Global impact of mineral dust on cloud droplet number concentration 1 2 Vlassis A. Karydis¹, Alexandra P. Tsimpidi¹, Sara Bacer¹, Andrea Pozzer¹, Athanasios Nenes^{2,3,4} and Jos Lelieveld^{1,5} 3 4 5 ¹Max Planck Institute for Chemistry, Mainz, 55128, DE 6 ²Georgia Institute of Technology, Atlanta, GA, 30332, USA 7 ³National Observatory of Athens, Palea Penteli, 15236, GR 8 ⁴Foundation for Research and Technology Hellas, Patras, 26504, GR 9 ⁵The Cyprus Institute, Nicosia, 1645, CY 10 11 Abstract 12 13 14 The importance of wind-blown mineral dust for cloud droplet formation is studied by considering *i*) the adsorption of water on the surface of insoluble particles, *ii*) the 15 16 particle coating by soluble material (atmospheric aging) which augments cloud condensation nuclei (CCN) activity, and *iii*) the effect of dust on inorganic aerosol 17 concentrations through thermodynamic interactions with mineral cations. The 18 ECHAM5/MESSy Atmospheric Chemistry (EMAC) model is used to simulate the 19 20 composition of global atmospheric aerosol while the ISORROPIA-II thermodynamic equilibrium model treats the interactions of K⁺-Ca²⁺-Mg²⁺-NH₄⁺-Na⁺-SO₄²⁻-NO₃⁻-Cl⁻-21 H₂O aerosol with gas-phase inorganic constituents. Dust is considered a mixture of 22 inert material with reactive minerals and its emissions are calculated online by taking 23 into account the soil particle size distribution and chemical composition of different 24 deserts worldwide. The impact of dust on droplet formation is treated through the 25 "unified dust activation parameterization" that considers the inherent hydrophilicity 26 from adsorption and acquired hygroscopicity from soluble salts during aging. Our 27 simulations suggest that the presence of dust increases cloud droplet number 28 concentrations (CDNC) over major deserts (e.g., up to 20% over the Sahara and 29 Taklimakan Deserts) and decreases CDNC over polluted areas (e.g., up to 10% over 30 southern Europe and 20% over northeastern Asia). This leads to a global net decrease 31 of CDNC by 11%. The adsorption activation of insoluble aerosols and the mineral 32 dust chemistry are shown to be equally important for the cloud droplet formation over 33 the main deserts, e.g., these effects increase CDNC by 20% over the Sahara. Remote 34 35 from deserts the application of adsorption theory is critically important since the increased water uptake by the large aged dust particles (i.e., due to the added 36 hydrophilicity by the soluble coating) reduce the maximum supersaturation and thus 37 cloud droplet formation from the relatively smaller anthropogenic particles (e.g., 38

CDNC decreases by 10% over southern Europe and 20% over northeastern Asia by applying adsorption theory). The global average CDNC decreases by 10% by considering adsorption activation, while changes are negligible when accounting for the mineral dust chemistry. Sensitivity simulations indicate that CDNC is also sensitive to the mineral dust mass and inherent hydrophilicity, and not to the chemical composition of the emitted dust.

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46 **1. Introduction**

47 Atmospheric aerosols from anthropogenic and natural sources adversely affect human health and influence the Earth's climate, both directly and indirectly 48 (Haywood and Boucher, 2000; Lohmann and Feichter, 2005; Andreae and Rosenfeld, 49 2008; IPCC, 2013; Kushta et al., 2014; Lelieveld et al., 2015). The direct climate 50 effect refers to the influence of aerosols on the radiative budget of Earth's atmosphere 51 by scattering and absorbing solar radiation (Seinfeld and Pandis, 2006). The indirect 52 effects include the ability of aerosols to affect the cloud optical thickness and 53 scattering properties of clouds (Twomey, 1974) as well as the cloud lifetime and 54 precipitation (Albrecht, 1989). The scientific interest in aerosol-cloud-climate 55 56 interactions initially focused on anthropogenic pollutants (e.g., sulfate) and to a lesser extent on naturally emitted aerosols (e.g., sea salt). However, among atmospheric 57 58 aerosols, mineral dust is of particular importance since it is globally dominant in terms of mass concentration in the atmosphere (Grini et al., 2005; Zender and Kwon, 59 60 2005) and can influence cloud and precipitation formation (Levin et al., 2005; Yin and Chen, 2007; Karydis et al., 2011a; Rosenfeld et al., 2011; Kallos et al., 2014). 61 62 Additionally, dust alone is responsible for more than 400,000 deaths attributable to air pollution per year (Giannadaki et al., 2014). 63

