

High Gas-Phase Methanesulfonic Acid Production in the OH-Initiated Oxidation of Dimethyl Sulfide at Low Temperatures

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ABSTRACT: Dimethyl sulfide (DMS) influences climate via cloud condensation nuclei (CCN) formation resulting from its oxidation products (mainly methanesulfonic acid, MSA, and sulfuric acid, H_2SO_4). Despite their importance, accurate prediction of MSA and H₂SO₄ from DMS oxidation remains challenging. With comprehensive experiments carried out in the Cosmics Leaving Outdoor Droplets (CLOUD) chamber at CERN, we show that decreasing the temperature from +25 to -10 °C enhances the gas-phase MSA production by an order of magnitude from OH-initiated DMS oxidation, while H₂SO₄ production is modestly affected. This leads to a gas-phase H₂SO₄-to-MSA ratio (H_2SO_4/MSA) smaller than one at low temperatures, consistent with field observations in polar regions. With an updated DMS oxidation mechanism, we find that methanesulfinic acid, CH₃S(O)OH, MSIA, forms large amounts of MSA. Overall, our results reveal that MSA yields are a factor of 2–10 higher than



those predicted by the widely used Master Chemical Mechanism (MCMv3.3.1), and the NO_x effect is less significant than that of temperature. Our updated mechanism explains the high MSA production rates observed in field observations, especially at low temperatures, thus, substantiating the greater importance of MSA in the natural sulfur cycle and natural CCN formation. Our mechanism will improve the interpretation of present-day and historical gas-phase H₂SO₄/MSA measurements.

KEYWORDS: dimethyl sulfide (DMS), OH-initiated oxidation, methanesulfonic acid (MSA), methanesulfinic acid $(CH_3S(O)OH, MSIA)$, low temperatures

INTRODUCTION

Dimethyl sulfide (DMS) is emitted into the atmosphere by marine bacteria and as a result of the degradation of dimethylsulfoniopropionate (DMSP) produced from phytoplankton.¹⁻³ These emissions are the most abundant biological source of sulfur,³ contributing between 18 and 42% of the global atmospheric sulfate aerosol.⁴ Sulfur-containing oxidation products from DMS-specifically, sulfuric acid (H₂SO₄) and

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methanesulfonic acid (CH₃S(O)(O)OH, MSA) that formed in the gas phase and condensed phase—take part in the formation of sulfur-containing aerosols and cloud condensation nuclei (CCN) in the marine atmosphere.^{3,5–7} They initially form small (ca. nanometer-sized) molecular clusters through nucleation,^{8–13} after which they grow by further condensation to CCN sizes. They thus affect the formation, optical properties, and lifetime of marine clouds,⁸ influencing cloud radiative properties and therefore climate.^{5,14–16}

The ratio of particulate MSA to non-sea-salt sulfate varies between 0.05 and 0.75 and is usually below 0.5.^{6,7,17,18} Understanding how gaseous MSA contributes to this ratio is important in understanding the impact of DMS oceanic emissions leading to the formation of low-volatility species via oxidation. The multiphase chemical mechanism is complex, and the yields of H₂SO₄ and MSA depend on temperature¹⁹ as well as atmospheric composition. Moreover, field measurements of gas-phase MSA and H₂SO₄ show a wide range of concentrations.^{12,20–23}

Atmospheric MSA is formed primarily from DMS oxidation, with high concentrations observed in polar regions.¹² Atmospheric H_2SO_4 is formed from sulfur dioxide (SO₂) oxidation, which comes from anthropogenic and volcanic emissions as well as DMS oxidation. It follows that the ratio of H_2SO_4 to MSA (H_2SO_4/MSA) is an important indicator of both regional variations and anthropogenic perturbation in both current and historical samples. Furthermore, both species may participate in new particle formation^{8–13} and CCN formation processes together with iodine oxoacids and ammonia in polar regions.^{24,25} Consequently, a quantitative, mechanistic understanding of the DMS oxidation mechanism(s), especially the resulting H_2SO_4/MSA , is essential to further our understanding of their roles in climate change.

DMS oxidation has been studied extensively with a wide array of instrumentation. However, chamber and flow-tube experiments have had difficulties reproducing DMS oxidation under atmospherically representative conditions, leaving us with significant gaps in our chemical understanding. Recent experimental work and in situ field observations have shown that autoxidation plays an important role in the oxidation of DMS at higher temperatures (>0 $^{\circ}$ C), proceeding through the formation of hydroperoxymethyl thioformate (HPMTF; HOOCH₂SCHO) via isomerization of the methylthiomethylperoxy radical (CH₃SCH₂OO, MSP), which is the primary product of DMS hydrogen abstraction.²⁹⁻³² However, subzero conditions (<0 °C) are challenging to investigate experimentally and so have received less attention. Hydrogen abstraction, both by OH and via isomerization (autoxidation), is suppressed at a lower temperature, and the OH addition pathway is known to dominate, as shown in previous studies.³³⁻³⁵ Finally, wall reactions can be problematic in the laboratory, and their effects on the DMS oxidation experiments have been unclear so far.²⁷

Model studies have addressed the importance of halogen chemistry^{36–39} and aqueous-phase processes^{28,37,38,40} to close the gaps between the modeled and measured oxidation products, suggesting reduced importance for OH-initiated DMS oxidation. However, even with recent revisions based on experimental findings, models with current state-of-the-art gas-phase mechanisms underestimate ambient MSA.^{37,40} Here, we show that those mechanisms overestimate H₂SO₄/MSA by a factor of 2–10 throughout the ambient temperature range. The mechanism adopted in this study is similar to the gas-

phase mechanism in the study of Wollesen de Jonge et al.,⁴¹ which revised the gas-phase chemistry (MCM) to explain their high particulate MSA.

We performed experiments in the CLOUD chamber at CERN¹³ under conditions that closely match the marine boundary layer (MBL) with an extensive suite of instruments. We investigated the gas-phase OH-initiated DMS oxidation over a wide temperature range $(-10 \text{ to } +25 \text{ }^{\circ}\text{C})$ and present a revised mechanism describing gas-phase DMS oxidation. In addition to H_2SO_4 and MSA, we quantified numerous intermediate products, including the key intermediate methanesulfinic acid ($CH_3S(O)OH$, MSIA), and used them to evaluate and constrain the mechanism. In addition to the gas-phase chemistry, we quantified semiempirical rate coefficients for the heterogeneous formation of the important products dimethyl sulfoxide (CH₃S(O)CH₃, DMSO) and dimethyl sulfone $(CH_3S(O)_2CH_3, DMSO_2)$. We designed our experiments such that the particle condensation sink (CS) is negligible. In CLOUD and most experiments, these multiphase processes occur on the chamber walls,^{27,41} but they can also occur on or within aerosols and cloud droplets in the atmosphere. We investigated the effects of temperature and NO_x on gas-phase OH-initiated DMS oxidation. Both our experiments and our revised oxidation mechanism closely match ambient gas-phase H_2SO_4/MSA , and together they provide a quantitative understanding of this critical natural biogeochemical process.

