed in the condensed phase. Under the assumption that 50-80% of MAE is observed as SOA, they estimated MAE yields are 18-32% from MPAN. We demonstrate that MAE is not wellconverted to SOA and, in particular, not via the mechanism that forms the dominantly-organic particles in the MPAN experiments. It instead appears that MAE is a remarkably stable and volatile gas-phase species, with similar volatility to MPAN, which by itself does not form SOA through equilibrium partitioning. Notably, when $\sim 12~(\pm 50\%)$ ppbv of MAE is injected into the chamber following the reaction period (Fig. 4B), no change to particle size or mass concentration can be observed.

To further evaluate the inertness of MAE, we injected a few hundred ppbv of MAE into a clean 1000 L teflon bag alongside 500 ppbv of HNO $_3$ (\sim 300 ppbv of which remained in the gas phase, and the rest, presumably, coated the chamber walls). No observable wall loss of MAE occurred over the course of 2.5 hours (Fig. S4, ESI†); this is identical to its behavior in the absence of acid. It appears that the small quantity of MAE formed from MACR oxidation is not an eager participant in the surface- and/or water-induced partitioning that often leads to SOA formation. The inertness we observe provides insight into

why 300 ppbv (\sim 1250 µg m $^{-3}$) of MAE produced only \sim 10 µg m $^{-3}$ of SOA through reactive uptake in the Lin $et~al.^{18}$ work, even when using seed particles that are exceptionally acidic (\sim 50 wt% H $_2$ SO $_4$ at RH < 10%). The evidence suggests that the low (<2%) gasphase mixing ratios observed in this work and by Lin $et~al.^{18}$ (from MACR) indeed represent a measure of the entire yield of MAE. Thus, it appears that the disagreement between this work and that of Lin $et~al.^{18}$ lies in the assumption of MAE's ability to form SOA, and not in the observations of MAE itself. As MAE formation is negligible both from MPAN and MACR, mechanism 1 is ruled out as a significant contributor to MPAN-derived SOA.

A key element to understanding the MPAN + OH reaction lies in the measurement of hydroxyacetone (HAC). The HAC yield determined here (\sim 25 \pm 7%) is consistent with the upper limit determined by Orlando et al. (<40%), ²¹ but more than a factor of two lower than the yield determined by Grosjean et al. $(\sim 60\%)$. Both of the previous works were performed under high-NO conditions using ethyl nitrite as an OH precursor. The source of discrepancy with regards to the Grosjean work is not clear; however, a similar trend in data agreement can be observed in the determination of rate coefficients of OH + MPAN. We measured $k_{288 \, \text{K}} \sim 3 \, (\pm 1) \times 10^{-11} \, \text{cm}^3 \, \text{mol}^{-1} \, \text{s}^{-1}$, which compares well with Orlando et al. ²¹ $(k_{277 \, \text{K}} \sim 3.2 \, (\pm 0.8) \, \times$ 10^{-11} cm³ mol⁻¹ s⁻¹), but is an order of magnitude higher than the value reported by Grosjean et al. 20 $(k_{298\,\mathrm{K}} \sim 3.6 ~(\pm 0.4) \times$ 10⁻¹² cm³ mol⁻¹ s⁻¹). The coefficient by Orlando et al. is preferred by IUPAC33 due to its consistency with the OH reaction with structurally-similar APNs, 47 and our data is in agreement with this recommendation. Thus, we only consider the comparison between this work and that of Orlando et al. 21

If the production of HAC occurs via the lactone-production pathway (as the data seem to indicate) then the yield would be independent of NO mixing ratio. The agreement in the HAC yield obtained under low-NO conditions (this work) and high-NO conditions²¹ suggests that HAC is not formed from the alkylperoxyl radical B in the MPAN + OH reaction. Finally, the experimentally-determined HAC yield (and, consequently the remainder HMML yield) is comparable to those theoretically predicted by Kjaergaard et al. 17 (\sim 36% of HAC). In contrast, Lin et al. 18 suggest a low HAC yield (\sim 3%). The current work cannot differentiate the 2% of thermalized radical A that is predicted to form, 17 ultimately generating the alkoxyl radical under high NOx conditions, because the products would likely include HAC and NO₃. Together, the experimental data support the HMML mechanism from the MPAN + OH reaction. We estimate approximately 25% of the reaction yields HAC + CO + NO₃ and 75% of the reaction yields HMML + NO3, with 75-100% of HMML leading to the formation of SOA under dry conditions.