Freshly emitted dust is considered insoluble. Reports of hygroscopic growth 64 measurements of dust particles indicate solubility to be very low, so that activation of 65 observed cloud condensation nuclei (CCN) has been attributed to soluble ions present 66 in the particles (Gustafsson et al., 2005; Herich et al., 2009; Koehler et al., 2009; 67 Garimella et al., 2014). Chemistry - climate models (CCMs) typically use Köhler 68 theory to describe droplet formation from dust, which assumes that the CCN activity 69 depends solely on their curvature effect and the fraction of soluble material on the 70 particle (Smoydzin et al., 2012). However, mineral dust can adsorb water which 71 72 results in a surface film of water with reduced activity (Sorjamaa and Laaksonen,

73 2007), and promote the formation of cloud droplets at cloud-relevant supersaturation, even of freshly emitted and chemically unprocessed dust particles (Sorjamaa and 74 Laaksonen, 2007; Kumar et al., 2009a). Kumar et al. (2009a) emphasized the 75 importance of including water adsorption effects in describing the hygroscopic growth 76 of mineral aerosols, which was then included in a droplet formation parameterization 77 78 (Kumar et al., 2009b) for use in models. Evidence on the importance of adsorption activation of dust particles is discussed in Kumar et al. (2011b; 2011a) for dry- and 79 wet-generated clays and mineral dusts representative of major regional dust sources 80 81 (North Africa, East Asia and North America). Adsorption activation was also found to be important for volcanic ashes (Lathem et al., 2011). The observed hygroscopicity 82 could not be attributed to the soluble ions present, but rather to the strong water vapor 83 adsorption on the particle surface. Furthermore, the surface fractal dimension derived 84 from dust and ash critical supersaturation data agrees well with previous methods 85 based on measurements of nitrogen adsorption, which contribute strong evidence for 86 adsorption effects on water activity and droplet activation (Laaksonen et al., 2016), 87 despite concerns raised by Garimella et al. (2014) on multiple charging effects on the 88 work of Kumar et al. (2011b). 89

Hatch et al. (2014) provided an alternative approach for parameterizing CCN activation of fresh atmospheric mineral aerosol. This approach was based on experimental water adsorption measurements on mineral clays compared to CCN measurements used by Kumar et al. (2011b), which require corrections for multiply charged particles and non-sphericity. Despite differences in the adsorption parameters reported from the above two studies, the adsorption derived CCN activities were quite similar and in excellent agreement.

97 Based on these findings, Karydis et al. (2011a) integrated the Kumar et al. (2009b) 98 parameterization into the Global Modeling Initiative (GMI) chemical transport model 99 (Considine et al., 2005) and found that insoluble mineral dust can contribute up to 100 24% of the cloud droplet number downwind of arid areas. Subsequently, the Kumar et 101 al. (2009b) parameterization has been integrated in a number of global and regional 102 models and applied to investigate the impact of mineral dust on warm cloud formation 103 (Bangert et al., 2012; Karydis et al., 2012; Gantt et al., 2014; Zhang et al., 2015).

Soluble inorganic ions like Ca^{2+} , Mg^{2+} , Na^+ , and K^+ that exist on the surface of mineral dust particles can participate in heterogeneous chemical reactions with acids such as HNO₃ and HCl. Furthermore, dust particles can provide reaction sites for the

107 SO_2 oxidation into H_2SO_4 . These processes result in the coating of dust particles by soluble material, which augments the hygroscopicity of dust and therefore its ability 108 to act as CCN (Kelly et al., 2007). On the other hand, highly oxidized, soluble organic 109 species, particularly including carboxylic acid groups (e.g., oxalic acid), can interact 110 with particles dominated by di-valent salts (e.g., CaCl₂) and strongly decrease their 111 hygroscopicity (Drozd et al., 2014). Due to their relatively large size, chemically aged 112 dust particles can act as giant CCN, enhancing precipitation as they efficiently collect 113 moisture and grow at the expense of smaller droplets (Feingold et al., 1999; Levin et 114 115 al., 2005). In addition, giant CCN compete with the submicron particles for water vapor, potentially reducing supersaturation and cloud droplet formation (Barahona et 116 al., 2010; Betancourt and Nenes, 2014b; Betancourt and Nenes, 2014a). 117

