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Formation of highmolecular-weight peroxides

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The relative importance of competing pathways for the formation of high-molecular-weight peroxides in the ozonolysis of organic aerosol particles

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Abstract

High-molecular-weight (HMW) organic compounds are an important component of atmospheric particles, although their origins, possibly including in situ formation pathways, remain incompletely understood. This study investigates the formation of HMW

- ⁵ organic peroxides through reactions involving stabilized Criegee intermediates (SCI's). The model system is methyl oleate (MO) mixed with dioctyl adipate (DOA) and myristic acid (MA) in submicron aerosol particles. Criegee intermediates are formed by the ozonolysis of the double bond in methyl oleate. An aerosol flow tube coupled to an Aerodyne quadrupole aerosol mass spectrometer is employed for the studies. The
- relative importance of different HMW organic peroxides, including secondary ozonides (SOZ's), α -acyloxyalkyl hydroperoxides and α -acyloxyalkyl alkyl peroxides (α AAHP-type compounds), diperoxides, and monoperoxide oligomers, is determined for the ozonolysis of different mixing mole fractions of MO in DOA and MA. Two SOZ's are identified as the HMW products of the ozonolysis of pure methyl oleate. Changes in
- ¹⁵ the mobility and vacuum aerodynamic diameters of the particles indicate that up to 67% of the product mass remains in the particles, with the balance evaporating. In mixed particles, SOZ's form within an inert matrix of DOA to as low as 0.04 mole fraction MO. In comparison, in mixed particles of MO and MA, α AAHP-type compounds form in high yields for initial MO mole fractions under 0.5, suggesting that SCI's efficiently
- ²⁰ attack the carboxylic acid group of myristic acid. The reactions of SCI's with carboxylic acid groups to form α AAHP-type compounds therefore compete with those of SCI's with aldehydes to form SOZ's, provided that both types of functionalities are present at significant concentrations. The results suggest that SCI's formed by the ozonolysis of unsaturated organic molecules in atmospheric particles could lead to the transforma-
- tion of carboxylic acids and other protic groups into HMW organic peroxides.



1 Introduction

Gas-phase ozonolysis of unsaturated organic molecules has been intensively investigated for decades (Atkinson and Arey, 2003). Ozonolysis is an important pathway for the degradation and the transformation of compounds in the atmosphere and is also

- ⁵ a potential night-time source of radical species (Paulson and Orlando, 1996). Furthermore, some of the reactions yield products having sufficiently low vapor pressure to form secondary-organic-aerosol (SOA) particles (Griffin et al., 1999). Knowledge of the further chemical reactions inside the particle phase is, however, limited. Investigations of particle-phase ozonolysis have been accelerating recently (e.g., Rudich, 2002). Katrib et al., 2004. Ziemenn, 2005). Although azonchusia reactions have been
- 2003; Katrib et al., 2004; Ziemann, 2005). Although ozonolysis reactions have been studied in depth by organic chemists for some time (Bailey, 1978), the reactions inside aerosol particles have important differences because of the mixture of components, the possible presence of water, and the coupling to the gas phase through evaporation and condensation. Ozonolysis of organic molecules inside particles can lead to the
- formation of increasingly polar compounds that alter physicochemical properties, such as particle hygroscopicity, CCN activity, and light scattering and extinction. The process by which these properties change during the residence time of a particle in the atmosphere is called "particle aging" (Kanakidou et al., 2005). In addition to reactions with ozone, particles also age by reaction with hydroxyl radical (Bertram et al., 2001;
- Rudich, 2003; Eliason et al., 2004), nitrate radical (Hung et al., 2005; Docherty and Ziemann, 2006), and internally mixed chemical species, including acid-catalyzed and uncatalyzed pathways (Jang et al., 2002; Kalberer et al., 2004; linuma et al., 2004; Tolocka et al., 2004; Gao et al., 2004).

A model system recently employed by numerous investigators in studies of particle aging is the ozonolysis of oleic acid in both pure and mixed systems. Ozonolysis reaction kinetics have been studied, including the gas-phase loss rate of ozone, the condensed-phase loss rate of the organic compound, and the effects of chemical mixtures, particle morphology, and physical state on those loss rates (Moise and Rudich,

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2002; Morris et al., 2002; Smith et al., 2003; Thornberry and Abbatt, 2004; Hearn et al., 2005; Hearn and Smith, 2005; Katrib et al., 2005a, 2005b; Knopf et al., 2005; Ziemann, 2005). Product identification and yield and the reaction mechanisms of ozonolysis have been investigated (Katrib et al., 2004; Zahardis et al., 2005, 2006; Hung et al., 2005; Ziemann, 2005;; Reynolds et al., 2006¹). Ozone-induced changes in particle properties including hydroscopicity (Asad et al., 2004). CCN activity (Broekhuizen et al.)

erties, including hygroscopicity (Asad et al., 2004), CCN activity (Broekhuizen et al., 2004), and density (Katrib et al., 2005b) have been quantified.

Several of these studies (Hung et al., 2005; Zahardis et al., 2005, 2006; Ziemann, 2005; Hearn et al., 2005; Reynolds et al., 2006¹) suggest the occurrence of ozone-induced oligomerization inside atmospheric aerosol particles as the ultimate fate of a

- ¹⁰ Induced oligomerization inside atmospheric aerosol particles as the ultimate fate of a portion of the stabilized Criegee intermediates (SCI's). SCI's are primary products of the ozonolysis of unsaturated organic molecules. In situ oligomerization is one possible explanation for the large quantities of high-molecular-weight (HMW) organic compounds that have recently been measured in atmospheric aerosol particles in urban
- ¹⁵ and rural locations (Zappoli et al., 1999). To further assess the likelihood of ozoneinduced oligomerization as an important source of HMW organic compounds in the atmosphere, the scientific community requires more quantitative information concerning the effects of several key differences between atmospheric processes and laboratory studies, both of which are tied to the chemistry of mixed particles compared to pure ones. These differences include the following:
 - The concentration of double bonds in particles of pure model compounds differs from that of atmospheric particles. This difference could affect the relative concentration of SCI's and thus the relative importance of competing pathways and product yields (including oligomerization) in a nonlinear kinetic system. Competing pathways identified in the ozonolysis of oleic acid include the formation of secondary ozonides (Hung et al., 2005; Zahardis et al., 2005; Ziemann, 2005;



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¹Reynolds, J. C., Last, D. J., McGillen, M., Nijs, A., Horn, A. B., Carpenter, L. J., Lewis, A. C.: Structural analysis of oligomeric molecules formed from the reaction products of oleic acid ozonolysis, submitted, 2006.

Reynolds et al., 2006¹), α -acyloxyalkyl hydroperoxides (i.e., ester hydroperoxides) (Hung et al., 2005; Ziemann, 2005; Zahardis et al., 2006; Reynolds et al., 2006¹), and diperoxides (Zahardis et al., 2005; Reynolds et al., 2006¹), all of which form HMW organic peroxides. Monoperoxide oligomers of the type (-CCOOC) are also possible (Nishikawa et al., 1995), although not observed by Reynolds et al. (2006)¹.

