Kinetics and Product Yields of the OH Initiated Oxidation of Hydroxymethyl Hydroperoxide


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Abstract

Hydroxymethyl hydroperoxide (HMHP), formed in the reaction of the C₁ Criegee intermediate with water, is among the most abundant organic peroxides in the atmosphere. Although reaction with OH is thought to represent one of the most important atmospheric removal processes for HMHP, this reaction has been largely unstudied in the laboratory. Here, we present measurements of the kinetics and products formed in the reaction of HMHP with OH. HMHP was oxidized by OH in an environmental chamber; the decay of the hydroperoxide and the formation of formic acid and formaldehyde were monitored over time using CF₃O⁻ chemical ionization mass spectrometry (CIMS) and laser induced fluorescence (LIF). The loss of HMHP by reaction with OH is measured relative to the loss of 1,2-butanediol \[ k_{\text{1,2-butanediol+OH}} = (27.0 \pm 5.6) \times 10^{-12} \text{ cm}^3 \text{ molecule}^{-1} \text{s}^{-1} \]. We find that HMHP reacts with OH at 295 K with a rate coefficient of \((7.1 \pm 1.5) \times 10^{-12} \text{ cm}^3 \text{ molecule}^{-1} \text{s}^{-1}\), with the formic acid to formaldehyde yield in a ratio of \(0.88 \pm 0.21\) and independent of NO concentration \((3 \times 10^{10} - 1.5 \times 10^{13} \text{ molecule cm}^{-3})\). We suggest that, exclusively, abstraction of the methyl hydrogen of HMHP results in formic acid while abstraction of the hydroperoxy hydrogen results in formaldehyde. We further evaluate the relative importance of HMHP sinks and use global simulations from GEOS-Chem to estimate that HMHP oxidation by OH contributes 1.7 Tg yr⁻¹ (1-3%) of global annual formic acid production.
Introduction

Hydroperoxides significantly contribute to the chemistry of the atmosphere due to their high reactivity. These species alter the atmosphere’s oxidative potential by acting as a reactive sink and transported reservoir of HO$_x$.\textsuperscript{1–3} They act as oxidants of SO$_2$ in the aqueous phase to produce SO$_4^{2–}$, thereby reducing air quality and visibility.\textsuperscript{4,5} In addition, hydroperoxides have been implicated in the inhibition of certain peroxidase enzymes essential to plant function,\textsuperscript{6,7} although some studies note that under certain conditions exposure to ozone can increase plant resistance to oxidative stress from hydroperoxides.\textsuperscript{8,9}

Hydroxymethyl hydroperoxide (HOCH$_2$OOH, HMHP) is among the hydroperoxides observed in significant abundance in the atmosphere. Reported concentrations of HMHP vary considerably, but typically fall in the low ppbv range during the summer and have been reported up to 5 ppbv over forested regions.\textsuperscript{10–13} Recently, HMHP concentrations were measured during the SEAC4RS (Studies of Emissions, Atmospheric Composition, Clouds, and Climate Coupling by Regional Surveys) flight campaign traversing across the southeastern United States between August 6 and September 23, 2013. HMHP mixing ratios varied considerably depending on location and altitude, but within the boundary layer (as determined by the NASA Airborne UV Differential Absorption Lidar (DIAL) instrument) the average HMHP mixing ratio was 0.25 ppbv with a maximum of 4.0 ppbv (Figure 1).

HMHP forms when terminal alkenes react with ozone in the presence of water vapor.\textsuperscript{14–20} Upon attack by O$_3$, the alkene fragments into a carbonyl and an energy-rich intermediate, which may be collisionally stabilized to form the C$_1$ Criegee intermediate (CH$_2$OO). The C$_1$ Criegee intermediate then reacts primarily with water vapor monomer or dimer (n=1,2) to form HMHP as the dominant product:

$$\text{CH}_2\text{OO} + (\text{H}_2\text{O})_n \rightarrow \text{HOCH}_2\text{OOH} + (\text{H}_2\text{O})_{n-1}$$ (1)

Atmospherically prevalent alkenes with terminal double bonds such as isoprene and β-pinene
as well as simpler alkenes such as ethene, propene, 1-butene, etc. contribute to HMHP formation (e.g. Figure 1). Reported maximum yields of HMHP from O₃ + alkene reactions typically fall around 0.4 for ethene, 0.14 for propene, 0.15 for 1-butene, and 0.15 for 1-pentene under humid conditions.²¹,²² Nguyen et al.²³ found that the maximum yield from isoprene is 0.44 at an RH of ∼40% and that the yield of HMHP from CH₂OO + H₂O is strongly dependent on RH; they attribute this finding to differences in the product distribution between reaction with water monomer and water dimer. Further investigation into the kinetics and products of CH₂OO with water dimer by Sheps et al.²⁴ shows that HMHP is formed in greater than 50% yield from this reaction.

