



Impact of mineral dust on the global nitrate aerosol direct and indirect radiative effect

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14 Abstract

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15 Nitrate (NO₃⁻) aerosol is projected to increase dramatically in the coming decades and may become the dominant inorganic particle species. This is due to the continued strong decrease in SO₂ emissions, which 16 is not accompanied by a corresponding decrease in NO_x and especially NH_3 emissions. Thus, the radiative 17 effect (RE) of NO₃⁻ aerosol may become more important than that of SO₄²⁻ aerosol in the future. The 18 19 physicochemical interactions of mineral dust particles with gas and aerosol tracers play an important role 20 in influencing the overall RE of dust and non-dust aerosols but can be a major source of uncertainty due to 21 their lack of representation in many global climate models. Therefore, this study investigates how and to 22 what extent dust affects the current global NO₃⁻ aerosol radiative effect through both radiation (RE_{ari}) and 23 cloud interactions (REaci) at the top of the atmosphere (TOA). For this purpose, multi-year simulations 24 nudged towards the observed atmospheric circulation were performed with the global atmospheric 25 chemistry and climate model EMAC, while the thermodynamics of the interactions between inorganic aerosols and mineral dust were simulated with the thermodynamic equilibrium model ISORROPIA-lite. 26 The emission flux of the mineral cations Na⁺, Ca²⁺, K⁺ and Mg²⁺ is calculated as a fraction of the total 27 aeolian dust emission based on the unique chemical composition of the major deserts worldwide. Our 28 29 results reveal positive and negative shortwave and longwave radiative effects in different regions of the 30 world via aerosol-radiation interactions and cloud adjustments. Overall, the NO₃⁻ aerosol direct effect 31 contributes a global cooling of -0.11 W/m², driven by coarse-mode particle cooling at short wavelengths. Regarding the indirect effect, it is noteworthy that NO_3^- aerosol exerts a global mean warming of +0.17 32 33 W/m^2 . While the presence of NO₃ aerosol enhances the ability of mineral dust particles to act as cloud 34 condensation nuclei (CCN), it simultaneously inhibits the formation of cloud droplets from the smaller 35 anthropogenic particles. This is due to the coagulation of fine anthropogenic CCN particles with the larger 36 nitrate-coated mineral dust particles, which leads to a reduction in total aerosol number concentration. This 37 mechanism results in an overall reduced cloud albedo effect and is thus attributed as warming.

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³⁹ Keywords: direct radiative effect, indirect radiative effect, nitrate aerosols, mineral dust





41 **1.** Introduction

Atmospheric aerosols are among the most complex components of the Earth's climate system. 42 This is due not only to the diversity of their origins, with many natural and anthropogenic emission 43 sources, but also to their extremely varied chemical composition and properties. The many 44 45 mechanisms by which they interact with each other and with physical entities such as radiation, clouds, land, and oceans add to their complexity and play a critical role in the energy balance of 46 the planet (Arias et al., 2021). The most direct way in which aerosols affect the Earth's energy 47 balance is through their interactions with solar shortwave (SW) and terrestrial longwave (LW) 48 49 radiation (IPCC, 2013). Overall, the radiative effect due to aerosol-radiation interactions (REari) is mainly dominated by the scattering of SW radiation back to space (negative radiative effect, 50 generating a cooling of the climate system) and the absorption of LW radiation (positive radiative 51 effect, generating a warming of the climate system) (Gao et al., 2018; Tsigaridis and Kanakidou, 52 2018). Aerosols belonging to the black and/or brown carbon family, together with mineral dust 53 54 particles, contribute to absorption (Kanakidou et al., 2005; Zhang et al., 2017; Wong et al., 2019), while the main inorganic aerosol components, such as sulfate and nitrate, as well as a significant 55 56 amount of organic carbon contribute mainly to scattering (Kirchstetter et al., 2004; (Bond and Bergstrom, 2006; Klingmüller et al., 2019; Zhang, 2020). However, mineral dust can also 57 influence the behavior of the RE_{ari} of anthropogenic pollution. Dust particles alter the 58 anthropogenic radiative effect of aerosol-radiation interactions by reducing the loading of 59 anthropogenic aerosols (either by coagulating with them or by adsorption of their precursor 60 inorganic trace gases), leading to less scattering of solar radiation and thus a warming effect (Kok 61 et al., 2023). 62

Atmospheric aerosols can also indirectly affect the Earth's energy balance by forming clouds, 63 controlling cloud optical thickness and scattering properties, and altering their precipitation and 64 lifetime (IPCC, 2013). Atmospheric aerosols act as cloud condensation nuclei (CCN), providing a 65 suitable surface for water vapor to condense, leading to the formation of liquid droplets that 66 67 develop into a corresponding liquid cloud (Lance et al., 2004). Such clouds are referred to as warm 68 clouds and are typically found in the lower troposphere (Khain and Pinsky, 2018). However, there is constant competition between small and large particles for the available amount of water vapor 69 70 (Barahona et al., 2010; Morales and Nenes, 2014). Under the same humidity conditions, the 71 presence of small particles will lead to the formation of small droplets with high number concentrations, while the presence of larger particles will lead to the formation of large droplets 72 but with lower number concentrations. Depending on the size characteristics of its particle 73 population, a warm cloud will exhibit different optical properties, with a population dominated by 74 smaller particles generally being more reactive in the SW spectrum. The change in cloud 75 reflectivity due to the presence of aerosols is referred to as the first radiative effect due to aerosol-76 cloud interactions (RE_{aci}) and was first described by Twomey (1977). The small size of 77 78 anthropogenic aerosols results in an overall smaller cloud droplet size, which reduces precipitation 79 efficiency and thus increases cloud lifetime. This contributes to cloud reflectivity and is referred to as the second radiative effect of aerosol cloud-interactions, first described by Albrecht (1989). 80 81 These two indirect effects are considered equally important for the total indirect radiative effect of aerosols (Lohmann and Feichter, 2005). Atmospheric aerosols exert a net cooling effect that can 82 83 partially mask the warming effect of greenhouse gases, therefore, the recent decline in





anthropogenic aerosol concentrations may accelerate global warming (Urdiales-Flores et al.,
2023). Overall, the radiative effect due to aerosol-cloud interactions is considered the main source
of existing uncertainty in the effective (total) radiative effect of aerosols in the atmosphere (Myhre
et al., 2014; Seinfeld et al., 2016).

