# **Opinion:** A research roadmap for exploring atmospheric methane

# 2 removal via iron salt aerosols

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20 Abstract. The escalating climate crisis requires rapid action to reduce the concentrations of atmospheric greenhouse gases and 21 lower global surface temperatures. Methane will play a critical role in near-term warming due to its high radiative forcing and 22 short atmospheric lifetime. Methane emissions have accelerated in recent years and there is significant risk and uncertainty associated with the future growth in natural emissions. The largest natural sink of methane occurs through oxidation reactions 23 24 with atmospheric hydroxyl and chlorine radicals. Enhanced atmospheric oxidation could be a potential approach to remove 25 atmospheric methane. One method proposes the addition of iron salt aerosols (ISA) to the atmosphere, mimicking a natural 26 process proposed to occur when mineral dust mixes with chloride from sea spray to form iron chlorides, which are photolyzed by sunlight to produce chlorine radicals. Under the right conditions, lofting ISA into the atmosphere could potentially reduce 27 atmospheric methane concentrations and lower global surface temperatures. Recognizing that potential atmospheric methane 28 29 removal must only be considered as an additive measure - in addition to, not replacing, crucial anthropogenic greenhouse gas 30 emission reductions and carbon dioxide removal - roadmaps can be a valuable tool to organize and streamline interdisciplinary 31 and multifaceted research to efficiently move towards understanding whether an approach may be viable and socially 32 acceptable, or if it is nonviable and further research should be deprioritized. Here we present a five-year research roadmap to 33 explore whether ISA enhancement of the chlorine radical sink could be a viable and socially acceptable atmospheric methane 34 removal approach.

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#### 35 1 Introduction

36 Driven by anthropogenic greenhouse gas emissions, Earth's average surface temperature has increased by 1.1 °C since 1850 37 (Forster et al., 2021). As global temperature increases, irreversible changes in the Earth system will likely occur, such as ice 38 sheet collapses in Greenland and Antarctica, coral reef die-off, abrupt permafrost thaw, mountain glacier loss, and Amazon 39 rainforest dieback (Lenton et al., 2008; McKay et al., 2022). To mitigate both near-term and long-term warming we must 40 undertake rapid, sustained action to establish a diverse portfolio of approaches to slow and then reverse the increase of 41 atmospheric greenhouse gas concentrations, ideally reducing them to preindustrial levels. Emissions reductions must be 42 prioritized. In addition, as near-term warming threatens to trigger climate tipping points, negative emissions approaches may 43 be used to remove greenhouse gases already in the atmosphere and counter rising natural and uncontrollable emissions.

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45 Methane (CH<sub>4</sub>) emissions have contributed roughly 0.5 °C to global warming relative to preindustrial times, second only to 46 carbon dioxide (CO2) (IPCC, 2023). Methane lasts roughly a decade in the atmosphere (UNEP, 2021), with a Global Warming 47 Potential 83 times that of carbon dioxide over 20 years and 30 times that of carbon dioxide over 100 years (Forster et al., 48 2021). The concentration of methane in the atmosphere is over 2.5 times pre-industrial levels, and the growth rate has 49 accelerated since 2006, with record increases in 2020 and 2021 (NOAA, 2023). Methane emissions come from anthropogenic 50 sources (fossil fuel use, agriculture, waste and wastewater, biomass burning, etc.) and natural sources (wetlands, oceans, 51 freshwaters, termites, permafrost, etc.), both of which are increasing (Jackson et al., 2020; Nisbet et al., 2023, Saunois et al., 52 2020). As the planet continues to warm and precipitation patterns change, natural methane emissions are expected to increase 53 from wetlands, as well as from permafrost due to abrupt thaw, thermokarst lake formation and expansion, and bacterial 54 processes (Dean et al., 2018; Neumann et al., 2019; Paudel et al., 2016; Peng et al., 2022; Zhang et al., 2023). On our current trajectory, natural emissions are estimated to increase by ~30-200 Tg CH4/yr by 2100 (Zhang et al., 2023; Kleinen et al., 55 56 2021). There is evidence that wetland emissions increases are already underway, with roughly half of the 2020 atmospheric 57 methane increase attributed to wetlands (Qu et al., 2022; Peng et al., 2022; Nisbet et al., 2023; Zhang et al., 2023).

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The natural sinks for atmospheric methane are oxidation by gas phase radicals (~95 %) and uptake into soil by methanotrophic bacteria and archaea (~5 %) (Saunois et al., 2020). The atmospheric lifetime of methane is thus mainly determined by the oxidative capacity of the atmosphere. Approximately 550 Tg methane per year is removed by reactions with tropospheric hydroxyl radicals (OH.); tropospheric chlorine radicals (Cl·) destroy ~11 Tg methane per year (Saunois et al., 2020). However, the tropospheric chlorine radical sink estimate is poorly constrained, ranging from 1 to 35 Tg methane per year (Saunois et al., 2020). Recent research suggests the tropospheric chlorine radical sink could be underestimated (van Herpen et al., 2023).

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66 Complex and non-linear atmospheric chemistry dictates the oxidation of methane, which results in the production of 67 formaldehyde and carbon monoxide which is further oxidized to carbon dioxide. Hydroxyl radical oxidation of methane can 68 result in the formation or loss of ozone, depending on the nitrogen oxide (NOx) concentrations. Chlorine radicals react to 69 remove both methane and ozone, the principal precursor of hydroxyl radical production (Lelieveld et al., 2002; Seinfeld and 70 Pandis, 2016). Therefore, chlorine radicals may either increase or decrease the atmospheric lifetime of methane depending on 71 the concentration of chlorine radicals and atmospheric conditions (Li et al., 2022, van Herpen et al., 2023, Li et al., 2023). 72 Volatile organic compounds and NOx will change the methane response since they directly and indirectly affect the 73 concentrations of hydroxyl and chlorine radicals. Furthermore, since reactions with carbon monoxide and methane are the 74 primary sinks for hydroxyl radicals, the abundance of methane impacts the oxidative character of the atmosphere in a self 75 feedback process (Lelieveld et al., 2002; Staniaszek et al., 2022; Holmes, 2018).

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77 Enhanced atmospheric oxidative sinks could increase the rate of atmospheric methane removal and therefore reduce near-term 78 warming. While not a replacement for much needed anthropogenic emissions reductions, enhancing atmospheric oxidative 79 sinks may be an important negative emission approach considering projected ongoing elevated natural methane emissions. 80 One untested proposal involves iron salt aerosols (ISA). This potential approach involves lofting iron-based particles into the 81 troposphere (e.g., from ships or towers) to catalytically produce chlorine radicals (Oeste, 2009; Oeste et al., 2017), mimicking 82 a natural phenomenon proposed to occur when mineral dust combines with sea spray aerosols (van Herpen et al., 2023). 83 Discussing natural analogues of this process and the current state of research, this paper presents a roadmap for research and 84 development that is needed to understand whether ISA enhancement of the chlorine radical sink may be a feasible, scalable, 85 and safe approach for atmospheric methane removal.

#### 86 2 State of the research

87 Currently, research into ISA falls into three categories: laboratory experiments to quantify the details of the mechanism, 88 observational analysis of the natural analogue of ISA, known as mineral dust sea spray aerosol (MDSA), and numerical 89 modeling evaluating the potential impacts of ISA.

### 90 2.1 Laboratory experiments

91 In a series of papers, Wittmer et al. demonstrated the production of chlorine atoms from iron-doped salts and aerosols (Wittmer 92 et al., 2015a; Wittmer et al., 2015b; Wittmer et al., 2017). Reproducing and expanding upon these laboratory studies is of the highest importance, since the mechanism of chlorine generation is poorly understood (van Herpen et al., 2023; Wittmer et al., 93 94 2015; Wittmer et al., 2017; Zhu et al., 1997). The ISA mechanism is catalytic in iron and may be catalytic in chlorine in natural 95 environments (Wittmer et al., 2017). Understanding of catalytic efficiency requires study of dependencies on conditions such 96 as aerosol size distribution and number density; humidity; rate of supply of acidity to the system; rate of coagulation of 97 aerosols; effects of organic chelating agents on iron activity; changes in chlorine escape probability due to aqueous chemical 98 conditions; changes in iron activity due to aerosol chemistry; and, understanding how the atmosphere behaves under high CIOx

- 99 conditions (Pennacchio et al., submitted). Furthermore, studies of real atmospheric conditions are needed to better understand
- 100 suppression of the ISA mechanism in the presence of sulfate and oxalate (Wittmer et al., 2015; Wittmer et al., 2017).