MATERIALS AND METHODS

CLOUD Chamber and Experiments. The CLOUD chamber is an electropolished stainless steel cylinder with a volume of 26.1 m³. The chamber is surrounded by an insulated thermal housing, which maintains high-temperature uniformity and stability, and it can be operated over a wide range of temperatures (from -70 to +100 °C) and relative humidity (from below 1% to above 90%). Ultrapure synthetic, humidified air (cryogenic 79% N₂ and 21% O₂) and a slight overpressure minimize contaminants in the chamber. All trace gases (e.g., SO₂, O₃, CO, NH₃, NO_x, and DMS) have independent gas lines connected to the chamber to avoid reactions caused by mixing in the gas lines before injecting into the chamber. Dry nitrogen was used to dilute DMS, NO₂, and O₃ from standard high-concentration gas bottles before reaching the chamber to obtain close to atmospheric concentrations.

Four 200 W Hg-Xe UV lamps (UVH LC8, Hamamatsu Photonics K.K., Japan) at wavelengths between 250 and 450 nm with adjustable power and a xenon fluoride excimer laser (UVX) at 248 nm were used to generate OH radicals via photolysis of ozone (O_3) . The distribution of UV lamps and laser in the chamber make the gas-phase oxidation products (e.g., OH) form uniformly in a few minutes. An LED sabre (LS3) at 385 nm was used to photolyze NO_2 producing NO. Sulfur dioxide (SO_2) was measured with a high-sensitivity pulse fluorescence analyser (model 43i-TLE; Thermo Fisher Scientific), ozone with a UV photometric ozone analyser (model 49C; Thermo Environmental Instruments), carbon monoxide using a nondispersion cross-modulation infrared analysis method (model APMA-370; Horiba), and nitrogen oxides NO with an ECO Physics CLD 780TR and NO₂ with a Cavity Enhanced Differential Optical Absorption Spectroscopy (CE-DOAS). The SO₂ monitor employed at CLOUD was unfortunately not able to measure the formed SO₂ from DMS

oxidation because of its high detection limit (0.5 ppb). A chilled dew-point mirror (EdgeTech Instruments) and a direct tunable diode laser absorption spectrometer (TDL hygrometer, Werle et al.⁴²) were used to continuously monitor the water vapor concentration.

The OH-initiated DMS oxidation experiment was conducted as follows: Before the oxidation experiments, the mixing fan speed was set to 100% to enhance turbulent mixing in the chamber, thus reducing the concentrations of oxidation products due to faster wall loss. The injection rates of ozone, DMS, and NO_x were set to reach the desired values. The experiment was then initiated by switching the UV lights on and setting the fan speed to 12% to initiate DMS oxidation (by OH radicals) and to reduce wall loss rates of DMS oxidation products to establish a new steady-state condition. Temperature-ramping experiments were conducted at two temperature ranges (+25 to +10 and +10 to -10 °C) with different initial concentrations of DMS, OH, and CO. The temperature ramps took a few hours each, which is only slightly longer than the residence time in the chamber. Therefore, unlike in a typical oxidation experiment, steady states were not reached for each temperature in the temperature-ramping experiments. As such, H₂SO₄-to-MSA ratios in these experiments should not be used directly in other contexts because they are not measured in steady state. They can, however, be used for investigating the temperature dependence by simulating the full concentration profile at each point in time and then comparing the model results with the experimental results.

Measurements of DMS and its Oxidation Products. State-of-the-art instruments were operated simultaneously to measure the gas-phase concentrations of DMS and its oxidation products. They were measured by a suite of advanced mass spectrometers including a nitrate-ion-based chemical ionization mass spectrometer (NO₃⁻-CIMS), a bromide chemical ionization mass spectrometer coupled with a multischeme chemical ionization inlet (Br⁻-MION-CIMS), the gas-phase channel of a bromide chemical ionization mass spectrometer equipped with a Filter Inlet for Gases and AEROsols inlet (Br⁻-FIGAERO-CIMS), selective reagent ionization mass spectrometers (SRI-TOF-MS), and protontransfer reaction time-of-flight mass spectrometer (H₃O⁺-CIMS, NH_4^+ -CIMS). The experiments were grouped into three sets depending on the availability of instruments and temperature. We conducted experiment set 1 at -10 °C, experiment set 2 at +10 and -10 °C, and experiment set 3 at +25 and +10 °C, with relative humidity ranging from 20 to 70%. NO3--CIMS and PTR3 worked during all of the experiments. However, the mode of PTR3 was changed from a regular H_3O^+ mode to NH_4^+ mode in experiment set 3. Br⁻-MION-CIMS measurements were only available in experiment set 1 and Br--FIGAERO-CIMS measurements were only in experiment set 2. The conditions of the experiments and relevant instruments are listed in Table S1.

DMS. A proton-transfer reaction time-of-flight mass spectrometer (H_3O^+ -CIMS) provided the DMS concentration. The instrument is based on the design of the proton-transfer reaction time-of-flight mass spectrometer (PTR3) described in Breitenlechner et al.⁴³ DMS concentrations were calibrated with a gas standard at specific conditions between experiments. The concentrations of DMS for the second experiment set were provided by the selective reagent ionization mass spectrometer (SRI-TOF-MS) described in detail by Canaval et al.⁴⁴ We calibrated both instruments regularly between

experiments with a standard gas mixture containing multiple volatile organic compounds to account for any possible drifts in transmission efficiency or absolute humidity dependency.