Understanding the relative rates of production and removal mechanisms of HMHP is key to assessing its lifetime and importance in the atmosphere. HMHP undergoes three major atmospheric removal processes: photolysis, deposition, and reaction with the hydroxyl radical. HMHP photodissociation likely behaves like that of other hydroperoxides, proceeding via cleavage of the O–O peroxy bond and thereby recycling the oxidant pool. However,
numerous studies investigating the spectroscopic properties of HMHP have concluded that photolysis is likely to be less significant than other atmospheric loss processes (photolysis rate of $J \sim 1 \times 10^{-6}$ s$^{-1}$ under typical atmospheric conditions).\textsuperscript{25–29} On the other hand, deposition contributes to efficient removal of HMHP. HMHP is highly water-soluble (Henry’s Law constant of $H \sim 10^6$ M atm$^{-1}$),\textsuperscript{30} and is therefore highly susceptible to rainout and other wet deposition processes. Nguyen et al.\textsuperscript{13} estimate an HMHP lifetime of 10 hours against dry deposition at a forested site in the southeastern United States and suggest that deposited HMHP may react heterogeneously to produce formic acid.

Similar to other peroxides, reaction with OH is expected to be an important sink of HMHP. However, the rate coefficient for this reaction has not been previously reported. Three possible H-abstraction pathways exist for HMHP oxidation: abstraction of (a) the hydroperoxidic hydrogen, (b) the alkyl hydrogen, and (c) the alcoholic hydrogen. These channels lead to the formation of formic acid (HCOOH) or formaldehyde (HCHO) with OH or HO$_2$ radicals as byproducts, respectively. Francisco and Eisfeld\textsuperscript{31} performed a theoretical calculation of HMHP + OH and concluded that pathway (b) dominates the reactivity as they find that this pathway has the lowest reaction barrier. However, no experimental evidence has been reported to test these conclusions.

In this study, we investigate the reaction of HMHP with OH. HMHP was oxidized by OH in an environmental chamber, and the decay of the hydroperoxide was monitored over time using CF$_3$O$^-$ chemical ionization mass spectrometry (CIMS). Product yields of formic acid and formaldehyde were characterized by CIMS and by laser induced fluorescence (LIF), respectively. The yield of formic acid was further characterized under varying NO concentrations to assess the HMHP + OH oxidation pathways. HMHP sinks and the impact of HMHP oxidation on global formaldehyde and formic acid concentrations are interpreted in the context of simulations using GEOS-Chem to evaluate the global importance of HMHP oxidation.
Experimental Methods

Instrumentation

Chemical Ionization Mass Spectroscopy (CIMS) is a versatile and robust technique for detecting a variety of atmospheric compounds, including hydroperoxides. Reagents and oxidation products in this work were monitored using a compact time-of-flight CIMS (ToF-CIMS, Tofwerk/Caltech) that employs a CF$_3$O$^-$ reagent ion for sensitive detection of gas-phase organic acids and multifunctional organic compounds. The CF$_3$O$^-$ CIMS technique has been described in detail in Crounse et al.$^{32}$, Paulot et al.$^{33}$, and St. Clair et al.$^{34}$.

Briefly, the reagent ions form by passing 380 sccm of 1 ppm CF$_3$OOCF$_3$ in N$_2$ through a cylindrical ion source containing a layer of radioactive polonium-210 (NRD LLC, $\leq$10 mCi). The sample air is diluted with dry N$_2$ (1750 sccm) in a Pyrex glass flow tube with a hydrophobic coating (Fluoropel 801A, Cytonix) that is maintained at a pressure of 35 mbar. The diluted sample air then mixes with the reagent ions, which selectively ionize analytes by forming ion clusters ($m/z =$ analyte mass + 85) or fluoride transfer ions ($m/z =$ analyte mass + 19), the dominance of which depends on the acidity and fluoride affinity of the target analyte. Product ions are transferred through a pinhole orifice and a conical hexapole ion guide to the time-of-flight mass spectrometer chamber. Compounds are separated in the mass spectrometer based on differences in their mass-to-charge ratio as they accelerate through the instrument.

The ToF-CIMS provides 10 Hz resolution data for masses between $m/z$ 19 and $m/z$ 396. In this study, HMHP was monitored at $m/z$ 149 (HMHP·CF$_3$O$^-$), the relative rate partner, 1,2-butandiol, was monitored at $m/z$ 175 (1,2-butandiol·CF$_3$O$^-$), formic acid (FA) was monitored at $m/z$ 65 (FA$_{H^+}$HF), and bis-HMP (bis-hydroxymethyl peroxide, HOCH$_2$OOCH$_2$OH) was monitored at both $m/z$ 113 (bis-HMP$_{H^+}$HF) and $m/z$ 179 (bis-HMP·CF$_3$O$^-$). All observed ion signals were normalized to the sum of the reagent anion signal ($^{13}$CF$_3$O$^-$ isotope at $m/z$ 86) and the water signal ($m/z$ 104, H$_2$O·$^{13}$CF$_3$O$^-$ isotope) to
account for fluctuations in the reagent ion concentration. Detection limits for the ToF-CIMS are typically 10 pptv for a 1 second integration period.

