Unanticipated Side Effects of Stratospheric Albedo Modification Proposals Due to Aerosol Composition and Phase

Daniel J. Cziczo¹,²,³*, Martin J. Wolf³, Blaž Gasparini⁴,⁵, Steffen Münch⁴ & Ulrike Lohmann⁶

The Earth has now warmed ~1.0 °C since the period 1850–1900, due in large part to the anthropogenic addition of greenhouse gases to the atmosphere. Most strategies to address this warming have called for a reduction of emissions and, often, accompanying removal of greenhouse gases. Other proposals suggest masking the increased radiative forcing by an increase in particles and/or clouds to increase scattering of incoming solar radiation. Two related recent proposals have suggested addition of calcite particles to the stratosphere, which one model suggests may enhance ozone. Here we show that the interaction of calcite with acidic materials in the stratosphere results in a more complex aerosol than has been previously considered, including aqueous and hydrate phases that can lead to ozone loss. Our study suggests particle addition to the stratosphere could also perturb global radiative balance by affecting high altitude cloud formation and properties. Experimental and modeling results suggest particles will act as the nucleation sites for polar stratospheric cloud ice and, after sedimentation into the troposphere, impact cirrus clouds in the absence of other efficient ice nucleating particles. These results show that an overly simplistic set of assumptions regarding intentional particle emissions to the atmosphere can lead to incorrect estimates of the radiative effect and fail to identify unintended consequences.

It has been established that the anthropogenic emissions of greenhouse gases have warmed the planet by ~1.0 °C since pre-industrial times¹,². There have been proposals to intentionally alter the atmospheric abundance of greenhouse gases both to use their warming potential to raise temperature for the benefit of colder climates³ and, more recently, to lessen the detrimental effects of increased global temperature⁴. The latter is termed ‘carbon capture and storage’ and falls within the concept of ‘geoengineering’, commonly defined as the intentional manipulation of planetary processes for a desired climatic effect.

Since at least the 1960’s there have been proposals to mask the increased radiative forcing associated with anthropogenic greenhouse gases by increasing planetary albedo⁵–⁷. Land and ocean albedo enhancement and, more recently, the addition of light-scattering particles to the atmosphere and/or manipulation of cloud properties have been proposed⁸. Highlighting the uncertainty surrounding these actions, a recent report by the National Academy of Sciences⁹ suggests such processes should be described as ‘climate interventions’ rather than ‘climate engineering’ or ‘climate management’, terms which imply a level of certainty that is not supported by experimental evidence. These albedo modification strategies do not address the other effects of greenhouse gases, such as ocean acidification⁵, and are predicted to have significant side effects, including changes in plant growth, precipitation, stratospheric heating and ozone loss and reduced concentrated solar power generation⁹,¹⁰. Increased scattering of solar radiation, termed shortwave radiation (SW), the cornerstone of albedo modification, also has side effects such as changes in precipitation and atmospheric chemistry⁵,¹⁰.
A number of proposals for climate intervention have suggested augmentation of the natural stratospheric aqueous sulfuric acid aerosol layer. Figure 1 shows this concept in relation to an unperturbed atmosphere. Volcanic eruptions have been observed to inject sulfuric acid precursors and water vapor into the stratosphere, leading to enhanced particle formation. Mixing and sedimentation processes result in particle lifetimes on the order of a few years. Volcanic enhancement of the concentration of light-scattering particles in the stratosphere causes increased scattering of SW, and lower global temperatures, for up to a few years. Volcanic effects are therefore transient, and not analogous to albedo modification proposals, which require a sustained and increasing input to mask a continually increasing concentration of greenhouse gases.