H₂SO₄, MSA, and CH₃S(O)₂OOH. The concentrations of H_2SO_4 , MSA, and $CH_3S(O)_2OOH$ were measured with a nitrate-ion-based chemical ionization mass spectrometer (NO₃⁻-CIMS; Tofwerk AG, Thun, Switzerland; Jokinen et al.⁴⁵). The specially designed inlet for chemical ionization at the ambient pressure system and its calibration and quantification procedures are well described by previous studies.^{46,47} We applied the same calibration coefficient $C_{\rm H,SO_4}$ = 4.13×10^{10} cm⁻³ per normalized signal (cps cps⁻¹; cps denotes counts per second) for the experiments carried out at +10 and -10 °C since charging efficiency does not vary significantly in this temperature range. The uncertainty for NO₃⁻-CIMS is mainly caused by H₂SO₄ calibration, below \sim 50%; this uncertainty includes the systematic error from calibration setup and statistic error. All uncertainties for each species from the different instruments are listed in Table S2. We estimate that MSA has a collision-limited charging efficiency and strong binding energy with less fragmentation than H₂SO₄ based on the cluster binding enthalpy calculated by quantum chemical methods (see Text S1.3 and Tables S3 and S4 for details). The measurement sensitivity of methanesulfonic peroxide $(CH_3S(O)_2OOH)$ is lower than the maximum sensitivity, and the given concentrations represent its lower limits. The detailed estimation for MSA and $CH_3S(O)_2OOH$ can be found in Text S1.3.

DMSO, DMSO₂, CH₃SCHO, and CH₃SOH. The concentrations of DMSO, DMSO₂, methyl thioformate (CH₃SCHO), and methanesulfenic acid (CH₃SOH) were measured either by an H₃O⁺- or NH₄⁺-CIMS.^{48°} Different ionization schemes $(H_3O^+- \text{ or } NH_4^+-\text{chemical ionization})$ were used in different experiments as shown in Table S1. We determined the instrument's collision limit calibration factor by ionizing 1 ppbv of hexanone from a gas standard diluted in air. Applying the collision limit calibration factor to all compounds ensures lower-limit estimates for their concentrations. As shown in Figure S2, the independence of the concentration ratios determined by both ionization methods (H₃O⁺- and NH₄⁺chemical ionization) with the collision limit calibration factor on sample gas humidity and temperature suggests that both DMSO and DMSO₂ are very likely ionized at the collision limit in both ionization modes. The lower-limit estimates are therefore most likely their true concentrations. The comparison between the two ionization methods at the same time can be realized with additional experiments performed with high ammonia concentration in the chamber. These additional experiments have much higher DMS, O₃, and NH₃ concentrations than the OH-initiated DMS oxidation experiments to investigate the formed particles at larger sizes of around 100 nm. In contrast to the OH oxidation experiments whose goal is to investigate the chemical mechanism, the goal of the additional experiments with high NH₃ is to investigate the CCN potential of the formed particles. That is, the much higher concentrations of DMS, O3, and NH3 facilitate sustained condensation that leads to larger particles. Additionally, the conducted voltage scan experiment in Figure S3 shows that the DMSO and DMSO₂ hydronium-water and ammonium clusters are stable against fragmentation at the chosen settings and that their proton affinity is high enough to even keep the charge upon collision-induced fragmentation at

higher voltages. Their uncertainties are estimated at around 20% in H_3O^+ -CIMS and 50–80% in NH_4^+ -CIMS with the larger uncertainty of the collision limit sensitivity in the latter. The reported concentrations of CH_3SCHO and CH_3SOH are lower-limit estimates also based on the results of the voltage scans. See Text S1.4 for the detailed description.

HPMTF and MSIA. A bromide chemical ionization mass spectrometer coupled with a multischeme chemical ionization inlet (Br⁻-MION-CIMS) and the gas-phase measurement of bromide chemical ionization mass spectrometer equipped with a Filter Inlet for Gases and AEROsols (Br--FIGAERO- $\ensuremath{\text{CIMS}}\xspace^{49}$ were the primary instruments to detect MSIA and HPMTF. The peaks of MSIA and HPMTF measured by Br--MION-CIMS can be found in Figure S4. If neither instrument was available due to instrument malfunction or absence in some experiments, NO3⁻-CIMS was used to measure MSIA and HPMTF. The instrument setup and operation of Br--MION-CIMS and FIGAERO are described in Rissanen et al.⁵⁰ and Lopez-Hilfiker et al.,⁴⁹ respectively. In addition to ionization with Br-, we also detected both MSIA and HPMTF with NH₄⁺ and H₃O⁺-CIMS. While direct calibrations of MSIA and HPMTF were not available, the intercomparison between the three ionization modes and the use of quantum chemical calculations provide reasonable constraints on the ionization efficiency of the two species and hence their concentrations.

We first derived MSA calibration coefficients for Br⁻-MION-CIMS and Br⁻-FIGAERO(g)-CIMS using the MSA concentration measured by NO₃⁻-CIMS. Afterward, MSA calibration factors in both instruments are used to estimate the lower limits of HPMTF and MSIA concentrations with high uncertainty. The traces shown in Figure S7 for MSIA from NH₄⁺- and H₃O⁺-CIMS are calibrated using the collision limit assumption. It is very likely that if ionization does not occur at the collision limit with all primary ion clusters with water, a strong water dependence would be observed between the different modes. The apparent humidity independence of the ratio of the MSIA concentration derived from the three different ionization schemes (NH₄⁺, H₃O⁺, and Br⁻) suggests that the compound is detected at the collision limit in these ionization schemes. However, the fragmentation caused by the electric field in the vacuum chamber also affects the detection efficiency. The signals of MSIA and HPMTF in Br⁻-MION-CIMS might underly some fragmentation or other unaccounted losses in the instrument or inlet. Yet, the voltage scan results in Figure S8 show that MSIA is detected at maximum sensitivity in both H₃O⁺ and NH₄⁺-CIMS. Therefore, we use the MSIA concentration from H₃O⁺-CIMS with around 28% uncertainty due to the calibration and inlet and instrumental loss correction. The signal of HPMTF is influenced by neighboring peaks and is divided into many water clusters in H_3O^+ -CIMS, while this is not the case in NH_4^+ -CIMS. Unfortunately, NH4⁺-CIMS was not available for many of the experiments, and the limit of detection in H₃O⁺-CIMS was too high to use the data during the herein analyzed experiments with low concentrations. As such, we use the lower-limit estimation for HPMTF concentrations from Br⁻-FIGAERO(g)-CIMS. See Section S1.5 for the detailed description and estimations.