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 In atmospheric particles, numerous oxygenated organic compounds may react with the SCI's, again affecting the relative importance of competing pathway, including oligomerization. These reactions imply the decomposition of saturated organic compounds and the formation of a variety of compounds, including other types of oligomers.

The present paper provides a report concerning the formation of HMW organic peroxides from the reactions of stabilized Criegee intermediates. Ozonolysis reactions are carried out for variable matrix substances and double-bond concentrations. The model system is methyl oleate in mixed submicron aerosol particles. Unlike oleic acid, methyl oleate does not have a free carboxylic acid group, which when present can propagate polymerization in a myriad of pathways. Hence the reactions of methyl oleate are expected to be simpler than those of oleic acid, which aids greatly in sorting through the importance of competing pathways. Methyl oleate mixed at systematic

- mole fractions with nonreactive dioctyl adipate (DOA) is an approach for varying the concentration of double bonds. Similarly, methyl oleate mixed at systematic mole fractions with myristic acid (MA), which bears a reactive –COOH group, is an approach for investigating the reactivity of SCI's towards this pathway. Comparison of products formed for these different conditions is employed as the experimental strategy
- to elucidate the relative contribution of reaction pathways leading to several classes of peroxides, including secondary ozonides (SOZ's as C(OO)OC), α -acyloxyalkyl peroxides (α AAHP-type compounds as C(O)OC(OOH)), diperoxides (DP's as C(OO)₂C), and monoperoxide oligomers (CCOOC) as product pathways in the formation of HMW

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organic peroxides. Ozonolysis products are identified using an Aerodyne aerosol mass spectrometer (AMS) coupled to an aerosol flow tube.

2 Experimental procedures

- 2.1 Chemicals
- Aerosol particles of pure methyl oleate and mixtures of it with other organic solutes were generated by atomizing solutions in organic solvents (HPLC-grade, TSI 3076 atomizer). To examine a possible influence of unevaporated solvent remaining in the aerosol particles, experiments were carried out in duplicate with ethyl acetate and methanol as solvents. Within experimental uncertainty, identical results were obtained in most experiments, and the few exceptions are discussed later in the text. The figures show data recorded from ethyl acetate solutions. Organic solutes used included methyl oleate (MO) (99%), ethyl oleate (EO) (98%), azelaic acid monomethyl ester (AAME) (85%), dioctyl adipate (DOA) (bis(2-ethylhexyl) adipate, 99%), and myristic acid (MA) (>99%). These compounds were purchased from Aldrich.
- 15 2.2 Instrumental setup

A diagram of the experimental setup is presented in Fig. 1. The aerosol passed through a diffusion charcoal scrubber and a bipolar charger. The aerosol flow was then introduced into a differential mobility analyzer (DMA) (TSI 3071A). 500-nm particles were classified (see Sect. 2.3). The particle loading ranged from 100 to 1000 μ g m⁻³. The

flow rates of the sample and sheath flows entering the DMA were 0.3 and 3 L min⁻¹, respectively. The monodisperse aerosol flow was introduced to a flow tube reactor (20-mm inner diameter and 60-cm length) at a rate of 0.15 L min⁻¹ and 1 atm. The remainder of the monodisperse aerosol was vented (0.15 L min¹ regulated by a critical orifice purchased from O'Keefe Controls Co.). The sheath flow of 0.75 L min⁻¹ for the



reactor was a dry O₂/N₂ mixture having 0 to 350 ppm O₃(0 to 8.8×10¹⁵ molec cm⁻³). Ozone was generated in ultrapure air by ultraviolet irradiation from a low-pressure mercury lamp (Jelight Model 600). The interaction time between the aerosol particles and the ozone was 6s when the injector was fully withdrawn, which was the condition for
 ⁵ most experiments described in this report. Experiments were at room temperature and 0% relative humidity. The integrated number concentration was measured with a Condensation Particle Counter (TSI 3022A, 0.30 L min⁻¹). Alternatively, a Scanning Mobility Particle Sizer (SMPS) (TSI 3936, 0.30 L min⁻¹) was used for mobility diameter measurements. The sheath flow in the tube reactor was regulated using a critical orifice (0.52 L min⁻¹).

The chemical composition of the aerosol particles was analyzed using a quadrupole aerosol mass spectrometer (AMS, 0.08 Lmin^{-1}) (Aerodyne Research, Inc.). Three different operating modes of the AMS were employed during the experiments, namely the mass spectrum (MS) scanning mode, the particle time-of-flight (TOF) mode, and the jump mass spectrum (JMS) mode (Jayne et al., 2000; Jimenez et al., 2003). The MS mode was used to obtain mass spectra from m/z 0 to 390, where m/z denotes the mass-to-charge ratio of the fragment. The TOF mode was used to measure the vacuum aerodynamic diameter (d_{va}) of particles. The JMS mode provided increased analytical sensitivity for monitoring intensities of high-mass ions (m/z>202).

20 2.3 Determination of aerosol particle size

In the absence of O_3 , the diameter of MO particles decreased by ca. 4 nm s^{-1} because of evaporation in the flow tube. The evaporation rate was determined by the following procedure. (1) The injector was fully inserted into the flow tube, thus giving a residence time of 0 s. The DMA voltage was adjusted to a mobility diameter (d_m) of

²⁵ 500 nm. The volume-equivalent diameter equaled the mobility diameter because MO was liquid at room temperature and because the aerosol particles were therefore non-porous and spherical. When the injector was fully withdrawn so that the residence time



of the particles in the flow tube was 6s, a decrease in d_{va} of approximately 20 nm was observed as a result of evaporation (Table 1). The d_m values of 500 and 477 nm at 0 and 6s injector positions, respectively, were calculated from the relationship of $d_m = (\rho_0/\rho_{MO})d_{va}$ (DeCarlo et al., 2004; Katrib et al., 2004b), where the densities ρ_{MO} and ρ_0 were 0.874 and 1.000 g cm⁻³, respectively. The mass spectrum verified no change

in particle composition for different injector positions, as expected.

In addition to evaporation, particle diameter also changed by the addition of O_3 to the flow. Factors contributing to the change in diameter included (1) the reactive uptake of O_3 by particles (i.e., increasing particle mass by adding oxygen atoms to molecules),

- ¹⁰ (2) increases or decreases in the particle density due to alterations of the chemical composition, (3) the suppression of MO evaporation as a result of MO consumption during ozonolysis, and (4) the formation and the evaporation of volatile products such as nonanal. As shown in Table 1, d_{va} of particles after the ozonolysis was 446 nm for an ozone exposure $P_{O3} \times t$ of 3×10^{-4} atm s, where P_{O3} and *t* are the partial pressure of O_3 and the interaction time in the flow tube, respectively. In a separate measurement
- (see also Sect. 3.2), d_m of 438 nm for the reacted particles was measured, from which a density of 1.02 g cm⁻³ was calculated by $\rho = \rho_0 d_{va}/d_m$ (DeCarlo et al., 2004).