Soluble coatings on dust are mostly evident in the atmosphere after long-range 118 transport of dust plumes. Anthropogenic NO_3^- and SO_4^{2-} mainly contribute to the 119 chemical aging of dust over continents while sea salt derived Cl⁻ is more important 120 over oceans (Sullivan et al., 2007; Fountoukis et al., 2009; Dall'Osto et al., 2010; 121 Tobo et al., 2010; Karydis et al., 2011b; Bougiatioti et al., 2016b; Weber et al., 2016). 122 Apart from the gas phase composition, the chemical processing of dust also depends 123 124 on its chemical composition and thus on the source region (Sullivan et al., 2009; Karydis et al., 2016). Several studies have revealed that Saharan dust can be 125 126 efficiently transported over the Mediterranean basin where it can acquire significant soluble coatings (mostly sea salt and sulfate) resulting in the enhancement of its 127 hygroscopicity and CCN activity (Wurzler et al., 2000; Falkovich et al., 2001; 128 Smoydzin et al., 2012; Abdelkader et al., 2015). Twohy et al. (2009) have shown that 129 Saharan dust often acts as CCN over the eastern North Atlantic and significantly 130 contributes to cloud formation west of Africa. Begue et al. (2015) analyzed a case of 131 possible mixing of European pollution aerosols with Saharan dust transported over 132 northern Europe, and found that aged Saharan dust was sufficiently soluble to impact 133 the hygroscopic growth and cloud droplet activation over the Netherlands. Asian dust 134 has also been reported to have a considerable impact on cloud formation after being 135 transported over long distances and mixed with soluble materials (Perry et al., 2004; 136 Roberts et al., 2006; Sullivan et al., 2007; Ma et al., 2010; Stone et al., 2011; 137 Yamashita et al., 2011). 138

139 Despite the importance of mineral dust aerosol chemistry for accurately predicting 140 the aerosol hygroscopicity changes that accompany these reactions, most

141 thermodynamic models used in global studies lack a realistic treatment of crustal species, e.g., assuming that mineral dust is chemically inert (Liao et al., 2003; Martin 142 et al., 2003; Koch et al., 2011; Leibensperger et al., 2011). Few global studies have 143 accounted for the thermodynamic interactions of crustal elements with inorganic 144 aerosol components (Feng and Penner, 2007; Fairlie et al., 2010; Xu and Penner, 145 2012; Hauglustaine et al., 2014; Karydis et al., 2016). Most of these models either 146 neglect the impact of dust on cloud droplet formation or apply simplified assumptions 147 about the CCN activity of dust, e.g., they convert "hydrophobic" dust to "hydrophilic" 148 149 dust by applying a constant κ -hygroscopicity (e.g., 0.1) and use Köhler theory to describe cloud droplet activation. However, accounting for both the inherent 150 hydrophilicity of dust and the acquired hygroscopicity from soluble salts could 151 improve the predictive capability of CCMs. For this purpose, Kumar et al. (2011a) 152 presented a "unified dust activation framework" (UAF) to treat the activation of dust 153 with substantial amounts of soluble material by considering the effects of adsorption 154 (due to the hydrophilicity of the insoluble core) and absorption (due to the 155 hygroscopicity of the soluble coating) on CCN activity. Karydis et al. (2011a) 156 provided a first estimate of aged dust contribution to global CCN and cloud droplet 157 158 number concentration (CDNC) by using the UAF. They found that coating of dust by hygroscopic salts can cause a twofold enhancement of its contribution to CCN. On the 159 160 other hand, aged dust can substantially deplete in-cloud supersaturation and hence reduce the CDNC. Bangert et al (2012) investigated the impact of Saharan dust on 161 cloud droplet formation over western Europe and found only a slight increase in 162 calculated CDNC. However, these studies did not include thermodynamic interactions 163 of mineral dust with sea salt and anthropogenic pollutants. Instead, a prescribed 164 fraction of mineral dust that is coated with ammonium sulfate was used to represent 165 the aged dust. 166