The increase in particles after volcanic eruptions has been observed to lead to a depletion of ozone in the stratospheric layer that absorbs ultraviolet SW before it reaches the surface. Two acids, HNO₃ and HCl, act as stratospheric reservoirs of nitrogen and chlorine radicals (termed NOₓ and ClOₓ, respectively) that catalytically destroy ozone. Particles in the stratosphere, natural or otherwise, can act as sites for heterogeneous chemistry involving these ozone-destroying radicals. Reactivity varies with particle phase: aqueous solutions, hydrates and ices generally have higher reactivity than anhydrous surfaces. Proposals for albedo modification suggesting augmenting sulfate aerosols would therefore lead to a destruction of ozone. There have been several comprehensive studies of the intricacies of stratospheric ozone depletion by particles naturally or anthropogenically added to the stratosphere. These consider polar and lower latitudes, dynamics, and the amount of injected material.

Some recent albedo modification proposals have suggested addition of light scattering but less chemically reactive (with respect to ozone depletion) particles such as alumina and calcite. Calcite (CaCO₃) addition has been suggested since it could scatter SW and might reduce stratospheric aerosol acidity and sequester NOₓ and ClOₓ, leading to increased ozone:

\[
\text{CaCO}_3(s) + \text{H}_2\text{SO}_4(g,aq) \rightarrow \text{CaSO}_4 + \text{H}_2\text{O}(l) + \text{CO}_2(g)
\]  
(1)

\[
\text{CaCO}_3(s) + 2 \text{HNO}_3(g,aq) \rightarrow \text{Ca(NO}_3)_2 + \text{H}_2\text{O}(l) + \text{CO}_2(g)
\]  
(2)

\[
\text{CaCO}_3(s) + 2 \text{HCl}(g) \rightarrow \text{CaCl}_2 + \text{H}_2\text{O}(l) + \text{CO}_2(g)
\]  
(3)

The underlying assumption in reactions (1)–(3) is that anhydrous salts are produced and that their surfaces are less reactive than aqueous, hydrate and ice surfaces. Moreover, eventual particle sedimentation would effectively remove part of the stratospheric NOₓ and ClOₓ burden. Keith et al. considered an addition of 2.1 and 5.6 Tg calcite per year in the form of 275 nm radius particles, resulting in Ca(NO₃)₂, CaSO₄ and CaCl₂ due to reaction with nitric, sulfuric and hydrochloric acid, respectively. This order corresponds to the assumed production due to the vapor pressure of each species: production of CaCl₂ is least favorable due to the high vapor pressure and low abundance of HCl. The estimated radiative impact ranged from 1–2 W/m² between the cases. Based on an
assumption of all products being anhydrous and inactive, a 3.8% increase in stratospheric ozone was estimated for the 2.1 Tg case.\(^6\)

Reactions (1)–(3), although chemically balanced, do not account for the correct form of the products under stratospheric conditions. The basis for Eq. (1) is that the ubiquitous stratospheric aqueous sulfuric acid layer, or sulfuric acid vapor, will react with the injected calcite particles via coagulation or uptake, respectively, and that all reactions will proceed until the calcite is converted (Fig. 2). Full conversion is inconsistent with the literature since it has been assumed that CaCO\(_3\) will fully react with stratospheric acids to produce similarly unreactive anhydrous salts.\(^6\) The literature does not support this assumption since sulfuric acid produces an unreactive CaSO\(_4\) ‘armor’ on CaCO\(_3\) upon reaction with H\(_2\)SO\(_4\) (Panel C)\(^18\). Regardless of the acid-CaCO\(_3\) reaction, the ultimate product under stratospheric conditions is not anhydrous but instead reactive hydrates and soluble salts (Panel D)\(^20,25,26\). Unlike aqueous H\(_2\)SO\(_4\), CaCO\(_3\) and the solid and hydrate products can nucleate ice heterogeneously (Fig. 3), thereby introducing effective ice nucleating particles, and impacting cloud formation, to the stratosphere and troposphere after sedimentation. Injected particle composition will evolve over time and single particles will most likely form a mixture of these.