Steady-State Measurement and Production Rate Calculation. The average residence time in the CLOUD chamber was \sim 1.3 h for experiment sets 1 and 2. The reaction rates of oxidation products were derived from steady-state

measurements, which means that the inflow of all gases, the concentration of oxidants, and oxidation products were kept constant. Under steady-state conditions, the rate of change of oxidation products equals zero, and we can write

$$\frac{\mathrm{d}[X]}{\mathrm{d}t} = p - [X] \times k_{\mathrm{loss}} - \sum_{i} k_{Y_{i}} \times [Y_{i}] \times [X] = 0$$
⁽¹⁾

Therefore, we can get the production rate p

$$p = [X] \times k_{\text{loss}} + \sum_{i} k_{Y_i} \times [Y_i] \times [X]$$
(2)

where X is the oxidation product, Y typically is OH or O_{3} , k_{Y_i} are the reaction rate coefficients for the reaction of X and Y, and k_{loss} is either the ventilation loss (2.1 × 10⁻⁴ s⁻¹ in experiment sets 1 and 2) or wall loss that is listed in Table S2. All of the oxidation product concentrations were measured at the steady-state condition. The OH concentration used in this study is estimated from our box model, and its reactions involved in forming OH radicals are listed in Table S7.

Modeling. In this study, a numerical model with zero dimension was set up and used to simulate DMS oxidation processes at different initial conditions (Table S1). We combine the simulation and experimental results to evaluate and constrain the role of different DMS gas-phase oxidation pathways in MSA and H₂SO₄ formation. The chemistry mechanism was generated with the Kinetic PreProcessor (KPP)⁵¹ and solved with the ordinary differential equation solver RODAS3 (Rosenbrock method of order 3).⁵² The concentrations of oxidation products are determined by the chemical production and loss from gaseous chemistry, wall reactions, ventilation loss, wall loss, and the condensation sink (CS) to generate particles in the CLOUD chamber. The measured temperature, relative humidity, DMS, O₃, CO, NO_x concentrations (Table S1), and O_3 photolysis rate with high time resolution were used as input to the model. The photolysis rate of O3 for UVH at full intensity or 5 W of UVX is around 6×10^{-5} s⁻¹. The OH concentrations are adjusted by changing the UV light intensity, O3, and H2O concentrations. The CLOUD chamber ventilation loss is mainly determined by the flush-out flow (~330 lpm in experiment sets 1 and 2, 250 lpm in experiment set 3). Species like H₂SO₄, MSA, and hydrogen oxide radicals are readily lost to the wall surface. We measured the lifetime of all species (Table S2) in the dark decay (lights off, fan 12%) and cleaning stage (lights off, fan 100%) to estimate the losses and applied them to our box model. With the increase of fan speed from 12 to 100%, the wall losses increased by a factor of 1.5-4.6 for different species. The measured loss of species like methyl thioformate (CH₃SCHO) in the cleaning stage is slightly larger than the ventilation loss caused by the flush-out flow. It suggests that the loss of CH₃SCHO is dominated by ventilation whose loss increases mildly with the increased fan speed. Therefore, we added extra loss terms for dimethyl sulfoxide $(CH_3S(O)CH_3, DMSO)$, dimethyl sulfone $(CH_3S (O)_2CH_3$, DMSO₂), and CH₃SCHO to fit their time evolutions in the cleaning stage. The particle condensation sink measured in this study is smaller than 1×10^{-5} s⁻¹, which is negligible compared to other loss processes. The detailed description of chemistry mechanisms used in the box model can be found in Text S2.



Figure 1. Schematic representation of DMS oxidation with OH radicals in this study. Most reactions are taken from MCMv3.3.1, Hoffman et al., and recent publications.^{29,31,32,41} The widths of the arrows indicate production rates (s⁻¹) on a linear scale, which arecalculated at -10 °C with 100 pptv DMS, 7×10^6 cm⁻³ OH, 40 ppbv O₃, 2 pptv NO, and 200 pptv NO₂. We set the same width for reaction rates when they are below 1.8×10^4 s⁻¹. The percentage given for each pathway indicates the branching ratio of the production rates at this condition. The precentage of MSP leads to pathways 1b is temperature-dependent. At +22 °C, the reaction follows pathway 1b to 97%. The reaction rate coefficients of the isomerization of MSP are discussed in Text S2. Red arrows highlight the important reactions proposed in this study. The mechanism of MSIA reacts with OH surrounded by the thin, dashed gray line is the possible alternative formation mechanism we suggest but their reaction rate coefficients are not fully studied. Therefore, we treat the MSIA converting to CH₃S(O)₂ in our box model. $k_{\text{DMS + O}_{3wall}}$ and $k_{\text{DMS + O}_{3wall}}$ are the semiempirical reaction rate coefficients for heterogeneous wall reactions (see Text S4 for details). The species in colored boxes in this figure are those we quantify, and the color of each box outline matches the color of each time trace in Figure 2. Species CH₃S(OH)CH₃, CH₃S(O₂)(OH)CH₃, CH₃SCH₂O, CH₃SO, and CH₃SO₂ have not been measured before but are included here based on modeling studies.^{33,35,40}

RESULTS

Identification of Oxidation Products from OH-Initiated DMS Oxidation. Figure 1 shows an oxidation mechanism initiated by the DMS + OH reaction and informed by the experiments we describe here, and Figure 2 shows an example time series of measured species for an experiment conducted at -10 °C with NO_x below the instrument detection limit, ≤ 2 pptv. With sensitive chemical ionization mass spectrometers, we are able to identify and quantify many intermediates and terminal products in this mechanism. The species we measure include MSA and H₂SO₄, as well as DMSO, DMSO₂, MSIA, and HPMTF. These are the major sulfur-containing products also previously identified and measured from DMS oxidation.^{26,27,29,32,53} We also identify $CH_3S(O)_2OOH$, CH_3SCHO , and CH_3SOH , as shown in the figures. Some species are quantified with proper calibration factors, while species like $CH_3S(O)_2OOH$, HPMTF, CH₃SOH, and CH₃SCHO (by NH₄⁺-CIMS) are lower-limit estimates due to the lack of authentic standards or generation methods. A more detailed description is given in the Materials and Methods section and Text S1.