The present work aims at advancing previous studies of dust influences on cloud 167 droplet formation by comprehensively considering *i*) the adsorption of water on the 168 surface of insoluble dust particles, *ii*) the coating of soluble material on the surface of 169 mineral particles which augments their CCN activity, and, *iii*) the effects of dust on 170 the inorganic soluble fraction of dust through thermodynamic interactions of semi-171 volatile inorganic species and sulfate with mineral cations. The ECHAM5/MESSy 172 Atmospheric Chemistry (EMAC) model (Jöckel et al., 2006) is used to simulate 173 aerosol processes, while the "unified dust activation framework" (Karydis et al., 174

175 2011a; Kumar et al., 2011a) is applied to calculate the CCN spectra and droplet number concentration, by explicitly accounting for the inherent hydrophilicity from 176 adsorption and acquired hygroscopicity from soluble salts by dust particles from 177 atmospheric aging. Mineral dust chemistry has been taken into account by using the 178 thermodynamic equilibrium model ISORROPIA II (Fountoukis and Nenes, 2007). 179 Dust emissions are calculated online by an advanced dust emission scheme that 180 accounts for the soil particle size distribution (Astitha et al., 2012) and chemical 181 composition (Karydis et al., 2016) of different deserts worldwide. The sensitivity of 182 183 the simulations to the emitted dust aerosol load, the mineral dust chemical composition and the inherent hydrophilicity of mineral dust is also considered. 184

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186 2. Model Description

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188 **2.1 EMAC Model**

We used the ECHAM5/MESSy Atmospheric Chemistry (EMAC) model (Jöckel et 189 190 al., 2006) which uses the Modular Earth Submodel System (MESSy2) (Jöckel et al., 2010) to connect submodels that describe the lower and middle atmosphere processes 191 with the 5th generation European Centre - Hamburg (ECHAM5) general circulation 192 model (GCM) as a dynamical core (Röckner et al., 2006). EMAC has been 193 194 extensively described and evaluated against in-situ observations and satellite retrievals (de Meij et al., 2012; Pozzer et al., 2012; Karydis et al., 2016; Tsimpidi et 195 al., 2017). The spectral resolution of the EMAC model used in this study is T63L31, 196 corresponding to a horizontal grid resolution of approximately $1.9^{\circ} \times 1.9^{\circ}$ and 31 197 vertical layers between the surface and 10 hPa (i.e. 25 km altitude). The model 198 dynamics has been weakly nudged (Jeuken et al., 1996) towards the analysis data of 199 the European Centre for Medium-Range Weather Forecasts (ECMWF) operational 200 model (up to 100 hPa) to represent the actual day-to-day meteorology in the 201 troposphere. EMAC is applied for 2 years covering the period 2004-2005 and the first 202 year is used as spin-up. 203

EMAC simulates the gas phase species through the MECCA submodel (Sander et al., 2011). Aerosol microphysics are calculated by the GMXe module (Pringle et al., 206 2010). The organic aerosol formation and chemical aging are calculated by the 207 ORACLE submodel (Tsimpidi et al., 2014). The CLOUD submodel (Röckner et al., 208 2006) calculates the cloud cover as well as cloud microphysics and precipitation of large scale clouds (i.e., excluding convective clouds). The cloud microphysical processes are computed by using the detailed two-moment liquid and ice-cloud microphysical scheme described in Lohmann and Ferrachat (2010), which enables a physically based treatment of aerosol–cloud interactions. The wet and dry deposition are calculated by the SCAV (Tost et al., 2006) and the DRYDEP (Kerkweg et al., 2006) sub-models.