**Figure 2.** The natural background stratosphere contains aqueous H\(_2\)SO\(_4\) particles which act as sites for reactions that destroy ozone (Panel A). Proposals for injection of additional aqueous H\(_2\)SO\(_4\) particles would scatter more solar radiation but promote additional ozone loss.\(^2\) Aqueous H\(_2\)SO\(_4\) particles do not nucleate ice heterogeneously, instead requiring temperatures below \(-38^°C\) and RH with respect to ice in excess of 145%.\(^30\) Recent proposals suggest addition of scattering materials that are less reactive, such as CaCO\(_3\) (Panel B).\(^6\) It has been assumed that CaCO\(_3\) will fully react with stratospheric acids to produce similarly unreactive anhydrous salts.\(^6\) The literature does not support this assumption since sulfuric acid produces an unreactive CaSO\(_4\) ‘armor’ on CaCO\(_3\) (Panel C)\(^18\). Regardless of the acid-CaCO\(_3\) reaction, the ultimate product under stratospheric conditions is not anhydrous but instead reactive hydrates and soluble salts (Panel D)\(^20,25,26\). Unlike aqueous H\(_2\)SO\(_4\), CaCO\(_3\) and the solid and hydrate products can nucleate ice heterogeneously (Fig. 3), thereby introducing effective ice nucleating particles, and impacting cloud formation, to the stratosphere and troposphere after sedimentation. Injected particle composition will evolve over time and single particles will most likely form a mixture of these.

**Figure 3.** The temperature and relative humidity required for ice nucleation by the particles proposed for addition to the stratosphere and the products expected from acid reactions. Water saturation (solid line) and homogeneous freezing (dashed line)\(^30\) are shown for reference. Conditions required for ice nucleation are shown for 275 and 550 nm diameter particles. For comparison, an effective tropospheric ice nucleating particle (INP), kaolinite mineral dust, and an ineffective INP, elemental carbon (EC), are shown. The proposed injection material, CaCO\(_3\), and the anhydrous and hydrate forms of CaSO\(_4\) were found to nucleate ice with moderate effectiveness. Calcium nitrate, incorrectly assumed to be an anhydrous salt by Keith et al.,\(^6\) is not an effective INP, since it is either a hydrate or an aqueous solution under these conditions.

assumption of all products being anhydrous and inactive, a 3.8% increase in stratospheric ozone was estimated for the 2.1 Tg case.\(^6\)

Reactions (1)–(3), although chemically balanced, do not account for the correct form of the products under stratospheric conditions. The basis for Eq. (1) is that the ubiquitous stratospheric aqueous sulfuric acid layer, or sulfuric acid vapor, will react with the injected calcite particles via coagulation or uptake, respectively, and that all reactions will proceed until the calcite is converted (Fig. 2). Full conversion is inconsistent with the literature since
CaSO₄ forms a surface layer that acts as ‘armor’ that prevents further reaction; this reaction has been extensively studied for use of limestone mitigation of acid mine drainage. Any incomplete conversion of CaCO₃ results in a smaller ozone increase than suggested by Keith et al. CaSO₄ exists in several forms not realized in Eq. (1) that have been extensively studied for industrial purposes. These include CaSO₄ dihydrate (gypsum), hemihydrate (CaSO₄·½H₂O), dihydrate (CaSO₄·2H₂O) and both soluble and insoluble anhydrites. Laboratories studies show production of the soluble anhydrite CaSO₄ at stratospheric temperatures (~200 K) and low relative humidity (RH), transitioning to the dihydrate at ~40% RH. Room temperature studies show that the hemihydrate is produced between these two phases, at ~20%, and the hexahydrate forms at 70–80% RH. Ozone depleting reactions of these compounds have not been studied, but other hydrates, such as nitric acid trihydrate, effectively promote ozone loss.

Keith et al. only discuss formation of an anhydrite (the difference between the soluble and insoluble forms is not noted), but at the mean RH with respect to liquid water of the lower stratosphere in the extratropics (~12 km altitude) of ~12% aqueous particles are the thermodynamically favorable form. The focus of most stratospheric aerosol augmentation studies to increase global albedo is on a mid-stratospheric layer, between ~20–25 km, where the mixing ratio of water vapor is 3–7 parts per million by volume (ppmv). There is a strong water vapor gradient from the tropopause to mid-stratosphere, with a decreasing RH as altitude increases. Values of 40% RH are not uncommon in the lower stratosphere, decreasing to 3% above 20 km. Particles are more likely to exist in anhydrous and lower hydrate forms at higher altitudes and as higher hydrates or in aqueous states closer to the tropopause. The exact phase state will depend on the specific particle type and water vapor and temperature profile. Overall, our analysis suggests that current assumptions of CaSO₄ particle phase are oversimplified and sulfate sequestration and ozone depletion impacts need to be reconsidered.