In addition to the ToF-CIMS, other instruments were employed to assess concentrations of formaldehyde, NO, and O\textsubscript{3} over the course of the experiment. Formaldehyde product yields were characterized by the NASA In Situ Airborne Formaldehyde (ISAF) instrument.\textsuperscript{35} Briefly, the ISAF instrument employs a pulsed tunable fiber laser for LIF detection of HCHO. The laser operates at 353 nm, exciting a single rotational transition of the A–X band in HCHO. The instrument has a 10 Hz sampling frequency that is averaged to 1 second, at which the precision is typically better than 20\% above 100 pptv. NO\textsubscript{x} and O\textsubscript{3} concentrations throughout the experiment were monitored with a NO\textsubscript{x} monitor (Teledyne 200EU) and an O\textsubscript{3} monitor (Teledyne 400E). All instruments sampled the chamber from an approximately 1 m loop of 0.635 cm OD PFA tubing.

**Synthesis**

A new method for synthesizing HMHP was performed in this study. The method is based on a technique described in Bauerle and Moortgat\textsuperscript{26}, in which formaldehyde vapor is passed through hydrogen peroxide to generate HMHP. Here, HCHO was prepared by gently heating crystalline paraformaldehyde (Sigma-Aldrich) and passing the resulting vapor through two successive cryotrapes at -65 °C and -196 °C to collect impurities and condense HCHO, respectively. A small (∼10 sccm) flow of N\textsubscript{2} was then passed over the collected HCHO, held at -65 °C, and bubbled through urea hydrogen peroxide (Sigma-Aldrich, 97\%, 1100 mg) in dichloromethane (DCM, 30 mL) held in a -30 °C cold bath. After approximately 5 minutes of flow, the -30 °C bath was removed and the reaction mixture allowed to warm to room temperature. HMHP formed from this method in a relative yield of approximately 4:1 HMHP to bis-HMP and with small amounts of HCHO and H\textsubscript{2}O\textsubscript{2} also present (as determined by gas-phase analysis of an evaporated droplet of the synthetic mixture). Note that previous synthesis using a similar method found significant safety hazards upon concentration of the
hydroperoxide product (e.g. Fry et al. 27).

Chamber Experiments

Experiments on HMHP + OH oxidation were conducted in a small environmental chamber. The chamber consisted of a 1 m$^3$ fluorinated ethylene propylene copolymer bag (Teflon-FEP, DuPont) within an enclosure equipped with UV lights (8 Sylvania 350 blacklights), and has been described previously. 36–38 The chamber was prepared by multiple flushes of dry air between successive experiments. Experiments were performed at ambient laboratory temperature (295 ± 2 K) and pressure (745 Torr). Table 1 provides a description of experimental conditions.

Table 1: Summary of conditions used in HMHP oxidation experiments. Mixing ratios are given in ppbv.

<table>
<thead>
<tr>
<th>Exp.</th>
<th>[HMHP]$_0$</th>
<th>[OH source]$_0$*</th>
<th>[1,2-BD]$_0$</th>
<th>initial [NO]$_0$</th>
<th>% HMHP Ox.</th>
<th>Objective</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>150</td>
<td>190</td>
<td>80</td>
<td>475</td>
<td>35%</td>
<td>Kinetics</td>
</tr>
<tr>
<td>2</td>
<td>90</td>
<td>200</td>
<td>15</td>
<td>460</td>
<td>40%</td>
<td>Kinetics</td>
</tr>
<tr>
<td>3</td>
<td>35</td>
<td>190</td>
<td>15</td>
<td>160</td>
<td>55%</td>
<td>Kinetics</td>
</tr>
<tr>
<td>4</td>
<td>165</td>
<td>100</td>
<td>—</td>
<td>445</td>
<td>30%</td>
<td>Yields</td>
</tr>
<tr>
<td>5</td>
<td>55</td>
<td>45</td>
<td>—</td>
<td>20</td>
<td>45%</td>
<td>NO$_x$ dep.</td>
</tr>
<tr>
<td>6</td>
<td>70</td>
<td>30</td>
<td>—</td>
<td>500</td>
<td>30%</td>
<td>NO$_x$ dep.</td>
</tr>
<tr>
<td>7</td>
<td>20</td>
<td>20</td>
<td>—</td>
<td>20</td>
<td>35%</td>
<td>NO$_x$ dep.</td>
</tr>
<tr>
<td>8</td>
<td>10</td>
<td>60</td>
<td>—</td>
<td>25</td>
<td>50%</td>
<td>NO$_x$ dep.</td>
</tr>
<tr>
<td>9</td>
<td>20</td>
<td>120</td>
<td>—</td>
<td>530</td>
<td>40%</td>
<td>NO$_x$ dep.</td>
</tr>
</tbody>
</table>

*OH source was methyl nitrite for kinetics experiments and isopropyl nitrite for the yields experiment.