The inorganic aerosol composition is computed with the ISORROPIA-II 215 (http://isorropia.eas.gatech.edu) thermodynamic equilibrium model (Fountoukis and 216 Nenes, 2007) with updates as discussed in Capps et al. (2012). ISORROPIA-II 217 calculates the gas-liquid-solid equilibrium partitioning of the K^+ - Ca^{2+} - Mg^{2+} - NH_4^+ -218 Na⁺-SO₄²⁻-NO₃⁻-Cl⁻-H₂O aerosol system. Potassium, calcium, magnesium, and 219 sodium are assumed to exist in the form of Ca(NO₃)₂, CaCl₂, CaSO₄, KHSO₄, K₂SO₄, 220 KNO₃, KCl, MgSO₄, Mg(NO₃)₂, MgCl₂, NaHSO₄, Na₂SO₄, NaNO₃, NaCl in the solid 221 phase and Ca^{2+} , K^+ , Mg^{2+} , Na^+ in the aqueous phase. More details about the EMAC 222 model set up used in this study can be found in Karydis et al. (2016). 223

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5 2.2 CCN Activity and Cloud Droplet Formation Parameterization

The cloud droplet formation parameterization is triggered only when warm clouds are present (i.e., cloud water is present and temperature exceeds 269 K). The equilibrium supersaturation, *s*, over the surface of a water droplet containing a solute particle (i.e., without any insoluble material present) is calculated using the hygroscopicity parameter, κ , based on κ -Köhler theory (Petters and Kreidenweis, 2007):

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$$s = \frac{4\sigma M_w}{RT\rho_w D_P} - \frac{D_{dry}^3 \kappa}{D_P^3} \quad (1)$$

where D_{dry} is the dry CCN diameter, D_P is the droplet diameter, σ is the CCN surface tension at the point of activation, ρ_w is the water density, M_w is the molar mass of water, *R* is the universal gas constant, and *T* is the average column temperature.

For insoluble particles (e.g., pristine mineral dust), the multilayer Frenkel-Halsey-Hill (FHH) adsorption isotherm model (Sorjamaa and Laaksonen, 2007) is used, which contains two adjustable parameters (A_{FHH} and B_{FHH}) that describe the contribution of water vapor adsorption on CCN activity. In this case, the equation describing the equilibrium supersaturation over the surface of a water droplet is given
by (Kumar et al., 2009b):

$$s = \frac{4\sigma M_w}{RT\rho_w D_P} - A_{FHH} \left(\frac{D_P - D_{dry}}{2D_w}\right)^{-B_{FHH}}$$
(2)

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where D_w is the diameter of a water molecule. The adsorption parameter A_{FHH} represents the interactions between the first water monolayer and the dust surface. B_{FHH} expresses the long range interactions of additional adsorbed water layers with the dust surface. Kumar et al. (2011b) tested a wide range of fresh unprocessed regional dust samples and minerals and found that one set of the FHH parameters ($A_{FHH} = 2.25 \pm 0.75$, $B_{FHH} = 1.20 \pm 0.10$) adequately reproduces the measured CCN activity for all dust types considered.

To account for the coating of soluble material on the surface of mineral dust, the "unified activation framework" (Karydis et al., 2011a; Kumar et al., 2011a) is used, which describes the water vapor supersaturation over an aerosol particle consisting of insoluble core with a soluble coating:

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$$s = \frac{4\sigma M_w}{RT\rho_w D_P} - \frac{\varepsilon_s D_{dry}^3 \kappa}{\left(D_P^3 - \varepsilon_i D_{dry}^3\right)} - A_{FHH} \left(\frac{D_P - \varepsilon_i^{1/3} D_{dry}}{2D_w}\right)^{-B_{FHH}}$$
(3)

where ε_i is the insoluble volume fraction and ε_s is the soluble volume fraction. Eq. 3 255 takes into account both the inherent hydrophilicity from adsorption expressed in the 256 third term of the equation and the acquired hygroscopicity from soluble salts by dust 257 particles expressed in the second term of the equation. The first term accounts for the 258 Kelvin effect. Noting that for a complete insoluble dust particle, i.e., as $\varepsilon_s \rightarrow 0$ and 259 $\varepsilon_i \rightarrow 1$, the UAF approaches FHH theory (Eq. 2). Black carbon (BC) is not expressed 260 by the FHH terms in eq. 3. Instead, BC is assumed to have zero hygrscopicity and 261 affects κ in eq. 3 according to the simple mixing rule. 262