The time traces in Figure 2 show the sequence of a typical experiment, starting with the characterization of wall losses in the dark, followed by the initiation of photochemistry. UV

lights, the trigger of OH radicals, were switched on at midnight (00:00) after DMS and ozone (O_3) concentrations stabilized. The time sequences of different DMS oxidation products show different behaviors. The evolution of H₂SO₄, MSA, MSIA, HPMTF, $CH_3S(O)_2OOH$, and CH_3SOH is consistent with the expected time response of gas-phase production through OH oxidation (with lights on) and wall loss. The measured concentration reaches their maxima in the order of a few minutes rather than a few hours like aqueous-phase reactions. The CH₃SCHO evolution is consistent with gas-phase production and loss to ventilation rather than the walls, which has a timescale of ~ 1.3 h (26.1 m³/330 lpm). DMSO shows a small fractional, but large absolute, increase with photochemistry, yet only a modest fractional wall loss. DMSO rapidly reaches a steady state, while DMSO₂ decreases slowly and continuously from the onset of the run, associated with the slowly decreasing product of the DMS and O₃ concentrations.

Mechanism Treatment via the Box Model. Figure S11 reveals that DMSO and DMSO₂ have a source from the wall, which has also been shown in a previous study.²⁷ As described in the Methods section, we first constrain wall effects to delineate gas-phase chemistry; however, the gas-phase photochemistry is dominant for all species other than DMSO₂ when the UV lights are on. In turn, DMSO₂ is long-lived in the gas



Figure 2. Experiment of OH-initiated DMS oxidation at -10 °C. (A) Time series of ultraviolet light intensity (blue line, left axis; used to photolyze O₃ to produce OH radicals) and mixing fan speed to accelerate wall losses (orange line, right axis). (B) O₃ (red line, left axis) and DMS (navy line, right axis) concentrations. (C) Measured DMS oxidation products (solid dots, colors correspond to box outlines in Figure 1). After a brief period ended at 21:48, UV lights were turned off and the fan speed was increased to full intensity to encourage wall losses. Most species were removed efficiently and showed a sharp drop. However, the loss of CH₃SCHO was mainly due to the chamber ventilation (with a dilution lifetime of 1.3 h). DMSO₂ was relatively insensitive to fan change since it was primarily produced from heterogeneous production on the walls, while DMSO was less affected by multiphase reactions (primarily from gas-phase reactions). When the fan speed was lowered and UV lights were turned on, the concentrations of most species increased due to OH oxidation of DMS. The shade for each time series presents the uncertainty of each species except for the species (HPMTF, CH₃S(O₂)OOH, and CH₃SOH) whose concentrations are lower-limit estimates; their uncertainty only includes the instrumental loss without correction from sensitivity.

phase and so has minimal influence on the chemistry. DMSO is also produced on the walls, but once OH is present, both gas-phase production and loss of DMSO via OH far exceed the wall production rate (see Text S4 and Figure S11).

As shown in Figure 2, the terminal products MSA and H_2SO_4 are far less abundant than the first-generation oxidation product CH_3SCHO , but MSA and H_2SO_4 have rapid wall loss, as do their precursors, such as MSIA, CH_3SOH , and HPMTF. Considering the wall loss rate, we find that the production rates for MSA ($3.2 \times 10^4 \text{ s}^{-1}$) and CH_3SCHO ($4.5 \times 10^4 \text{ s}^{-1}$) are similar, with uncertainties at around 50 and 20%, mainly caused by their measured concentrations. We seek these production rates along with molar yields and branching ratios. To constrain the production rates, we need a photochemical model of the chamber and the DMS oxidation sequence, including the walls.

The measured wall losses in Table S2 show that MSA and H_2SO_4 have the same values, allowing a direct comparison of the two species' concentrations. As shown in Figure 2, the measured MSA is high and 3.7 times larger than H_2SO_4 , which cannot be explained when assuming that MSA is exclusively formed from the hydrogen abstraction channel (pathway 1 in Figure 1). We observe a positive relationship between MSIA and MSA, shown in Figures S12 and S13. Previous studies^{54,55} have concluded that SO₂ is the lone major product of the reaction of OH with MSIA. However, the abstraction of an acidic H-atom by OH radicals is typically slow because acidity implies a deficiency in electron density.⁵⁶ By contrast, OH addition to the S-atom in MSIA is more likely to produce an

intermediate product that may decompose to form sulfurous acid (H₂SO₃) and CH₃,⁵⁵ but may also react with O₂ to produce MSA. This pathway and its reaction rate coefficients are not yet well understood. Therefore, in the box model, we add the pathway proposed by Yin et al.,⁵⁷ Lucas et al.,⁵⁸ and Hoffmann et al.⁴⁰ that MSIA reacts with OH or NO₃ radicals forming methyl sulfonyl radicals (CH₃S(O)₂) as an intermediate, which then either decomposes to SO₂ or reacts with O₃ to form MSA (via CH₃SO₃ reaction with HO₂) and SO₃, instead of producing SO₂ only. A recent study also applied this pathway to a multiphase model⁴¹ to explain the observed high particulate MSA/sulfate at 0 and 20 °C.

Lv et al.⁵⁵ suggested that the reaction of MSIA and O₃ is unlikely to be competitive with MSIA and OH because of the large discrepancy between these two reaction rate coefficients. Even at high O₃ concentrations (ca. 100 ppbv), the first-order reaction rate coefficient of MSIA by O₃ is 4.4×10^{-10} s⁻¹, which is much smaller than the first-order reaction rate coefficient of MSIA by OH at 2.8×10^{-4} s⁻¹ (with lowest OH concentrations, ~1 × 10⁶ cm⁻³). Therefore, the critical conclusion is that we would suggest that MSIA reacts with OH, producing both MSA and H₂SO₄. Thus, both the addition and abstraction pathways of DMS oxidation form MSA in the gas phase, explaining the higher MSA concentrations observed in field observations and CLOUD experiments.

To treat wall effects, we add wall reactions as pathway (3) to our DMS oxidation mechanism (Figure 1) by directly applying two semiempirical rate coefficients $k_{\text{DMS} + O_{3\text{wall}}}$ and $k_{\text{DMSO} + O_{3\text{wall}}}$



Figure 3. Measured and modeled gas-phase concentrations of identified species at an OH-initiated DMS oxidation experiment. Circles are the identified species measured by NO_3^- -CIMS, H_3O^+ -CIMS, and Br^- -FIGAERO_(g). Solid lines represent simulation results, including wall reactions of DMSO and DMSO₂ and dashed lines represent simulation results excluding wall reactions. Wall loss is included in our box model; therefore, we can compare the measured and simulated values directly. The differences between the modeled and measured HPMTF and CH₃SCHO are mainly because they are lower-limit estimates (see Texts S1.4 and S1.5 for quantitative measurement), which means their concentrations are underestimated in this case. The shadow area in the subplot of HPMTF presents the variation caused by applying different reaction rate coefficients of isomerization of MSP. Since the production rate of CH₃SOH is unclear, the simulated CH₃SO₄ concentrations (lower-limit estimates), but we do not investigate it in detail because it is not important for understanding MSA and H₂SO₄ formation. The shade for each time series presents the uncertainty of each species except for the species (HPMTF, CH₃S(O₂)OOH, and CH₃SOH) whose concentrations are lower-limit estimates; their uncertainty only includes the instrumental loss without correction from sensitivity.