88 Mineral dust influences the anthropogenic radiative effect through aerosol-cloud interactions in several ways that can result in either a net warming or net cooling effect. Dust particles can increase 89 the number of CDNC in remote areas since through chemical aging by pollutants (Nenes et al., 90 91 2014; Karydis et al., 2017), dust particles become more hygroscopic and require lower supersaturation thresholds for activation (Karydis et al., 2011). This is caused by the transfer of 92 93 anthropogenic pollutants towards remote desert regions which enhances the solubility of dust 94 particles. In such regions, this mostly results in increased cloud albedo and a net cooling effect. 95 However, dust particles also tend to reduce the availability of smaller anthropogenic CCN. This is due to intrusions of aged dust particles into polluted environments which reduce the numbers of 96 97 smaller aerosols through increased coagulation with them. This results in lower cloud reflectivity 98 (albedo) and thus a net warming effect (Klingmüller et al., 2020). Furthermore, when dust is above or below low-level clouds, the resulting effect of local heating is an increase in total cloud cover 99 100 due to enhanced temperature inversion or enhanced upward vertical motion, respectively (Kok et al., 2023). On the other hand, when dust is present inside low-level clouds, local heating enhances 101 in-cloud evaporation, resulting in an overall decrease in cloud cover. Kok et al. (2023) showed that 102 the amount of desert dust in the atmosphere has increased since the mid-19th century, causing an 103 overall cooling effect on the Earth that masks up to 8% of the warming caused by greenhouse 104 gases. If the increase in dust were halted, the previously hidden additional warming potential of 105 greenhouse gases could lead to slightly faster climate warming. 106

 NO_3 is expected to dominate the global aerosol composition in the coming decades due to the 107 predicted limited availability of SO_4^{-2} following the abrupt decline in SO_2 emissions, which will 108 not necessarily be accompanied by proportional reductions in NOx and NH₃ emissions (Bellouin 109 et al., 2011; Hauglustaine et al., 2014). Excess NO_3^{-1} is expected to exert a cooling RE_{ari} by 110 scattering SW radiation (Bauer et al., 2007a; Xu and Penner, 2012; Myhre et al., 2013; IPCC, 111 112 2013; Li et al., 2015), but the RE_{aci} is much more complex and complicated and can lead to both cooling and warming. Mineral dust thus becomes a key factor, as it is one of the main promoters 113 114 of NO_3^- aerosol formation, providing a very suitable surface for gaseous HNO₃ condensation to 115 the aerosol phase (Karydis et al., 2011; Trump et al., 2015). This affects not only the optical properties of dust aerosols, which will influence their overall RE_{ari}, but also how they can alter 116 cloud formation and microphysics. NO_3^- aerosols increase the hygroscopicity of mineral dust 117 (Kelly et al., 2007) by providing layers of soluble material on their surface, thus increasing their 118 119 ability to act as CCN (Karydis et al., 2017). In doing so, they also increase the size of dust particles through hygroscopic growth and therefore their coagulation efficiency. Thus, nitrate-dust 120 121 interactions are a complex mechanism that ultimately affects climatology in a variety of ways. The 122 role of mineral dust in modifying the influence of NO3⁻ aerosols in the global RE_{aci} is not yet well understood. This study aims to focus on the extent of the RE_{ari} and RE_{aci} of NO₃⁻ aerosols and on 123 how interactions with mineral dust regulate both on a global scale. 124

This study is organized as follows: in Section 2, details of the modeling setup for conducting the global simulations as well as the treatment of dust-nitrate interactions in the model are





- discussed and the methodology for calculating the global RE_{ari} and RE_{aci} of NO_3^- aerosols is explained. Section 3 presents the main results for the global RE_{ari} for coarse and fine NO_3^- aerosols
- 129 for the base case simulation and the sensitivity cases listed in Table 1. Section 4 presents the results
- 130 for the global RE_{aci} of total NO_3^- aerosols, while section 5 includes the feedback mechanism of
- 131 dust-nitrate interactions with cloud microphysics. Finally, the main conclusions and a general
- discussion on the scope of the study are presented in section 6.
- 133 <u>**Table 1:**</u> Differences between base case and sensitivity simulations performed.

Simulation Name	Conditions Applied
Base Case	Mineral dust ion composition according to Karydis et al. (2016)
Sensitivity 1: Chemically Inert Dust	Mineral dust emitted exclusively as a chemically inert bulk particle
Sensitivity 2: Homogeneous Ion Composition	Global homogeneous ionic composition of mineral dust particles according to Sposito (1989)
Sensitivity 3: Half Dust Scenario	50% reduced dust emission flux.
Sensitivity 4: Increased Dust Scenario	50% increased dust emission flux.

134 2. Methodology

135 2.1 Model Setup

The simulations were performed with the global atmospheric chemistry and climate model 136 EMAC (ECHAM/MESSy) (Jockel et al., 2006), which includes several submodels describing 137 138 atmospheric processes and their interactions with oceans, land, and human influences. These submodels are linked through the Modular Earth Submodel System (MESSy) (Jockel et al., 2005) 139 to a base model, the 5th Generation European Center Hamburg General Circulation Model 140 (ECHAM) (Roeckner et al., 2006). The submodel system used in this work includes the MECCA 141 submodel, which performs the gas phase chemistry calculations (Sander et al., 2019). The SCAV 142 submodel is responsible for the in-cloud liquid-phase chemistry and wet deposition processes (Tost 143 et al., 2006; Tost et al., 2007b), while DRYDEP and SEDI are used to compute the dry deposition 144 of gases and aerosols and gravitational settling, respectively (Kerkweg et al., 2006). All aerosol 145





146 microphysical processes are calculated by the GMXe submodel (Pringle et al., 2010a; Pringle et al., 2010b), where aerosols are divided into 4 lognormal size modes (nucleation, Aitken, 147 accumulation and coarse). Each mode is defined in terms of aerosol number concentration, number 148 mean dry radius, and geometric standard deviation (sigma). The mean dry radius for each mode is 149 allowed to vary within fixed bounds $(0.5 \text{ nm} - 6 \text{ nm} \text{ for nucleation}, 6 \text{ nm} - 60 \text{ nm} \text{ for Aitken}, 60 \text$ 150 nm - 700 nm for accumulation, and above 700 for coarse) and the sigma is fixed and equal to 1.59 151 for the first three size modes and 2 for the coarse mode. The coagulation of aerosols is also handled 152 153 by GMXe, following Vignati et al. (2004) and the coagulation coefficients for Brownian motion are calculated according to Fuchs and Davies (1964). The partitioning between the gas and aerosol 154 155 phases is calculated using the ISORROPIA-lite thermodynamic module (Kakavas et al., 2022) as 156 implemented in EMAC by Milousis et al. (2024). The optical properties of the aerosols and the radiative transfer calculations are simulated by the submodels AEROPT (Dietmuller et al., 2016) 157 and RAD (Dietmuller et al., 2016), respectively. AEROPT can be called several times within a 158 model time step with different settings for the aerosol properties. More details are given in section 159 2.3.1. All cloud properties and microphysical processes are simulated by the CLOUD submodel 160 (Roeckner et al., 2006) using the two-moment microphysical scheme of Lohmann and Ferrachat 161 (2010) for liquid and ice clouds. The activation processes of liquid cloud droplets and ice crystals 162 follow the physical treatment of Morales and Nenes (2014) and Barahona and Nenes (2009), 163 164 respectively, as described by Karydis et al. (2017) and Bacer et al. (2018). More details are given 165 in Section 2.3.2.

The meteorology for each of the simulations was nudged by ERA5 reanalysis data (C3S, 2017), thus this study estimates the radiative effect of nitrate aerosols with respect to RE_{ari} and RE_{aci} separately, rather than the effective (total) radiative effect, as this would require multiple free-run simulations with prescribed sea surface temperatures for each case separately. The spectral resolution used for each simulation was T63L31, which corresponds to a grid resolution of 1.875° x 1.875° and 31 vertical layers up to 25 km in height. The period covered by the simulations is from 2007 to 2018, with the first year representing the model spin-up period.

