Experimental reaction rates constrain estimates of ozone response to calcium carbonate geoengineering

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Stratospheric solar geoengineering (SG) would impact ozone by heterogeneous chemistry. Evaluating these risks and methods to reduce them will require both laboratory and modeling work. Prior model-only work showed that CaCO₃ particles would reduce, or even reverse ozone depletion. We reduce uncertainties in ozone response to CaCO₃ via experimental determination of uptake coefficients and model evaluation. Specifically, we measure uptake coefficients of HCl and HNO₃ on CaCO₃ as well as HNO₃ and ClONO₂ on CaCl₂ at stratospheric temperatures using a flow tube setup and a flask experiment that determines cumulative long-term uptake of HCl on CaCO₃. We find that particle ageing causes significant decreases in uptake coefficients on CaCO₃. We model ozone response incorporating the experimental uptake coefficients in the AER-2D model. With our new empirical reaction model, the global mean ozone column is reduced by up to 3%, whereas the previous work predicted up to 27% increase for the same SG scenario. This result is robust under our experimental uncertainty and many other assumptions. We outline systematic uncertainties that remain and provide three examples of experiments that might further reduce uncertainties of CaCO₃ SG. Finally, we highlight the importance of the link between experiments and models in studies of SG.
The deliberate introduction of aerosols into the stratosphere as a means of solar geoengineering (SG) might reduce some climate risks. Yet physical risks and efficacy of stratospheric aerosol geoengineering are poorly constrained due to uncertainty in the climate’s response to aerosol radiative forcing and to local stratospheric processes that include stratospheric chemistry, heating, and aerosol microphysics. Uncertainties about local process can be reduced by experiments. A prior modeling study from our group suggested that calcium carbonate (CaCO$_3$) might enable stratospheric geoengineering with reduced ozone loss or even ozone increase, but that study lacked measurements of important CaCO$_3$-specific reaction rates. It assumed that uptake coefficients (the probability that a gas-phase molecule is reactively bound following collision with a solid aerosol surface) of ozone-relevant gas species on CaCO$_3$ is independent of the fraction of CaCO$_3$ that been consumed in the reaction. This uncertainty needs to be resolved by empirical methods.

The use of empirical tools to study solar geoengineering has been limited. Field experiments are contentious. Modeling studies use existing experimental results for sulfates but there have been very few laboratory studies to address uncertainties of SG using non-sulfate particles. This work is among the first efforts to measure uptake coefficients on solid particles at stratospheric temperatures. Our purpose is reducing uncertainty about SG.

Sulfate is the most widely-studied material for stratospheric SG because of the prominence of natural sulfate aerosols in the stratosphere and the long history exploring their links to climate. In addition, there appear to be ways to inject SO$_2$ and likely H$_2$SO$_4$ based on existing technologies. However, sulfate aerosols pose well-known risks such as ozone depletion and stratospheric heating. CaCO$_3$ particles have been proposed as an alternative to sulfate, and major motivations include the potential to reduce both risks. This work addresses the ozone depletion aspect. Many modeling studies that explore chemical impacts of potential SG deployment have shown ozone depletion effects from a range of sulfate distribution scenarios, chiefly due to their impacts on NOX (NO and NO$_2$) and halogen species. These effects could counteract the recovery of ozone under the Montreal Protocol and increase risks of SG scenarios designed to mitigate impacts of climate change.

Non-sulfate materials have been proposed to address various risks of SG. Nearly two decades ago, Teller and Wood proposed the use of alumina nanoparticles based on scattering considerations while cautioning about environmental impacts. Other solid particles including TiO$_2$, SiC, ZrO$_2$, diamond, and CaCO$_3$ have been explored. Diamond and alumina might reduce ozone depletion compared with sulfate when producing the same radiative forcing.

CaCO$_3$ produces much less stratospheric heating than does sulfate, and is a plausible material given the extensive industrial-scale manufacture of micron-scale calcite particles. CaCO$_3$ could also address the risks of ozone depletion by reactive uptake of acidic species containing halogens, sulfur, and nitrogen, effectively scrubbing ozone-depleting species from the stratosphere. However, due to the competition between denoxification and dehalogenation, it is difficult to know the net effect on ozone a priori.