Reagents were added to the chamber sequentially after flushing the chamber bag with dry air and filling it to near 50% with zero air. NO (1993 ± 20 ppmv in N$_2$, Matheson) was prepared by filling an evacuated 500 cm$^3$ glass bulb to the desired pressure and backfilling with N$_2$ before adding to the chamber. Next, 1,2-butandiol (1,2-BD, ≥ 98%, Sigma-Aldrich) was added as a relative rate partner by flowing 20 L min$^{-1}$ dry air over a small drop of the diol placed in a glass vial. 1,2-butandiol was chosen as a relative rate partner because it is detectable by the CF$_3$O$^-$ CIMS technique and has a known OH reaction rate constant
that is expected to be similar to that of HMHP + OH. A method similar to that outlined in Taylor et al. was used to synthesize the HO source used in this study, methyl nitrite (CH$_3$ONO). Approximately 200 ppbv of CH$_3$ONO was added to the chamber via serial dilution in a 500 cm$^3$ glass bulb. Finally, HMHP was added to the chamber by first cryo-collecting the sample to remove high volatility impurities generated from the synthesis. Zero air was passed for 10-30 seconds over a three way vial containing approximately 0.5–1.2 mL of synthesized HMHP in DCM and HMHP was subsequently trapped in a cold bath at -80 °C. Upon removal of the bath, the cryotrapped sample was flushed into the chamber with zero air for approximately 20 minutes until the remainder of the chamber volume was filled. H$_2$O$_2$ and HCHO were present in the chamber in minor amounts (∼5% and ∼2% relative to HMHP, respectively) from the HMHP synthetic mixture described above.

Photooxidation was initiated after stabilization of the CIMS signals (15 to 50 min). The UV lights were turned on to generate OH radicals via

$$\text{CH}_3\text{ONO} + h\nu \rightarrow \text{HO}_2 + \text{NO} + \text{HCHO}$$

This process produced OH concentrations that were typically ∼50 times greater than average atmospheric levels. Oxidation lasted until the OH precursor was depleted (∼1 hour), utilizing 3 of the chamber’s UV lights. Approximately 30–50% of HMHP was oxidized.

The chamber was prepared in a very similar manner for all experiments, with a few notable exceptions. For Exps. 4–9, 1,2-butanediol was not added to the chamber to minimize any interference in the product yield due to oxidation of this species. For Exp. 4, in which HCHO was measured, isopropyl nitrite was used as the OH source to preclude HCHO interference from CH$_3$ONO photolysis. In addition, a series of experiments were conducted to assess the NO$_x$ dependence of the formic acid yield. In these experiments, a further step was taken to purify HMHP from the reaction mixture. The solvent and high volatility
impurities were removed first by either flowing zero air over the reaction mixture at -80 °C or by placing the reaction mixture under vacuum. To isolate HMHP from the lower volatility bis-HMP synthetic byproduct, the remaining reaction mixture was collected in a cold trap and HMHP was eluted at a temperature of -15 °C.

**Calibration**

A gravimetric technique was used to calibrate the ToF-CIMS for formic acid. A commercially available formic acid standard (Sigma-Aldrich, 98%) was dissolved in water (1% w/w solution) and volatilized into the chamber by flowing a known amount of dry air over the sample until it had completely evaporated. The dry and water-dependent sensitivities were determined by addition of varying concentrations of water vapor to the sample before it entered the CIMS instrument. This water-vapor calibration was applied to the ToF-CIMS formic acid signal during analysis. Because there is no commercially available standard, the absolute sensitivity of HMHP could not be determined. Using the synthetic sample, a calibration for the change in HMHP sensitivity based on water vapor was performed in a manner similar to that of formic acid. This relative calibration was used in analysis of the ToF-CIMS HMHP signals. The bis-HMP ToF-CIMS sensitivity relative to HMHP was estimated from ion-molecule collision rates, which were parameterized from the calculated dipole moment and polarizability of the neutral molecules. See the Supplemental Information for further details of the ToF-CIMS calibration procedures. HCHO instrument sensitivity for ISAF was determined via calibration against standard additions of a commercially available HCHO standard to zero air. See Cazorla et al. for further details of the ISAF calibration procedure.

**Theoretical Methods**

Reaction rate constants for the unimolecular reactions are calculated with the approach by Møller et al. using multi-conformer transition state theory with Eckart tunneling. The
electronic energies are calculated using coupled cluster methods in Molpro2012, while zero-
point vibrational energy corrections and partition functions are calculated using density
functional theory in Gaussian 09. Rice-Ramsperger-Kassel-Marcus (RRKM) modeling of
selected reactions is done using the Master Equation Solver for Multi-Energy well Reactions
(MESMER) and the MultiWell program suite. See Supplemental Information for details.

Results and Discussion

HMHP + OH Rate Coefficient

The HMHP + OH reaction rate coefficient ($k_{\text{HMHP}}$) relative to that of 1,2-butanediol + OH
($k_{\text{diol}}$) was determined at ambient temperature using data from experiments 1–3 (Table 1).
The rate coefficient of 1,2-butanediol with OH is $(27.0 \pm 5.6) \times 10^{-12}$ cm$^3$ molecule$^{-1}$s$^{-1}$
at $T = 296$ K. To obtain the rate constant for HMHP + OH relative to that of 1,2-
butanediol + OH, the natural logarithm of the HMHP mixing ratio (normalized to the
initial concentration) was plotted as a function of the natural logarithm of the normalized
1,2-butanediol mixing ratios over the course of oxidation (Figure 2). The slope of a linear
regression analysis incorporating error in both dimensions gives the reaction rate of HMHP
relative to 1,2-butanediol ($k_{\text{HMHP}[\text{OH}]} / k_{\text{diol}[\text{OH}]}$) for each experiment. Table 2 lists the
relative rates $k_{\text{HMHP}} / k_{\text{diol}}$ and gives a recommended rate constant for the OH oxidation of
HMHP (calculated by taking a mean of all experimental runs weighted by their respective
uncertainties).