For the organic compounds used in the mixed particle experiments, control studies employing pure particles showed that MA (ca. 1 nm s^{-1}), AAME (ca. 3 nm s^{-1}), and EO (ca. 2 nm s^{1}) also evaporated in the flow tube. No evaporation of DOA was detected.

2.4 Mole fraction of methyl oleate in mixed particles

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Prior to the ozonolysis of particles of MO mixed with a matrix substance such as DOA or MA, the mole fraction of methyl oleate (x_{MO}) was determined, as follows. The weight percent of MO (w_{MO}) was related to the relative contribution of m/z 296 of the MO



molecular ion to the mass spectrum, as follows:

$$w_{\rm MO} = \frac{100 R/E_{\rm matrix} \left(l_{\rm mix}^{296} / \sum_{m/z} l_{\rm mix}^{m/z} \right)}{R/E_{\rm MO} \left(l_{\rm MO}^{296} / \sum_{m/z} l_{\rm MO}^{m/z} \right) + (R/E_{\rm matrix} - R/E_{\rm MO}) \left(l_{\rm mix}^{296} / \sum_{m/z} l_{\rm mix}^{m/z} \right)}$$
(1)

where RIE_{MO} and RIE_{matrix} are the ionization efficiencies of MO and of the matrix substance, respectively, relative to nitrate as ammonium nitrate (Jayne et al., 2000). The derivation of Eq. (1) is provided in the appendix. The *RIE* values were determined using particles of pure compounds. The measurements were made at the 6-s injector position, thereby including the effect of volatilization in the flow tube. In Eq. (1), I_{MO}^{296} and I_{mix}^{296} are the m/z 296 intensities recorded for pure MO and mixed MO/matrix particles, respectively. $\Sigma I_{MO}^{m/z}$ and $\Sigma I_{mix}^{m/z}$ denote intensity sums for m/z 0 to 390, omitting m/z values of air beams (e.g., m/z 28). The matrix substances were verified to have negligible intensity at m/z 296.

The mole fraction of MO (x_{MO}) was calculated from its weight percent, as follows:

$$x_{\rm MO} = \frac{w_{\rm MO}}{MW_{\rm MO}} \left/ \left(\frac{w_{\rm MO}}{MW_{\rm MO}} + \frac{100 - w_{\rm MO}}{MW_{\rm matrix}} \right) \right.$$
(2)

where MW_{MO} and MW_{matrix} were the molecular weights of MO and the matrix substance, respectively. The mole fractions determined by Eq. (2) differed by up to 50% from those of the reservoir solutions used in atomization, presumably because of different evaporation rates of MO and the matrix substances in the scrubber and elsewhere in the apparatus.

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3 Results and discussion

Reaction pathways of the ozonolysis of pure MO particles (Fig. 2) have similarities with those of pure oleic acid particles (Ziemann, 2005), although several of the pathways for polymerization are absent (Hung et al., 2005). Methyl oleate reacts with O_3 diffusing

into the particles and forms a primary ozonide (1,2,3-trioxolane). Because this primary ozonide is unstable, it readily decomposes and forms Criegee intermediates (CI) and aldehydes. In one reaction pathway, the CI's isomerize to form carboxylic acids, including *n*-nonanoic acid (NA) and azelaic acid monomethyl ester (AAME). The CI's may also relax from their high vibrational states to form SCI's. For reasons outlined in
 Sect. 3.4, the evidence indicates that relaxation to form SCI's, rather than isomerization to NA or AAME, is the dominant fate of the CI's.

The SCI's can react by several pathways to lead to molecules of MW greater than that of MO. Reaction mechanisms are presented in Fig. 2. Peroxides can form by the reaction of an SCI with aldehydes or carboxylic acids or with another SCI, all of

- which are products of MO ozonolysis. SCI's react with aldehydes to form SOZ's (1,2,4trioxolanes, SOZ1 to 3) or monoperoxide oligomers (CCOOC). The self-reactions of SCI's form diperoxides (1,2,4,5-tetroxane, DP1 to 3) (Zahardis et al., 2005). SCI's also react with protic or nucleophilic molecules to form hydroperoxides, in particular α acyoxyalkyl hydroperoxide products (α AAHP1 to 4) through the reactions of the SCI's
- with the carboxylic acids formed as first-generation products in MO ozonolysis. The hydroperoxides can react further with aldehydes to form α -acyloxyalkyl hydroxyalkyl peroxides (AAHAP's), which have MW's greater than those of the α AAHP's. Rebrovic (1992) proposes that AAHAP's can continue to react with carboxylic acids to lose water and form *bis*(acyloxy-1-alkyl) peroxides (BAAP's), although Nishikawa et al. (1995)
- ²⁵ suggest that a terminal product of AAHAP is more favored. α AAHP, AAHAP, and BAAP compounds are hereafter grouped together under the label " α AAHP-type compounds."

The products can be categorized as having low or high MW's based on the highest molecular weight expected by the products of primary ozonolysis, namely m/z 202

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of azeleic acid monomethyl ester. Low-molecular-weight (LMW) products therefore include *n*-nonanal (NN), 9-oxononanoic acid monomethyl ester (OAME), *n*-nonanoic acid, and azelaic acid monomethyl ester. HMW products are composed of these molecules as subunits and include SOZ's, diperoxides, monoperoxide oligomers, and
 ⁵ αAAHP-type compounds. These HMW products are all part of the peroxide family.

In the following sections, we present results on the ozonolysis and loss of methyl oleate in pure particles (Sect. 3.1), on the effects of ozonolysis on the rates of particle evaporation (Sect. 3.2), on the connections between evaporation and the formation of HMW products (also Sect. 3.2), and on the relative importance in mixed particles of the competing pathways that form SOZ's, α AAHP-type compounds, diperoxides, and monoperoxide oligomers (Sects. 3.3 to 3.5). The relative importance of the competing pathways is assessed by comparing product yields across a suite of experiments in which methyl oleate is mixed at systematic mole fractions with dioctyl adipate (DOA) and myristic acid (MA). Section 3.6 addresses the relative rates of the competing path-

3.1 Ozonolysis and loss of methyl oleate

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Figure 3 shows the decrease in the concentration of methyl oleate (m/z 296) when pure MO aerosol particles are exposed to increasing O_3 . For an O_3 exposure above 1.2×10^{-4} atm s¹, MO disappears nearly completely. The slope of the decay in Fig. 3 yields a reactive uptake coefficient of $1.1 \pm 0.2 \times 10^{-3}$ (Katrib et al., 2005). This value is consistent with a literature value of $1.23 \pm 0.10 \times 10^{-3}$ (Hearn et al., 2005).