## 101 2.2 The natural analogue of ISA: mineral dust sea spray aerosol

102 The natural mineral dust sea spray aerosol (MDSA) mechanism is proposed to occur when iron from mineral dust mixes with 103 the chlorine in sea spray, forming iron chlorides which are photolyzed to produce chlorine radicals (van Herpen et al., 2023). 104 ISA could mimic this natural MDSA phenomenon by only aerosolizing what is currently believed to be the key component of 105 the mineral dust: the photoactive iron.

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107 van Herpen et al. (2023) provided the first evidence that methane may be removed over the North Atlantic by chlorine radicals 108 produced by the MDSA mechanism. North African semi-arid regions are the dominant source of iron-containing mineral dust, 109 with frequent transport over the North Atlantic (Prospero et al., 2021). Using air samples collected in Barbados during North 110 African dust events (Mak et al., 2003), a model parameterized with the MDSA mechanism of chlorine radical production 111 produced results consistent with a previously unexplained 13C-depletion in the reaction product carbon monoxide (CO) (van Herpen et al., 2023). Carbon monoxide produced from chlorine radical oxidation of methane is extremely depleted in 13C 112 113 which makes  $\delta 13C(CO)$  a very sensitive indirect detection method of chlorine radicals (Röckmann et al., 1999). The variability 114 of  $\delta 13C(CO)$  in atmospheric air is the main evidence that the MDSA process is occurring, as there is no other mechanism 115 proposed that can explain the carbon monoxide isotope signature (Mak et al., 2003).

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117 As a proxy for methane oxidation by chlorine radicals, studies of the isotopic composition of carbon monoxide in the mid-118 Atlantic boundary layer are underway to further confirm the MDSA mechanism. This includes a regular air sampling program 119 at four ground-based stations in the North Atlantic (located in Barbados, Canary Islands, Cape Verde, and Brazil), as well as 120 Atlantic transect sampling onboard commercial vessels. Time series at the ground-based stations provide high-resolution 121 observations at the longitudinal margins of trans-Atlantic dust transport, including the site where seasonal depletions in 122  $\delta 13C(CO)$  were previously observed (Mak et al., 2003). Air samples collected in north-south ship transects may allow spatial 123 correlation of the  $\delta 13C(CO)$  excursions with North African dust plumes. Together, these samples provide an opportunity to 124 investigate the formation process of MDSA, as well as the seasonal and spatial influence of North African dust on tropospheric 125 chlorine radical oxidation.

# 126 2.3 Modeling

Early modeling studies indicate that atmospheric chlorine additions can increase or decrease methane concentrations, depending on the concentration of chlorine that is present in the atmosphere (Horowitz et al., 2020; Li et al., 2022; Saiz-Lopez et al., 2023). Chlorine radicals readily oxidize ozone, the main precursor of hydroxyl radicals, which results in less methane oxidation via hydroxyls; at low atmospheric concentrations, chlorine radicals reduce ozone concentrations without having a 131 commensurate impact on methane. Even though chlorine radicals react 16x faster with methane than the reaction of hydroxyl 132 radicals with methane (Atkinson, 2006), hydroxyl radicals dominate the methane oxidation sink because they are much more 133 abundant than chlorine radicals. As more chlorine is emitted, ozone concentrations will be reduced so that proportionally more 134 chlorine radicals react with methane. The increased methane destruction by chlorine radicals will eventually outcompete 135 decreased destruction by hydroxyl. In an initial, highly simplified model scenario, Li et al. (2023) found that a reduction in 136 methane concentration could be achieved if more than 90 Tg Cl/yr (three times the estimated present-day emission rate) was 137 added evenly to the atmosphere over all ocean surfaces, and lowering the global methane burden by 2,000 Tg would require 138 the emission of an additional 1,000 Tg Cl2 /yr. However, assuming a uniform increase of chlorine over all ocean surfaces may 139 underestimate the potential effectiveness of local chlorine radical generation where efficiency may be condition-dependent 140 (e.g., NO<sub>x</sub>, CO, and chlorine concentrations, humidity, temperature, altitude, etc.) (Meidan et al., submitted). For example, 141 considering the MDSA natural analogue of ISA, van Herpen et al (2023) found that high dust concentrations in the North 142 Atlantic corresponded with net methane removal, while globally lower dust concentrations led to a net increase of methane. 143

144 Current models may not accurately capture the speed and efficiency of producing chlorine radicals via the ISA mechanism due 145 to assumptions of the percentage of photoactive iron in the emitted iron, aerosol pH, aerosol mixing rates, and more. Another 146 challenge with Earth system models is that they instantaneously dilute emissions to model grid dimensions which could lead 147 to underestimates or overestimates of the effectiveness of the ISA mechanism, especially when considering iron emission 148 additions from point sources like ships (e.g. Meidan et al., submitted). For example, the mixing of the iron and sea salt within 149 the aerosol is modeled to occur instantaneously (Meidan et al., submitted; van Herpen et al., 2023); however, in reality it would 150 likely take hours to days, leading the global model to overestimate the rate of chlorine radical production. Furthermore, the 151 ISA mechanism is likely to occur faster in high NOx environments (Oeste et al., 2017) but could be less efficient in high sulfate 152 environments (Bondy et al., 2017; Chen et al., 2020; Legrand et al., 2017; Pio et al., 1998), and both NOx and sulfate may be 153 co-emitted with iron (e.g. from a ship plume). Thus, models that instantaneously dilute emissions across the grid dimensions 154 may misrepresent the ISA mechanism. Overall, it is unclear whether current Earth system models overestimate or 155 underestimate the efficiency of the oxidation mechanism. Additional detailed box modeling focusing on deployment sites and 156 constrained by field observations are necessary to assess the effectiveness of the mechanism. However, local box models are 157 less reliable over timescales where mixing between different air masses is relevant. This motivates the need for high-resolution 158 regional and seasonal modeling over ocean basins and variable resolution configurations embedded in global models.

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160 Considering the difficulty of simulating variable atmospheric chemical conditions (e.g., atmospheric chemical composition, 161 solar radiation, wind mixing, etc.) across different geographic locations, it is important to develop an ensemble of models that 162 enable uncertainty assessment. Such a model ensemble will allow a comprehensive exploration of different iron salt aerosol 163 deployment scenarios (magnitude each year), aerosol particle sizes, and deployment location and timing.

#### 165 3 Roadmap

## 166 3.1 Roadmap framework

Roadmaps are used in climate research to define knowledge gaps, needs, and associated outputs as they relate to interdependencies and timelines, particularly in instances that benefit from integrated, interdisciplinary research. Recent examples include geochemical carbon dioxide removal (Masano et al., 2022), ocean-based carbon dioxide removal (Ocean Visions, 2023), ice sheet contributions to sea level rise (Aschwanden et al., 2021), and solar radiation management (Wanser et al., 2022). A coordinated, thorough, and science-based approach is needed to ensure that resources are used efficiently, stakeholders and interdisciplinary teams are engaged on appropriate timelines, and efforts are focused towards sequenced research questions and milestones.

#### 174 3.2 Viability assessment

175 The viability of an atmospheric methane removal approach can be assessed by considering its potential for feasibility, 176 scalability, and social license to operate. A feasible approach must be climate beneficial, safe, acceptable for its side effects, 177 and cost-plausible. Determination of scalability will be approach-specific, acknowledging that the scale of increased natural 178 emissions is anticipated to be tens of millions of tons of methane per year (Kleinen et al., 2021).