Anthropogenic aerosol and trace gas emissions were taken from the CMIP6 database (O'Neill 173 174 et al., 2016) according to the SSP370 scenario. Natural NH₃ emissions (from land and ocean) were based on the GEIA database (Bouwman et al., 1997), and natural volcanic SO₂ emissions were 175 taken from the AEROCOM database (Dentener et al., 2006). Biogenic NO emissions from soils 176 were calculated online according to the algorithm of Yienger and Levy (1995), while lightning-177 produced NO_x was also calculated online by the LNOx submodel (Tost et al., 2007a) using the 178 179 parameterization of Grewe et al. (2001). DMS emissions from the oceans are calculated online by the AIRSEA submodel (Pozzer et al., 2006). Sea salt emissions are based on the AEROCOM 180 database (Dentener et al., 2006) following the chemical composition reported by Seinfeld and 181 Pandis (2016), i.e. 30.6% Na⁺, 3.7% Mg⁺, 1.2% Ca²⁺, 1.1% K⁺, and 55% Cl⁻. Dust emissions are 182 calculated online using the parameterization of Astitha et al. (2012). In this scheme, while the 183 184 surface friction velocity is the most important parameter for the amount of the emitted dust flux, the meteorological information for each grid cell is also taken into account. Dust particles are 185 emitted in the accumulation and coarse size modes of the insoluble fraction, but can be transferred 186 187 to the soluble fraction after either coagulation with other soluble species and/or by condensation 188 of soluble material on their surface. Both processes are treated and calculated by GMXe and





ISORROPIA-lite. The emissions of mineral ions (Ca²⁺, Mg²⁺, K⁺, and Na⁺) are estimated as a 189 fraction of the total dust emission flux based on the soil chemical composition of each grid cell. 190 This is done using desert soil composition maps from Klingmüller et al. (2018) which are based 191 on the mineral ion fractions from Karydis et al. (2016). To assess the impact of changes in mineral 192 dust chemistry and emissions on the global NO3⁻ aerosol RE_{ari} and RE_{aci}, four additional sensitivity 193 simulations were performed (Table 1). In the first sensitivity simulation, mineral dust is described 194 only by a bulk, chemically inert species. In the second sensitivity case, the chemical composition 195 196 of the mineral dust was assumed to be spatially uniform, with a percentage distribution for bulk dust, Na^+ , K^+ , Ca^{2+} and Mg^{2+} particles assumed to be 94%, 1.2%, 1.5%, 2.4% and 0.9% 197 respectively according to Sposito (1989). Finally, two additional simulations were performed to 198 199 assess the impact of the global mineral dust budget on the results, where the dust emission fluxes were first halved and then increased by 50% to account for the historical increase in global dust 200 mass load since pre-industrial times, as reconstructed by Kok et al. (2023). 201

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203 2.2 Treatment of Dust-Nitrate Interactions

204 The interactions between mineral dust and nitrate aerosols play a crucial role in altering the size 205 distribution and optical properties of both species and can also strongly influence cloud 206 microphysical processes (Fig. 1). Therefore, these interactions affect both the RE_{ari} and the RE_{aci} of both nitrate and dust aerosols. First, the adsorption of HNO₃ onto the surface of dust particles 207 208 is a process that strongly promotes the formation of nitrate aerosols on dust (Karydis et al., 2016). 209 We treat this condensation process using the GMXe submodel. Specifically, the amount of gas phase species that kinetically condenses within a model time step (equal to 10 minutes in this 210 study) is calculated according to the diffusion-limited condensation theory of Vignati et al. (2004). 211 This information is then passed to the ISORROPIA-lite thermodynamic module to calculate the 212 213 gas/aerosol partitioning.

Specifically, the module receives as input the ambient temperature and humidity along with the 214 diffusion-limited concentrations of H₂SO₄, NH₃, HNO₃, and HCl, the concentrations of the non-215 volatile cations (NVCs) Na⁺, K⁺, Ca²⁺ and Mg²⁺, and the concentrations of the ions SO_4^{2-} , NO_3^{-} , 216 NH₄⁺, and Cl⁻ present in the aerosol phase from the previous time step. The module then calculates 217 the equilibrium reactions of the NO_3^- anion with the NVCs, depending on their abundance with 218 respect to the SO_4^{2-} anion, taking into account mass conservation, electroneutrality, water activity 219 equations and precalculated activity coefficients for specific ionic pairs (Fountoukis et al., 2007; 220 221 Kakavas et al., 2022). Therefore, in all cases where mineral dust is considered chemically active, all reactions of nitrate aerosols with NVC are treated. The salts that may be formed are assumed 222 223 to be completely deliquesced as follows:

224
$$Ca(NO_3)_2 \rightarrow Ca^{2+}_{(aq)} + 2NO^-_{3(aq)}$$

225 $NaNO_3 \rightarrow Na^+_{(aq)} + NO^-_{3(aq)}$

226
$$KNO_3 \rightarrow K^+_{(aq)} + NO^-_{3(aq)}$$

227
$$Mg(NO_3)_2 \rightarrow Mg^{2+}_{(aq)} + 2NO^-_{3(aq)}$$





More information on equilibrium reactions and equilibrium constants as well as the corresponding thermodynamic equilibrium calculations can be found in Fountoukis and Nenes (2007).

The coating of dust particles by nitrate aerosols during gas/aerosol partitioning calculations is 231 an important process that leads to an increase in dust solubility and hygroscopicity (Laskin et al., 232 233 2005). Therefore, after these processes have taken place, a large fraction of the originally insoluble 234 dust particles has become soluble (Fig. 1a), which leads to changes in their optical properties, as their increased ability to absorb water makes them more efficient in extinguishing SW radiation 235 236 and absorbing and emitting LW radiation (Fig. 1a, 1b) (Kok et al., 2023). The transfer to the soluble fraction after coating with soluble material is handled by the GMXe submodel, which also provides 237 238 key aerosol attributes necessary for the calculation of the dust optical properties (see Section 2.3). 239



240 Figure 1: Conceptual illustration of how dust-nitrate interactions affect the total NO₃- (left) RE_{ari} and 241 (right) RE_{aci}. a) In dust-rich environments, nitric acid transported from anthropogenic pollution and biomass 242 burning regions interacts with mineral cations to form a soluble coating on the surface of dust particles. The 243 dominant effect of these interactions is an enhanced LW absorption (warming RE_{ari}) by the coarse dust 244 particles. b.) In nitrate-rich environments, the intrusion of dust particles and their subsequent interaction 245 with freshly emitted nitric acid leads to an overall increase in aerosol hygroscopicity and thus a stronger SW reflection (cooling RE_{ari}). c.) In dust-rich environments, the number of ice crystals in cirrus clouds is 246 247 reduced while their size is increased due to the interaction of dust particles with the transported HNO₃. This 248 results in an optical thinning of the ice clouds, which leads to less trapping of outgoing LW radiation 249 (cooling RE_{aci}). d.) In nitrate-rich environments, the increased wet radius of aged dust particles leads to 250 enhanced coagulation with smaller particles, resulting in a decrease in the number of smaller aerosols and, in turn, a decrease in the number of activated particles in cloud droplets by smaller aerosols, which 251 252 ultimately leads to a reduction in the backscattering of SW radiation by warm clouds (warming RE_{ari}).