Calculation of CDNC is carried out in two conceptual steps, one involving the determination of the "CCN spectrum" (i.e., the number of CCN that can activate to form droplets at a certain level of supersaturation), and another one determining the maximum supersaturation, s_{max} , that develops in the ascending cloudy air parcels used to represent droplet formation in EMAC. The CDNC is then the value of the CCN spectrum at s_{max} . The "CCN spectrum", $F^{s}(s)$, is computed following Kumar et al. (2009b) and assumes that particles can be described either by KT or FHH theory. $F^{s}(s)$ for an external mixture of lognormal particle size distributions is given by:

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$$F^{s}(s) = \int_{0}^{s} n^{s}(s) ds = \sum_{i=1}^{n_{m}} \frac{N_{i}}{2} \operatorname{erfc} \left[-\frac{\ln\left(\frac{s_{g,i}}{s}\right)}{x\sqrt{2}\ln(\sigma_{i})} \right] \quad (4)$$

where s is the level of water vapor supersaturation, $n^{s}(s)$ is the critical 273 supersaturation distribution, $s_{g,i}$ is the critical supersaturation of the particle with a 274 diameter equal to the geometric mean diameter of the mode i, σ_i is the geometric 275 standard deviation for the mode i, and x is an exponent that depends on the 276 activation theory used. For modes following Köhler theory, $x = -\frac{3}{2}$ (Fountoukis and 277 Nenes, 2005), while for insoluble particles following FHH theory, x depends on 278 A_{FHH} and B_{FHH} (Kumar et al., 2009b) (i.e., x=0.86 for $A_{FHH}=2.25$ and B_{FHH} 279 =1.20 used here). In the case of UAF, x lies between the KT and FHH-AT limits, and 280 is determined from Eq. (3) by performing a power law fit between s_g and D_{dry} as 281 described in Kumar et al. (2011a). The calculation of s_g involves determining the 282 maximum of the relevant equilibrium curve in equilibrium with the surrounding water 283

284 vapor
$$\left(\frac{ds}{dD_p}\right|_{D_p=D_g} = 0$$
 in Eqs. 1-3). Once D_g is determined, it can be substituted in Eqs.

1-3 to obtain s_g .

The maximum supersaturation, s_{max} , in the ascending parcel is calculated from an equation that expresses the supersaturation tendency in cloudy air parcels, which at the point of maximum supersaturation becomes (Nenes and Seinfeld, 2003; Barahona and Nenes, 2007)

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$$\frac{2aV}{\pi\gamma\rho_w} - Gs_{\max}I(0, s_{\max}) = 0 \quad (5)$$

where V is the updraft velocity (i.e., not including convection) calculated online by assuming that the sub-grid vertical velocity variability is dominated by the turbulent transports and by choosing the root-mean-square value of the GCM model-generated

turbulent kinetic energy (TKE) as a measure. Based on this assumption, the in-cloud 294 updraft velocity can be expressed as $V = \overline{V} + 0.7\sqrt{TKE}$, where \overline{V} is the GCM-295 resolved large scale updraft velocity (Lohmann et al., 1999a; Lohmann et al., 1999b). 296 Following Morales and Nenes (2010), V can be considered as a "characteristic updraft 297 velocity" which yields CDNC value representative of integration over a probability 298 density function (PDF) of updraft velocity. Morales and Nenes (2010) have shown 299 that this assumption applies well to large scale clouds (i.e., stratocumulus), which are 300 the type of clouds described by the CLOUD sub-model in EMAC. a, γ, G in Eq. (5) 301 are parameters defined in Nenes and Seinfeld (2003). $I(0, s_{max})$ is the "condensation 302 integral" which expresses the condensational depletion of supersaturation upon the 303 growing droplets at the point of s_{max} in the cloud updraft. It is expressed as the sum of 304 305 two terms:

$$I(0, s_{\max}) = I_K(0, s_{\max}) + I_{FHH}(0, s_{\max})$$
(6)

The first term on the right hand side of Eq. (6), $I_K(0, s_{\text{max}})$, describes the contribution 307 from particles that follow the Köhler theory and is calculated using the revisited 308 population splitting approach of Betancourt and Nenes (2014a). The second term, 309 $I_{FHH}(0, s_{max})$, represents the contribution of freshly emitted or aged dust particles to 310 the condensation integral and is represented in Kumar et al. (2009b) and Karydis et al. 311 (2011a). Once s_{max} is determined by numerically solving Eq. (5), the number of 312 cloud droplets that form in the parcel, N_d , is obtained from the "CCN spectrum" (Eq. 313 (4)) computed for s_{max} , i.e., $N_d = F(s_{\text{max}})$. The cloud droplet formation 314 parameterization presented here has been extensively evaluated by comparing 315 computations of N_d and s_{max} and their sensitivity to aerosol properties against 316 detailed numerical simulations of the activation process by a parcel-model 317 (Betancourt and Nenes, 2014a). 318