One important shortcoming of Keith et al. is uncertainty of the uptake coefficients due to lack of experimental data. Here we measure the uptake coefficients of four reactions at stratospheric temperature:

$$\text{CaCO}_3 + \text{HCl}.$$  \hspace{1cm} (1)

$$\text{CaCO}_3 + \text{HNO}_3.$$  \hspace{1cm} (2)

$$\text{CaCl}_2 + \text{HNO}_3.$$  \hspace{1cm} (3)

$$\text{CaCl}_2 + \text{ClONO}_2.$$  \hspace{1cm} (4)

These reactions are important as they directly impact the chlorine and nitrate budget in the stratosphere, which critically affect stratospheric ozone concentration. Eqs. (1) and (2) determine the capacity of CaCO$_3$ to sequester NO$_x$ and halogen species, and Eq. (3) and Eq. (4) release reactive halogen species back to the stratosphere. We did not include heterogeneous reactions of gaseous HBr, BrONO$_2$, and H$_2$SO$_4$ with CaCO$_3$ particles because Keith et al. showed that these reactions caused little change in background mixing ratios of BrO$_x$ or H$_2$SO$_4$. Uptake coefficients of Eqs. (1) and (2) have been experimentally determined, but not under stratospheric conditions. These coefficients of Eqs. (3) and (4) have not been studied previously. In addition, Keith et al. did not include heterogeneous catalysis on CaCO$_3$ surfaces such as ClONO$_2$ + HCl. Here we assess the potential impacts of these reactions. We find that exposure to gas-phase acids causes CaCO$_3$ particles to age rapidly, reducing their reactivity. This is in stark contrast to liquid sulfuric acid particles with surfaces that are unaffected by aging. We use these data to constrain a chemical transport model of stratospheric chemistry and aerosol microphysics and find that the prior model results overestimated ozone response.

### Results

#### Experimental determination of uptake coefficients

Flow tube experiments have been widely used to study heterogeneous uptake coefficients for gas-solid reactions relevant to the stratosphere. For example, Molina et al. studied the uptake of ClONO$_2$ on silica and alumina in the presence of HCl. Davies and Cox studied the reaction between HONO and NaCl. Schmidt et al. studied reaction between acetone and TiO$_2$. Figure 1a shows the schematic of our flow tube experiment, it is similar to Molina et al. with slight modifications. We synthesized ClONO$_2$ inline using plasmonic dissociation of Cl$_2$ and cryogenically purified it inline (more details in the method section). As there are no literature values for the uptake coefficient of ClONO$_2$ on CaCl$_2$, we first compare our results for the uptake coefficients of ClONO$_2$ on NaCl with existing literature values to validate our synthesis and measurement methods. The results confirm that our experimental and analysis approach as well as our ClONO$_2$ synthesis method produce valid results: the uptake coefficient of ClONO$_2$ on NaCl was determined to be $5 \pm 3 \times 10^{-3}$ at room temperature, which is close to the values reported in literature (Gebel et al. reported $6.5 \pm 3 \times 10^{-3}$).

Uptake coefficients were calculated using the method of Knopf (details in the “method” section). Typical time series for an uptake experiment for Eqs. (1) and (4) are shown in Fig. 2a, b and more detailed uptake coefficients are measured in Table 1. Consistent with previous literature, the reactions all show large initial uptake which gradually decays to a slowly varying semi steady state value (uptake signal changes by $<$5%/min). This process demonstrates a slow deactivation of the particles’ reactivity for gaseous uptake reactions.

Uptake coefficients at the semi steady state are shown in Table 1. For Eq. (2), this level occurs after about 10 min. Immediately after the insert is retracted upstream of the injector following the uptake experiment, a small increase in the HCl signal corresponding to reversibly adsorbed HCl on the surface is observed, consistent with the uptake experiment results in literature and the observation that HCl is reversibly adsorbed on the calcite surface before irreversible uptake occurs. For Eq. (2),
the semi steady state level occurs after about 3 min. In contrast to Eq. (2), there is no desorption peak after the insert is pulled upstream of the injector, signaling that this reaction proceeded without the adsorption step as in Eq. (2). For the HNO₃ uptake reactions, the initial rapid changes in uptake could not be observed because there is a prominent time-varying background uptake by the quartz insert, while the background uptake of HCl and ClONO₂ are smaller and did not show time-dependent changes.