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The key milestone questions below can help determine the viability of ISA and whether it should continue to be prioritized.
The research that informs the key milestones questions should be pursued in parallel (Table 1).

- Is enhancement of the chlorine radical oxidative sink of methane via the ISA mechanism effective and climate beneficial? At what scale?
- What impacts could the ISA approach have on Earth systems and human systems, both positive and negative? Is there
   a cost-plausible ISA deployment method?
- 3. What is needed to advance a structure of ethical governance and social license for utilizing atmospheric intervention
   to reduce atmospheric methane concentrations?

## 188 3.2.1 Milestone question #1: Is the ISA mechanism effective and climate beneficial and scalable?

189 The complexity and nonlinearity of atmospheric chemistry and meteorology requires laboratory, field, and plume and global 190 modeling studies of the efficiency of chlorine radical production, its dependence on atmospheric conditions and other gases, 191 and the impact on methane removal.

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- An important assumption in previous studies is that only 1.8 % of iron is photoactive (van Herpen et al., 2023; Meidan et al., submitted). However, the amount of ISA that is photoactive may vary by emission source (e.g. ship emissions may have more
- 195 photoactive iron relative to mineral dust; Ito, 2013; Rodriguez et la., 2021), geographical location, aerosol pH, the presence of

other chemical constituents (e.g. sulfate and NOx), and altitude (Mahowald et al., 2018). Furthermore, the rate of chlorine production – and subsequent rate of methane oxidation – per mass of photoactive iron is estimated to result in the removal of 45 methane molecules per iron atom per day (van Herpen et al., 2023), but has many uncertainties including the time that iron remains in the atmosphere which may be impacted by large regional variability in deposition rates (Meidan et al., submitted). The efficiency, cost, safety (e.g. air quality), and net radiative forcing of ISA will depend on the percentage of iron that is photoactive, the rates of chlorine production and methane oxidation per mass of photoactive iron, and the lifetime of photoactive-iron based aerosol.

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Current studies assume that the chlorine radicals released from the photochemical reaction with iron will react (e.g., with methane) to form hydrochloric acid, which will then be reabsorbed back into the aerosol and thus recycled (van Herpen et al., 2023; Oeste et al., 2017). It is unclear under which atmospheric conditions this cycle occurs, but if some chlorine radicals are lost then the cycling would be less efficient. One potential mechanism by which the cycling efficiency could be reduced is if hydrochloric acid is produced and deposited into the ocean (Wang et al., 2019). Therefore, further laboratory measurements and detailed box models are needed to further study hydrochloric acid recycling efficiency by ISA.

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Better understanding of how sulfur dioxide and NOx concentrations impact the ISA mechanism is also needed (Oeste et al., 2017). Sulfur dioxide and NOx concentrations vary regionally and locally and their emissions may sometimes coincide with those of iron.

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Smaller ISA particles have greater surface area to mass ratios and stay longer in the atmosphere, likely increasing the efficiency of the ISA mechanism. Furthermore, smaller particles tend to be more acidic (Pye et al., 2020), which is required for the mechanism to be active (Wittmer et al., 2015; 2017). However, smaller aerosols may be transported further to coastal or inland locations where these particles could contribute directly to negative human health effects or deposit on terrestrial or ice-covered surfaces with unintended consequences. The aerosol size may also affect marine cloud cover, thereby influencing local radiative forcing. In addition, there tends to be more sulfate in smaller aerosols, which may reduce the effectiveness of the ISA mechanism (Bondy et al., 2017; Chen et al., 2020; Legrand et al., 2017; Pio et al., 1998).

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Studying MDSA (the natural analogue of ISA) through field studies is essential to understand the effectiveness of this mechanism under different atmospheric conditions and its geographical extent, thereby better constraining atmospheric and Earth system models. Early MDSA field studies are underway to explore the seasonality and spatial extent of methane oxidation by chlorine radicals that may occur in natural dust plumes through proxy measurements of  $\delta 13C(CO)$ . Further studies – both natural analogue and *in-situ* ISA enhancement studies – would benefit from alternative ISA detection and quantification approaches, including direct chlorine measurements or additional proxy measurements to reinforce existing observations.

Ideally, models will be developed to include the entire MDSA mechanism, including implementation of the isotope effect in the chlorine radical reaction with methane, thus enabling direct comparison of model results to observations of  $\delta$ 13C(CO). At present, some models (EMAC; Gromov 2014) include complete carbon monoxide isotope representation, but not the MDSA mechanism, whereas other models (CAM-CHem; van Herpen et al., 2023) include an initial representation of the MDSA mechanism but do not incorporate the isotopic effects.

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Isotopic signatures and dust from ice core paleo records may elucidate evidence of historical MDSA activity. Methane isotope measurements of air trapped in polar ice cores have been used to constrain the methane budget in the past (Bock et al. 2017; Sapart et al., 2012; Fischer et al., 2008; Ferretti et al., 2005), but possible variations in the chlorine-based methane sink have not been taken into account in these studies. Dust levels have undergone strong changes in the past (e.g. Fischer et al., 2007; Han et al., 2018; Yuan et al., 2020; Yue et al., 2023), and associated changes in MDSA may have affected the paleo records of  $\delta$ 13C(CH4).

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243 If modeling, laboratory studies, and natural analogue field studies prove that a promising, safe mechanism exists to produce 244 chlorine radicals at sufficient scale and under diverse atmospheric conditions, it may be appropriate to consider field studies 245 with intentional enhancement of ISA, using a broad suite of atmospheric measurements to understand how the mechanism 246 performs in the real atmosphere (see Section 3.2.3). Prior to performing any ISA enhancement field studies - even at a small 247 and controlled scale - it is essential to engage and work collaboratively with potentially impacted communities, policy and 248 science leaders, governmental bodies, NGOs, media, and other stakeholders to ensure that actions are conducted with social 249 license and an appropriate governance framework with free, prior, and informed consent (FAO, 2016). For example, a study 250 could involve controlled enhancement of dust or emitted aerosolized iron, or could investigate existing anthropogenic 251 emissions of iron (e.g. from a ship plume, power plant, iron foundry, etc). The scale of the study should be suitable to 252 accommodate a likely non-linear atmospheric response, where substantial increases in chlorine - thus iron emissions - may 253 be needed before there is a decrease in methane.

# 3.2.2 Milestone question #2: What are the potential Earth system and human systems impacts of ISA, and is it costplausible?

Lifecycle analyses are necessary to assess the potential benefits, tradeoffs, risks, uncertainties, and costs of ISA. As part of this analysis, understanding the potential impacts of ISA enhancement on the Earth system and human system is imperative before considering large scale deployment. Human system impacts may include human health outcomes, as well as indirect human impacts from Earth system changes. For example, if ISA resulted in ocean acidification there could be marine life implications resulting in economic, biodiversity, and cultural impacts for coastal communities. Furthermore, if chlorine drifts into urban areas it could stimulate ozone formation and cause negative human health impacts (Wang et al., 2020). Earth system modeling must be conducted to understand the impact of atmospheric conditions, aerosol size, and release locations, magnitudes, and timing, on ocean systems, terrestrial systems, the cryosphere, the stratosphere, and clouds and precipitation. Results from atmospheric modeling (plume and global), laboratory studies, and initial field studies will inform priorities and research directions for Earth system studies and could potentially inform the design of field studies to verify model results.

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269 The potential broader impacts of ISA deployment beyond removing methane are not well understood. These may include co-270 benefits such as reductions in atmospheric black carbon (Li et al., 2021, Oeste et al., 2017) and ozone (Li et al., 2023; van 271 Herpen et al., 2023), both of which are climate warming agents. However, there are also potential negative impacts that must 272 be further explored including ocean acidification, stratospheric ozone loss, adverse chemical side reactions (such as COCl2 273 formation), and albedo changes from deposition on ice surfaces (Li et al., 2023). Iron aerosols absorb light and thus tend to 274 warm the planet, offsetting some of the lowered radiative forcing from oxidized methane (Matsui et al., 2018; Li et al., 2021; 275 Meidan et al., submitted). There are also potential side effects such as indirect radiative forcing due to marine cloud brightening 276 and carbon dioxide absorption by ocean iron fertilization (Emerson, 2019) that could be either favorable or unfavorable.