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320 **2.3 Aerosol Precursor Emissions**

Dust emission fluxes are calculated online by an advanced dust flux scheme developed by Astitha et al. (2012). This scheme uses an explicit geographical representation of the airborne soil particle size distribution based on soil characteristics in every grid cell. Emissions of crustal species (Ca^{2+} , Mg^{2+} , K^+ , Na^+)

are estimated as a fraction of mineral dust emissions based on the chemical 325 composition of the emitted soil particles in every grid cell (Karydis et al., 2016). 326 Emissions of sea spray aerosols are based on the offline monthly emission data set of 327 AEROCOM (Dentener et al., 2006) assuming a composition of 55% Cl⁻, 30.6% Na⁺, 328 7.7% SO_4^{2-} , 3.7% Mg^{2+} , 1.2% Ca^{2+} , 1.1% K^+ (Seinfeld and Pandis, 2006). The 329 CMIP5 RCP4.5 emission inventory (Clarke et al., 2007) is used for the anthropogenic 330 primary organic aerosol emissions from fossil fuel and biofuel combustion sources. 331 The open biomass burning emissions from savanna and forest fires are based on the 332 333 GFED v3.1 database (van der Werf et al., 2010). More details about the aerosol phase emissions used by EMAC can be found in Karydis et al. (2016) and Tsimpidi et al. 334 (2016). 335

Related anthropogenic emissions of NO_x, NH₃, and SO₂, which represent the 336 gaseous precursors of the major inorganic components, are based on the monthly 337 emission inventory of EDGAR-CIRCE (Doering, 2009) distributed vertically as 338 presented in Pozzer et al. (2009). The natural emissions of NH₃ are based on the 339 GEIA database (Bouwman et al., 1997). NO_x produced by lightning is calculated 340 online and distributed vertically based on the parameterization of Grewe et al. (2001). 341 342 The emissions of NO from soils are calculated online based on the algorithm of Yienger and Levy (1995) as described in Ganzeveld et al. (2002). Eruptive and non-343 344 eruptive volcanic degassing emissions of SO₂ are based on the AEROCOM data set (Dentener et al., 2006). The oceanic DMS emissions are calculated online by the 345 AIRSEA submodel (Pozzer et al., 2006). More details about the gas phase emissions 346 used by EMAC can be found in Pozzer et al. (2012) and Karydis et al. (2016). 347

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349 **3. Model Results and Evaluation**

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351 **3.1 Model Predictions**

The annual and seasonal (during DJF and JJA) mean CDNC, calculated by EMAC with UAF implementation for the lowest model level at which clouds are formed (centered at 940 hPa), are shown in Figure 1. In this study, CDNC is referred to the number concentration of droplets nucleated in-cloud and represents an upper limit since droplet depletion by collision, coalescence and collection are not taken into account. Therefore, the calculated CDNC is mostly sensitive to the cloud updraft velocity and the total aerosol number concentration (Karydis et al., 2012), which are

the main drivers of the $s_{\rm max}$ calculations. The annual mean aerosol number 359 concentration, updraft velocity, and $s_{\rm max}$ at 940 hPa, as well as the low-level 360 cloudiness calculated by EMAC are shown in Figure 2. The calculated CDNC is also 361 sensitive to the fraction of mineral dust present in the aerosol since it can affect the 362 aerosol-water vapor interactions by changing the exponent x in Eq. 4. The annual 363 mean insoluble fraction of the particles in the accumulation and coarse mode (where 364 mineral dust exists) are shown in figure 3. The calculated global annual mean CDNC 365 at 940 hPa is 231 cm⁻³. 366