The mass spectra recorded for pure MO aerosol particles prior to ozonolysis, as well as those of the particle-phase products after ozonolysis, are shown in Fig. 4. Methyl oleate has characteristic peaks for the parent ion $[MO]^+$ at m/z 296 and for the fragment ions at m/z 206 at m/z 206

²⁵ ment ions at m/z 222 and 264 (Fig. 4a). The peak at m/z 264 corresponds to $[MO-32]^+$, which is the ion formed by the removal of methanol from the methyl ester group. The $[MO-74]^+$ ion at m/z 222 results from a loss of $CH_2=C(OH)OCH_3$ via McLafferty rearrangement of the methyl ester group in the molecule (McLafferty and Tureek, 1993).



The intensities at m/z 222, 264, and 296 decrease and eventually disappear for increasing O₃ exposure (data not shown). Concomitantly, the intensities of the product peaks increase, notably at m/z 141, 143, 152, 155, 171, and 185 (Fig. 4b). In addition, small but non-negligible peaks at m/z 279, 291, 311, 323, and 355 are present above m/z
⁵ 202. These peaks are not attributable to MO fragmentation. The data in Fig. 4b therefore implicate the presence of ozonolysis products having molecular weights greater than 202 (see Sect. 3.3).

- 3.2 Decreased rates of particle evaporation: connections to the formation of High-Molecular-Weight Products
- ¹⁰ The formation of HMW products is inferred from the relative evaporation rates of reacted and unreacted aerosol particles. Prior to ozonolysis, MO particles evaporate at a rate of 4 nm s¹ (Table 1). In comparison, the rate for ozone-reacted particles is 0.15 nm s¹. This evaporation rate is measured after complete MO reaction; therefore, the change results from the evaporation of semivolatile products from the particles.
- ¹⁵ This evaporation rate is twenty times slower than that of AAME (3 nm s⁻¹), which, as a diacid compared to NN, NA, and OAME, is the LMW product having the lowest expected volatility and hence the slowest rate of evaporation. The possible explanation that the slow evaporation rates result from the decreased activities and hence the lower vapor pressures of a mixed chemical system appears insufficient to explain the large
- ²⁰ magnitude of the reduced rate of evaporation. The evaporation rate of 0.15 nm s⁻¹ therefore implies significant inhibition of evaporation, which can be explained by the formation of HMW products having very low vapor pressures. A related inference is that the sum of the volume fractions of the four LMW products in the particles has an upper limit of 5%. This conclusion is based upon an assumed lower limit of 3 nm s¹
- ²⁵ for the rate of evaporation of the LMW products when pure, an assumed behavior of an ideal solution in the organic mixture of products, and an absence of any effects of diffusion or altered partial molar volumes.

Further analysis suggests that more than 60 wt% of the ozonolysis products have

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high molecular weights, as follows. The mass m_p of a spherical particle can be calculated by $m_p = (\pi/6)d_m^3 \rho$ and $\rho = \rho_0 d_{va}/d_m$ (Katrib et al., 2005b). The ratio of particle mass before and after the O₃ exposure is then 0.78, according to the data shown in Table 1 for 0 and 6 s. In this calculation, the density of products is 1.02 g cm^{-3} by use of by $\rho = \rho_0 d_{va}/d_m$, although no correction to d_m is made for up to 1% diameter change arising from the evaporation of MO ozonolysis products between the flow tube and the SMPS. For comparison, by similar analysis, Katrib et al. (2005b) report that the density of products from oleic acid ozonolysis is 1.05 g cm^{-3} . The ratio of 0.78 increases to 0.80 if MO is taken to react completely at 2 s based upon the reaction kinetics so

- that a correction can be made from 2 to 6 s for the evaporation of products; however, this correction is not used in our treatment. The mass ratio of 0.78 can be compared to the expected ratio of 1.16 arising from the stoichiometric addition of ozone in the absence of volatilization. The product mass yield is therefore 67 wt%. We infer that HMW products constitute over 95% of this mass because MO has reacted completely
- and because LMW products constitute an upper limit of 5% by volume of the particles, at least according to our previous inference. A product mass yield of 67 wt% in turn suggests that 33 wt% of the ozonolysis products evaporates in the flow tube. These products may include the LMW products (NN, NA, OAME, and AAME) as well as some higher vapor pressure HMW products.
- 20 3.3 Pathways to secondary ozonides

Formation of SOZ's during MO ozonolysis is supported by the presence of fragment ions at m/z 279, 291, 311, 323, and 355 (Fig. 5a), which arise from SOZ2 (MW 344) and SOZ3 (MW 388) (cf. Fig. 2). M-33 of SOZ2 at m/z 311 forms from the loss of HO₂ (MW 33), which is a common fragmentation pathway for electron-impact ionization of SOZ's (Ziemann, 2005). Methanol (MW 32) is eliminated from methyl esters, yielding a peak at M-32. The peak at m/z 279 is therefore attributed to M-33-32 of SOZ2. By analogy, SOZ3 has peaks at m/z 355 (M-33) and m/z 323 (M-33-32). In addition, SOZ3



has a peak at M-33-32-32 because of the elimination of two methyl-ester groups (cf. SOZ2 and SOZ3 in Fig. 2).

The peak assignments of SOZ's are further confirmed by the ozonolysis of aerosol particles of ethyl oleate. The expected peaks at m/z 279, 291, 325, 337, and 383

- ⁵ are apparent in Fig. 5b, corresponding to ethyl oleate analogues of SOZ2 and SOZ3 (dubbed SOZ2_EO and SOZ3_EO). The loss of ethanol (MW 46) instead of methanol (MW 32) is observed. Fragment ion peaks of SOZ2_EO occur at m/z 279 (M-33-46) and 325 (M-33), and those of SOZ3_EO occur at m/z 291 (M-33-46-46), 337 (M-33-46), and 383 (M-33).
- ¹⁰ The ozone-profile concentrations of SOZ2 and SOZ3, reflected in Fig. 6 by the increasing intensities of peaks at m/z 311 and 323 for higher ozone partial pressure, behave as expected, lending further support to the accuracy of the peak assignments. The leveling off of peak intensity above 2.0×10^{-5} atm O₃ (1.5×10^{-4} atm s⁻¹) corresponds to the reactive depletion of MO (Fig. 3), indicating that further ozone exposure does not affect the SOZ yields. The signal intensity at m/z 185 (Fig. 6b), which arises from the acylium ion [CH₃OCO(CH₂)₇CO]⁺, similarly increases and then levels off for higher ozone concentrations. The acylium ion forms by electron-impact ionization of

SOZ's and other oxygenated products (Ziemann, 2005).