To investigate whether there is a route to HMML from thermalized alkyl radicals, we performed experiments with saturated APNs that are analogous to MPAN. The photooxidation of the saturated analogues occurs *via* H-abstraction instead of OH-addition, which generates lower-energy alkyl radicals. For example, we synthesized A in the chamber *via* the OH-initiated photochemistry of 3-hydroxy-2-methylpropanoyl peroxynitrate (HMPPN). HMPPN is a major product of the high-NO₂ OH-initiated oxidation

of hydroxymethyl propanal (HMP), which was produced by dark ozonolysis of 231MBO (Section S1 and Fig. S5 and S6A, ESI†). Expected gas-phase products arising from RO₂ and RO radicals were observed from the chemistry of A but not A* (Fig. S7, ESI†), *e.g.*, the hydroxynitrate, the hydroxyhydroperoxide, hydroxyacetone, and 2-oxopropanoyl peroxynitrate. ²¹ Qualitatively compared to MACR, the SOA formation from the photochemistry of HMP (Exp. #11, Fig. S6B, ESI†) and isobutyraldehyde (Exp. #10, not shown) is negligible (\ll 1%). Chan *et al.* ¹³ first suggested the importance of the β-unsaturation in MPAN, following the observation that α , β -unsaturated aldehydes produced the largest SOA yields. Our results are consistent with their explanation.

Additionally, we use the potential energy surfaces from the Kjaergaard $et~al.^{17}$ work to calculate the relative fates of A. Compared to a typical effective rate for the reaction of alkyl radicals with O_2 at 1 atm air $(O_2 = 21\%)$ and 298 K $(k_{\rm eff} \sim 1 \times 10^7~{\rm s}^{-1})$, the thermalized unimolecular decomposition of A to HMML is much slower $(k_{\rm (A\rightarrow HMML)} < 10^3~{\rm s}^{-1})$. This translates to a calculated 0% yield of HMML in the HMPPN + OH reaction (100% reacts with O_2). In comparison, Kjaergaard $et~al.^{17}$ estimated the decomposition of A* to HMML is fast under the conditions they studied $(k_{\rm (A^*\rightarrow HMML)} \sim 4 \times 10^9~{\rm s}^{-1})$, translating to a 61–74% HMML yield from MPAN (2% is stabilized to A). The theoretical-derived kinetic results are consistent with observations for these C_4 APN-derived alkyl radicals.

Fig. 6 summarizes the chemical mechanisms in the OH-initiated oxidation of MPAN and HMPPN to reflect the current scientific knowledge. It is also possible that larger APNs behave differently than the C₄ APNs discussed here. For example, it is not clear if APNs from the monoterpene aldehydes such as pinonaldehyde (saturated) and limonaldehyde (unsaturated, in the gamma position) are able to produce SOA or undergo a different photochemical fate. These larger APN systems are intriguing because they can be produced in the laboratory²² but have not been observed in the ambient atmosphere.

Atmospheric fate of HMML

To study the SOA formation from HMML under conditions more relevant to the atmosphere, we performed MACR high-NO₂ photooxidation experiments with hydrated ammonium sulfate (AS) seed particles at 40 and 85% RH (efflorescence and deliquescence RH of AS are ~ 30 and $\sim 80\%$, respectively).48 Fig. 7 shows that the SOA production is lower and gas-phase 2MGA yields are higher when the reaction occurs in deliquesced AS particles (Exp. #13, liquid water \sim 75 vol%) compared to primarily organic particles (under dry conditions, Exp. #9). The small MAE signal did not increase under humid conditions, suggesting that the formation of this compound is likely not important in the ambient environment. The production of HAC is observed to slightly increase toward the end of the experiment. However, this change is within the uncertainty of the HAC determination ($\pm 30\%$). If the difference in HAC signal between dry and humid is real, the underlying mechanism is unclear. The systematic uncertainty in calculating mixing ratio for 2MGA is approximately $\pm 50\%$. However, if all of the ~12-13 ppb of 2MGA in the gas phase is converted to particle mass, the volatilization of 2MGA may explain almost 100% of the $\sim 60 \ \mu g \ m^{-3}$ SOA mass discrepancy between the two experiments. 2MGA is typically used as a tracer for isoprene SOA; however, it appears that its accommodation in the condensed phase is low at typical aerosol-phase pH. Recent observations in the Southeast U.S. and Amazon forests support the idea that 2MGA may exist primarily in the gas phase in humid biogenic environments.49