Using a numerical model (see Material and Methods for the configuration and mechanism), we then evaluate the proposed mechanism herein. We compare the measured oxidation products (circles) with their modeled values, including wall reactions (solid lines) in Figure 3 for the same OH-initiated DMS oxidation experiment as Figure 2.

The modeled MSA, H_2SO_4 , $CH_3S(O)_2OOH$, and DMSO concentrations in Figure 3 are within 50% uncertainty of the measured values. The time evolution of HPMTF and MSIA in the simulation follows the same trend as the measured values. The modeled MSIA concentrations exceed the measured value (within an uncertainty of 28%) by a factor of 3, which is believed to be reasonable due to the uncertainties in DMS chemistry, chamber loss parameterizations, and systematic measurement errors. The smallest discrepancy between the modeled and measured HPMTF is a factor of 6. This difference can be explained by the uncertainty and the reduced

sensitivity of the gas-phase measurement of Br⁻-FIGAERO-CIMS to HPMTF due to a stronger declustering process compared to the maximum sensitivity compounds (see Text S1.5 for details). The discrepancy between the modeled and measured HPMTF can be varied from 6 to 14 by applying different reaction rate coefficients of the isomerization of MSP. As shown in Figure 3, the variation of HPMTF at low temperatures has a limited effect on DMS oxidation.

When we exclude the wall reaction from the model, the resulting vast difference between the modeled and measured $DMSO_2$ concentrations and their qualitative behaviors (Figure 3) indicates that wall reactions are essential to understanding the $DMSO_2$ formation. On the other hand, the simulation without wall interactions captures the time evolution of the other species except for DMSO, which indicates that OH-initiated DMS oxidation is their dominant source, and DMSO₂ is unlikely to be their primary precursor. The model suggests

that DMSO is formed for the most part from OH-initiated DMS oxidation in the gas phase but interacts strongly with the walls. The modeled DMSO concentrations are 54 and 46 pptv, with and without wall reactions, respectively. However, the temporal evolution of the concentration cannot be reliably predicted without wall interactions. As shown in Table S2, the uncertainties of measured MSA, H₂SO4, MSIA, CH₃SCHO, DMSO, and DMSO₂ are below 50%. Although the uncertainties of HPMTF, CH₃SOH, and CH₃S(O)₂OOH are high, their influences on our main conclusion—that MSA likely forms substantially via gas-phase oxidation of MSIA by OH—are negligible since they are not the determining species in this argument.

Overall, the underestimated calibration factor can explain the discrepancy between the modeled and simulated HPMTF. Based on the close agreement between the model and measurements for major oxidation products (DMSO, DMSO₂, MSIA, MSA, and H_2SO_4), we conclude that the mechanism presented here can represent DMS oxidation by OH radicals in the marine atmosphere.

Temperature Dependence for OH-Initiated DMS **Oxidation.** Temperature has a strong effect on (1) the branching ratio of addition/abstraction at the first step of DMS oxidation by OH radicals, (2) thermal decomposition of CH_3SO_2 and CH_3SO_3 in abstraction pathway 1a, (3), degradation of MSIA (which is not fully understood), and (4) peroxy radical isomerization of MSP in abstraction pathway 1b. All of these, especially the thermal decomposition, influence H₂SO₄/MSA. We plot H₂SO₄/MSA versus temperature in Figure 4 to present the temperature dependence for OH-initiated DMS oxidation. The green and red circles are from our CLOUD experiments with and without NO_x, respectively. The purple symbols are ambient data, and the black lines are box-model simulations with various mechanisms. The CLOUD data come from two separate temperature-ramping experiments, one cooling from +25 to +10 °C and the other from +10 to -10 °C, both without NO_x. In Figure 4, H_2SO_4/MSA ranges from 0.3 to 1.3 (-10 to +10 °C) and 1.4 to ~ 11 (+10 to +25 °C), decreasing exponentially with decreasing temperature. Ambient observations of gas-phase H₂SO₄/MSA vary from 1 to 17 at Mace Head, Ireland (during inferred experimental temperature dependence. The ambient concentrations of DMS, O₃, OH, and NO_x during the measurements of H₂SO₄/MSA from Beck et al., presented in Figure 4, were, unfortunately, unknown and likely differed from our experimental conditions. The differences will inevitably influence the absolute MSA and H₂SO₄ concentrations reported in Beck et al.'s observation and our experiments. However, we find that the ratio of H₂SO₄/MSA is mainly determined by temperature (Figure 4) instead of precursor vapor concentrations, and therefore the H₂SO₄/ MSA from our experiments is nonetheless used to compare with field observations. MSA formation has a stronger temperature dependence than H₂SO₄. A detailed discussion of the temperature dependence for all of the oxidation products is given in Text S5 and Figure S14.

We modeled the temperature dependence of H_2SO_4/MSA with the OH-initiated DMS oxidation mechanism developed here (solid line) and compare it to the gas-phase mechanism from MCMv3.3.1⁵⁹ (dashed line) and Hoffmann et al.⁴⁰ (dotted line) in Figure 4. These model results differ by up to



Figure 4. Temperature dependence of the H₂SO₄-to-MSA ratio. Green circles represent the H_2SO_4/MSA ratio without NO_x (green) from two temperature-ramping experiments measured by NO3--CIMS; red circles are the experimental H₂SO₄/MSA with NO_x. The NO2 and NO differences between the red and green circles are around 400 and 8 pptv. The error bars represent the standard deviation of temperature and H₂SO₄/MSA. Purple square symbols are the daytime average of ambient measurement. They are the ambient measurements from 4 and 5 May, 5 and 6 August 2017 in Ny-Ålesund station, 20 August 2015 in Villum,¹² 16 and 17 December 2014 in ABOA station,⁶⁸ and 17 June 1999 in Mace head.²¹ We decreased the temperature to study the temperature dependence, in contrast to increasing the temperature, to avoid the emission of contamination from the wall. Lines represent the simulation results using the OHinitiated gas-phase oxidation mechanism from MCMv3.3.1 (dashed line, with constant NO_x concentration) and Hoffman et al.⁴⁰ (dotted line). The simulation results in this study are presented as gray rectangles by varying the reaction rate coefficient of the isomerization of MSP.