253 In general, the changes in the properties of dust particles through their interactions with nitrate aerosols will result in more efficient removal rates, mainly through wet deposition, due to their 254 higher hygroscopicity and increased size (Fan et al., 2004). The reduced number of dust particles 255 that can act as ice nuclei (IN) and their increased size can lead to an optical thinning of cirrus 256 257 clouds (Fig. 1c) (Kok et al., 2023). Furthermore, the changes induced by dust-nitrate interactions reduce the activation of smaller aerosols in warm clouds (Fig. 1d). In particular, the enhanced 258 259 hygroscopicity of dust particles will lead to a faster depletion of the available supersaturation, as 260 they act as giant CCN that absorb large amounts of water vapor to activate into cloud droplets (Karydis et al., 2017). In addition, the population of smaller aerosols will also be depleted by 261 262 increased coagulation with the large dust particles. As a consequence of the different degrees of complexity of the dust-nitrate interactions, it is very important to note that they do not always 263 result in a linear response in terms of how they affect climate through their subsequent interactions 264 with radiation, clouds, or both. 265

266

267 2.3 **Radiative Effect Calculation**

To calculate the global RE_{ari} and RE_{aci} of NO_3^- aerosols, the optical properties from the 268 AEROPT submodel and the radiative transfer calculations from the RAD submodel were used. 269 First, AEROPT provides the aerosol extinction (absorption and scattering) coefficients, the single 270 271 scattering albedo, and the aerosol asymmetry factor for each grid cell with a vertical distribution 272 analogous to the vertical resolution used. The GMXe submodel is used to provide input of aerosol attributes for the calculation of aerosol optical properties, which is done online using 3D look-up 273 tables. The tables provide information on the real and imaginary parts of the refractive index and 274 the Mie size parameter per size mode (Dietmuller et al., 2016). Then, the radiative scheme of RAD 275 uses the particle number weighted average of the extinction cross section, the single scattering 276 277 albedo, and the asymmetry factor as input for the radiative transfer calculations. In addition to AEROPT, RAD takes input from the submodels ORBIT (Earth orbital parameters), CLOUDOPT 278 279 (cloud optical properties) (Dietmuller et al., 2016), and IMPORT (import of external datasets) to 280 calculate the radiative transfer properties for longwave and shortwave radiation fluxes separately. Both the AEROPT and RAD submodels can be invoked multiple times within a model time step, 281 each time with different settings for the aerosol optical properties, allowing radiative transfer 282 estimates for identical climatological conditions. This is of paramount importance for the 283 calculation of the REari of aerosols since any effects due to possibly different climatological 284 conditions must be eliminated. Henceforth, all references to RE estimates, as well as net, 285 longwave, and shortwave flux quantities, will refer to the top of the atmosphere (TOA) only. 286

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2.3.1 Radiative Effect from Aerosol-Radiation Interactions (REari)

To estimate the global RE_{ari} of all aerosols as well as that of total, coarse, and fine NO₃⁻ aerosols,</sup> 288 289 3 simulations were performed for each sensitivity case in Table 1. In the first simulation all aerosol species are present, in the second simulation NO3⁻ aerosols are completely removed by turning off 290 291 their formation, and in the third simulation coarse mode NO_3^{-1} aerosols are removed by forcing 292 HNO₃ to condense only on the fine mode (i.e., sum of the three smaller lognormal size modes: 293 nucleation, Aitken, and accumulation). For each of these three simulations, the radiative transfer 294 routines are called twice for each time step. One call uses the normal aerosol optical properties, 295 and the other call uses an aerosol optical depth equal to 0 to emulate an atmosphere without





296 aerosols. Essentially, the global RE_{ari} of each simulation can be calculated by taking the difference 297 between the net fluxes between the two calls. More specifically, the first simulation will yield the REari of the total aerosol load (F1,ari hereafter), the second simulation will yield the REari of all 298 aerosols except NO₃⁻ (F_{2.ari} below), and the third simulation will yield the RE_{ari} of all aerosols 299 except the coarse mode NO_3^- (F_{3,ari} below). Since the above estimates of the radiative effect were 300 computed using the exact same climatology, its effect was effectively eliminated. However, in 301 302 order to isolate the NO₃⁻ aerosol radiative effect, it is also essential to disable any aerosol-cloud 303 interactions, otherwise the cooling effect would be severely underestimated because cloud scattering would make aerosol scattering less relevant (Ghan et al., 2012). For this purpose, the 304 305 simplest cloud scheme available in the EMAC model is used, which calculates cloud microphysics according to Lohmann and Roeckner (1996) and cloud coverage according to Tompkins (2002). 306 307 To disable aerosol-cloud interactions, no aerosol activation routines are used to avoid coupling with the activation schemes. Overall, the global RE_{ari} of total, coarse, and fine NO_3^- aerosols are 308 obtained as follows: 309

310 •
$$F_{NO3,ari}(F_{N,ari}) = F_{1,ari} - F_{2,ari}$$

311 •
$$F_{coarseNO3,ari}(F_{cN,ari}) = F_{1,ari} - F_{3,ari}$$

312 •
$$F_{fineNO3,ari}(F_{fN,ari}) = F_{3,ari} - F_{2,ari}$$

313 2.3.2 Radiative Effect from Aerosol-Cloud Interactions (REaci)

In this work we estimate the effect of total NO_{3} - aerosols on the calculated global RE_{aci}. 314 Climatology plays a crucial role in aerosol-cloud interactions and simulating a "fine-only NO₃-315 atmosphere", as done for the REari calculations, would produce an unrealistic climatological 316 scenario, since coarse-mode NO₃⁻ is strongly associated with cations in mineral dust particles 317 318 (Karydis et al., 2016), making them quite effective as CCN (Karydis et al., 2017). Therefore, the RE_{aci} calculations require only 2 simulations for each sensitivity case separately: one with all 319 aerosols present and one with the entire NO₃⁻ aerosol load removed by turning off their formation. 320 321 The global RE_{aci} is then given by:

$$\bullet \quad F_{NO3,aci}(F_{N,aci}) = FF_N - F_{N,ari}$$

where FF_N is the total NO₃⁻ aerosol feedback radiative effect. Since $F_{N,ari}$ is calculated using the 323 324 methodology described in Section 2.3.1, it is only necessary to estimate FF_N . This is equal to the 325 difference in net fluxes between the two simulations. There is no need to emulate an aerosol-free 326 atmosphere here since any differences induced by different climatologies must be included. The two simulations performed for the calculation of RE_{aci} use the cloud formation scheme as described 327 in Lohmann and Ferrachat (2010), which uses prognostic equations for the water phases and the 328 329 bulk cloud microphysics. In addition, the empirical cloud cover scheme of Sundqvist et al. (1989) is used. For aerosol activation, the CDNC activation scheme of Morales and Nenes (2014) is used, 330 331 which includes the adsorption activation of mineral dust as described in Karydis et al. (2017). The 332 effect of dust-nitrate interactions on clouds presented here refers to the lowest level of cloud formation at 940 hPa. For the ICNC activation, the scheme of Barahona and Nenes (2009) is used, 333 334 which calculates the ice crystal size distribution through heterogeneous and homogeneous freezing 335 as well as ice crystal growth.