Calcite, which is expected to remain a solid particle, and anhydrous and the dihydrate of CaSO₄ act as moderately effective INPs. Ca(NO₃)₂, which exists as higher order hydrates and in solution under these conditions, is only observed to nucleate ice homogeneously. This finding also reinforces the reactivity of Ca(NO₃)₂ for ozone depleting reactions. The ability of a fraction of injected particles to act as INPs is important in two regimes: upper tropospheric cirrus ice and polar stratospheric clouds. The former case is further discussed in the next paragraphs. The role of heterogeneous nucleation on polar stratospheric clouds, sites on which ozone-depleting reactions occur, is not resolved. Thus, the impact on cloud formation and ozone depletion from addition of anthropogenic INPs to this region of the atmosphere is currently unknown.

In order to estimate the impact of injection of calcite into the stratosphere we use a general circulation model which can simulate aerosol transport, evolution, and radiative effects as well as aerosol-cloud interactions in both liquid and ice clouds. We simulate a continuous stratospheric injection of calcite particles with a modal radius of 275 nm with the calcite refractive indices consistent with Keith et al. The calcite burden required ~3 model years to equilibrate at about 5 Mt, consistent with a particle lifetime of just under 1 year (Fig. S1). The simulated radiative forcing from calcite aerosols alone is ~1.5 W/m² averaged for years 4–10 of the simulation (Fig. 4, panel A). This result is comparable to the value of ~2 W/m² reported by Keith et al., the variance is due to differences in aerosol treatment (modal vs. sectional aerosol schemes) and the interactive simulation of stratosphere-to-troposphere transport. Sedimentation processes are normally well resolved in models, however, coarse vertical resolution in the stratosphere compared to thin aerosol or cloud layers may lead to numerical diffusion and be a limitation for properly resolving circulation. This can affect correctly simulating the residence time of particles in the stratosphere.

The decreased SW radiation flux that reaches tropospheric clouds reduces their ability to scatter solar radiation, implying a positive (warming) cloud radiative effect of 0.5 W/m² (Fig. 4, panel B) with the net radiative effect of the calcite injection at ~1.1 W/m² (Fig. 4, panel C). It is unclear if Keith et al. considered this “cloud shielding” effect but it has been previously shown for sulfate and more complex injection simulations. Our simulations also allow sedimentation of particles across the tropopause and their ability to either form new cloud particles or shift their formation mechanisms based on the laboratory measurements. The treatment of ice cloud formation mechanism (homogeneous versus heterogeneous nucleation) varies between models and has been shown...
to result in substantial radiative difference33–35. Our model simulations only show small changes in cirrus clouds in case of abundant natural INPs and larger changes if natural INPs are rare (Supplementary Materials). These results show that a full consideration of the chemical, physical and radiative impacts of albedo modification proposals is necessary to provide an understanding of the impact on the planet.

Methods

Particles were produced from aqueous solutions, anhydrous and hydrate crystals, depending on their predicted phase state at tropopause conditions. Ice nucleation onset was determined within a liter-sized cloud chamber36. Also known as an ‘ice cloud chamber’, tropopause conditions of temperature and relative humidity at which cirrus clouds form can be controlled to test INP properties. For the global climate simulations, the ECHAM-HAM general circulation model was used. ECHAM-HAM includes a two-moment aerosol scheme, capable of simulating aerosol emissions, growth, coagulation, and sinks and a two-moment cloud microphysics scheme with prognostic equations for cloud liquid and ice, suitable for simulations of aerosol-cloud interactions37,38.

The Supplementary Information contains a full discussion of the methods.