an order of magnitude compared to the CLOUD measurements, but all show a similar exponential temperature dependence. This is because they all include thermal decomposition for $CH_3S(O)_2$ and CH_3SO_3 in abstraction pathway 1a, which contains an exponential temperature dependence that largely determines H₂SO₄/MSA. However, our mechanism shows much better agreement with the observed H₂SO₄/MSA compared to the other two mechanisms. This is because the reaction of MSIA with OH radicals contributes more to MSA production and consequently reduces H₂SO₄/MSA rather than only contributing to SO₂ formation, as e.g., in MCMv3.3.1. Overall, the observed temperature effect suggests a potentially greater role for MSA in colder regions in important processes such as new particle formation and growth, as our mechanism and observations show that the H_2SO_4/MSA ratio is smaller than 0.5 when the temperature is below roughly +2 $^{\circ}$ C.

Effect of NO_x on OH-Initiated DMS Oxidation. In addition to temperature, NO_x (i.e., NO₂ and NO) influences OH-initiated DMS oxidation, for example, through bimolecular reactions between peroxy radicals (RO₂) and NO. As shown in Figure 4, H₂SO₄/MSA from two experiments with NO_x (red circles) agrees with the temperature-ramping experimental results (green circles), indicating that NO_x has a negligible effect compared to the temperature sensitivity over this same range. The NO_x-ramping experiments in Figure S15 confirm that NO₂ and NO only slightly enhance H₂SO₄/MSA in contrast to the strong effect from temperature change. The measured H₂SO₄/MSA at +10 °C (Figure S15A) and -10 °C (Figure S15B) increase by 54 and 100% as NO₂ increases from 0 to 400 pptv. The effect of NO_x on HPMTF formation is not clear (Figure S15). We observe a slight drop of HPMTF (17%) and almost constant DMS in Figure S15A. However, Figure S15B shows a 100% increase in HPMTF and a 20% drop in DMS because DMS reacts with NO₃ radicals. We cannot quantify the effect of NO_x on HPMTF formation because of the interference from N₂O₅ formed from the reaction of O₃ and NO₂, which has a close molecular weight to HPMTF.

In MCMv3.3.1, the modeled H₂SO₄/MSA (Figure S16) is very sensitive to NO₂ concentration varies dramatically as NO₂ increases from 0 to 160 pptv. However, the modeled H₂SO₄/ MSA from the other two mechanisms (Figure 4) are insensitive to NO_x. In MCMv3.3.1, NO₂ constrains the formation of MSA through the reaction of CH₃SO with NO₂, which forms CH₃S(O)₂, because it excludes the isomerization reaction of CH₃SO₂ producing CH₃S(O)₂ (Hoffmann et al.⁴⁰). The isomerization reaction is estimated with a constant value of ~1 s⁻¹ in previous studies,^{60,61} which is higher than the bimolecular reaction rate of CH₃SO with NO₂ (~1.2 × 10⁻¹ s⁻¹ with a maximum 400 pptv NO₂), although it lacks the temperature dependence.

Overall, the suite of simultaneously measured key intermediates helped us to adjust the previous DMS oxidation schemes. Our mechanism has two major improvements. First, the influence of NO_x on product distribution is greatly diminished compared to the MCMv3.3.1. Second, the key ratio H_2SO_4/MSA is substantially reduced (MSA is increased), with the absolute ratio and temperature dependence in the mechanism agreeing with both the experimental data and ambient observations.

DISCUSSION AND ATMOSPHERIC IMPLICATION

DMS oxidation in the atmosphere is affected by temperature, NO_{xy} the distribution of oxidants including OH, NO_{3} , halogen compounds, ^{36,37,62} and available water for multiphase oxidation in aerosols and droplets.^{40,63} In this study, we followed the kinetics of a suite of gas-phase reaction intermediates and final products in DMS oxidation experiments by OH to investigate the effects of temperature and NO_{x} . Our results show a strong temperature dependence for DMS oxidation, especially the yields of oxidation products such as MSA.

Our experimental results show high MSA concentrations and reduced H_2SO_4/MSA from OH-initiated DMS oxidation, especially at low temperatures (<0 °C). From the box-model results, it is clear that the inclusion of the gas-phase MSA formation mechanism via oxidation of MSIA by OH improves the agreement between modeled and experimental results at low temperatures. When MSIA reacts with OH, it may produce $CH_3S(O)_2$ or another adduct intermediate; regardless, both $CH_3S(O)_2$ and the adduct intermediate decompose to SO_2 or SO_3 to form H_2SO_4 or react with O_3 or O_2 to form MSA. For both pathways, we can expect a similar temperature dependence.

At low temperatures, the oxidation of DMS with OH proceeds more via the OH addition channel, forming abundant DMSO and MSIA, and consequently, a large amount of MSA. At high temperatures, the reaction rate of the OH addition channel decreases, which lowers MSIA and MSA formation, while the hydrogen abstraction channel becomes more important. Also, the predominance of MSP isomerization (\geq 95, at 295 K)²⁹ leads to the HPMTF formation, which

increases H_2SO_4 production substantially through oxidation of HPMTF.^{29,30}

Therefore, we propose that MSIA is important in MSA formation at low temperatures, which can also be connected to the observed high particulate MSA/sulfate in laboratory⁴¹ and field observations^{7,17} since both MSA and H₂SO₄ participate in particle growth. Interestingly, NO_x is far less influential than previously thought. At -10 °C, H₂SO₄/MSA increases by 100% after ramping up NO₂ (from 0 to 400 pptv) and NO (from 0 to 8 pptv). But the observed change (red circles) is small (less than 50%) compared to the effect of changing temperature (about an order of magnitude when the temperature decreases from +10 to -10 °C).