The formation of SOZ3, a cross ozonide, demonstrates that SCI's have a lifetime sufficiently long to diffuse and react with other molecules. A normal ozonide forms from a single parent molecule, as exemplified by SOZ2 from SCI1 + OAME or SCI2 + NN. A cross ozonide, however, requires two parent molecules. For example, the precursors to SOZ3 are SCI2 and OAME, each of which is exclusionary to the formation of the other from a single molecule. Based upon possible molecular combinations, the predicted

relative molar yields of SOZ's are SOZ1:SOZ2:SOZ3 = 1:2:1. This analysis, however, omits any preference towards SCI1 or SCI2 in the cleavage of the primary ozonide and assumes that the reactions of SCI's with NN and OAME are equally favorable. In comparison to the 2:1 prediction, the data instead show that the relative intensities of the SOZ2 peaks compared to those of SOZ3 are 3:1 (Fig. 6a), indicating a correspondingly



higher yield of SOZ2, at least for the assumption of similar ionization efficiencies. In a bulk ozonolysis study of MO, the product yields of SOZ1, 2, and 3 are 9.9%, 66%, and 13%, respectively, based upon HPLC analysis (Wu et al., 1992). Caging of the SCI/aldehyde pair by solvent molecules leads to a greater yield of the normal ozonide 5 SOZ2 by increasing the effective rate constant of SCI1 with OAME and SCI2 with NN

(Loan et al., 1965). The formation of SOZ3 shows that the caging effect is overcome in many reactions, which implies that SCI's diffuse through the matrix prior to reacting with other molecules.

The absence of a change in the relative yield of the cross ozonide compared to the normal ozonide for increasing MO concentration in an inert matrix of dioctyl adipate (DOA) provides further evidence that the caging effect is weak for this system (Fig. 7a). The effectiveness of the caging of the SCI's and the aldehydes formed by the scission of the primary ozonide weakens for increasing solvent polarity (del Rio et al., 1997; Ziemann, 2005). For instance, the normal ozonide from 2-pentene is preferred in the

- ¹⁵ nonpolar solvent pentane because of a strong caging effect (Loan et al., 1965). The relative yields of SOZ2 and SOZ3 are shown in Fig. 7a for an increasing mole fraction of MO in DOA. The yields are calculated from the ratio of the intensities of m/z 311 for SOZ2 and m/z 323 for SOZ3 to the initial intensity of m/z 296 for MO. The SOZ yields and their ratio are constant within measurement uncertainty for MO mole frac-
- tions ranging from 0.04 to 1. The constant SOZ3/SOZ2 ratio (Fig. 7b) suggests that the formation of normal ozonide SOZ2 via caging is of limited importance in a polar solvent of mixed MO and DOA and, moreover, that any changes in this caging effect for changes in the MO/DOA solvent composition are not detectable as variance in the measured product yields. An implication of the experiment is that the yields of SOZ's
- ²⁵ under diluted conditions are as high as in pure particles. We therefore conclude that potential competing pathways, such as unimolecular rearrangement of diffusing SCI's to carboxylic acids, are not significant.

Absent in the results is any evidence for the occurrence of SOZ1 (MW 310; M-33 at m/z 267). This negative finding is consistent with the low product yield reported in



bulk ozonolysis (Wu et al., 1992), suggesting that SOZ1 could be present below our detection limit. An additional possibility is that the products of aerosol particles and bulk studies differ: the small system size of the former allows volatile reactants and products to evaporate. The absence of carboxylic acid groups in SOZ1 implies a higher volatility for it compared to SOZ2 and SOZ3 (Ziemann, 2005). Moreover, volatilization of NN, which is the precursor to SOZ1, could be important, as supported in studies of oleic acid ozonolysis (Moise and Rudich, 2002; Thornberry and Abbatt, 2004). Although Zahardis et al. (2005) report that the SOZ1 forms as an ozonolysis product of oleic acid aerosol particles, Reynolds et al. (2006)¹ do not. Ziemann (2005) suggests that SOZ1 is absent because of high volatility.

3.4 Pathways to α -Acyloxyalkyl-1-Hydroperoxides

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Although Fig. 2 presents a pathway that α AAHP-type compounds could form via the reactions of SCI with the carboxylic acid groups of LMW products, the results shown in Fig. 5 indicate that the pathway to α AAHP-type compounds is in fact insignificant for the case of ozonolysis of pure MO. $[M-33]^+$ is a characteristic fragment ion of α AAHP's 15 (Ziemann, 2005). Peak intensities in Fig. 5a are insignificant for m/z 293 of α AAHP1-33, m/z 327 of α AAHP2a-33 or α AAHP2b-33, and m/z 371 of α AAHP3-33. Moreover, BAAP's and AAHAP's, which are also α AAHP-type compounds, have substructures identical to those of α AAHP's and, if present, should contribute peak intensity at the same m/z values. The accuracy of these peak assignments for α AAHP-type compounds is confirmed by analysis of the products formed by the ozonolysis of mixed AAME/MO particles. AAME, which is one of the LMW products of MO ozonolysis, promotes the formation of α AAHP-type compounds (Fig. 2). For an AAME mole fraction of 0.50 prior to ozonolysis, strong peaks appear at m/z 295, 327, and 371 upon ozonolysis (mass spectra not shown). The intensity of m/z 311 of SOZ2 also decreases, as 25 expected. The peaks are assigned as $[\alpha AAHP2a, b - HO_2 - CH_3OH]^+$ for m/z 295, $[\alpha AAHP2a, b - HO_2]^+$ for m/z 327, and $[\alpha AAHP3 - HO_2]^+$ for m/z 371. The strength of



yield of α AAHP-type compounds under these conditions, establishes an upper limit of 7 wt% yield of α AAHP-type compounds for pure MO particles based upon the minimum detection threshold. The absence of α AAHP-type compounds as products in the ozonolysis of pure MO particles is consistent with previous bulk MO ozonolysis studies ⁵ using ¹H-NMR (Rebrovic, 1992) and HPLC (Nishikawa et al., 1995) for product characterization. In contrast to MO ozonolysis, α AAHP-type compounds are the dominant products in the ozonolysis of oleic acid (Hung et al., 2005; Ziemann, 2005; Zahardis et al., 2006; Reynolds et al., 2006¹). This result can be explained by the chemical differences of MO and oleic acid as solvents, namely the SCI-inert ester group of MO 10 compared to the SCI-scavenging carboxylic acid group of oleic acid.

The formation of α AAHP-type compounds during MO ozonolysis in the presence of carboxylic acids is further investigated by studying product formation in mixed MA/MO aerosol particles (Fig. 8). Myristic acid (MA) is a C₁₄ saturated hydrocarbon terminated by a carboxylic acid group. α AAHP_MA1 (MW 386) and α AAHP_MA2 (MW 430) form ¹⁵ by the reactions SCI1 + MA and SCI2 + MA, respectively. For initial mole fractions of MO less than 0.5, strong peak intensities occur at m/z 335, 347, 353, 365, and 379, corresponding to [α AAHP_MA1 - HO₂ - H₂O]⁺ (e.g., 335 = 386 - 51), [α AAHP_MA2 - HO₂ - CH₃OH - H₂O]⁺, [α AAHP_MA1 - HO₂]⁺, [α AAHP_MA2 - HO₂ - CH₃OH]⁺, and [α AAHP_MA2 - HO₂ - H₂O]⁺, respectively (Fig. 8a). These structures may also result from the breakdown of AAHAP's and BAAP's of MO and MA during AMS analysis. Figure 8b further shows that when α AAHP-type compounds occur in high yield, the yield of SOZ's decreases (Fig. 8b), which can be expected for competitive parallel