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The potential effects of ISA on air quality and human health are also poorly understood. Enhanced chlorine could lead to beneficial reductions in tropospheric ozone. However, the reduced hydroxyl radical production may increase lifetimes of atmospheric trace species that may be detrimental to human health. Moreover, iron aerosols themselves present a human health risk, especially when small (O'Day et al., 2022). Therefore, more research is required to determine under which conditions (including deployment location/timing, particle composition, and ISA size) the air quality impact is beneficial or detrimental.

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Engineering deployment modalities and implementation scenarios is one of the later steps in this roadmap, only to be pursued if earlier dependencies are addressed and ISA proves effective and climate beneficial with acceptable side effects. Nevertheless, to avoid delaying potential future deployment-readiness and to iteratively refine design in advance of any *in-situ* field studies, development and engineering of a nozzle sprayer delivery system could begin in parallel with early research activities.

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As part of a lifecycle analysis, the cost of materials (e.g., iron), infrastructure, and other implementation resources must be assessed. To be cost-plausible, the cost per ton of methane removed must have a viable path to becoming lower than the social cost of methane, a monetary valuation that estimates the socioeconomic impact caused by an additional metric ton of methane (Azar et al., 2023).

#### 295 3.2.3 Milestone question #3: What is needed to advance ethical governance and social license?

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298 It is imperative that governance and social impacts be considered in parallel and iteratively with the development of any 299 atmospheric methane removal approach. Addressing the climate crisis requires engagement beyond technical solutions; 300 collaboration and transparency between physical scientists, behavioral scientists, media, the public, policy-makers, NGOs, Indigenous peoples, and other stakeholders is essential to ensure that an ethical governance framework is established (Dowell 301 302 et al., 2020; Diamond et al., 2022; Data for Progress, 2023; Carbon180, 2022). There must be effective engagement and 303 education to co-create research questions and iteratively communicate research findings and results, as well as risks and co-304 benefits. Failure to do so jeopardizes the trust and sound decision making of communities and governments (The Arctic 305 Institute, 2021), threatening our ability to critically and openly assess potential climate solution approaches through a scientific 306 process. Ideally, an external governance framework would be developed which is enforceable and legally binding; however, 307 there is also value in internal governance frameworks which may, for example, be based around a code of conduct, advisory 308 or review boards, or other non-binding structures.

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Table 1: Key research and development activities for a five-year timeline beginning in 2023. Funding is needed for all research and development activities, including those that are already underway.

	Research and development activity question #1: Is enhancement of the chlorine ial? At what scale?	Year in which work becomes a funding priority oxidative sink of	Anticipated duration (variable depending on outcomes) methane via the ir	Academia on salt aeros	Non- profits ol (ISA) me	Government echanism effect	Engineering consultant ive and
Modeling studies	Atmospheric plume and high-resolution regional modeling to understand if ISA- driven chlorine radical production results in net methane loss using various ISA sizes and concentrations of iron, NO., and other chemical species.	2023	3 years	Ø	Ø	Ø	

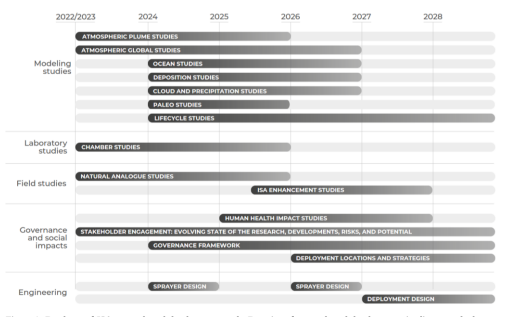
	Global atmospheric modeling to understand if ISA-driven chlorine radical production results in net cooling, considering radiative forcing from methane, ozone, and iron aerosols, the photoactivity of the iron, meteorological and atmospheric chemistry conditions, and ISA size.	2023	4 years	Ŋ	Ŋ	Q	
	Paleo modeling to understand variability in the chlorine-based methane sink that may have been associated with past dust changes.	2024	2 years	Ŋ	Ŋ	Ŋ	
Laboratory studies	Chamber studies to understand the photoactive iron fraction, chlorine radical production efficiencies, methane oxidation efficiencies, and reactions with other species.	2023	3 years	Ŋ	$\Sigma$	Ŋ	
Field studies	Natural analogue studies to (a) better constrain the atmospheric radical sinks; (b) understand the spatial extent and magnitude of the natural analogue, MDSA; and (c) understand the sources and distribution of photoactive iron (from MDSA as well as combustion iron sources).	2023	3 years	Ø	Ŋ	Q	
	In-situ ISA enhancement field studies (e.g. ship plume and/or ground based) at a controlled experiment site to measure iron speciation and changes in chlorine, methane, and other species.	2025	2+ years	Ŋ	Ŋ	Ø	
Engineering	Initial design study for ISA sprayer system.	2024	1 year				$\bigtriangledown$
	question #2: What impacts could the ISA app leployment method?	roach have on Ea	arth systems and h	uman system	s, both po	sitive and negat	ive? Is there a
Modeling studies	Modeling of ocean system impacts considering different release locations/timing, ISA sizes, and atmospheric conditions.	2024	3+ years	Ŋ	Ŋ	Ŋ	

	Modeling of ISA deposition impacts on different land and ice surfaces considering different release locations/timing, ISA sizes, and atmospheric conditions.	2024	3+ years	Ŋ	Ŋ	Ŋ	
	Modeling of atmospheric impacts including clouds, precipitation, and stratospheric ozone considering different release locations/timing, ISA sizes, and atmospheric conditions.	2024	3+ years	Ø	Ŋ	Ø	
	Lifecycle analysis to quantify ISA climate benefits, tradeoffs, and cost.	2024	4+ years	Ø	Ŋ	Ø	
Governance and social impacts	Study of potential human health impacts considering different release locations/timing, ISA sizes, and atmospheric conditions.	2025	3 years	Ø	Ŋ	ß	
	Advanced design study for ISA sprayer system.	2026	1 year				
Engineering	Design study of deployment modalities and implementation scenarios.	2027	2 years	Ø			Ø
	43: What is needed to advance a structure of ethane concentrations?	ethical governa	nce and social licen	se for utilizin	g atmosph	eric interventio	n to reduce
	Engage with stakeholders regarding the research findings and results, risks, and potential.	2023	4+ years		Ø	Ŋ	
Governance and social impacts	Develop a collaborative governance framework to monitor and report on environmental and social impacts.	2024	4+ years		Ŋ	Ŋ	
	Stakeholder engagement for identification of potential deployment locations and strategies.	2026	3 years			Ŋ	V

# 316 3.3 Priorities and timeline

317 It may be advantageous to pursue multiple research questions in parallel because the output from one research question may 318 inform the inputs for other research questions. As such, activities can be sequenced using a prioritized timeline (Table 1), 319 where later research activities and action areas often have multiple dependencies on earlier activities. For example, the 320 engineering design study is suggested to begin in year 4 (2027) because it can start prior to having complete Earth system 321 modeling results, human health study outcomes, or conclusions from ISA enhancement field studies. However, the engineering 322 design study cannot advance to its later stages until earlier activities have been thoroughly addressed. This expedited schedule 323 advances possible timelines to avoid delaying potential deployment-readiness, but is not meant to accelerate the timeline 324 beyond appropriate caution and due diligence. Setbacks in addressing early research questions and action areas will likely 325 result in timeline delays.





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Figure 1: Roadmap of ISA research and development needs. Duration of research and development timelines may be longer or shorter than depicted and/or exceed the five-year horizon.