The uncertainty for all experiments is dominated by the 20% uncertainty in the 1,2-
butanediol rate constant. Other appreciable sources of error arose from the loss of HMHP
on the walls of the chamber, equilibration of the ToF-CIMS signals, and the error in the
linear fit. To account for uptake of HMHP to the chamber walls, a first-order loss rate of
$(0.8 \pm 0.2) \times 10^{-5}$ s$^{-1}$ was used in the data analysis (5% correction to the HMHP data).
This rate was determined by filling the chamber with 12 ppbv of HMHP and monitoring
the decay of the signal in the dark. The signal from 1,2-butanediol was also corrected for minor wall loss (0.5% correction to data). In addition, HMHP mixing ratios were corrected for a minor loss due to photolysis \((J = 8.5 \times 10^{-7} \text{ s}^{-1})\), calculated from measured light flux in the chamber and from quantum yields and cross sections in Sander et al.\(^5\); total correction of \(<1\%\)). Data from \(t<15\) and \(t>45\) minutes into the oxidation was not used in Figure 2 or to evaluate the kinetics of HMHP + OH (5% correction to the rate constant). The first few minutes of oxidation were disregarded to minimize error due to equilibration of sampling surfaces, such as chamber and tubing walls. At long oxidation times, most of the OH precursor had been depleted, causing photochemical reactions to slow and making relative loss to the chamber walls substantial.

Figure 2: Fit (black) of the natural logarithmic decay of HMHP vs 1,2-butanediol (diol) during oxidation. HMHP and 1,2-butanediol data are measured at ToF-CIMS signals \(m/z\) 149 and \(m/z\) 175, respectively, and averaged over 30 s between 15 and 45 min into the oxidation. The relative decay is used to determine the HMHP + OH oxidation rate.

**HMHP Oxidation Products**

Experiments to determine the yields of formic acid and HCHO from HMHP oxidation were conducted in a manner similar to that of the kinetics experiments. A constant correction factor of \(0.10\times[\text{HMHP}]\) was subtracted from all HCHO signals to account for an estimated
Table 2: Relative rate of HMHP + OH to 1,2-butanediol + OH \((k_{\text{HMHP}}/k_{\text{diol}})\) at 295 K for Experiments 1–3 and derived absolute HMHP + OH rate coefficient \((10^{-12} \text{ cm}^3 \text{ molecule}^{-1} \text{s}^{-1})\).

<table>
<thead>
<tr>
<th>Experiment 1</th>
<th>Experiment 2</th>
<th>Experiment 3</th>
<th>Rate Coefficient</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.262 ± 0.008</td>
<td>0.275 ± 0.011</td>
<td>0.253 ± 0.015</td>
<td>7.1 ± 1.5</td>
</tr>
</tbody>
</table>

Uncertainties are 1σ standard deviations from measurement uncertainties; the rate coefficient also includes error in the 1,2-butanediol + OH rate coefficient.

HMHP decomposition in the ISAF HCHO instrument (see the Supplemental Information for more details). Note that a more thorough investigation of the conversion of HMHP to HCHO on ISAF instrument surfaces, as was done for ISOPOOH conversion to HCHO, would provide more confidence in the magnitude of the conversion rate. HMHP and formic acid were corrected for wall loss by applying the experimentally-derived wall loss rates to the data. Under the experimental conditions, formic acid wall loss is comparable to wall production, the sum of which is minor compared to total formic acid production from HMHP (~ 2.5%). The loss of formic acid and HCHO due to the reaction with OH was accounted for as described by Eq. VI of Atkinson et al. (1.5% for FA and up to 15% for HCHO; see the Supplemental Information). HCHO was also corrected for loss due to photolysis \((J = 1.7 \times 10^{-5} \text{ s}^{-1}\), calculated from measured light flux in the chamber and from quantum yields and cross sections in Sander et al.; up to 5% correction). As previously discussed, bis-HMP was present in the chamber during the oxidation experiment. Observed formic acid mixing ratios were corrected for bis-HMP + OH production of formic acid, assuming that bis-HMP oxidation produces 2 equivalents of formic acid. This process is calculated to produce up to 40% of total formic acid, using a bis-HMP + OH rate coefficient that is 35% that of HMHP + OH as determined from the kinetics experiments (see the Supplemental Information).

The ratio of the formic acid to the HCHO yield was found in Exp. 4 by comparing the change in these species during the oxidation period and using a linear regression that accounts for error in both dimensions. These yields are assessed from data taken between
Figure 3: Production of formic acid (FA) compared with that of HCHO. Formic acid was measured on the ToF-CIMS at $m/z$ 65 and HCHO data are from ISAF during Experiment 4. The signals are corrected for losses outlined in the text, and averaged over 30 s between 15 and 45 min into the oxidation. The black line indicates the best fit to the data.

t=15 minutes and t=45 minutes into the oxidation, to minimize error due to equilibration of sampling surfaces and to loss on chamber walls. The results of the linear regression analysis is presented in Figure 3. The reaction of HMHP with OH produces formic acid and HCHO in comparable amounts with a formic acid to HCHO product ratio of 0.88 ± 0.21. The reported uncertainty arises from the errors in the calibration of the ToF-CIMS data and from corrections due to HMHP interference in ISAF, product loss due to reaction with OH, and bis-HMP + OH production of formic acid as outlined above, as well as from uncertainty in the line of best fit.