Received: 9 May 2018; Accepted: 27 October 2019;
Published online: 11 December 2019

References

1. IPCC, Global warming of 1.5°C. An IPCC Special Report on the impacts of global warming of 1.5 °C above pre-industrial levels and related global greenhouse gas emission pathways, in the context of strengthening the global response to the threat of climate change, sustainable development, and efforts to eradicate poverty [Masson-Delmotte, V. et al (eds)] (2018).
2. McNutt, M. et al. Climate Intervention: Reflecting Sunlight to Cool Earth. (The National Academies Press, 2015).
3. Arrhenius, S. A. Worlds in the making: The evolution of the universe (Harper Brothers, 1908).
4. McNutt, M. et al. Climate Intervention: Climate Intervention: Carbon Dioxide Removal and Reliable Sequestration. (The National Academies Press, 2015).
5. Dykema, J. A., Keith, D. W. & Keutsch, F. N. Improved aerosol radiative properties as a foundation for solar geoengineering risk assessment. Geophys. Res. Lett. 43, 7758–7766 (2016).
6. Keith, D. W., Weisenstein, D. K., Dykema, J. A. & Keutsch, F. N. Stratospheric solar geoengineering without ozone loss. Proc. Natl. Acad. Sci. 113, 14910–14914 (2016).
7. Crutzen, P. J. Albedo enhancement by stratospheric sulfur injections: A contribution to resolve a policy dilemma? Clim. Change 77, 211–220 (2006).
8. Horing, D. F. et al. Restoring the quality of our environment: Report of the environmental pollution panel. President's Science Advisory Committee (White House Press, 1965).
9. Murphy, D. M. Effect of Stratospheric Aerosols on Direct Sunlight and Implications for Concentrating Solar Power. Environ. Sci. Technol. 43, 2784–2786 (2009).
10. Robock, A. 20 reasons why geoengineering may be a bad idea. Bull. Atomic Sci. 64, 14–18 (2008).
11. Robock, A. Volcanic eruptions and climate. Geophys. Res. 38, 191–219 (2000).
12. Hamill, P., Jensen, E. J., Russell, P. B. & Bauman, J. J. The life cycle of stratospheric aerosol particles. Bull. Am. Met. Soc. 78, 1395–1410 (1997).
13. WMO (World Meteorological Organization), Assessment for Decision-Makers: Scientific Assessment of Ozone Depletion: 2018, Global Ozone Research and Monitoring Project—Report No. 58 (2018).
14. Solomon, S. Stratospheric ozone depletion: A review of concepts and history. Rev. Geophys. 37, 275–316 (1999).
15. Rolf Müller, R., Grooß, J.-U., Zafar, A. M., Robrecht, S. & Lehmann, R. The maintenance of elevated active chlorine levels in the Antarctic lower stratosphere through HCl null cycles. Atmos. Chem. Phys. 18, 2985–2997 (2018).
16. Richter, J. H. et al. Stratosynamical dynamical response and ozone feedbacks in the presence of SO2 injections. J. Geophys. Res. 122(12), 557–12,573 (2017).
17. Tilmes, S. et al. Effects of different stratospheric SO2 injection altitudes on stratospheric chemistry and dynamics. J. Geophys. Res. 123, 4654–4673 (2018).
18. Fusi, L., Primicerio, M. & Monti, A. A model for calcium carbonate neutralization in the presence of armoring. Appl. Math. Model. 39, 348–362 (2015).
19. van der Merwe, E. M. & Strydom, C. A. Purification of South African phosphogypsum for use as Portland cement retarder by a combined thermal and sulphuric acid treatment method. So. African J. Sci. 100, 411–414 (2004).
20. Chou, I.-M. & Seal, R. R. Magnesium and calcium sulfate stabilities and the water budget of Mars. 18. Fusi, L., Primicerio, M. & Monti, A. A model for calcium carbonate neutralization in the presence of armoring.
21. Gierens, K., Schumann, U., Helten, M., Smit, H. & Marenco, A. A distribution law for relative humidity in the upper troposphere and lower stratosphere derived from three years of MOZAIC measurements. Atmos. Chem. Phys. 17, 1218–1226 (1999).