# 330 4 Conclusion

The activities outlined in this roadmap require coordinated efforts across multiple government agencies to financially support 331 332 research and development, to ensure robust assessment and governance processes, and to foster international engagement. This 333 work is valuable for multiple reasons: 334 1. We need to understand if ISA is a feasible, scalable, and safe methane removal method, or if it is nonviable and 335 further research should be deprioritized. 336 2. Though this roadmap is ISA-specific, the research and development needs identified here contribute to 337 fundamental understanding of processes and mechanisms that are broadly applicable to exploration of other 338 methane removal approaches. 339 3. The research outlined in this roadmap will contribute to constraining the global methane budget and oxidative 340 character of the atmosphere, which will improve our understanding of atmospheric chemistry, Earth system 341 dynamics, and air quality. 342 343 Addressing the climate crisis requires a diverse portfolio of climate solutions. It is essential that aAtmospheric methane 344 removal approaches are should only be researched in addition to, not replacing, crucial anthropogenic greenhouse gas emission 345 reductions and carbon dioxide removal. Atmospheric methane removal approaches could play a future role in overall climate

change mitigation alongside aggressive anthropogenic emissions reductions, for example by dampening the impacts of anthropogenically-amplified natural methane emissions (e.g., from wetlands or permafrost thawing), particularly if they become uncontrollable due to climate change. All atmospheric methane removal approaches are at a very early stage (Jackson et al., 2021; Ming et al., 2022; Spark, 2023); all require further research and none are ready for deployment. We hope that this ISA roadmap, and other atmospheric methane removal roadmaps that follow, will help accelerate, prioritize, and parallelize research that is essential to understanding which climate solutions to pursue.

# 352 Author contribution

KG and SA wrote the manuscript draft. TJ, PH, NM, DM, MJ, MvH, YX, AS-L, TR, CB, ER, and DM contributed to, reviewed,
 and edited the manuscript.

# 355 Competing Interests

356 Spark Climate Solutions has in the past and anticipates in the future making research grants in alignment with this roadmap.

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#### 357 Acknowledgements

358	We thank two anonymous reviewers whose thoughtful and constructive comments improved the paper. We also thank Romany
359	Webb, Sabine Fuss, Jean-François Lamarque, Paige Brocidiacono, Eric Davidson, Rob Jackson, and Celina Scott-Buechler

360 for helpful discussions during preparation of the manuscript.

#### 361 References

Aschwanden, A., Bartholomaus, T. C., Brinkerhoff, D. J., and Truffer, M.: Brief communication: A roadmap towards credible
 projections of ice sheet contribution to sea level, The Cryosphere, 15, 5705–5715, https://doi.org/10.5194/tc-15-5705-2021,
 2021.

365

Atkinson, R., Baulch, D. L., Cox, R. A., Crowley, J. N., Hampson, R. F., Hynes, R. G., Jenkin, M. E., Rossi, M. J., Troe, J.,
 and IUPAC Subcommittee: Evaluated kinetic and photochemical data for atmospheric chemistry: Volume II – gas phase
 reactions of organic species, Atmos. Chem. Phys., 6, 3625–4055, https://doi.org/10.5194/acp-6-3625-2006, 2006.

- Azar, C., Martin, J.G., Johansson, D.J.A., and Sterner, T.: The social cost of methane, Climatic Change, 176,
   https://doi.org/10.1007/s10584-023-03540-1, 2023.
- 372

369

Bock, M., Schmitt, J., Beck, J., Seth, B., Chappellaz, J., and Fischer, H.: Glacial/interglacial wetland, biomass burning, and
geologic methane emissions constrained by dual stable isotopic CH4 ice core records, P. Natl. Acad. Sci. USA, 114, E5778E5786, https://doi.org/10.1073/pnas.1613883114, 2017.

376

380

381Carbon180,SettingDAConTrack:StrategiesforHubImplementation:382https://static1.squarespace.com/static/5b9362d89d5abb8c51d474f8/t/6261d1890b76863f1047a2dd/1650577901659/Carbon138380-SettingDAConTrack.pdf, last access: 7 December 2023, 2022.

- 384
- Chen, Y., Cheng, Y., Ma, N., Wei, C., Ran, L., Wolke, R., Größ, J., Wang, Q., Pozzer, A., Denier van der Gon, H. A. C.,
  Spindler, G., Lelieveld, J., Tegen, I., Su, H., and Wiedensohler, A.: Natural sea-salt emissions moderate the climate forcing of
  anthropogenic nitrate, Atmos. Chem. Phys., 20, 771–786, https://doi.org/10.5194/acp-20-771-2020, 2020.
- 388

<sup>Bondy, A.L., Wang, B., Laskin, A., Craig, R.L., Nhliziyo, M.V., Bertman, S.B., Pratt, K.A., Shepson, P.B., and Ault, A.P.:
Inland Sea Spray Aerosol Transport and Incomplete Chloride Depletion: Varying Degrees of Reactive Processing Observed
during SOAS, Environ. Sci. Technol., 51, 9533-9542, https://doi.org/10.1021/acs.est.7b02085, 2017.</sup> 

389	Data for Progress, Advancing Equitable Deployment of Regional DAC Hubs:
390	https://www.filesforprogress.org/memos/advancing-equitable-deployment-of-regional-dac-hubs.pdf, last access: 7 December
391	2023, 2023.
392	
393	Dean, J.F., Middelburg, J.J., Röckmann, T., Aerts, R., Blauw, L.G., Egger, M., Jetten, M.S.M., de Jong, A.E.E., Meisel, O.H.,
394	Rasigraf, O., Slomp, C.P., in't Zandt, M.H., and Dolman, A.J.: Methane Feedback to the Global Climate System in a Warmer
395	World, Rev. Geophys., 56, 207-250, https://doi.org/10.1002/2017RG000559, 2018.
396	
397	Diamond, M.S., Gettelman, A., Lebsock, M.D., McComiskey, A., Russell, L.M., Wood, R., and Feingold, G.: To assess
398	marine cloud brightening's technical feasibility, we need to know what to study - and when to stop, P. Natl. Acad. Sci. USA,
399	119, https://doi.org/10.1073/pnas.2118379119, 2022.
400	
401	Dowell, G., Niederdeppe, J., Vanucchi, J., Dogan, T., Donaghy, K., Jacobson, R., Mahowald, N., Milstein, M., and Zelikova,
402	T.J.: Rooting carbon dioxide removal research in the social sciences, Interface Focus, 10,
403	https://doi.org/10.1098/rsfs.2019.0138, 2020.
404	
405	Emerson, D.: Biogenic Iron Dust: A Novel Approach to Ocean Iron Fertilization as a Means of Large Scale Removal of Carbon
406	Dioxide From the Atmosphere, Front. Mar. Sci., 6, https://doi.org/10.3389/fmars.2019.00022, 2019.
407	
408	Fischer, H., Behrens, M., Bock, M., Richter, U., Schmitt, J., Loulergue, L., Chappellaz, J., Spahni, R., Blunier, T., Leuenberger,
409	M., and Stocker, T.F.: Changing boreal methane sources and constant biomass burning during the last termination, Nature,
410	452, 864-867, https://doi.org/10.1038/nature06825, 2008.
411	
412 413	Fischer, H., Siggaard-Andersen, M.L., Ruth, U., Rothlisberger, R., and Wolff, E.: Glacial/interglacial changes in mineral dust and sea-salt records in polar ice cores: sources, transport, and deposition. Rev. Geophys., 45,
413	https://doi.org/10.1029/2005RG000192, 2007.
415	
416	Ferretti, D.F., Miller, J.B., White, J.W., Etheridge, D.M., Lassey, K.R., Lowe, D.C., Meure, C.M., Dreier, M.F., Trudinger,
417	C.M., Van Ommen, T.D., and Langenfelds. R.L.: Unexpected changes to the global methane budget over the past 2000 years,
418	Science, 309, 1714-1717, https://doi.org/10.1126/science.1115193, 2005.
419	
420	Forster, P., T. Storelvmo, K. Armour, W. Collins, JL. Dufresne, D. Frame, D.J. Lunt, T. Mauritsen, M.D. Palmer,
421	M. Watanabe, M. Wild, and H. Zhang: The Earth's Energy Budget, Climate Feedbacks, and Climate
422	Sensitivity, Climate Change 2021: The Physical Science Basis, Contribution of Working Group I to the Sixth