Fig. 8A shows the CIMS gas-phase measurement of 2MGA (from a 0.075 M solution of the synthesized standard) as the solution is increasingly acidified. 2MGA behaves similarly to a semivolatile compound such as isoprene epoxydiol in the CIMS instrument, *i.e.*, it takes approximately 10 minutes to equilibrate

Fig. 6 The OH-initiated oxidation mechanism of MPAN and HMPPN derived from the low-RH experiments conducted in this work. The HMML formation from MPAN arises from a chemically-activated mechanism while the product formation from HMPPN arises from a thermalized alkyl radical + O_2 reaction.

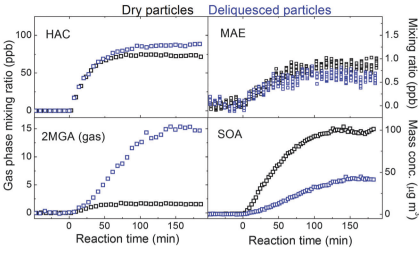


Fig. 7 The formation of gas-phase species and secondary organic aerosols under dry conditions (black markers) and under humid conditions (85% RH) with deliquesced ammonium sulfate seeds (blue markers).

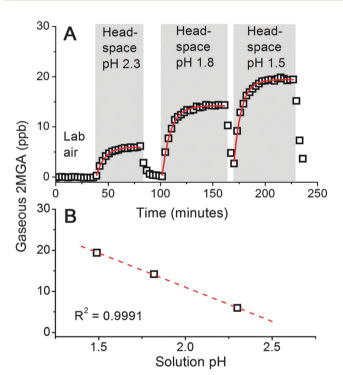


Fig. 8 CIMS measurements of the headspace air above a 0.075 M 2-methylglyceric acid (2MGA) solution: (A) the gaseous 2MGA signal when switching from lab air to solutions with different pH and (B) the headspace 2MGA mixing ratio (obtained through exponential fitting) is a linear function of pH in the measurement range.

with teflon tubing compared to a more-volatile compound such as hydroxyacetone (<1 min). The gas-phase mixing ratio linearly increases with decreasing solution pH (Fig. 8B) in the measurement range, as expected for a semi-volatile organic acid. This pH-dependent volatilization is likely to be important for understanding the SOA formation from not only MPAN oxidation, but other types of atmospheric reactions that produce organic acids in the condensed phase, *e.g.*, cloud-processing oxidation, 50,51 ozonolysis of alkenes, 52,53 among other reactions.

Interestingly, gas-phase 2MGA is not significantly enhanced and aerosol mass is not significantly decreased in the 40% RH experiment (Exp. #12) compared to the dry experiments. It is possible that the liquid water content under these conditions (\sim 30 vol%) is not high enough to dominate the reaction with

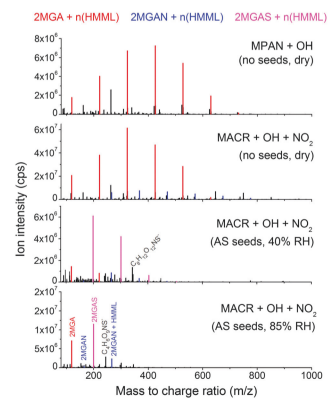


Fig. 9 Negative mode ESI-HRMS analysis of select SOA samples, integrated for retention times 2.5–3.5 min. The inorganic constituents have been separated by HPLC. Oligomer families comprised of HMML units are color-coded based on the monomer compounds (2MGA = 2-methylglyceric acid, 2MGAN = 2-methylglyceric acid nitrate, and 2MGAS = 2-methylglyceric acid sulfate).