The reactive loss of MA is also observed (Fig. 8c). MA itself does not react with ozone so that its loss is explained by reaction with SCI's. For initial MO mole fractions less than 0.4, the magnitude of the decrease in MA appears fully explained by assuming that 100% of SCI's formed by MO ozonolysis reacts with the –COOH group of MA (dashed line in Fig. 8c). The molar product yield of α AAHP-type compounds from MO is therefore also expected to be nearly 100%, and the SOZ molar product yield could

pathways.



approach 0%. For MO mole fractions greater than 0.4, however, MA does not react completely. Furthermore, its fractional loss decreases slightly in tandem with the decreasing yields of α AAHP-type compounds (Figs. 8a and 8c). The implication could be that as the concentration of C=O species increases, such as by increased formation

- ⁵ of LMW products OAME and NN, and as the concentration of COOH species concurrently decreases, there may be a crossover in the fate of the SCI's from α AAHP to SOZ pathways for mixed MA/MO particles of increasing MO mole fraction (Fig. 2). For bulk ozonolysis products of pure MO and MO/NA 1:1 (w/w) mixture, Nishikawa et al. (1995) measure a 89.7 t% yield of α AAHP-type compounds. Such a high yield is consistent with the results presented in Figs. 8a and 8c, although the carboxylic acid used (NA) is
 - different from that used in this study (MA).

The high yield of SOZ's and the absence of α AAHP-type compounds in the ozonolysis of pure MO particles suggest that NA and AAME do not form in high yield. Isomerization of CI's is therefore a minor pathway. Hung et al. (2005) reach the same

- ¹⁵ conclusion for the ozonolysis of oleic acid. The CI's thermalize to form SCI's, which react as shown in Fig. 2 to form the measured products. The present study with MO provides further evidence that CI isomerization is a minor pathway by demonstrating that α AAHP-type compounds readily form when –COOH bonds are made available by preparing mixed AAME/MO or MA/MO. Therefore, if NA or AAME could form from
- ²⁰ isomerization of Cl's in pure MO particles, rapid reaction with available SCl's would take place and α AAHP-type compounds would be observed. Cl isomerization is presumably more favored in the gas phase, for which it is widely reported, than in the condensed-phase because of the longer lifetime for collision deactivation in the case of the former.
- ²⁵ SCI reactions with carboxylic acid groups are an example of a broader reaction mechanism of SCI's with protic or nucleophilic molecules to form hydroperoxides. For example, ozonolysis of MO particles prepared by atomization from methanol and without a charcoal scrubber in place leads to a change in the mass spectra (data not shown). Peaks appear at m/z 157 and 201, which are attributed to the $[M - HO_2]^+$ ions



of $CH_3(CH_2)_7CH(OOH)OCH_3$ and $CH_3OCO(CH_2)_7CH(OOH)OCH_3$. These alkoxy hydroperoxides form by the reaction between SCI's and residual methanol.

3.5 Pathways to diperoxides and monoperoxide oligomers

The detected high-mass fragments provide no direct evidence for the presence of diperoxides or monoperoxide oligomers (CCOOC) (Fig. 5a). For example, DP1, 2, and 3 have molecular weights of m/z 325, 360, and 404, yet no significant intensity is observed at these m/z values or at other m/z values that might correspond to the loss of methanol, water, or other small fragments during electron-impact ionization.

The absence of direct evidence notwithstanding, diperoxides or monoperoxide oligomers could form but go undetected. They are thermally unstable and therefore may pyrolyze to LMW molecules on the 250°C vaporizer surface of the AMS (Leiva et al., 2004; Ziemann, 2005). In this case, the LMW molecules, rather than the diperoxides or monoperoxide oligomers, would be detected by the subsequent electron-impact ionization and quadrupole mass filtering of the AMS. Another possibility is that the DP's may form but volatilize from the particles and therefore go undetected, which is an explanation analogous to that put forward for the absence of SOZ1 in the detected products (Sect. 3.3).

In the MA/MO experiments, for increasing MO mole fraction there is a corresponding positive linear trend in the yield of SOZ's and a negative nonlinear trend in the yield of α AAHP-type compounds (Fig. 8). This relationship implies the formation of additional products from the SCI's. Under the assumption that the mechanism shown in Fig. 2 is complete, possible fates of the SCI's include diperoxides, monoperoxide oligomers, and the LMW products NA and AAME. Our analysis of the collected mass spectra does not further discriminate among these possibilities.

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3.6 Assessment of the relative importance of competing reactions

The relative importance of competing pathways in the ozonolysis of mixed MA/MO particles is assessed using a model. An assumption is made of steady-state concentrations for the SCI's. The modeled reactions are as follows:

$${}_{5} SCI's + \{LMW aldehydes\} \xrightarrow{k_{1}} SOZ's$$
 (R1)

SCI's + MA
$$\xrightarrow{\kappa_2}$$
 α AAHP-type compounds (R2)

The reaction mechanism omits a pathway for the formation of LMW carboxylic acids (AAME and NA) because the data imply that these products are minor (Sect. 3.2). Possible formation of diperoxides and monoperoxide oligomers is also not directly modeled (see below).

The yields of α AAHP-type compounds in this competitive kinetics scheme are shown in Fig. 9 as curves corresponding to different assumed ratios $k_2:k_1$. The superimposed solid squares show the sums of the signal intensities m/z 335, 347, 353, 365, and 379, which arise from α AAHP-type compounds. The intensities are scaled to 100% yield for an MO mole fraction of 0.15. Comparison of the solid squares to the curves constrains $k_2:k_1$ between 0.3 and 3 for MO mole fractions less than 0.5. For higher mole fractions, the apparent decrease in $k_2:k_1$ may occur because the reaction mechanism of R1 and R2 is incomplete. For example, the formation of diperoxides is second-order in SCI's.

4 Conclusions and atmospheric implications

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²⁰ The ozonolysis of methyl oleate in pure and mixed submicron aerosol particles to form high-molecular-weight organic peroxides is investigated. Two secondary ozonides are identified as major HMW ozonolysis products in pure MO particles and in particles of MO diluted in an inert matrix of DOA. α -acyloxyalkyl peroxides are the dominant



products when MO is ozonolyzed in mixed particles having high concentrations of – COOH groups (e.g., myristic acid in this study). The reactions of the SCI's for carboxylic groups to form αAAHP-type compounds compete with those of the SCI's for LMW aldehydes to form SOZ's, provided that both types of functionalities are present at significant concentrations. Although the mass spectra do not indicate the presence of diperoxides or monoperoxide oligomer products, their formation is nevertheless not ruled out because pyrolysis or evaporation is possible prior to analysis, and, given the constraints of mass balance, their formation is inferred as plausible.