To assess carbon closure from HMHP + OH, an upper limit to the HMHP sensitivity was estimated using the yield experiment. As discussed previously, the absolute sensitivity of the ToF-CIMS to HMHP is not well-known. Assuming that formic acid and HCHO are the only two products from HMHP + OH, the total change in HMHP mixing ratios over the experiment was assumed to be equal to the change in the sum of the two products. This mixing ratio was then used to derive an upper limit to the expected HMHP sensitivity on the ToF-CIMS. Using this sensitivity, we calculate HMHP yields from ethene ozonolysis.
experiments conducted in the 1 m³ chamber (see the Supplemental Information). Within error, these yields are the same as those reported by Hasson et al.²¹, consistent with formic acid and HCHO as the only major products from the reaction of HMHP with OH.

Figure 4: Mechanism of HMHP gas-phase oxidation by OH. HMHP oxidation may proceed via one of three pathways: (a) abstraction of the hydroperoxidic hydrogen, (b) abstraction of the methyl hydrogen, and (c) abstraction of the alcoholic hydrogen. In theory, pathway (a) may further bifurcate depending on the NO concentration.

The formic acid yield was also evaluated as a function of NO mixing ratio, which was varied between a few and more than 500 ppbv [\(\sim (0.003-1.5) \times 10^{13}\) molecules cm\(^{-1}\)]. As shown in Figure 4, formic acid can form following abstraction of the methyl, alcoholic, or hydroperoxy hydrogens. Abstraction of the hydroperoxide H (pathway (a) in Figure 4) leads to the formation of the hydroxy peroxy radical (HOCH\(_2\)OO). This radical likely forms with excess energy from the abstraction and may react unimolecularly to lose HO\(_2\) before undergoing collisional thermalization. Following stabilization, the radical has two possible subsequent reaction paths: unimolecular thermal decomposition to HO\(_2\) and HCHO or bimolecular reaction with NO forming the hydroxy alkoxy radical (HOCH\(_2\)O), which then decomposes to H and formic acid.²⁵,³¹,⁵³,⁵⁴ The 1,4-H shift of HOCH\(_2\)OO leading to a hydroperoxy alkoxy radical (OCH\(_2\)OOH) has a calculated rate constant of only 3 \(\times 10^{-7}\) s\(^{-1}\) and is therefore disregarded (see Supplemental Information). Like pathway (a), abstraction of the alcoholic
H of HMHP (pathway (c) in Figure 4) leads to the formation of the HOCH$_2$OO radical due to a favorable 1,4-H shift; however, this pathway is expected to be minor in comparison to pathways (a) and (b) due to the difference in known ROH vs. ROOH abstraction rates.$^{55}$

Results from these experiments indicate that there is no obvious dependence of the formic acid yield on the amount of NO present (Figure 5). The initial concentration of NO used in these experiments is listed in Table 1; note that NO concentrations in the chamber generally decrease from this value as the oxidation proceeds. The imprecision in the measured yield is a result of the experimental challenges described above. These include the need to accurately describe the wall loss of the hydroperoxides and formic acid, as well as errors in accounting for formic acid production from the oxidation of bis-HMP.

![Figure 5: Correlation of experimental formic acid (FA) yields with derived uncertainty against initial NO mixing ratio. The formic acid yields do not appear to exhibit a dependence on NO, indicating that the unimolecular HO$_2$-loss rate is faster than expected from literature reports of thermalized HOCH$_2$OO. These yields are compared with those predicted by a box model using a range of unimolecular HOCH$_2$OO decomposition rates.](image)

The lack of dependence of the formic acid yield on [NO] shows some discrepancy with the predicted yield from reported unimolecular HOCH$_2$OO decomposition rates. The decompo-
position rate of HO\(_{\text{CH}_2\text{OO}}\) has been measured to be quite slow, between 1.5 – 140 s\(^{-1}\) at T = 298 K (see Table 3). Even at the fastest experimentally-determined decomposition rate, reaction of the peroxy radical with NO would be expected to be competitive (pseudo first-order rate of 120 s\(^{-1}\) at highest NO concentrations) with the unimolecular decomposition. As a result, the formic acid yield would be expected to depend on NO, such that higher NO concentrations would predict greater formic acid yield. Shown in Figure 5 are kinetic box model calculations of the predicted formic acid yield when using unimolecular HO\(_2\)-loss rates of 25 s\(^{-1}\), 50 s\(^{-1}\), 100 s\(^{-1}\), 200 s\(^{-1}\), 400 s\(^{-1}\), and 800 s\(^{-1}\) (see Supplemental Information). The lack of clear dependence on [NO] suggests that the unimolecular decomposition in these experiments occurs at a rate of greater than a few hundred per second at T = 295 K.