Fig. 1 Schematics of the experimental setup and some potentially important reaction processes. a Schematics of the flow tube and mass spectrometer used for the flow tube experiments. b Schematics of the important physical and chemical interactions during heterogeneous uptake of gaseous molecules.

Fig. 2 Results from uptake experiments. a, b Typical uptake signal from the flow tube experiments for HCl + calcite and ClONO₂ + CaCl₂ reactions respectively. The baseline was subtracted and signal normalized to the background level. c, d Results from long-term uptake experiments and parametric fit results for HCl + calcite and ClONO₂ + CaCl₂ respectively. Thickness-dependent uptake coefficients listed in Table 1 are extracted from these fits. Error bars in the data points in c represent systematic error from the experiment. Results from the mass spectrometer measurements are overlaid on the plots. The shadowed regions show 95% confidence intervals of the parametric fit extracted using bootstrapping as outlined in the experimental section. The standard errors for fitting parameters A and n (defined in the text) are 0.18 and 0.05, and the covariance is −0.08.
characteristics. We therefore did not extract the time-dependent uptake signal for these reactions as the case for Eq. (2) and (3), and calculated the semi-steady-state uptake coefficient based on the difference in mass spectrometer (MS) signal between the background signal and after the signal reaches a steady value.

Flow tube experiments measure rates of surface reactions but are not suited to explore continuous changes in the uptake coefficients after exposures on the order of days or weeks as the uptake signal decreases below the detection limit. To explore the long-duration regime we conducted a set of long-term uptake “flask experiments” to measuring reaction rates between CaCO₃ particles and HCl.

Flask experiments are conducted for Eq. (1) with three HCl concentrations, and the results are shown in Fig. 2c. The result from one flow tube experiment is also displayed for comparison. The concentrations span six orders of magnitude, and for each HCl concentration, we see that the reacted layer thickness growth rates decrease with reaction time, representing decreasing reactive uptake coefficients as a function of reaction extent.

### Table 1 Parameterization of thickness-dependent uptake coefficients.

| Reaction                        | γ value          | Thickness-dependent parameterization | Notes                                                      |
|---------------------------------|------------------|--------------------------------------|------------------------------------------------------------|
| HCl + Calcite -> CaCl₂         | (7.2 ± 5) x 10⁻⁵| 2.66 x 10⁻³ ÷ c₁d                      | Three flow tube experiment measurement cycles              |
|                                 | (3.3 ± 0.6) x 10⁻⁷| or 3.41 x 10⁻⁴ ÷ c₁D                  | 16 long term uptake measurements at 3 different HCl       |
|                                 | to (3.0 ± 0.6) x 10⁻¹² b | or 3.41 x 10⁻⁴ ÷ c₁D                  | concentrations                                           |
| HNO₃ + Calcite -> Ca(NO₃)₂      | (2.5 ± 1.8) x 10⁻⁴| 1.43 x 10⁻⁴ ÷ c₁                      | Three flow tube experiment measurement cycles, error      |
|                                 |                  | or 3.41 x 10⁻⁴ ÷ c₁D                  | increased by background adsorption from HNO₃              |
| HNO₃ + CaCl₂ -> Ca(NO₃)₂        | (2.5 ± 1.5) x 10⁻⁴| 3.41 x 10⁻⁴ ÷ c₁                      | Three measurement cycles by MS, error increased by        |
|                                 |                  | or 3.41 x 10⁻⁴ ÷ c₁D                  | background adsorption from HNO₃                           |
| Cl(NO₃)₂ + CaCl₂ -> Ca(NO₃)₂   | (9.8 ± 9) x 10⁻⁵ | 6.26 x 10⁻⁴ ÷ c₁                      | Three measurement cycles by MS                            |

All flow tube experiments were done at 215 K and long-term uptake experiments were done at 200 K. Mass spectrometer (MS) experiments were done at 1.6 Torr and long-term uptake experiments were done at ambient pressure.