336 3. Radiative Effect from Aerosol-Radiation Interactions (RE_{ari})

337 3.1 Base Case

The global average RE_{ari} of total NO_3^- aerosols at the top of the atmosphere was found to be -338 0.11 W/m², which is within the reported range of the estimated present day all-sky direct radiative 339 effect of total NO₃⁻ aerosols by other studies (Liao et al., 2004; Bauer et al., 2007a; Bauer et al., 340 2007b; Xu and Penner, 2012; Myhre et al., 2013; IPCC, 2013). The NO₃⁻ cooling of the RE_{ari} 341 342 calculated by EMAC is driven by the scattering of SW radiation (equal to -0.34 W/m^2), which outweighs the warming due to absorption of LW radiation (equal to $+0.23 \text{ W/m}^2$) (Table 2). The 343 344 RE_{ari} of the total NO_3^- aerosol is dominated by the coarse particles, as they account for 82% of the calculated SW cooling and all the LW warming (Table 2; Figure 2). The warming effect of the 345 coarse particles is strongest over the dust belt zone and especially over the regions of the Middle 346 East and the northern face of the Himalayan plateau. These regions are characterized by high 347 concentrations of coarse NO₃⁻ aerosols due to the adsorption of HNO₃ on desert soil particles 348 (Karydis et al., 2016; Milousis et al., 2024). Therefore, the warming due to absorption of terrestrial 349 LW radiation by coarse-mode nitrates interacting with mineral dust is the strongest over these 350 areas (Fig. 1a), reaching up to +4 W/m² (Fig. 2iii). On the other hand, the scattering of solar SW 351 radiation is higher over regions with higher concentrations of total NO₃⁻ aerosols. Over the USA 352 and Europe, the SW RE_{ari} is -1.5 W/m² (Fig. 2v). However, the cooling exerted by nitrate aerosol 353 through the SW RE_{ari} is more profound over areas where it interacts strongly with high 354 concentrations of mineral dust particles (Fig. 1b). Such areas include the Congo Basin, where 355 HNO₃ from tropical forest biomass burning interacts with Saharan mineral dust particles; the North 356 Indian region, where anthropogenic HNO3 emissions interact with mineral dust particles from the 357 Taklimakan desert; and the East Asian region, where HNO₃ emissions from Chinese megacities 358 359 interact with mineral dust particles from the Gobi Desert. These regions can result in an average cooling of up to -4 W/m^2 (Fig. 2v). 360

Interestingly, cooling through SW interactions is not evident over the Sahara Desert, most likely 361 362 due to the minimal presence of nitrate aerosols in the $PM_{2.5}$ size range (not shown). The high concentrations of nitrate aerosols in the PM_{2.5-10} range over the Sahara contribute to LW 363 absorption, but SW scattering is almost exclusively associated with the regions dominated by high 364 concentrations in the PM_{2.5} (and smaller) size range (i.e. East Asia). Moreover, nitrate aerosols 365 366 over the Sahara are exclusively associated with dust particles which have a relatively high imaginary part of the refractive index, with typical values for SW wavelengths being ~ 0.005 (Di 367 Biagio et al., 2019). As a result, nitrate aerosols over the Sahara exhibit relatively strong absorption 368 369 in the SW spectrum as well, and the combination of the bright surface of the desert below leads to a weakened cooling effect and even sometimes in localized warming. 370

In contrast to the radiative effect of coarse NO_3^- particles, the RE_{ari} of fine NO_3^- particles due to interactions with LW radiation is negligible (Fig. 2iv), while its effect on SW radiation is about an order of magnitude smaller than that induced by the coarse NO_3^- aerosols (Fig. 2vi). The cooling induced by fine NO_3^- aerosols from the scattering of SW radiation is stronger over regions of high anthropogenic activity, namely the East Asian and Indian regions, which are characterized by high concentrations of fine NO_3^- aerosols, and peaks over eastern China (-0.25 W/m²). Conversely, a weak warming effect (+0.15 W/m²) is calculated over deserts (e.g. Gobi, Sahara).







Figure 2: Global mean TOA net RE_{ari} for (i) coarse and (ii) fine NO3 aerosols; longwave RE_{ari} for (iii)
 coarse and (iv) fine NO3 aerosols; shortwave RE_{ari} for (v) coarse and (vi) fine NO3 aerosols, as calculated
 by EMAC from the base case simulation.





- 382 **<u>Table 2</u>**: Net, longwave, and shortwave global mean TOA RE_{ari} of total, coarse, and fine NO₃⁻
- aerosols for the base case and each sensitivity case simulations.
- 384

385

Simulation	Aerosol Component	T	OA RE _{ari} (W/n	n ²)
	-	Net	LW	SW
	Total NO ₃ -	- 0.11	+ 0.23	- 0.34
Base Case	Coarse NO ₃ -	- 0.09	+ 0.23	- 0.32
	Fine NO ₃ ⁻	- 0.02	~ 0	- 0.02
	Total NO ₃ ⁻	- 0.09	+ 0.11	- 0.20
Chemically Inert Dust	Coarse NO ₃ -	- 0.07	+ 0.11	- 0.18
	Fine NO ₃ ⁻	- 0.02	~ 0	- 0.02
	Total NO ₃ ⁻	- 0.09	+ 0.18	- 0.27
Homogeneous Ion Composition	Coarse NO ₃ -	- 0.08	+ 0.18	- 0.26
	Fine NO ₃ -	- 0.01	~ 0	- 0.01
	Total NO ₃ -	- 0.08	+ 0.19	- 0.27
Half Dust Scenario	Coarse NO ₃ -	- 0.06	+ 0.19	- 0.25
	Fine NO ₃ -	- 0.02	~ 0	- 0.02
	Total NO ₃ -	- 0.10	+ 0.27	- 0.37
Increased Dust Scenario	Coarse NO ₃ -	- 0.08	+ 0.27	- 0.35
	Fine NO ₃ -	- 0.02	~ 0	- 0.02





387 3.2 Sensitivity of REari Estimates

388 The comparison of the calculated total NO_3^- radiative effect due to interactions with net, LW, 389 and SW radiation for the sensitivity cases listed in Table 1 can be found in Table 2, which shows each of the estimates. Consideration of nitrate interactions with mineral dust cations can greatly 390 affect the NO3⁻ REari estimates. Assuming that mineral dust particles are inert, the estimated 391 392 warming due to LW radiation interactions is 52% weaker than in the base case where dust 393 reactivity is considered. Similarly, the cooling effect exerted by nitrate aerosols through interactions with SW radiation is estimated to be 41% weaker under the assumption that mineral 394 dust is non-reactive. Both estimates are lower when mineral dust is assumed to be chemically inert, 395 since HNO₃ is no longer effectively adsorbed on dust particles and therefore the RE_{ari} by coarse 396 NO_3^- aerosol is significantly weaker compared to the base case where it dominates the total NO_3^- 397 398 effect (see Sect. 3.1). However, since both the estimated warming and cooling are weaker, the 399 effects partially cancel each other out, resulting in a net cooling effect (-0.09 W/m^2) that is 18% 400 weaker compared to the base case calculations. Assuming a homogeneous ionic composition for 401 the dust, results in SW cooling and LW warming being 21% and 22% lower, respectively, weakening the estimate for the net cooling RE_{ari} by 18% (-0.09 W/m²). The net direct radiative 402 403 effect of total NO_3^- is the same for the cases where dust is assumed to have a homogeneous 404 chemical composition and where it has no chemical identity, indicating the importance of both 405 aspects for the impact of dust-nitrate interactions on the direct radiative effect.