HMML or there is a barrier to 2MGA volatilization. For example, if organic-inorganic phase separation occurs at RH between efflorescence and deliquescence, 54,55 the reaction of gas-phase HMML with water and solvated ions and the volatilization of aqueous 2MGA may be hindered by an organic shell. Additionally, when liquid water content is lower (and thus, ionic strength is higher), organic acids such as 2MGA may form involatile salts with NH₄⁺ and other cations.⁵⁶ Salt formation is not expected to be significant for deliquesced particles, but can be considerably enhanced for drier metastable particles at 30-50% RH.57

Fig. 9 shows the results of high-resolution mass spectrometry (HR-MS) analyses of SOA collected from MPAN and MACR photooxidation. When the photooxidation of MACR is performed without inorganic seeds and under dry conditions, long oligomer families of 2MGA that are separated by 102.0317 Da (the exact mass of HMML) are observed, consistent with earlier mass spectrometry observations. 12,14,58 The SOA composition from MPAN + OH appears almost identical to that of MACR + OH + NO₂. In addition to the 2MGA + HMML oligomers, the MACR high-NO_x photooxidation produces an organic nitrogen oligomer family (Fig. 9, blue peaks). The organic nitrogen family of 2MGA-nitrate + HMML elutes primarily at later retention times (8-20 min), likely due to their lower polarity compared to organic sulfates and acids, but is absent in other chemical systems, including the MPAN + OH low-NO_x reaction. These N-containing SOA compounds are likely formed from the reaction of HNO₃ (from NO₂ + OH reaction in the MACR high-NO_r experiments) with HMML in the condensed phase or interfacial region, which generate the nitrate ester of 2MGA through nucleophilic ring opening.

When aqueous sulfate is present, the sulfate ester of 2MGA is produced (Fig. 9, magenta peaks), which react with HMML similarly to 2MGA and 2MGA-nitrate. In ammonium sulfate particles at 40% RH, the oligomer families of 2MGA-sulfate + n(HMML) and 2MGA + n(HMML) are observed up to the trimer and dimer, respectively. In the deliquesced AS seeds at 85% RH, the monomer compounds 2MGA, 2MGA-sulfate, and the sulfate nitrate (C₄H₆O₉NS⁻ ion) dominate the mass spectra. An exception is the dimer 2MGA-nitrate + HMML that has higher signal than the monomer. This may be attributed to the fact that nitrate ion is a poor nucleophile and nucleophilic substitution with water and sulfate may readily occur, so the monomer nitrate has a short condensed-phase lifetime. Thus, the organonitrate may be more stable in the dimer form. However, it is also possible the nitrate dimer is more ionizable in negative mode ESI compared to the monomer. We do not expect a large ionization difference between 2MGA-nitrate and 2MGA as the carboxyl moiety is believed to be the charge carrier for this compound in negative mode ESI with aqueous-based solvents.⁵⁹

Fig. 10 Mechanism of SOA production from HMML (the blue dashed area represents the condensed phase). When particle liquid water is high, the main fate of HMML is reaction with water to produce monomeric 2MGA or sulfate and nitrate ions to produce 2MGA-sulfate and 2MGA-nitrate, respectively. The 2MGA monomer is volatile enough to re-partition to the gas phase. When the SOA composition is higher in organics than water, the main fate of HMML is reaction with its condensed-phase derivatives, producing low-volatility polyesters that increase SOA mass. Hydrated particles with higher free acidity may favour the monomeric form of 2MGA by neutralizing the carboxylate and hydrolyzing the sulfate and nitrate esters, possibly suppressing SOA growth

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The combined mass of the SOA from these organic acid and hetero-atom compounds in the 85% RH experiment, however, is lower than the mass of 2MGA in the gas phase (Fig. 7).