*fit to equation γ = A(t)^n, with the measurements from long-term uptake experiments. For the parameterizations, the dimensions for the concentration (C) and product layer thickness (d) are /cm³ and Å, respectively, n is the area-normalized product amount and t is time.

*The values listed were calculated from experiments with C = 5.4 x 10³/cm³, d = 0.05 Å and C = 3.6 x 10⁻⁷/cm³, d = 33.3, respectively.

Estimating thickness-dependent uptake coefficients. Unlike liquid sulfate particles, whose surface chemical composition are not affected by gas-phase uptake, CaCO₃ particles age under gaseous exposure as products accumulate on their surfaces. As a result, the uptake coefficients change with exposure. We incorporate these changes by formulating uptake coefficients that depend on the product layer thickness and refer to them as thickness-dependent uptake coefficients. This is challenging because uptake coefficient changes are affected by many processes including the changes of available CaCO₃ reaction sites as a function of reaction extent, adsorption and equilibration of water on the particle surface, diffusion of gaseous species to available reaction sites, crystallization of reaction products, and steric changes that might cause fractures in the product layer (schematics shown in Fig. 1b). Various empirical models were used to account for these changes including a shrinking-core model, a Crystallization and Fracture modeling framework, a kinetic model, and a surface solvation model. None of these processes have been studied extensively under stratospheric conditions. We therefore posit a semi-empirical model that assumes that the gas phase uptake rate is proportional to the diffusion rate of reactive gaseous species through a surface product layer to reach available CaCO₃ reaction sites. The physical intuition is that the reaction rate is dominated by a diffusion-like process where the availability of the reactant plays a dominant role after the initial surface product forms and slows further reaction. The corresponding form of the cumulative product per unit area is n_p = A(C^n t)^3, where A is the fitting parameter to the empirical data, C is the concentration of the gas phase reactive species, t is time of reaction, and n is an empirical constant chosen differently for various reactions based on their uptake mechanisms. As discussed in the experimental result section, Eq. (2) proceeds following an initial reversible surface adsorption and dissociation of HCl on CaCO₃, and n was chosen as 1/2, consistent with the argument of Huthwelker et al. ; Eq. (4) proceeds without the initial reversible adsorption, and n was set to be 1. For Eq. (2), 1/2 is also the best fit for the data when n floats in the fitting routine. The uptake mechanisms for Eqs. (2) and (3) are unknown, and we formulated the reactions based on both the 1/2th and 1st order assumptions. This expression of n_p was chosen as it parsimoniously captures the reaction and diffusion processes. The corresponding thickness-dependent parameterizations are shown in Table 1. We selected this model because it fits our data well and is parsimonious (with only essential parameters to ensure low model variances) considering the high uncertainties due to various kinetic processes. We attempted a few other fitting models that are either more complicated or do not explain the experimental trends as well (Supplementary Fig. S1). The 95% confidence interval shown in Fig. 2c, d are calculated using bootstrapping and shows higher error where data are sparse.

Modeling global ozone impact. We use a slightly updated version of the atmospheric chemistry model used in Keith et al., adopting the highest injection rate of 5.6 Tg/yr CaCO₃ from that analysis (see “method” section). In addition to the scenario using the full thickness-dependent parameterization (thk-dep) and the 2040 baseline scenario without solar geoengineering, we studied five other scenarios: (i) thk-ind, with thickness-independent uptake coefficient (γ) values from the second column in Table 1, (ii) no psc, with the same γ values as in the thk-dep case but with polar stratospheric clouds (PSC) reactions turned off, (iii) 0 x γ, with γ values for Eqs. (1)–(4) set to 0, (iv) thk-dep + conc-dep, with thickness-dependent γ values using the alternative concentration-dependent γ values for Eqs. (2) and (3) (this captures uncertainties in the uptake mechanism of Eqs. (2) and (3) as described in the previous section), and (v) 10 x γ, with γ values set to ten times the thk-dep case. The 0 x γ and 10 x γ scenarios capture the uncertainties of calcite particle surface area density since changing the uptake coefficient is equivalent to changing the surface area density available for gas-phase uptake.
Figure 3 shows the impact of CaCO₃ emissions on ozone loss rates caused by important ozone-depleting species compared with Keith et al. The NO₃ cycles play a much more important role than ClO₂ catalytic cycles as a function of altitude, averaged from 60°S to 60°N. The thk-ind case shows an increase over the baseline in the NOₓ-related loss rates in contrast to the results from the thk-ind scenario and Keith et al. This is consistent with the decrease in ClO activation and ClO concentration, which in turn lead to increased NOx levels that drive hydrolysis of N₂O₅. Both lead to increased NOx levels that drive ozone loss around the altitude of CaCO₃ injection. Another impact of the decreased sulfate is a slight decrease in PSCs. However, the comparison between the thk-dep result and the no psc result shows that the column changes due to PSC variations do not offset the ozone decrease caused by NOₓ increase.