DAC

Hubs:

423 Assessment Report of the Intergovernmental Panel on Climate Change, edited by: Masson-Delmotte, V., P. Zhai, A. Pirani,

424	S.L. Connors, C. Péan, S. Berger, N. Caud, Y. Chen, L. Goldfarb, M.I. Gomis, M. Huang, K. Leitzell, E. Lonnoy,
425	J.B.R. Matthews, T.K. Maycock, T. Waterfield, O. Yelekçi, R. Yu, and B. Zhou, Cambridge University Press,
426	Cambridge, United Kingdom and New York, NY, USA, 923-1054, doi:10.1017/9781009157896.009, 2021.
427	
428	Gromov, S.S.: Stable isotope composition of atmospheric carbon monoxide: a modeling study: https://openscience.ub.uni-
429	mainz.de/handle/20.500.12030/4250, last access: 7 December 2023, 2014.
430	
431	Han, C., Do Hur, S., Han, Y., Lee, K., Hong, S., Erhardt, T., Fischer, H., Svensson, A.M., Steffensen, J.P., and Vallelonga, P.:
432	High-resolution isotopic evidence for a potential Saharan provenance of Greenland glacial dust, Sci. Rep., 8,
433	https://doi.org/10.1038/s41598-018-33859-0, 2018.
434	
435	Holmes, C.D.: Methane Feedback on Atmospheric Chemistry: Methods, Models, and Mechanisms, J. Adv. Model Earth Sy.,
436	10, 1087-1099, https://doi.org/10.1002/2017MS001196, 2018.
437	
438	Horowitz, H.M., Holmes, C., Wright, A., Sherwen, T., Wang, X., Evans, M., Huang, J., Jaegle, L., Chen, Q., Zhai, S., and
439	Alexander, B.: Effects of sea salt aerosol emissions for marine cloud brightening on atmospheric chemistry: Implications for
440	radiative forcing, Geophys. Res. Lett., 47, https://doi.org/10.1029/2019GL085838, 2020.
441	
442	IPCC, AR6 Synthesis Report: Climate Change 2023: https://www.ipcc.ch/report/sixth-assessment-report-cycle/, last access 7
443	December 2023, 2023.
444	
445	Ito, A.: Global modeling study of potentially bioavailable iron input from shipboard aerosol sources to the ocean, Global
446	Biogeochem. Cy., 27, https://doi.org/10.1029/2012GB004378, 2013.
447	
448	Jackson, R.B., Abernethy, S., Canadell, J.G., Cargnello, M., Davis, S.J., Feron, S., Fuss, S., Heyer, A.J., Hong, C., Jones, C.D.,
449	Matthews, H.D., O'Conner, F.M., Pisciotta, M., Rhoda, H.M., de Richter, R., Solomon, E.I., Wilcox, J.L., and Zickfeld, K.:
450	Atmospheric methane removal: a research agenda, Phil. Trans. R. Soc. A, 379, https://doi.org/10.1098/rsta.2020.0454, 2021.
451	
452	Jackson, R.B., Saunois, M., Bousquet, P., Canadell, J.G., Poulter, B., Stavert, A.R., Bergamaschi, P., Niwa, Y., Segers, A.,
453	and Tsuruta, A., Increasing anthropogenic methane emissions arise equally from agricultural and fossil fuel sources, Environ.
454	Res. Lett., 15, https://doi.org/10.1088/1748-9326/ab9ed2, 2020.
455	
456	Kleinen T. Gromov S. Steil, B. and Brovkin, V.: Atmospheric methane underestimated in future climate projections

en, T., Gromov, S., Steil, B., and Brovkin, V.: Atmospheric methane underes Environ. Res. Lett., 16, https://doi.org/10.1088/1748-9326/ac1814, 2021. 457

458	
459	Legrand, M., Preunkert, S., Wolff, E., Weller, R., Jourdain, B., and Wagenbach, D.: Year-round records of bulk and size-
460	segregated aerosol composition in central Antarctica (Concordia site) - Part 1: Fractionation of sea-salt particles, Atmos.
461	Chem. Phys., 17, 14039–14054, https://doi.org/10.5194/acp-17-14039-2017, 2017.
462	
463	Lelieveld, J., Peters, W., Dentener, F.J., and Krol, M.C.: Stability of tropospheric hydroxyl chemistry, J. Geophys. Res., 107,
464	https://doi.org/10.1029/2002JD002272, 2002.
465	
466	Lenton, T.M., Held, H., Kriegler, E., Hall, J.W., Lucht, W., Rahmstorf, S., and Schellnhuber, H.J.: Tipping elements in the
467	Earth's climate system, P. Natl. Acad. Sci. USA, 105, 1786-1793, https://doi.org/10.1073/pnas.0705414105, 2008.
468	
469	Li, Q., Fernandez, R.P., Hossaini, R., Iglesias-Suarez, F., Cuevas, C.A., Apel, E.C., Kinnison, D.E., Lamarque, J.F., and Saiz-
470	Lopez, A.: Reactive halogens increase the global methane lifetime and radiative forcing in the 21st century, Nat. Commun.,
471	13, https://doi.org/10.1038/s41467-022-30456-8, 2022.
472	
473	Li, L., Mahowald, N. M., Miller, R. L., Pérez García-Pando, C., Klose, M., Hamilton, D. S., Gonçalves Ageitos, M., Ginoux,
474	P., Balkanski, Y., Green, R. O., Kalashnikova, O., Kok, J. F., Obiso, V., Paynter, D., and Thompson, D. R.: Quantifying the
475	range of the dust direct radiative effect due to source mineralogy uncertainty, Atmos. Chem. Phys., 21, 3973-4005,
476	https://doi.org/10.5194/acp-21-3973-2021, 2021.
477	
478	Li, Q., Meidan, D., Hess, P., Anel, J.A., Cuevas, C.A., Doney, S., Fernandez, R.P., van Herpen, M., Hoglund-Isaksson, L.,
479	Johnson, M.S., Kinnison, D.E., Lamarque, J.F., Röckmann, T., Mahowald, N.M., and Saiz-Lopez, A.: Global environmental
480	implications of atmospheric methane removal through chlorine-mediated chemistry-climate interactions, Nat. Commun., 14,
481	https://doi.org/10.1038/s41467-023-39794-7, 2023.
482	
483	Li, Y., Song, Y., Li, X., Kaskaoutis, D.G., Gholami, H., and Li, Y: Disentangling variations of dust concentration in Greenland
484	ice cores over the last glaciation: An overview of current knowledge and new initiative, Earth-Science Reviews, 242,
485	https://doi.org/10.1016/j.earscirev.2023.104451, 2023.
486	
487	Maesano, C.N., Campbell, J.S., Foteinis, S., Furey, V., Hawrot, O., Pike, D., Aeschlimann, S., Reginato, P.L., Goodwin, D.R.,
488	Looger, L.L., Boyden, E.S., and Renforth, P.: Geochemical Negative Emissions Technologies: Part II. Roadmap, Front. Clim.,
489	4, https://doi.org/10.3389/fclim.2022.945332, 2022.
490	