Although atmospheric particles are of much greater chemical heterogeneity than those of the model particles in our study, certain common elements remain, and the SCI reactivity observed in this study suggests possible reaction pathways of ozone with unsaturated organic molecules to yield HMW peroxides in atmospheric aerosol particles. Mainly, the intermolecular reaction of stabilized Criegee intermediates with carboxylic acid and other protic or nucleophilic moieties may compete with the forma-

- tion of secondary ozonides and other products. Furthermore, in regard to SCI reactivity, both the unreacted and the reacted model particles of our study have important chemical similarities to atmospheric particles (Table 2). The concentration of double bonds in atmospheric particles ranges from 12 to 28 mole percent, which is several times greater than the 0.2 to 5.4% of the unreacted model particles. The concentrations
- C=O and C-OH in atmospheric particles ranges from 1 to 8% to 1 to 5%, respectively. Carboxyl acids (–COOH) may account for a significant portion of these C=O and C-OH groups. The mixed MA/MO particles have comparable –COOH concentrations. For example, the concentration is 2.1% for MA/MO particles of 0.40 mole fraction. No-tably, αAAHP-type compounds become important HMW products for mole fractions of
- 0.40 and greater. An additional chemical factor is particle liquid water content, which is nearly zero in the experiments of this study at 0% relative humidity whereas it can be substantial in atmospheric particles (Dick et al., 1996). SCI's can react with water, having the effect of favoring LMW over HMW products. Pryer et al. (1991), for example, report that H₂O₂ and aldehydes evolve for MO reacted in aqueous emulsions: the SCI



- + H₂O yields a hydroxy hydroperoxide that hydrolyzes to form the observed products.
 Ozone concentration and particle residence time must also be considered in regard to extending the laboratory results to possible processes in the atmosphere. The ozone mixing ratio in the troposphere of 10 to 100 ppb is approximately 1000 times
- ⁵ less than that of 50–70 ppm employed in many parts of this study (Figs. 5 and 7–9). Slower ozonolysis rates in atmospheric particles could suggest that steady-state SCI concentrations are lower than in the model particles, which would imply that reactions second-order in SCI, such as the formation of diperoxides, are less favored. The formation of α AAHP-type compounds, secondary ozonides, and monoperoxide oligomers,
- ¹⁰ however, remains favorable provided that abundant COOH, –C-OH, and –C=O functionalities are present in atmospheric particles. Based upon analysis offered above, α AAHP-type compounds could become favored to secondary ozonides and monoperoxide oligomers once the ratio of concentrations of –COOH to –C=O exceeds a critical ratio. A related implication of these reaction pathways is that the direct ozonolysis of unsaturated organic molecules can lead to the indirect processing of saturated organic
- molecules having carboxylic, carbonyl, and hydroxyl groups.

The residence time of particles in the atmosphere, which is one to two weeks, is much longer than the 6s of most of our experiments, which has important implications for the steady-state concentration of possible HMW peroxides in atmospheric particles.

- Over a time period of days, some HMW peroxides could react further, including lysing into LMW products or alternatively further oligomerization to higher molecular weights. For instance, diperoxides thermally decompose to form LMW aldehydes (Leiva et al., 2004). Peroxides also react by photochemical and acid-catalyzed aqueous pathways (Ziemann, 2005). SOZ's undergo acid-catalyzed hydrolysis to form LMW aldehydes
- and carboxylic acids (Bernatek and Ledaal, 1960). Some reactions, however, can also increase MW. For example, as a result of thermal decomposition, SOZ's and diperoxides can initiate chain oligomerization by free-radical pathways (Ewing et al., 1989; Rimmer and Ebdon, 1995).

The results of this study inform on the factors affecting the relative importance of

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several reaction pathways that could yield HMW peroxides from the reactions of ozone with unsaturated organic molecules in atmospheric particles. Future investigations to further advance these results should focus on the chemical characteristics of atmospheric particles, such as the partitioning of organic molecules among phases (Griffin
 et al., 2003), the chemical morphology of particles (Katrib et al., 2005a), and the product yields affected by increased reaction times and lower ozone concentrations.

Appendix A

Derivation of Equation 1

¹⁰ The weight percent of MO (w_{MO}) is related to the relative contribution of m/z 296 of the MO molecular ion to the mass spectrum, as follows. The signal intensity m/z of mixed particles ($I_{mix}^{m/z}$) is the sum of the signal intensities from molecules of methyl oleate ($I_{MO/mix}^{m/z}$) and other molecules in the matrix ($I_{matrix/mix}^{m/z}$): $I_{mix}^{m/z} = I_{MO/mix}^{m/z} + I_{matrix/mix}^{m/z}$. This relationship extends to the sum of intensities across m/z 0 to 390, omitting m/z values 15 of air beams (e.g., m/z 28):

$$\sum_{m/z} I_{mix}^{m/z} = \sum_{m/z} I_{MO/mix}^{m/z} + \sum_{m/z} I_{matrix/mix}^{m/z}.$$
 (A1)

Under the assumption that all signals of a fragmentation pattern lie within the AMS measurement domain of m/z 0 to 390, the sum of AMS signal intensities are proportional to mass (Katrib et al., 2004). Therefore, an RIE-weighted mixing rule holds that $\left(\sum_{m/z} I_{MO/mix}^{m/z} / RIE_{MO}\right) : \left(\sum_{m/z} I_{matrix/mix}^{m/z} / RIE_{mat}\right) = w_{MO}:100 - w_{MO}.$ Rearrangment



of this equation leads to:

$$\sum_{m/z} I_{matrix/mix}^{m/z} = \frac{(100 - w_{MO}) R I E_{matrix}}{w_{MO} R I E_{MO}} \sum_{m/z} I_{MO/mix}^{m/z}$$
(A2)

Substituting Eq. (A2) into Eq. (A1), we obtain:

$$\sum_{m/z} I_{mix}^{m/z} = \frac{w_{MO}RIE_{MO} + (100 - w_{MO})RIE_{matrix}}{w_{MO}RIE_{MO}} \sum_{m/z} I_{MO/mix}^{m/z}$$
(A3)

5 which upon rearrangement leads to:

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$$I_{\rm mix}^{296} / \sum_{m/z} I_{\rm mix}^{m/z} = \frac{w_{\rm MO} R I E_{\rm MO}}{w_{\rm MO} R I E_{\rm MO} + (100 - w_{\rm MO}) R I E_{\rm matrix}} \left(I_{\rm mix}^{296} / \sum_{m/z} I_{\rm MO/mix}^{m/z} \right)$$
(A4)

The matrix substances have negligible intensity at m/z 296, which allows the accurate approximation $I_{\text{mix}}^{296} / \sum_{m/z} I_{\text{MO/mix}}^{m/z} \approx I_{\text{MO}}^{296} / \sum_{m/z} I_{\text{MO}}^{m/z}$. The right-hand quantity is a constant determined in a separate AMS measurement using pure MO particles. This substitution yields:

$$I_{\rm mix}^{296} / \sum_{\rm m/z} I_{\rm mix}^{\rm m/z} = \frac{w_{\rm MO} R I E_{\rm MO}}{w_{\rm MO} R I E_{\rm MO} + (100 - w_{\rm MO}) R I E_{\rm matrix}} \left(I_{\rm MO}^{296} / \sum_{\rm m/z} I_{\rm MO}^{\rm m/z} \right)$$
(A5)

Rearrangement of Eq. (A5) yields Eq. (1):

$$w_{\rm MO} = \frac{100 R/E_{\rm matrix} \left(I_{\rm mix}^{296} / \sum_{\rm m/z} I_{\rm mix}^{\rm m/z} \right)}{R/E_{\rm MO} \left(I_{\rm MO}^{296} / \sum_{\rm m/z} I_{\rm MO}^{\rm m/z} \right) + (R/E_{\rm matrix} - R/E_{\rm MO}) \left(I_{\rm mix}^{296} / \sum_{\rm m/z} I_{\rm mix}^{\rm m/z} \right)}$$
(A6)
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⁵ this material are those of the authors and do not necessarily reflect the views of the National Science Foundation.