This ozone change in the thk-dep scenario below the baseline are caused primarily by NOₓ. As Fig. 3 shows, the changes in the NOₓ and ClOₓ cycles are smaller and in the opposite direction compared with the Keith et al. estimates, and the overall effect is a decrease in the ozone column compared to the baseline. Similarity between results from the thk-dep scenario and the 0×γ results show that these impacts are not caused by Eqs. (2)–(4) but are predominantly caused by the decrease in background sulfate from neutralization by CaCO₃ particles. This causes a decrease in Cl activation and ClO concentration, which in turn results in less ClONO₂. It also leads to less heterogeneous hydrolysis of N₂O₅. Both lead to increased NOₓ levels that drive ozone loss around the altitude of CaCO₃ injection. Another impact of the decreased sulfate is a slight decrease in PSCs. However, the comparison between the thk-dep result and the no psc result shows that the column changes due to PSC variations do not offset the ozone decrease caused by NOₓ increase.

Discussion

Most narrowly, this study shows that our prior work overestimated the impact of CaCO₃ aerosol on ozone chemistry. The aged CaCO₃ aerosol is close to unreactive with respect to gas-phase uptake reactions. As a result, these uptake coefficients produce smaller ozone response in the opposite direction than predicted in Keith et al., and thus no contribution to the restoration of column ozone. However, since CaCO₃ reduces stratospheric heating and ozone depletion compared to sulfate aerosols, and since there may be methods of increasing the ozone benefit of CaCO₃ by increasing surface reactivity, it is still a viable candidate for SG.

The reduced ozone response is robust under a wide range of uncertainties. To assess the impact of uncertainties in the thickness-dependent γ parameterizations, we conduct model runs assuming upper and lower limits of the thickness-dependent uptake coefficient confidence intervals shown in Fig. 2c, d. In addition, we capture uncertainties in surface area density of the CaCO₃ particles with the 10-γ and 0-γ scenarios. Neither this scenario nor the inclusion of gas-phase concentration-dependent γ for Eqs. (2) and (3) (thk-dep + conc-dep) produce more than 1% ozone column differences anywhere as shown in Fig. 4. This is explained by the fact that the ozone response is dominated by changes in sulfate surface area, not uptake reactions of gas-phase species on CaCO₃.

More generally, these results highlight the importance of considering changes in uptake coefficients due to gas-phase exposure...
when determining the impacts of solid reactive particles. Further studies should explore the uptake coefficients of other ozone related gases. The details of the physical and chemical changes from reaction with sulfate aerosols, including those that contain organics\cite{34} are also important. Finally, there is uncertainty about how reactivity depends on the details of morphology and surface chemistry, which is important because CaCO$_3$ aerosol could be produced by a variety of technologies\cite{35,36}.