491	Mahowald, N.M., Hamilton, D.S., Mackey, K.R.M., Moore, J.K., Baker, A.R., Scanza, K.A., and Zhang, Y.: Aerosol trace
492	metal leaching and impacts on marine microorganisms, Nat. Commun., 9, https://doi.org/10.1038/s41467-018-04970-7, 2018.
493	
494	Mak, J., Kra, G., Sandomenico, T., and Bergamaschi, P.: The seasonality varying isotopic composition of the sources of carbon
495	monoxide at Barbados, West Indies, J. Geophys. Res., 108, https://doi.org/10.1029/2003JD003419, 2003.
496	
497	Matsui, H., Mahowald, N.M., Moteki, N., Hamilton, D.S., Ohata, S., Yoshida, A., Koike, M., Scanza, R.A., and Flanner, M.G.:
498	Anthropogenic combustion iron as a complex climate forcer, Nat. Commun., 9, https://doi.org/10.1038/s41467-018-03997-0,
499	2018.
500	
501	McKay, D.I.A., Stall, A., Abrams, J.F., Winkelmann, R., Sakschewski, B., Loriani, S., Fetzer, I., Cornell, S.E., Rockstrom, J.,
502	and Lenton, T.M.: Exceeding 1.5°C global warming could trigger multiple climate tipping points, Science, 377,
503	https://doi.org/10.1126/science.abn7950, 2022.
504	
505	Meidan, D.: Evaluating the potential of iron-based interventions in methane reduction and climate mitigation, submitted to
506	Environ. Res. Lett., https://doi.org/10.17632/v6mmbm75wd.1.
507	
508	Ming, T., Li, W., Yuan, Q., Davies, P., de Richter, R., Peng, C., Deng, Q., Yuan, Y., Caillol, S., and Zhou, N.: Perspectives
509	on removal of atmospheric methane., Advances in Applied Energy, 5, https://doi.org/10.1016/j.adapen.2022.100085, 2022.
510	
511	Neumann, R.B., Moorberg, C.J., Lundquist, J.D., Turner, J.C., Waldrop, M.P., McFarland, J.W., Euskirchen, E.S., Edgar,
512	C.W., and Turetsky, M.R.: Warming Effects of Spring Rainfall Increase Methane Emissions From Thawing Permafrost,
513	Geophys. Res. Lett., 46, 1393-1401, https://doi.org/10.1029/2018GL081274, 2019.
514	
515	Nisbet, E.G., Manning, M.R., Dlugokencky, E.J., Michel, S.E., Lan, X., Röckmann, T., Denier van der Gon, H.A.C., Schmitt,
516	J., Palmer, P.I., Dyonisius, M.N., Oh, Y., Fisher, R.E., Lowry, D., France, J.L., White, J.W.C., Brailsford, G., and Bromley,
517	T.: Atmospheric Methane: Comparison Between Methane's Record in 2006-2022 and During Glacial Terminations, Global
518	Biogeochem. Cy., 37, https://doi.org/10.1029/2023GB007875, 2023.
519	
520	NOAA Global Monitoring Laboratory, Earth System Research Laboratories, Trends in Atmospheric Methane:
521	https://gml.noaa.gov/ccgg/trends_ch4/, last access: 7 December 2023.
522	
523	Ocean Visions, Ocean-Based Carbon Dioxide Removal: Road Maps: https://www2.oceanvisions.org/roadmaps/, last access:

524 7 December 2023.

525								
526	O'Day, P.A., Pa	attammatt	el, A., Aronste	ein, P., Leppert, V	.J., Forman, H	.J.: Iron Speciation	in Respirable	Particulate Matter
527	and Implication	s for Hum	an Health, Env	viron. Sci. and Tech	nol., 56, 7006-	7016, https://doi.or	g/10.1021/acs.	est.1c06962, 2022.
528								
529	Oeste,	F.	D.,	Method	for	cooling	the	troposphere:
530	https://patentim	ages.stora	ge.googleapis.	.com/ac/64/72/49f3	3c79762e44f/A	U2010203265B2.p	df, last access:	7 December 2023,
531	2009.							
532								
533	Oeste, F. D., de	Richter, F	R., Ming, T., ar	nd Caillol, S.: Clim	ate engineering	g by mimicking nat	ural dust clima	te control: the iron
534	salt aerosol met	hod, Earth	n Syst. Dynam.	., 8, 1–54, https://d	oi.org/10.5194	/esd-8-1-2017, 201	7	
535								
536	Paudel, R., Mal	howald, N	I.M., Hess, P.O	G.M., Meng, L., ar	nd Riley, W.J.:	: Attribution of cha	anges in global	wetland methane
537	emissions from	n pre-indu	ustrial to pres	sent using CLM4	.5-BGC, Envi	ron. Res. Lett., 1	1, https://doi.	org/10.1088/1748-
538	9326/11/3/0340	20, 2016.						
539								
540	Pennacchio, L.,	, van Her	pen, M., Meid	lan, D., Saiz-Lope	z, A., and Joh	nnson, M.S.: Catal	ytic efficiencie	es for atmospheric
541	methane remova	al in the hi	igh-chlorine re	gime, submitted to	Environ. Res. 1	Lett., https://doi.org	g/10.26434/che	mrxiv-2023-3r8sf.
542								
543	Peng, S., Lin, X	., Thomps	son, R.L., Xi, Y	Y., Liu, G., Hauglu	staine, D., Lan	, X., Poulter, B., Ra	imonet, M., Sai	unois, M., Yin, Y.,
544	Zhang, Z., Zhen	ng, B., and	Ciais, P.: Wet	land emission and a	atmospheric sir	nk changes explain	methane growt	h in 2020, Nature,
545	612, 477–482, ł	nttps://doi.	.org/10.1038/s4	41586-022-05447-	w, 2022.			
546								
547	Pio, C.A., and L	Lopes, D.A	A.: Chlorine los	ss from marine aero	osol in a coasta	l atmosphere, J. Ge	ophys. Res., 10	)3, 25, 263 - 25, 272,
548	https://doi.org/1	0.1029/98	3JD02088, 199	18.				
549								
550	Prospero, J.M.,	Delany,	A.C., Delany	A.C., and Carlson	, T.N.: The Di	iscovery of African	n Dust Transpo	ort to the Western
551	Hemisphere and	d the Saha	aran Air Laye	r, B. Am. Meteoro	ol. Soc., 102, H	E1239-E1260, http:	s://doi.org/10.1	175/BAMS-D-19-
552	0309.1, 2021.							
553								
554	Pye, H. O. T., N	Nenes, A.,	Alexander, B.	, Ault, A. P., Barth	n, M. C., Clegg	, S. L., Collett Jr.,	J. L., Fahey, K.	M., Hennigan, C.
555	J., Herrmann, H	l., Kanakio	lou, M., Kelly,	, J. T., Ku, IT., M	cNeill, V. F., R	liemer, N., Schaefer	r, T., Shi, G., T	ilgner, A., Walker,
556	J. T., Wang, T.,	, Weber, F	R., Xing, J., Za	averi, R. A., and Z	uend, A.: The	acidity of atmosph	eric particles a	nd clouds, Atmos.
557	Chem. Phys., 20	0, 4809–48	888, https://doi	i.org/10.5194/acp-2	20-4809-2020,	2020.		