These aerosol phase HR-MS observations are consistent with earlier suggestions that relative humidity plays a role in modifying the composition and yield of isoprene-derived high-NO_x SOA. 60,61 However, previous works did not study 2MGA particle/ gas partitioning and discussions were framed with respect to poly-condensation reactions (e.g., a Fisher type esterification) – a hypothesis that is not consistent with the present data. Furthermore, it has been recently shown that Fisher type esterification is too slow to form 2MGA oligomers under standard conditions. 62 This does not, however, preclude condensation reactions from occurring when aerosol particles or cloud/fog droplets are evaporated. 63,64 In light of the gas-phase mechanism of MPAN + OH presented here, the molecular driving force behind the effect of RH on MPAN aerosol composition and yield can now be understood in terms of lactone polymerization.

Fig. 10 shows the proposed mechanism with which HMML is converted to SOA and gas-phase 2MGA. Lactones are wellknown to produce polyesters via polyaddition. 65 We suspect that, in nature, the anionic ring opening polymerization scheme is the active mechanism. Nucleophilic attack at the sp³ carbon is dominant for α -lactones, ⁶⁶ partly due to the stability of the carboxylate (estimated to be only $\sim 10 \text{ kcal mol}^{-1}$ higher in energy than the α-lactone).67 The propagated carboxylate can repeatedly add HMML under low RH conditions, where its main fate is reaction with organics that coat the particle. In ambient SOA, ring-opening initiators may be aqueous inorganic anions or myriad carboxylates that are present in the condensed phase. When liquid water is in excess (as is mainly the case in the lower troposphere), it appears the main fate of HMML is reaction with H₂O, aqueous sulfate, or aqueous nitrate, and not with organics. Because the organonitrate is more easily hydrolyzed, the major product of the reaction under typical atmospheric conditions is expected to be monomeric 2MGA and 2MGA sulfate. The 2MGA may re-partition into the gas phase, leading to significantly reduced SOA mass yields.

Atmospheric implications

The representation of MPAN oxidation in atmospheric models has important ramifications. Thus far, only the proposed MAE mechanism (i.e., reactive uptake) has been considered for modelling SOA formation and framing field observations.^{68,69} Previously modelled SOA formation from MAE increases with particle acidity, while simulations based on HMML are expected to exhibit a different temporal and spatial variability. Pye et al. 68 used the HMML and MAE yields suggested by Lin et al. 18 (57% and 21%, respectively); however, both compounds were assigned to have the acid-catalysed heterogeneous fate of MAE (i.e., essentially converting HMML mass to MAE). The observed and predicted SOA from MPAN deviates from the 1:1 relationship in that work, likely a result of the assumed

SOA mechanism. Fig. 8 and the mechanisms in Fig. 10 suggest a more complex impact of particle free acidity. For example, in conditions when particle free acidity is high, the neutral form of 2MGA will be favored and the organosulfate/organonitrate may hydrolyze more rapidly.⁷⁰ Both situations will enhance the abundance of 2MGA in the gas phase at the expense of lower-volatility derivatives that are important for SOA formation and growth.

The regional atmospheric importance of MPAN will vary with NO₂/NO ratio and isoprene emissions. Kjaergaard et al. 17 estimated that approximately 4 Tg of HMML is produced per year (up to $0.8 \mu g m^{-3} day^{-1}$ in the Southeast U.S.). However, whether the carbon from HMML is directed toward the formation of gaseous 2MGA or SOA will depend on atmospheric conditions and particulate matter characteristics. Typically, oligomers are not observed in ambient SOA and the mass concentrations of 2MGA and 2MGA-sulfate are low (e.g., 7-9 ng m⁻³ and 12 ng m⁻³ in Southeast U.S., respectively.)⁷¹ The low ambient observations in the Southeast US, a region characterized by higher particle liquid water and lower particle pH,72,73 are consistent with the HMML-derived SOA formation mechanism (Fig. 10). This work demonstrates that certain "SOA tracers" have more complex fates in nature than previously recognized. Interpreting the abundance of aqueous 2MGA and 2MGA-sulfate will require knowledge of physical parameters that may not be available for these compounds (e.g., Henry's Law coefficients, reaction coefficients, temperature and pH-dependent accommodation). We expect that integrating the lactone mechanism into atmospheric models will produce a more accurate representation of the magnitude, temporal variation, and spatial distribution of isoprenederived organic compounds (whether in the gas or condensed phase) near anthropogenically-influenced regions.

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