Our work illustrates how laboratory research can address uncertainties in stratospheric aerosol geoengineering. It is likely that there will be many other examples of novel measurements that can be accomplished with well-established experimental methods and are crucial in evaluating novel proposals that address risks posed by SG. As an illustration, the following experiments are straightforward applications of existing methods yet could be important to assessing SG risks:

**Other heterogeneous reactions on CaCO$_3$.** Heterogeneous reactions of ClONO$_2$ and HCl with water and HCl on particle surfaces present key uncertainties of ozone response to CaCO$_3$ geoengineering. Among these, reactions with HCl could be more important since the uptake coefficients of ClONO$_2$ in the presence of HCl was shown to be two orders of magnitude greater than in the presence of water on solid surfaces\cite{5,36}. Including an uptake coefficient of 0.02 for ClONO$_2$ + HCl (the value measured on alumina surface\cite{36}) for both the thk-dep and thk-ind cases decreases global ozone column by 2–3% compared to cases without this reaction (Supplementary Fig. S3). Ozone decreases up to 15% below the baseline over the south pole in the thk-dep case. Further including an uptake coefficient of 0.1 for HOCl + HCl did not change the ozone columns by more than 0.5%. Our results modified the ozone response estimate of Keith et al.\cite{9}, but some relevant reactions are still poorly constrained by experiment. A next step would be to measure reaction rates of ClONO$_2$ + HCl on fresh and aged CaCO$_3$ surfaces.

**Optical properties of CaCO$_3$ aerosol.** The major proposed advantage of CaCO$_3$ over sulfuric acid aerosol is that it causes less radiative cooling due to less solar and IR-band absorption and thus could pose lower risk with respect to changes in dynamics, which are poorly understood\cite{38}. Yet, this hypothesis is based on bulk optical properties of pure CaCO$_3$ that may differ from aged CaCO$_3$ aerosols in the stratosphere (e.g., surface salts could form hydrates that shift the absorption of terrestrial radiation and Ca(NO$_3$)$_2$ may absorb in the UV). Well-established methods could be used to measure the real and imaginary index of refraction of aged CaCO$_3$ powder. The aging process should include a range of stratospherically relevant processes including gas-phase reactions (as we explored in the flask experiments) and exposure to UV radiations.

**Ice nucleating properties.** Aerosols used in SG may impact cirrus cloud formation once they descend into the upper troposphere\cite{37,38}. The ice nucleating properties are important in determining impacts of the sedimented aerosols on cirrus clouds. In addition, ice may nucleate onto the CaCO$_3$ particles in the Antarctic winter under PSC conditions, affecting PSC formation. Experiments using a variety of laboratory experimental techniques\cite{39} could test ice nucleating properties on CaCO$_3$ powder that had been prepared under the range of conditions described in the prior paragraph. For example, Cziczo et al.\cite{4} has tested ice nucleating properties of CaCO$_3$, CaSO$_4$, and Na(NO$_3$)$_2$ and found that they can cause significant changes to cirrus clouds absent of natural ice nucleating particles\cite{4}.

This study may have lessons for management of solar geoengineering research. Sound estimates of SG’s risks and efficacy will require experiments and models. Current estimates are almost entirely based on large-scale modeling. Models are crucial, but overreliance on models may encourage overconfidence when users ignore the simplifications hidden in their assumptions. Empirical methods are needed to help reduce uncertainty. Models can explore how results depend on uncertainty in parameters to motivate experimental research which, in turn, can provide better constraints on parameters used in models. The back-and-forth between the experimental and modeling communities is essential for environmental science. The complexity of experimental results like these may serve as a useful reminder of the deep uncertainties underlying predictions about the consequences of solar geoengineering.

### Methods

**Gas flow system and flow tube.** All gas tanks are purchased from Airgas except for the SF$_6$ gas (Praxair, research grade). The nitrogen gas is ultra-high purity, and the HCl gas is a 1% blend in nitrogen. The nitric acid is generated from a permeation tube (KIN-TEK Trace Source Refillable permeation tube with a target emission rate of, ng/min at 50 °C). When doing experiments, we operate the permeation tube at 20 °C and calibrated the emission rate using the mass spectrometer.

The flow tube is jacketed with a length of 1 m and an inner diameter of 26 mm. The insert has an inner diameter of 22 mm, wall thickness of 1 mm, and length of 22 cm. The cooling was done by circulating an ethanol-dry ice mixture through the jacket. The temperature of the gas stream is monitored by three K-type thermocouples at the two ends and the middle of the flow tube. The temperature gradient of a typical experiment through the flow tube is 1–2 K. For all the flowtube experiments, the flowtube temperature was controlled at 215 K and the pressure was 1.7 Torr. Since only dry gas streams are used in the flowtube experiments, and the reactive gas streams pass through a ~20 cm quartz tubing at 215 K before reacting with particles, we treat the water vapor content in the experimental region to be 0.