406 In the Half Dust scenario, the LW warming estimate is 17% weaker than in the base case, while 407 the SW estimate is even more so (21%), resulting in a lower net cooling estimate of -0.08 W/m². Finally, the Increased Dust scenario shows the strongest LW warming effect (17% increase over 408 409 the base case) due to an increase in coarse mode nitrate. At the same time, the cooling effect due 410 to interactions with SW radiation shows a smaller increase of 9%. Thus, accounting for the historical increase in mineral dust emissions results in a net cooling estimate of -0.10 W/m^2 , which 411 is smaller than the base case. Interestingly, the behavior of the global total $NO_3^- RE_{ari}$ does not 412 exhibit linearity with respect to the global dust load. This is not surprising since the nitrate-dust 413 interactions themselves are not linearly correlated, and a given increase or decrease in dust 414 415 emissions does not lead to an analogous change in nitrate aerosol levels. For example, Karydis et al. (2016) have shown that moving from a scenario in which nitrate-dust chemistry is not 416 417 considered to one in which it is, but with half dust emissions, resulted in a 39% increase in the tropospheric burden of nitrate aerosols. However, moving from a scenario with half to full dust 418 419 emissions, the corresponding increase was only 9%. In our case, moving from the chemically inert dust scenario to the half dust scenario led to an 18% increase in atmospheric nitrate aerosol burden, 420 while moving from the half dust scenario to the base case led to an additional 8% increase, and 421 422 finally moving from the base case to the increased dust scenario led to an even smaller increase of 5%. 423

424 **4** Radiative Effect from Aerosol-Cloud Interactions (RE_{aci})

425 **4.1 Base Case**

The global average RE_{aci} of total NO_3^- aerosols at the top of the atmosphere was found to be +0.17 W/m². In contrast, an estimate of the RE_{aci} of nitrate aerosols by Xu and Penner (2012) showed only a trivial cooling effect for particulate NO_3^- (-0.01 W/m²). Similar to the RE_{ari}, the net





RE_{aci} estimated by EMAC is driven by the effect on the SW part of the spectrum, which causes a 429 warming effect of +0.27 W/m², while the effect on the LW radiation causes an average cooling of 430 -0.10 W/m² (Table 3). Overall, the net RE_{aci} of total NO₃⁻ aerosols is reversed compared to the net 431 REari, i.e. REaci exerts a strong cooling effect over regions where REari exerts a warming effect and 432 433 vice versa (Fig. 3i). The reason for this is that the regions contributing to a cooling RE_{ari} are dominated by smaller sized nitrate aerosols and vice versa. Therefore, the size characteristics of 434 the dominant nitrate aerosol population lead to different effects on the cloud optical properties as 435 436 discussed in section 1. For example, as the dominance of smaller nitrate aerosols decreases over a particular region, the optical thinning of low-level clouds will have an opposite effect on the REact 437 438 (Fig. 1d). Details of the mechanism by which nitrate-dust interactions affect cloud microphysical processes are discussed in section 5. Over North America and Europe, RE_{aci} causes a warming 439 effect of up to $+3 \text{ W/m}^2$, driven solely by the effect on SW radiation (Fig. 3iii). Over the regions 440 of East Asia and the Amazon and Congo basins, RE_{aci} reaches a maximum of +5 W/m², driven by 441 both the effect on the SW (up to $+4 \text{ W/m}^2$) and LW (up to $+1.5 \text{ W/m}^2$) parts of the radiation 442 spectrum. The cooling effect of RE_{aci} (up to -2 W/m²) extends mainly between the equatorial line 443 and the Tropic of Cancer, mainly due to the interaction of nitrate aerosols with desert dust particles 444 (e.g. from the Sahara) and their effect on the terrestrial spectrum (LW) (Figs. 1c & 3ii). The cooling 445 effect of dust interactions with anthropogenic particles in the LW spectrum corroborates the 446 447 findings of Klingmüller et al. (2020) and is attributed to the reduced ice-water path due to the 448 depletion of small aerosols, which in turn leads to less trapped outgoing terrestrial radiation. In addition, Kok et al. (2023) note how the presence of dust particles leads to an optical thinning of 449 cirrus clouds by reducing the number of ice crystals while increasing their size, which also leads 450 to less trapping of outgoing LW radiation and thus a cooling effect (Fig. 1c). On the other hand, 451 the warming effect of dust interactions with anthropogenic particles in the SW spectrum requires 452 further investigation and is therefore discussed in more detail in Section 5. 453







454 **Figure 3:** Global mean TOA RE_{aci} for total NO_3^- aerosols. Estimates for (i) net, (ii) longwave, and (iii) 455 shortwave, as calculated by EMAC from the base case simulation.





456	Table 3: Net, longwave, and shortwave global mean TOA REaci of total NO3 ⁻ aerosols for the
457	base case and each sensitivity case simulations.

Simulation		TOA REaci (W/m ²))
	Net	LW	SW
Base Case	+ 0.17	- 0.10	+ 0.27
Chemically Inert Dust	+ 0.11	- 0.06	+ 0.17
Homogeneous Ion Composition	+ 0.13	- 0.09	+ 0.22
Half Dust Scenario	+ 0.15	- 0.08	+ 0.23
Increased Dust Scenario	+0.14	- 0.11	+ 0.25

458

459 **4.2 Sensitivity of RE**aci Estimates.

Table 3 shows the comparison of the net, LW, and SW contributions of total NO_3^{-1} to the RE_{aci} 460 at the top of the atmosphere as calculated by the base case simulation and all sensitivity cases 461 considered. By assuming a chemically inert dust, the calculated net RE_{aci} of nitrate decreases by 462 35%, resulting in a net warming of +0.11 W/m². As with the RE_{ari} estimate, this sensitivity case 463 produces the largest deviation from the base case among all sensitivity simulations, for both the 464 SW (37% less warming) and LW (40% less cooling) estimates. This is due to the fact that the 465 absence of dust-nitrate interactions does not have such a large impact on the population of both 466 aerosols and activated particles (see also Section 5). The assumption of a homogeneous ionic 467 composition of the mineral dust leads to a weakened LW cooling estimate of 10% and a weakened 468 SW warming estimate of 19% resulting in a net NO₃⁻ RE_{aci} of +0.13 W/m² (24% lower than in the 469 470 base case).

The reduced dust emissions result in a 15% weaker warming in the SW spectrum and a 20% 471 weaker cooling in the LW spectrum, leading to an overall $NO_3^- RE_{aci}$ of +0.15 W/m² (12% weaker 472 than the base case scenario). This is because the reduced loading of nitrate aerosols, especially in 473 the coarse mode, in the half dust scenario results in less absorption of LW radiation (Fig. 1c) (hence 474 less cooling). Similarly, the effect of dust-nitrate interactions on the activation of smaller particles 475 476 (Fig. 1d) is less drastic and results in a weaker inhibition of SW radiation scattering (hence less warming, see also Section 5). Finally, increased dust emissions in the increased dust scenario show 477 a 10% increase in the LW cooling and an 8% decrease in the SW warming effect, surprisingly 478 resulting in a net warming $(+0.14 \text{ W/m}^2)$ that is lower than in the half dust scenario. The reason 479 480 that this scenario results in more LW cooling than the base case is that the increased amount of dust particles leads to even more optical thinning of the ice clouds, and therefore even less trapping 481 482 of LW radiation (more cooling). However, the reason why the SW warming estimate is lower than the base case is more complicated. First, the transition from the half dust scenario to the base case 483 and then to the increased dust scenario does not lead to an analogous increase in the nitrate aerosol 484





485 burden (see Section 3.2). Moreover, since the number of aerosols has increased from the increased dust scenario to the base case, but the relative humidity has remained largely the same, there is 486 more competition for water vapor because it is now distributed over a larger population. As a 487 result, the wet radius increase in the presence of nitrates is not as strong in the increased dust 488 scenario compared to the base case, and the depletion of smaller sized particles is also not as strong 489 (not shown). The implications of the depletion of the aerosol population in the presence of nitrate 490 aerosols on the microphysical processes of warm clouds, and consequently on SW warming, are 491 492 discussed in the next section.