**Table 3: Reported rates of HO\(_{\text{CH}_2\text{OO}}\) decomposition**

<table>
<thead>
<tr>
<th>Study</th>
<th>Method</th>
<th>Rate (s(^{-1}))</th>
<th>Notes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Su et al., 1979(^{25})</td>
<td>FTIR</td>
<td>1.5</td>
<td>Kinetic simulations with experimental data</td>
</tr>
<tr>
<td>Veyret et al., 1982(^{56})</td>
<td>FTIR</td>
<td>30</td>
<td>Kinetic simulations with experimental data</td>
</tr>
<tr>
<td>Barnes et al., 1985(^{57})</td>
<td>FTIR</td>
<td>20</td>
<td>Measured loss of HO(_2) by proxy (HO(_2)NO(_2))</td>
</tr>
<tr>
<td>Veyret et al., 1989(^{58})</td>
<td>UV</td>
<td>125</td>
<td>Kinetic fits to loss of HO(_2)</td>
</tr>
<tr>
<td>Burrows et al., 1989(^{59})</td>
<td>UV/FTIR</td>
<td>140</td>
<td>Kinetic simulations of measured K(_{eq})</td>
</tr>
<tr>
<td>Hermans et al., 2005(^{60})</td>
<td>MC-TST*</td>
<td>200</td>
<td></td>
</tr>
<tr>
<td>Morajkar et al., 2013(^{61})</td>
<td>cw-CRDS</td>
<td>55</td>
<td>Measured loss of HO(_2)</td>
</tr>
<tr>
<td>This study</td>
<td>MC-TST*</td>
<td>440</td>
<td>See Supplemental Information</td>
</tr>
</tbody>
</table>

*Multifconformer transition state theory*

The difference between the rate for unimolecular loss of HO\(_2\) inferred in this study with those previously reported (Table 3) likely reflects some combination of experimental error and differences in the initial energy distribution of the HO\(_{\text{CH}_2\text{OO}}\) radicals. Most of the measured rates shown in Table 3 were determined by observing the loss of HCHO, HO\(_2\), or the formation of the HO\(_{\text{CH}_2\text{OO}}\) radical in the presence of excess formaldehyde both with and without NO\(_x\) present. Morajkar et al.\(^{61}\), for example, invert the time dependence of HO\(_2\) to diagnose two rate coefficients: the initial rapid loss of HO\(_2\) is used to assess the rate of formation of HO\(_{\text{CH}_2\text{OO}}\) (the inverse of the unimolecular decomposition) and the
second subsequent and much slower loss is used to infer the equilibrium coefficient; the proper assignment of the HO₂ dynamics is thus complicated. Theoretical calculations of HOCH₂OO decomposition performed in this study suggest a significantly faster decomposition rate of 440 s⁻¹ at 298 K (see Supplemental Information).

The rate of decomposition of HOCH₂OO is also likely sensitive to how this species is formed. In the studies shown in Table 3, HOCH₂OO is produced cold via the association reaction of HCHO + HO₂. In contrast, when HOCH₂OO forms as a result of HMHP oxidation, some of the reaction exothermicity is likely deposited in the peroxy radical, speeding the rate of decomposition. RRKM simulations of the experimental system (see Supplemental Information) suggest that, provided that less than 8 kcal mol⁻¹ of the exothermicity deposits into H₂O, all of the HOCH₂OO will decompose to formaldehyde and HO₂ before collisional stabilization. Hence, we interpret the lack of NO-dependence of the formic acid yields as implying that the HOCH₂OO formed from HMHP oxidation by OH decomposes at a rate sufficiently high such that formaldehyde is the only product following abstraction of the hydroperoxide hydrogen, both for conditions of this experiment and those relevant in the atmosphere. Therefore formic acid production from HMHP + OH reaction in the atmosphere originates exclusively from abstraction of the methyl hydrogen.

**Atmospheric Implications**

**Atmospheric Fate of HMHP**

Consideration of the three major atmospheric loss processes for HMHP (deposition, OH reaction, and photolysis) allows for the estimation of its total lifetime and the relative contribution of each loss mechanism. For example, HMHP fluxes and OH concentrations were measured in the southeastern United States during the Southern Oxidant and Aerosol Study (SOAS) in summer 2013. During the campaign, the diurnal-average OH concentration was typically around 1×10⁶ molecules cm⁻³, but on some days peaked at levels more than twice as large.²² Our measured HMHP + OH rate coefficient produces a lifetime with respect to
oxidation by OH, $\tau_{OH}$, of between 15 and 40 hours. By comparison, the diurnal-average of the cloud-free atmospheric photodissociation rate at ground level is calculated by the Tropospheric Ultraviolet-Visible (TUV) radiation model (NCAR/ACD) to be $1.8 \times 10^{-6}$ s$^{-1}$. This value gives an HMHP lifetime of about one week against photolysis in the boundary layer. Using the dry deposition velocity of HMHP measured by Nguyen et al.\textsuperscript{13} during SOAS (4 cm s$^{-1}$) and an assumed mixed layer depth of 1.5 km, the lifetime of HMHP with respect to dry deposition is 10 hours. For these conditions, oxidation by OH accounts for between approximately 20–40% of HMHP loss.