**ClONO$_2$ Synthesis.** The ClONO$_2$ synthesis consists of three steps:

\begin{equation}
\text{Cl}_2 + \text{RF plasma} \rightarrow 2\text{Cl}.
\end{equation}

\begin{equation}
\text{Cl} + \text{O}_3 \rightarrow \text{ClO}.
\end{equation}

\begin{equation}
\text{ClO} + \text{NO}_2 \rightarrow \text{ClONO}_2.
\end{equation}

The gas tanks are all purchased from Airgas (Ar: HP grade, O$_2$: UHP grade, Cl$_2$ and NO$_2$: 1% blend in nitrogen). The RF plasma is generated by a Technical Research and Manufacturing Inc. DBS301 RF generator with a frequency range of 0.125–0.25 GHz. The Power output is measured by a Bird 43 General Purpose Wattmeter and is tuned to be between 15 and 40 W during operation. The ozone is generated from an A22 Ozone MP 1900 ozone generator. The ozone concentration as measured by an ozone meter is 1.5%.

The synthesis conditions were simulated in a box model to guide selections of design parameters that resulted in the highest ClONO$_2$ yield. A quartz tube acts both as an RF cavity for generating the chlorine radical and as a mixing region for the combination of ClO and NO$_2$. The length of the quartz tube is 70 cm. The distance between the two vertical flanges is 40 cm. The outer diameter is 1/2 in. and the inner diameter is 1.01 cm. We assumed no radicals remains after the cold trap because the quartz wool in the cold trap provides a large surface area for them to recombine or otherwise react.

After ClONO$_2$ is synthesized, it is collected and purified in a cold trap cooled by a cold jacket where chilled nitrogen gas flows through. The concentration of the effluent ClONO$_2$ is measured through absorption spectroscopy in a quartz cell with an absorption length of 99.14 cm. Absorption spectra in the 200–295 nm region were acquired using a fiber-coupled Ocean Optics USB4000 UV–Vis spectrometer (10-µm slit width, 200–400 nm 5 grating) illuminated by a Hamamatsu L2D2 deuterium lamp. The concentration of ClONO$_2$ is then calculated based on the absorption cross section data in the JPL database\cite{40}. It is determined to be about 75 ppm under typical synthesis conditions.

**Material and preparation for the flow tube experiment.** The calcite particles are purchased from American Elements with a manufacture-reported average diameter of 200 nm. The Brunauer–Emmett–Teller (BET) surface area of these particles is measured by a Quantachrome Monosorb instrument using N$_2$ as the adsorption gas. The BET surface area reported was 14.8 ± 0.5 m$^2$/g. We also measured the BET surface area of samples after processing by isopropyl alcohol (IPA) and found that IPA processing did not change the BET surface area. The NaCl particles (puriss. p.a., ≥99.5%, powder or crystals) used are purchased from Sigma Aldrich. To increase the amount of surface area per unit mass, the particles are then processed
in a Retsch PM100 planetary ball mill. The BET surface area of the resulting NaCl particles is measured by a Beckman Coulter SA 3100 Surface Area Analyzer using He as the absorption gas. The BET surface area is 0.051 ± 0.003 m²/g for NaCl before milling, and 0.38 ± 0.05 m²/g after milling.

Before the experiment, the CaCO₃ particles are deposited on the insert from a suspension of CaCO₃ in IPA. A known volume of a suspension with known mass concentration of CaCO₃ is pipetted into the insert. The insert is heated at roughly 60°C to spread the particles evenly over the insert while the IPA evaporates. This process is repeated until a desired amount of CaCO₃ is deposited on the insert. The insert is then heated in an oven at 250°C for roughly 30 min to drive off any excess IPA. A similar deposition method has been used to prepare inserts described in other experiments.4 The insert is then placed into the flow tube for at least 2 h before an uptake experiment starts. According to Santschi and Rossi,3 this is sufficient to drive off all but strongly bound surface water. It is possible that some IPA molecules could adsorb to the CaCO₃ surface even after the oven and vacuum processing. However, we expect this effect to be small since heating has been shown to be effective at removing adsorbed water from CaCO₃ surfaces by Santchi and Rossi, and IPA is more volatile than water.