493 **5** Effect Of NO₃⁻ Aerosols on Cloud Microphysics

494 5.1 Maximum Supersaturation, Hygroscopicity and Wet Radius

To further investigate the cause of the positive RE_{aci} induced by the NO₃⁻ aerosols, their effect 495 on the characteristics of the aerosol population as well as on the cloud microphysics is investigated, 496 with respect to the lowest forming cloud level of 940 hPa. For this purpose, a sensitivity simulation 497 is performed assuming a 'nitrate aerosol free' (NAF) atmosphere, in which the formation of NO₃-498 499 aerosols has been switched off, but an advanced cloud scheme is considered. This simulation is used to determine whether the presence of NO_3^- aerosols has a significant effect on the 500 501 hygroscopicity and size of atmospheric aerosols and ultimately on the maximum supersaturation 502 developed during cloud formation. Over polluted areas affected by transported dust air masses from surrounding arid areas, the presence of NO_3^{-1} aerosols can increase the CCN activity of the 503 504 large mineral dust particles, resulting in a reduction of the maximum supersaturation and inhibiting the activation of the small anthropogenic particles into cloud droplets (Klingmüller et al., 2020). 505 Results from the NAF sensitivity simulation support this hypothesis over parts of Eastern and 506 Central Asia, where the maximum supersaturation decreases by up to 0.05%. In contrast, the 507 508 presence of NO_3^- aerosols increases maximum supersaturation by up to 0.2% over North America, Europe, the Middle East, and parts of southern Asia (Fig. 4ii). Therefore, changes in maximum 509 supersaturation caused by the presence of NO₃⁻ aerosols cannot explain their warming effect 510 511 through the REaci.

The presence of NO_3^{-} has a significant effect on the hygroscopicity of both fine and coarse 512 aerosols and consequently on their wet radius, as shown in Figures 1a, b & 4. This is most evident 513 for coarse desert dust particles, which mix with NO_3^- aerosols from urban and forest regions, 514 increasing their hygroscopicity by an order of magnitude (up to 0.1), especially over the African-515 516 Asian dust belt and the Atacama Desert in South America (Fig. 4vi). Aerosol hygroscopicity is 517 similarly increased for the fine mode particles both near arid regions and over the highly industrialized region of Southeast Asia (Fig. 4iv). On the other hand, the aerosol hygroscopicity 518 for the two size modes is only slightly reduced, by up to 0.06 (or <10%) over the oceans and coasts 519 520 of Europe and East Asia, due to interactions of NO_3^- with sea salt particles, reducing their hygroscopicity. The increased ability of both coarse dust aerosols and smaller aerosols to absorb 521 water leads to an increase in their wet radius, but in different parts of the world. For example, fine 522 523 particle sizes increase by up to 0.04 µm (up to 40%) mostly over regions of high anthropogenic 524 activity (North America, Europe, and East Asia) (Fig. 4viii). On the other hand, coarse mode particle sizes are increased by up to 0.1 µm (up to 10%) over the forests of central Africa and the 525





- 526 African-Asian dust belt zone (Fig. 4x), while showing a similar decrease near the coasts of the
- 527 polluted northern hemisphere due to the effect of NO_3^- on the hygroscopicity of sea salt.













Change in Fine Aerosol Wet Radius











Figure 4: (i) Global mean maximum supersaturation, fine aerosol (iii) hygroscopicity and (v) wet radius, and coarse aerosol (vii) hygroscopicity and (ix) wet radius, as calculated by EMAC from the base case simulation at the altitude of 940 hPa. Absolute difference between base case and Nitrate Aerosol Free (NAF) sensitivity simulation in (ii) maximum supersaturation, fine aerosol (iv) hygroscopicity and (vi) wet radius, and coarse aerosol (viii) hygroscopicity and (x) wet radius at the altitude of 940 hPa. Red indicates higher values calculated by the base case simulation in the presence of NO₃⁻ aerosols.

537 5.2 Number Concentrations of Aerosol and Activated Particles.

Figure 5 shows the effect of NO_3^- on the number concentration of fine and coarse aerosols 538 between the base case and the 'NAF' sensitivity simulation, as well as the total aerosol population. 539 The presence of NO₃⁻ aerosols decreases the total aerosol number concentration over forests and 540 541 polluted regions (see also Fig. 1d). This behavior is driven solely by the decrease in smaller particle sizes, as the effect is minimal for the coarser particles (Figs. 5ii & 5iv). The largest decrease is 542 calculated over East and South Asia (up to 1000 cm⁻³ or 10%), while decreases of up to 200 cm⁻³ 543 544 on average (~10%) are found over Europe, the USA, and Central Africa. This effect is directly related to the increased wet radius of the aerosol population (Fig. 4viii) over these regions and thus 545 546 to its depositional efficiency. In addition, coarse dust particles become more hygroscopic due to interactions with NO_3^- aerosols that increase in size, resulting in increased coagulation with the 547 smaller anthropogenic particles, which reduces their abundance. 548

549 The reduced aerosol number concentration in the presence of NO₃⁻ can lead to a reduction of particles that are also activated into cloud droplets. Such behavior can be seen in Figure 6, which 550 shows the effect of NO_3^- on the number concentration of activated fine and coarse particles in 551 cloud droplets between the base case and the 'NAF' sensitivity simulation. The reduction in the 552 total number of activated cloud droplets is almost entirely due to the reduction in smaller size 553 particles (Figs. 6ii & 6iv). A reduction in the total number of activated droplets of up to 30 cm⁻³ or 554 10% is observed over the USA, Amazon, Europe, Central Africa, and parts of the Middle East, 555 while this reduction reaches up to 100 cm⁻³(10%) over Southeast Asia, where the largest reductions 556 in aerosol numbers are also calculated (Fig. 4ii). In turn, these are the regions where the warming 557 effect of NO₃⁻ aerosols on the calculated mean RE_{aci} is strongest (Figure 3i). The small increase in 558 activated droplets (~ 10 cm⁻³ or 1%) over Beijing, which concerns the fine mode particles, is most 559 likely because their number concentration decreases with increasing size. The high aerosol number 560 concentration there, which is the global maximum (Figure 5i), results in a hotspot of more readily 561 activated particles in the presence of NO_3^- . On the other hand, the CDNC decreases slightly over 562 563 the Sahara due to the more efficient deposition capacity of coarse dust particles due to their 564 interactions with nitrate aerosols, which is also reflected in the decrease in aerosol number (Fig. 6iv). Overall, the lower particle number in the presence of NO_3^- aerosols hinders the ability of the 565 smaller anthropogenic particles to activate into cloud droplets, leading to a reduced cloud cover 566 and thus a reduced cloud albedo effect. Therefore, not only less LW radiation is absorbed, but 567 568 more importantly, less SW radiation is scattered back to space, resulting in an overall warming of the net average RE_{aci} for total NO₃⁻ aerosols. 569







572 Figure 5: Global mean number concentration of (i) fine and (iii) coarse aerosols as calculated by EMAC
573 from the base case simulation at the altitude of 940 hPa. Absolute difference between the base case and the
574 Nitrate Aerosol Free (NAF) sensitivity simulation in the number concentration of (ii) fine and (iv) coarse
575 aerosols at the altitude of 940 hPa. Blue indicates that number concentrations are lower in the presence of
576 NO₃⁻ aerosols.