22. Hurst, D. F. et al. Recent divergences in stratospheric water vapor measurements by frost point hygrometers and the Aura Microwave Limb Sounder. Atmos. Meas. Tech. 9, 4447–4457 (2016).
23. Murphy, D. M. & Koop, T. Review of the vapour pressures of ice and supercooled water for atmospheric applications. Q. J. R. Meteorol. Soc. 131, 1539–1565 (2005).
24. Hurst, D. F. et al. Comparisons of temperature, pressure and humidity measurements by balloon-borne radiosondes and frost point hygrometers during MOHAVE-2009. Atmos. Meas. Tech. 4, 2777–2793 (2011).
25. Liu, Y. J., Zhu, T., Zhao, D. F. & Zhang, Z. F. Investigation of the hygroscopic properties of Ca(NO3)2, and internally mixed Ca(NO3)2/CaCO3 particles by micro-Raman spectrometry: Atmos. Chem. Phys. 8, 7205–7215 (2008).
26. Gough, R. V., Chevrier, V. F. & Tolbert, M. A. Formation of liquid water at low temperatures via the deliquescence of calcium chloride: Implications for Antarctica and Mars. Plan. Space Sci. 131, 79–87 (2016).
27. C. H. Pruppacher, H. R. & Klett, J. D. Microphysics of Clouds and Precipitation 2nd edn (Kluwer, 1997).
28. Köhler, H. The nucleus in and the growth of hygroscopic droplets. Trans. Faraday Soc. 32, 1152–1161 (1936).
29. Koop, T., Loo, R., Tsias, A. & Peters, T. Water activity as the determinant for homogeneous ice nucleation in aqueous solutions. Nature 406, 611–614 (2000).
30. Schmidt, H. et al. Solar irradiance reduction to counteract radiative forcing from a quadrupling of CO2: climate responses simulated by four earth system models, Earth Syst. Dynam. 3 (2012).
31. Kuebeler, M., Lohmann, U. & Feichter, J. Effects of stratospheric sulfate aerosol geo-engineering on cirrus clouds. Geophys. Res. Lett. 39, L23803 (2012).
32. Gasparini, B. & Lohmann, U. Why cirrus cloud seeding cannot substantially cool the planet. J. Geophys. Res. 121, 4877–4893 (2016).
33. Jensen, E. et al. On the Susceptibility of Cold Tropical Cirrus to Ice Nuclei Abundance. J. Atmos. Sci. 73, 2345–2464 (2016).
34. Storelvmo, T., Boos, W. R. & Herger, N. Cirrus cloud seeding: A climate engineering mechanism with reduced side effects? Philos. Trans. R. Soc. London 372, 20140116 (2014).
35. Garimella, S. et al. The SPectrometer for Ice Nuclei (SPIN): an instrument to investigate ice nucleation. Atmos. Meas. Tech. 9, 2781–2795 (2016).
36. Neubauer, D., Lohmann, U., Hoose, C. & Frondoso, M. G. Impact of the representation of marine stratocumulus clouds on the anthropogenic aerosol effect. Atmos. Chem. Phys. 14, 11997–12022 (2014).
37. Stevens, B. et al. Atmospheric component of the MPI-M Earth System Model: ECHAM6. J. Adv. Model. Earth Syst. 5, 146–172 (2013).

Acknowledgements
The authors thank J. Dykema for providing relevant particle refractive indexes for use in the radiative simulations and D. Murphy and J. Smith for discussions. S. Münch has been supported by the Swiss National Science Foundation under Project 200021E-166726.

Author contributions
D.J.C., M.J.W. Laboratory studies, data analysis, and paper writing; B.G. Performed and analyzed simulations, paper writing. Implementation of a new tracer particle into the model; S.M. Implementation of the calcite particles as INPs for the cirrus scheme; U.L. Paper writing.

Competing interests
The authors declare no competing interests.

Additional information
Supplementary information is available for this paper at https://doi.org/10.1038/s41598-019-53595-3.
Correspondence and requests for materials should be addressed to D.J.C.
Reprints and permissions information is available at www.nature.com/reprints.