**Global Modeling**

To investigate the global importance of HMHP chemistry, we simulate the production and fate of HMHP using the chemical transport model GEOS-Chem. GEOS-Chem is a three-dimensional model of tropospheric chemistry driven by assimilated meteorological observations from the NASA Goddard Earth Observing System (GEOS).\textsuperscript{63} The model includes isoprene oxidation chemistry,\textsuperscript{64} which has been extensively updated to reflect recent mechanistic studies.\textsuperscript{23,36,37,65–67} We have updated the GEOS-Chem mechanism to include HMHP yields from alkene ozonolysis taken from Neeb et al.\textsuperscript{68}, Hasson et al.\textsuperscript{22}, and Nguyen et al.\textsuperscript{23}, as well as HMHP loss due to deposition from Nguyen et al.\textsuperscript{13}, photolysis based on Roehl et al.\textsuperscript{28}, and OH oxidation from the results presented in this study. We have assumed that the HMHP + OH rate coefficient exhibits the same temperature dependence as the reaction of its homologue methyl hydroperoxide (MHP, CH$_3$OOH) with OH. The simulations reported here were conducted for the year 2014 on a global 4° × 5° latitude by longitude grid, following a 1-year model spin up, and use model version 10-01 with GEOS-FP meteorology.

HMHP forms in substantial quantities in regions with large biogenic VOC emissions. Globally, HMHP has a total annual production of 12.4 Tg yr$^{-1}$, with 8.6 Tg yr$^{-1}$ of that from the approximately 7.5% of isoprene that reacts with ozone. The annually averaged boundary layer ($z = 0–0.5$ km) HMHP mixing ratios are typically around 0.1 ppbv, but
reach up to 0.5 ppbv in the heavily forested regions of South America and Africa where isoprene emissions are largest (Figure 6A). In the southeastern United States, the GEOS-Chem predicted average HMHP mixing ratios for the summer of 2013 in the boundary layer are around 0.3 ppbv (see the Supplemental Information). By comparison, the average HMHP mixing ratio observed in the boundary layer in the southeastern United States during the SEAC4RS flight campaign in summer 2013 was 0.25 ppbv and reached as high as 4.0 ppbv
As products of HMHP oxidation, global concentrations and distributions of formic acid and HCHO are altered when HMHP is included in the model. The model predicts that 40% of HMHP is lost to OH, thereby producing 1.7 Tg yr$^{-1}$ of formic acid, with the remaining loss due to deposition (52%) and photolysis (7%). The magnitude of the OH oxidation pathway is substantial in the lower atmosphere; in certain locations up to 25% of local formic acid mixing ratios is simulated to arise from the oxidation of HMHP (Figure 6B). However, the global production of formic acid from HMHP + OH is small compared with previous GEOS-Chem budget estimates of 51 Tg yr$^{-1}$ of formic acid from photochemical oxidation and compared with estimates of 100-120 Tg y$^{-1}$ from observations.$^{69,70}$ Instead, HMHP acts as an intermediate species, producing formic acid further from emissions sources and resulting in higher formic acid mixing ratios in remote areas such as over the Atlantic Ocean. In contrast, the mixing ratios of HCHO are not significantly altered by inclusion of HMHP, as the model predicts only 0.1% of the total global annual HCHO production is due to HMHP oxidation, peaking at 1% of total local production in regions with high HMHP (see the Supplemental Information). Note that the mixing ratios of HMHP presented in Figure 6A are likely an underestimate as two important HMHP precursors, ethene and β-pinene, are not explicitly treated in the model. In addition, the contribution of HMHP oxidation to global formic acid concentrations is likely also a lower estimate. In these simulations, we assume that dry deposition represents a permanent loss of carbon from the atmosphere. If instead this process results in a flux of formic acid as suggested by Nguyen et al.$^{13}$, this would further increase the atmospheric concentrations of this carboxylic acid from HMHP.

Conclusions

The reaction rate constant from the HMHP + OH reaction shows that OH oxidation is a major loss process for HMHP in the atmosphere. Both OH reaction and, likely, dry deposition
of HMHP lead to the formation of formic acid, a notable point given that models currently underestimate the concentration of atmospheric formic acid compared with measurements. Studies such as Paulot et al.\textsuperscript{71} and Millet et al.\textsuperscript{70} indicate that measured summertime boundary layer concentrations of formic acid can be more than double the model predicted values. These authors suggest that the discrepancy necessitates a 2–3 times larger source of formic acid than models currently contain, most likely in the form of formic acid production from secondary chemistry of biogenic and other chemical precursors. The results of this study further constrain the formic acid budget from a hydroperoxide that forms in the oxidation of a variety of biogenic and anthropogenic precursors, and show that formic acid production from HMHP oxidation is not enough to account for the large discrepancy between models and observations. Additionally, given the importance of deposition to HMHP loss and potential for formic acid formation, this work highlights the need for improved understanding of surface chemistry.

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Supporting Information Available

The following files are available free of charge.

- Calibration procedures
- MC-TST and RRKM modeling details
- GEOS-Chem modeling details

This material is available free of charge via the Internet at http://pubs.acs.org/.

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Graphical TOC Entry