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Figure 6: Global mean number concentration of activated (i) fine and (iii) coarse aerosols as calculated by
 EMAC from the base case simulation at the altitude of 940 hPa. Absolute difference between the base case
 and the Nitrate Aerosol Free (NAF) sensitivity simulation in the number concentration of activated (ii) fine
 and (iv) coarse aerosols at the altitude of 940 hPa. Blue indicates that number concentrations are lower in
 the presence of NO₃⁻ aerosols.

590

591 6. Conclusions and Discussion

This study presents the effects of interactions between mineral dust and NO_3^- aerosols on the present-day global TOA radiative effect of the latter. We investigate how the presence of dust affects the radiative effect of NO_3^- aerosols, both through aerosol interactions with radiation and separately with clouds (RE_{ari} and RE_{aci}, respectively). Sensitivity simulations are also performed, varying both the mineral dust composition and its emissions, to assess their effect on the calculated NO_3^- aerosol radiative effect.

598 It was found that the global average net RE_{ari} of total NO₃⁻ aerosols is -0.11 W/m², which is 599 mainly due to the cooling from the shortwave part of the radiation spectrum due to scattering,





equal to -0.34 W/m². A warming from the longwave part of the spectrum due to absorption was 600 found to be +0.23 W/m² on global average and was mainly located over regions with high 601 concentrations of coarse NO₃⁻ aerosols. SW cooling was also observed in these regions, but also 602 over regions of high anthropogenic activity, mainly over the polluted northern hemisphere. The 603 overall sign of the net RE_{ari} for total NO₃⁻ aerosols was dominated by the behavior of the coarse 604 mode particles, which contributed on average -0.09 W/m^2 . Specifically, coarse nitrate particles 605 were responsible for 82% of the net cooling and 100% of the net warming, while the fine mode 606 607 played a minor role. The sensitivity experiments revealed that the chemistry of the mineral dust is the most important factor in changing the estimated RE_{ari} of the total NO_3^- aerosols. In particular, 608 LW warming is most affected by this assumption, being 52% weaker after assuming chemically 609 inert dust emissions, while the SW cooling is reduced by 41% compared to the base case 610 simulation, amounting to a net cooling of -0.09 W/m². A globally homogeneous ionic composition 611 for mineral dust had a smaller effect in LW (22% decrease) and SW (21% decrease) but resulted 612 in the same net estimate of -0.09 W/m^2 . Halving the dust emissions resulted in weaker estimates 613 for LW and SW by 17% and 21%, respectively, and the lowest overall net RE_{ari} of -0.08 W/m². 614 On the other hand, a 50% increase in dust emissions increased both LW warming and SW cooling 615 by 17% and 9% respectively, resulting in a net cooling RE_{ari} of -0.10 W/m², indicating the strong 616 non-linear relationship of nitrate-dust interactions and how they affect the radiative effect 617 estimates. 618

The global average net RE_{aci} of total NO₃⁻ aerosols was +0.17 W/m² due to the effect on the 619 shortwave portion of the spectrum. This was found to be $+0.27 \text{ W/m}^2$, while the cooling from the 620 longwave part was -0.10 W/m². Spatially, the net RE_{aci} is reversed compared to the net RE_{ari} for 621 total NO_3^- aerosols, where regions responsible for a strong SW cooling of the RE_{ari} contribute to a 622 strong SW warming of the REaci and vice versa. This is due to the fact that nitrate-dust interactions 623 624 challenge the dominance of smaller particles over heavily polluted regions, reducing the reflectivity of warm cloud and thus having an opposite effect on the RE_{aci}. The sensitivity 625 626 experiments again showed that the consideration of the chemistry of the mineral dust is important for the calculation of the RE_{aci} of the total NO_3^- aerosols. When the dust was assumed to be 627 chemically inert, the LW and SW estimates were up to 40% weaker, resulting in a warming of 628 +0.11 W/m². Assuming a homogeneous ion composition resulted in a smaller weakening of the 629 estimates (up to 18%) and a net warming of $+0.13 \text{ W/m}^2$. When dust emissions were halved, the 630 LW cooling was reduced slightly more than in the base case, resulting in a net warming of +0.15631 W/m². The 50% increase in dust emissions had the largest effect on LW behavior (10% increase), 632 but surprisingly the net estimate (+0.14 W/m²) was smaller than in the half-dust scenario. The 633 reason for this is that the SW estimate did not increase but decreased by 8% due to the fact that in 634 this scenario the increased nitrate burden causes increased competition for the available 635 supersaturation and the effect of dust-nitrate interactions on the smaller aerosol populations is not 636 as emphasized as in the base case. 637

The total NO_3^- aerosol RE_{aci} shows a positive sign, which is attributed to a reduced cloud albedo effect. More specifically, although the presence or absence of NO_3^- aerosol in the atmosphere did not significantly affect the total available maximum supersaturation, it did alter both the hygroscopicity and wet radii of the aerosols. In the presence of NO_3^- , the hygroscopicity of aerosols over deserts was increased by up to an order of magnitude, leading to an increase in their wet





643 radius of up to 10%, with an even larger increase of up to 40% for smaller particles over urban regions. Therefore, in the presence of NO_3^- aerosols, there is an increased depletion of fine particles 644 by coagulation with coarser particles (i.e. mineral dust) that have been further increased in size. 645 The reduction in the number of aerosols is as much as 10% in some regions, with maximum 646 reductions calculated over Southeast Asia. This reduction in the number of fine aerosols leads to 647 a reduction in the number of cloud droplets activated by fine aerosols (also up to 10%), which 648 649 would otherwise have absorbed more outgoing longwave radiation and, more importantly, 650 scattered more incoming shortwave radiation. Thus, the reduced cloud albedo effect leads to a 651 cooling in the longwave part of the spectrum, which is offset by a strong warming in the shortwave 652 part.





654 Code and Data Availability

The usage of MESSy (Modular Earth Submodel System) and access to the source code is licensed 655 to all affiliates of institutions which are members of the MESSy Consortium. Institutions can 656 become a member of the MESSy Consortium by signing the "MESSy Memorandum of 657 Understanding". More information can be found on the MESSy Consortium website: 658 http://www.messy-interface.org (last access: 22 May 2024). The code used in this study has been 659 based on MESSy version 2.55 and is archived with a restricted access DOI 660 661 (https://doi.org/10.5281/zenodo.8379120, The MESSy Consortium, 2023). The data produced in the study is available from the authors upon request. 662

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673

674 Competing Interests

At least one of the (co-)authors is a member of the editorial board of Atmospheric Chemistry andPhysics.

677 Author Contributions

AM and VAK wrote the paper with contributions from KK, APT, JFK, MK, and AN. VAK planned the research with contributions from APT, MK and AN. AM, KK and VAK designed the methodology for the radiative effect calculations. AM performed the simulations and analyzed the results, assisted by VAK and APT. All the authors discussed the results and contributed to the paper.

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