**Flow tube experiment.** The uptake experiment design follows Molina et al. but differs in two respects. First, instead of moving the injector with respect to the stationary insert and flow tube, the insert is moved relative to the stationary injector and flow tube. Four magnets attached to the injector rod and the insert allow for this configuration. This eliminates condensation on the injector rod, but it retracts from the low temperature region of the flow tube without having to heat the injector. It also eliminates the possibility of gas leaks into the flow tube due to mechanical movement of the injector relative to the vacuum seal. Second, instead of electronic ionization with electron ionization, ions are produced by chemical ionization by SF₆ as outlined in Huay et al. An alternative method used in the literature to measure heterogeneous uptake coefficients is the aerosol flow tube experiment, where the aerosols are dispersed in a gas stream and flow with the carrier gas. This method eliminates issues with gas-phase diffusion limitations and gas-transfer limitations within the particle layers. However, one key disadvantage of aerosol flow tube experiments compared to our coated wall flow tube experiment is that it is difficult to extend the residence time of particles in the reaction region beyond tens of seconds. As shown in the results section, reactive uptake coefficients on CaCO₃ particles continue to change for more than 10 min due to gradual surface saturation. Aerosol flow tube experiments cannot change the gas flow rate, so to overcome the issue with gas phase diffusion limitation with the coated wall flow tube experiments, we applied a correction to the uptake coefficients as outlined in the following section. This correction was small due to the low uptake coefficients measured in our experiments. In addition, to probe the effect of gas transfer limitations within the particle layers, we measured uptake coefficients of Eq. (2) for different CaCO₃ layer thickness and found that the uptake coefficients measured at the semi steady state were 2 ± 1.5 × 10⁻⁴, 4 ± 2 × 10⁻⁴, and 3 ± 2 × 10⁻⁴ for 0.5, 1.5, and 2.5 layers of coated CaCO₃ thickness on the insert. These are similar to values reported by Santschi and Rossi for cleaved CaCO₃ marbles and did not differ with layer thickness as expected from diffusion limitation. To overcome this problem, we modeled the diffusion limitation in the particle layer separately for each particle layer height. We calculated the effective Sherwood number, and the effective uptake coefficient y = C - γfl/ωg. The uptake signal of this baseline uptake at steady state was subtracted from these experiments and subtracted from the uptake signals when calculating the uptake coefficients. For the flow tube experiments, random errors come mainly from the precision of the ion-selective electrode used to measure the chloride ion concentration (2% of the measured voltage). Another source of error is the dead volume in the glass vessel, which takes time for the influent HCl–N₂ mixture to displace at the exit of the reaction. To account for this effect, we included an uncertainty term that is equivalent to the time that the influent gas fully displaces the dead volume (~10 s). This effect is negligible for longer period reactions (~1 min) but has a significant effect for the shorter-term reactions.

**Modeling method.** In order to probe the impact on global stratospheric ozone, we conduct similar simulations as Keith et al. using the AER 2-D chemical transport-aerosol model. All model specifications follow that study except for the uptake coefficients to isolate the impact of uptake coefficients measured in this work. We investigate the highest CaCO₃ emission rate in that study with 5.6 Tg yr⁻¹ of uniform CaCO₃ emission between 30°S and 30°N and between 20 and 25 km. This produces roughly 2 W m⁻² of global radiative forcing (RF). CaCO₃-related reactions that are included in the model consist of Eqs. (1)–(4) and coagulation of sulfate aerosols with CaCO₃ particles. The coagulation was treated in the same way as Keith et al., assuming liquid sulfate reacts with CaCO₃ to form CaSO₄ instantaneously once sulfate aerosols collide with CaCO₃ aerosols. The above trace gas emissions for 2040 conditions are taken from IPCC RCP 6.0, providing 2.3 ppbv of Cl₂ in the upper stratosphere but with temperature and circulation from a 1978–2004 climatology.