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Table 1. Mobility diameter (nm) for increasing time. Mobility diameters at 0 and 6 are estimated from measurements of the vacuum aerodynamic diameters (within parentheses) by the AMS and use of the relationship $d_m = (\rho_0 / \rho_{MO}) d_{va}$, whereas mobility diameters at 27 and 45 s are measured by the SMPS. Average values of three experiments, which agree within 1 nm, are shown for pure MO particles exposed to no ozone and for particles composed of products after complete MO ozonolysis. In the absence of ozone, the particle diameter decreases because of evaporation of MO. In the presence of ozone, MO is entirely ozonized by 27 s so that the diameter decreases recorded at 27 and 45 s arise from the evaporation of semivolatile ozonolysis products. ^aResidence times of 27 s or 45 s are based upon 6 s in the flow tube, 2 or 20 s in the tubing between the flow tube and the DMA, 5 s in the DMA inlet, and an effective 14 s within the electric field of the DMA. The actual residence time within the electric field is approximately 28 s, yet the particles are evaporating during this time so that the measured effective mobility diameter is a convolution of the entering and exiting diameters and the flow field within the DMA. We estimate that this temporal mean diameter corresponds to one-half the residence time (14 s).

	Time (s)			
	0	6	27 ^a	45 ^a
no ozone exposure	500	477	425	356
	(437)	(417)	(–)	(–)
ozone exposure of 5×10^{-5} atm O ₃	_	_	438	434
	(–)	(446)	(–)	(–)

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Table 2. Chemical functional groups present in atmospheric particles collected in East Asia compared to those present prior to ozonolysis in the model aerosol particles of this study. Values are shown in mole percent. In the last column, the alkene ozone exposure is given. For the ambient aerosol, 50 ppb O_3 is assumed. ^aRanges for ten samples presented by Maria et al. (2003) based upon infrared analysis. ^bLower limit of methyl oleate concentration of this study. ^cValues in parentheses are the fractions of the carbonyl bonds in carboxylic acid groups.

	[C=O] (%)	[C-OH] (%)	[Alkene] (%)	[Alkene]×P _{O3} (10 ⁻⁹ atm)
ambient aerosol	1–8 ^a	1–5 ^a	12–28 ^a	6–14 ^a
methyl oleate	2.7 (0) ^c	0	5.4	3.2×10 ³
4 mol% methyl oleate in DOA ^b	2.3 (0) ^c	0	0.2	1.3×10 ²
40 mol% methyl oleate in myristic acid	3.2 (2.1) ^c	2.1	0.5	3.2×10 ²

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Fig. 1. Schematic diagram of the experimental setup. Key: DMA, differential mobility analyzer; CPC, condensation particle counter; SMPS, scanning mobility particle sizer; TMP, turbo molecular pump; QMS, quadrupole mass spectrometer; MFC, mass flow controller; MFM, mass flow meter; CO, critical orifice; NV, needle valve.

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Fig. 2. Proposed reaction pathways for the ozonolysis of pure methyl oleate. Formation pathways of HMW organic peroxides are shown, including secondary ozonides (SOZ's), α -acyloxyalkyl peroxides (α AAHP's), diperoxides, and monoperoxide oligomers. Stereoisomers are not presented separately. The reaction mechanism follows that introduced by Ziemann (2005) for the ozonolysis of oleic acid. For reference, the figure also shows the structures of the matrix substances DOA and MA. Dilution of MO in DOA or MA affects the relative contribution of the each pathway to the product yields.

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Fig. 4. Mass spectra of methyl oleate aerosol particles (a) before and (b) after ozone exposure. The particle mobility diameter prior to ozonolysis is 500 nm. The O_3 exposure is 5×10^{-5} atm for 6 s. $I^{m/z}$ is the intensity recorded when tuned to m/z, and $\Sigma I^{m/z}$ denotes the intensity sum for m/z 0 to 390, omitting m/z values of air beams (e.g., m/z 28).





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Fig. 5. Mass spectra for m/z 270 through 390 amu after the ozonolysis of **(a)** methyl oleate and **(b)** ethyl oleate aerosol particles. The particle mobility diameter prior to ozonolysis is 500 nm. The ozone exposure is 5×10^{-5} atm for 6 s. The structures of the SOZ molecules are shown in Fig. 2.

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Fig. 6. Appearance of products for increasing ozone partial pressure in the ozonolsyis of pure methyl oleate. **(a)** Signal intensities m/z 311 and 323 of SOZ fragment ions. **(b)** Signal intensity m/z 185 of the acylium ion $[CH_3OCO(CH_2)_7CO]^+$. The peak intensities are normalized to the intensity of m/z 296 prior to ozonolysis. The O₃ concentration for which methyl oleate reacts to near completion is shown by the vertical line. Dashed lines are guides to the eye. The particle mobility diameter prior to ozonolysis is 500 nm. The exposure time is 6 s.

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Fig. 7. Signal intensities for increasing mole fraction of methyl oleate in DOA after O_3 exposure of 5×10^5 atm for 6 s. (a) Signal intensities m/z 311 and 323 of SOZ products normalized to the initial intensity of m/z 296 (i.e., the MO molecular ion). (b) Ratio of the signal intensity of m/z 323 to that of m/z 311. Dashed lines are guides to the eye. The particle mobility diameter prior to ozonolysis is 500 nm.





Fig. 8. Signal intensities for increasing mole fraction of methyl oleate in MA after O₃ exposure of 5×10^{-5} atm for 6 s. (a) Signal intensities m/z 335, 347, 353, 365, and 379 of α AAHP-type compounds normalized to the initial intensity of m/z 296 (MO molecular ion). (b) Signal intensities m/z 311 and 323 of SOZ products normalized to the initial intensity of m/z 296. (c) Signal intensity m/z 228 of the MA molecular ion normalized to its initial intensity. The dashed curve in *c* shows the calculated ratio I_{0}^{228}/I_{0}^{228} for the upper limit that 100% of SCI's formed by MO ozonolysis reacts with the –COOH group of MA. Dotted lines are guides to the eye. The particle mobility diameter prior to ozonolysis is 500 nm.