As shown in Table S5, our experimental MSA (0.2 to $1.5 \times 10^7 \text{ cm}^{-3}$) and H₂SO₄ (0.3 to $3 \times 10^6 \text{ cm}^{-3}$) concentrations formed from OH-initiated DMS oxidation experiments conducted at -10 °C are in the range of field measurements. For example, the concentrations of MSA and H₂SO₄ measured in Ny-Ålesund station (Figure S17, air temperature ranging from -10 to -7 °C) range from 0.1 to $4 \times 10^7 \text{ cm}^{-3}$ and 0.2 to $7 \times 10^6 \text{ cm}^{-3}$, respectively. Aerosol concentrations also affect gas-phase MSA and H₂SO₄ concentrations because of their different aerosol partitioning. However, the particle condensation sink in this study is smaller than $1 \times 10^{-5} \text{ s}^{-1}$, which is only slightly larger than the values at Ny-Ålesund (~4 $\times 10^{-4} \text{ s}^{-1}$) and Villum (~3 $\times 10^{-4} \text{ s}^{-1}$).

DMSO₂ and DMSO are important sulfur reservoirs in our study. They can also be formed from wall reactions without OH radicals. Our simulations show that wall reactions are the major source of DMSO₂ but have a limited role for other oxidation products compared to the gas-phase reactions (even DMSO). In the atmosphere, analogous to the wall of the chamber, cloud droplets and aerosol water could play a similar role. Therefore, the observed high concentrations of DMSO₂ (40–120 pptv) and DMSO (3–18 pptv) at night over the Arabian Sea⁶⁴ and Pacific Ocean⁶⁵ may be associated with multiphase DMS oxidation.

The DMS oxidation lasts only a few hours in the CLOUD chamber, while it can be for a few days in the atmosphere depending on the conditions. To gauge possible atmospheric evolution (a few days), we simulated OH-initiated DMS oxidation over a long timescale (~ 1 week) at -10 °C, excluding walls and ventilation. As shown in Figure S18, terminal products MSA and H₂SO₄ accumulate in parallel over time, ultimately constituting the dominant reaction products. DMSO, MSIA, CH₃SCHO, and CH₃SOH are the precursors for MSA and H₂SO₄, reaching a steady state during the first day of simulation. HPMTF is a precursor for SO₂, and after 1 week, both reach high concentrations and slowly approach steady state. Field measurements indicate high concentrations of HPMTF (up to 50 pptv in the marine atmosphere³²) and its possible participation in particle formation. Accounting HPMTF chemistry in a global chemistry transport model⁶⁶ results in a significant decrease in boundary layer levels of SO2 and H₂SO₄. Novak et al.⁶⁷ demonstrate that the rapid loss of HPMTF to clouds terminates DMS oxidation to SO₂. This is important at high temperatures. On the other hand, at low temperatures, HPMTF production is strongly reduced, and the main product becomes MSA via MSIA, so the fraction of oxidized DMS leading to SO2 is significantly lowered. The concentration of DMSO₂, a terminal product of OH-initiated oxidation of DMS, remains low in the absence of heterogeneous reactions. These simulations suggest that

MSA may be the dominant oxidation product of DMS oxidation for conditions found in the high-latitude marine atmosphere with low temperatures.

The agreement between the present chamber study, our model, and ambient observations over a wide temperature range is consistent with OH-initiated DMS oxidation governing the formation of gas-phase MSA and H_2SO_4 in the remote atmosphere. Our observation of many intermediate products closes gaps between the measured and modeled MSA. Specifically, MSIA and HPMTF are useful intermediates that serve as markers for OH addition and hydrogen abstraction channels, respectively. Therefore, measurement and proper calibration of these two species become important in future laboratory experiments and field observations for studying DMS. These findings improve our understanding of the atmospheric DMS oxidation process and its contribution to the natural sulfur cycle and potentially the formation of biogenic aerosol and thus CCN.

ASSOCIATED CONTENT

S Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acs.est.2c05154.

Additional description of quantitative measurement of DMS and its oxidation products; quantum chemical calculation method; detailed chemistry mechanism used in the box model; HO_2 and OH estimation from the box model; heterogeneous wall reactions of DMS and O_3 in the CLOUD chamber; temperature effect on the DMS oxidation process; and reactions of DMS oxidation chemistry (PDF)

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J.S., X.-C.H., and M.Sip. planned the experiments. J.S., W.S., X.-C.H., G.M., M.W., R.M., B.R., R.B., L.C., F.A., D.M.B., B.B., Z.B., D.C., B.C., L.D., J.D., H.F., M.G., R.G., M.H., V.H., W.K., J.E.K., A.K., H.L., C.P.L., B.L., N.G.A.M., H.E.M., D.M., R.L.M., B.M., T.M., T.P., J.P., M.P., A.A.P., S.S., M.Si., M.Su., Y.J.T., A.T., N.S.U., D.W., Y.Wang, S.K.W., A.W., Y.Wu., M.Z.-W., U.B., J.C., R.C.F., R.V., O.M., K.M., K.L., M.R., J.K., I.E.-H., F.B., and M.Sip. prepared the CLOUD facility or measuring instruments. J.S., W.S., X.-C.H., G.M., M.W., R.M., B.R., R.B., L.C., A.A., F.A., D.M.B., B.B., D.C., J.D., H.F., M.G., M.H., V.H., D.K., W.K., J.E.K., H.L., C.P.L., B.L., N.G.A.M., D.M., R.L.M., B.M., T.M., J.P., M.Sim., M.Su., Y.J.T., A.T., D.W., S.K.W., Y.Wu., M.Z.-W., J.C., R.C.F., R.V., and J.K. collected the data. J.S., W.S., X.-C.H., P.Z., G.M., M.W., R.M., B.R., L.C., S.I., J.K., M.Sim., M.Su., and S.K.W. analyzed the data. J.S., W.S., X.-C.H., P.Z., S.I., P.R., M.Sim., D.S., R.W.D.J., U.B., J.C., R.C.F., A.H., K.M., K.L., M.R., J.K., I.E.-H., F.B., M.Sip., N.M.D., and D.R.W. contributed to the scientific discussion. J.S. and N.M.D. wrote the manuscript with contributions from W.S., X.-C.H., M.R., J.K., I.E.-H., F.B., and D.R.W. J.S., W.S., X.-C.H., P.Z., R.B., Z.B., J.D., H.F., N.G.A.M., R.G., P.R., R.W.D.J., S.S., D.S., D.W., M.Z.-W., U.B., M.R., J.K., I.E.-H., F.B., M.S., N.M.D., and D.R.W. commented on and edited the manuscript.

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