- Qu, Z., Jacob, D.J., Zhang, Y., Shen, L., Varon, D.J., Lu, X., Scarpelli, T., Bloom, A., Worden, J., and Parker, R.J.: Attribution 560 of the 2020 surge in atmospheric methane by inverse analysis of GOSAT observations, Envion. Res. Lett., 17, https://doi.org/10.1088/1748-9326/ac8754, 2022. 561 562 563 Röckmann, T., Brenninkmeijer, C.A.M., Crutzen, P.J., and Platt, U.: Short term variations in the 13C/12C ratio of CO as a 564 measure of Cl activation during tropospheric ozone depletion events in the Arctic, J. Geophys. Res., 104, 1691-1697, 565 https://doi.org/10.1029/1998JD100020, 1999. 566 Rodriguez, S., Prospero, J.M., Lopez-Darias, J., Garcia-Alvarez, M.I., Zuidema, P., Nava, S., Lucarelli, F., Gaston, C.J., 567 568 Galindo, L., and Sosa, E.: Tracking the changes of iron solubility and air pollutants traces as African dust transits the Atlantic 569 in the Saharan dust outbreaks, Atmos. Environ., 246, https://doi.org/10.1016/j.atmosenv.2020.118092, 2021. 570 571 Saiz-Lopez, A., Fernandez, R.P., Li, Q., Cuevas, C.A., Fu, X., Kinnison, D.E., Times, S., Mahajan, A.S., Martin, J.C.G., 572 Iglesias-Suerez, F., Hossaini, R., Plane, J.M.C., Myhre, G., and Lamarque, J.F.: Natural short-lived halogens exert an indirect 573 cooling effect on climate, Nature, 618, 967-973, https://doi.org/10.1038/s41586-023-06119-z, 2023. 574 575 Sapart, C.J., Monteil, G., Prokopiou, M., van de Wal, R.S.W., Kaplan, J.O., Sperlich, P., Krumhardt, K.M., Van der Veen, C., 576 Houweling, S., Krol, M.C., Blunier, T., Sowers, T., Martinerie, P., Witrant, E., Dahl-Jensen, D., and Röckmann, T.: Natural 577 and anthropogenic variations in methane sources during the past two millennia, Nature, 490, 85-88, 578 https://doi.org/10.1038/nature11461, 2012. 579 580 Saunois, M., Stavert, A. R., Poulter, B., Bousquet, P., Canadell, J. G., Jackson, R. B., Raymond, P. A., Dlugokencky, E. J., 581 Houweling, S., Patra, P. K., Ciais, P., Arora, V. K., Bastviken, D., Bergamaschi, P., Blake, D. R., Brailsford, G., Bruhwiler, 582 L., Carlson, K. M., Carrol, M., Castaldi, S., Chandra, N., Crevoisier, C., Crill, P. M., Covey, K., Curry, C. L., Etiope, G., 583 Frankenberg, C., Gedney, N., Hegglin, M. I., Höglund-Isaksson, L., Hugelius, G., Ishizawa, M., Ito, A., Janssens-Maenhout, 584 G., Jensen, K. M., Joos, F., Kleinen, T., Krummel, P. B., Langenfelds, R. L., Laruelle, G. G., Liu, L., Machida, T., Maksyutov, 585 S., McDonald, K. C., McNorton, J., Miller, P. A., Melton, J. R., Morino, I., Müller, J., Murguia-Flores, F., Naik, V., Niwa, Y., Noce, S., O'Doherty, S., Parker, R. J., Peng, C., Peng, S., Peters, G. P., Prigent, C., Prinn, R., Ramonet, M., Regnier, P., Riley, 586 587 W. J., Rosentreter, J. A., Segers, A., Simpson, I. J., Shi, H., Smith, S. J., Steele, L. P., Thornton, B. F., Tian, H., Tohjima, Y.,
- 588 Tubiello, F. N., Tsuruta, A., Viovy, N., Voulgarakis, A., Weber, T. S., van Weele, M., van der Werf, G. R., Weiss, R. F.,
  - 589 Worthy, D., Wunch, D., Yin, Y., Yoshida, Y., Zhang, W., Zhang, Z., Zhao, Y., Zheng, B., Zhu, Q., Zhu, Q., and Zhuang, Q.:
- 590 The Global Methane Budget 2000–2017, Earth Syst. Sci. Data, 12, 1561–1623, https://doi.org/10.5194/essd-12-1561-2020,
- 591 2020.
- 592

593	Seinfeld, J. H. and Pandis, S. N.: Atmospheric chemistry and physics: from air pollution to climate change. John Wiley &
594	Sons, ISBN 0471720186, 2016.
595	
596	Spark Climate Solutions, Approaches to Atmospheric Methane Removal: https://www.sparkclimate.org/methane-
597	removal/101/approaches, last access: 7 December 2023.
598	
599	Staniaszek, Z., Griffiths, P.T., Folberth, G.A., O'Connor, F.M., Abraham, NL., and Archibald, A.T.: The role of future
600	anthropogenic methane emission in air quality and climate. NPJ Clim. Atmos. Sci, 5, https://doi.org/10.1038/s41612-022-
601	00247-5, 2022.
602	
603	The Arctic Institute, Sami Council resistance to SCoPEX highlights the complex questions surrounding geoengineering and
604	consent: https://www.thearcticinstitute.org/sami-council-resistance-scopex-highlights-complex-questions-geoengineering-
605	consent/, last access: 7 December 2023, 2021.
606	
607	FAO, Food and Agriculture Organization of the United Nations, Free Prior and Informed Consent: An indigenous peoples'
608	right and a good practice for local communities: https://www.fao.org/3/i6190e/i6190e.pdf, last access: 7 December 2023, 2016.
609	
610	UNEP, United Nations Environment Programme, Global Methane Assessment: Benefits and Costs of Mitigating Methane
611	Emissions: https://www.unep.org/resources/report/global-methane-assessment-benefits-and-costs-mitigating-methane-
612	emissions, last access: 7 December 2023, 2021.
613	
614	van Herpen, M.M.J.W., Li, Q., Saiz-Lopez, A., Liisberg, J.B., Röckmann, T., Cuevas, C.A., Fernandez, R.P., Mak, J.E.,
615	Mahowald, N.M., Hess, P., Meidan, D., Stuut, JB.W., and Johnson, M.S.: Photocatalytic Chlorine Atom Production on
616	Mineral Dust-Sea Spray Aerosols over North Atlantic, P. Natl. Acad. Sci. USA, 120,
617	https://doi.org/10.1073/pnas.2303974120, 2023.
618	
619	Wang, X., Jacob, D. J., Eastham, S. D., Sulprizio, M. P., Zhu, L., Chen, Q., Alexander, B., Sherwen, T., Evans, M. J., Lee, B.
620	H., Haskins, J. D., Lopez-Hilfiker, F. D., Thornton, J. A., Huey, G. L., and Liao, H.: The role of chlorine in global tropospheric
621	chemistry, Atmos. Chem. Phys., 19, 3981–4003, https://doi.org/10.5194/acp-19-3981-2019, 2019.
622	
623	Wang, X., Jacob, D.J., Fu, X., Wang, T., Le Breton, M., Hallquist, M., Liu, Z., McDuffie, E.E., and Liao, H.: Effects of
624	Anthropogenic Chlorine on PM2.5 and Ozone Air Quality in China, Environ. Sci. Technol., 54, 9908-9916,
625	https://doi.org/10.1021/acs.est.0c02296, 2020.

627	Wanser, K., Doherty, S.J., Hurrell, J.W., and Wong, A.: Near-term climate risks and sunlight reflection modification: a
628	roadmap approach for physical science research, Climatic Change, 174, https://doi.org/10.1007/s10584-022-03446-4, 2022.
629	
630	Wittmer, J., Bleicher, S., Ofner, J., and Zetzsch, C.: Iron (III)-induced activation of chloride from artificial sea-salt aerosol.
631	Environ. Chem., 12, 461-475, https://doi.org/10.1071/EN14279, 2015a.
632	
633	Wittmer, J., Bleicher, S., and Zetzsch, C.: Iron (III)-induced activation of chloride and bromide from modeled salt pans, J.
634	Phys. Chem. A, 119, 19, 4373-4385, https://doi.org/10.1021/jp508006s, 2015b.
635	
636	Wittmer, J., and Zetzsch, C.: Photochemical activation of chlorine by iron oxide aerosol. J. Atmos. Chem., 74, 187-204,
637	https://doi.org/10.1007/s10874-016-9336-6, 2017.
638	
639	Yuan, T., Yu, H., Chin, M., Remer, L.A., McGee, D., and Evan, A.: Anthropogenic decline of African dust: Insights from the
640	Holocene records and beyond. Geophys. Res. Lett., 47, 22, https://doi.org/10.1029/2020GL089711, 2020.
641	
642	Zhang, Z., Poulter, B., Feldman, A.F., Ying, Q., Ciais, P., Peng, S., and Li, X.: Recent intensification of wetland methane
643	feedback, Nat. Clim. Change, 13, 430-433, https://doi.org/10.1038/s41558-023-01629-0, 2023.
644	
645	Zhu, X. R., Prospero, J. M., and Millero, F. J.: Diel variability of soluble Fe(II) and soluble total Fe in North African dust in

646 the trade winds at Barbados, J. Geophys. Res., 102, 21297–21305, https://doi.org/10.1029/97JD01313, 1997.