Over the continents, the predicted annual mean CDNC is 546 cm⁻³ and exceeds 367 1000 cm⁻³ over the industrialized areas of Europe, central and eastern Asia, and North 368 America. In these areas, the aerosol number concentration is high (exceeding 10,000 369 cm^{-3} ; Figure 2a), while the calculated updraft velocities (0.5-1 m s⁻¹; Figure 2b) allow 370 the development of sufficiently high s_{max} (0.1-0.3%; Figure 2c) for the activation of 371 5% (over eastern China) to 15% (over central Europe) of the pollution aerosols into 372 cloud droplets. The simulated s_{max} is close to the estimated s_{max} (0.2%-0.5%) for 373 stratocumulus clouds based on data from continental air masses (Twomey and 374 Wojciechowski, 1968; Martin et al., 1993) indicating that the combination of aerosol 375 number concentration and updraft velocity in the model is realistic. 376

377 While the aerosol number concentration over the industrialized areas remains fairly constant throughout the year, the updraft velocity is higher during the boreal winter 378 (i.e., DJF) resulting in a seasonal peak of CDNC during DJF (exceeding 2,000 cm⁻³) 379 over North America, Europe and eastern Asia (Figure 1b). The highest annual mean 380 CDNC is calculated over northern India ($\sim 2,000 \text{ cm}^{-3}$) where the model simulates 381 highest aerosol concentrations (~30,000 cm⁻³). Over Southeast Asia and India, CDNC 382 peaks during JJA (exceeding 2,000 cm⁻³; Figure 1c), affected by the East Asian 383 Monsoon and the high updraft velocities developed during the wet season. Relatively 384 high CDNC (annual mean of 300-700 cm⁻³) are also calculated over the tropical 385 regions of the Southern Hemisphere which are influenced by biomass burning. 386 Relatively low values are calculated over the Congo Basin where the mean updraft 387 velocity is typically low (below 0.2 m s⁻¹) leading to low s_{max} (below 0.05%) and 388 cloud droplet activation ($\sim 300 \text{ cm}^{-3}$). 389

Downwind of deserts, the calculated CDNC varies between 100 cm⁻³ (e.g., Patagonia, and Australian deserts) to 1,000 cm⁻³ (e.g., Sahara, Arabian, Taklimakan, 392 Gobi and Atacama). In the vicinity of the Sahara and Arabian deserts, the mean updraft velocity is ~0.5 m s⁻¹. However, downwind of the western part of the Sahara 393 the aerosol number concentration is relatively low (~ 1.000 cm⁻³) leading to higher 394 s_{max} (~0.2%) but low CDNC (~200 cm⁻³). On the other hand, downwind of the 395 eastern Sahara and Arabian deserts the aerosol concentration is higher (2,000-3,000 396 cm⁻³). Over these areas the presence of a high number of coarse dust particles 397 significantly reduces s_{max} (~0.05%), but at the same time they efficiently activate into 398 cloud droplets (CDNC varies from 500 to 1,000 cm⁻³). Close to Patagonia and 399 Australia, despite the high updraft velocities ($\sim 1 \text{ m s}^{-1}$), the aerosol concentration is 400 low (below 500 cm⁻³) and also CDNC is relatively low (~100 cm⁻³). The highest 401 updraft velocities are calculated around the Atacama and Gobi deserts (over 1 m s⁻¹) 402 leading to both high s_{max} (over 0.3%) and CDNC (~1,000 cm⁻³). However, the central 403 Asian deserts (e.g., Gobi) are under the influence of the Siberian anticyclone during 404 winter (i.e., DJF) which causes katabatic winds (that inhibit the formation of positive 405 updraft velocities) and very low temperatures that prevent the formation of liquid 406 water clouds. 407

Over the oceans, the predicted annual mean CDNC is 113 cm⁻³ and exceeds 500 408 cm⁻³ along the coasts of Mediterranean countries, China, India, SE Asia, California, 409 the northeastern USA and western Africa (Fig. 1). Over many coastal regions aerosol 410 concentrations are relatively high (5,000-10,000 cm⁻³), however, the low updraft 411 velocities ($\sim 0.2 \text{ m s}^{-1}$) result in lower CDNCs than over land (Figure 1). The 412 Mediterranean and Yellow Seas are somewhat exceptional since the annual mean 413 updraft velocities are higher in these regions (~0.3 m s⁻¹), resulting in higher s_{max} 414 (~0.1% and ~0.3%, respectively) and therefore high CDNC (~800 cm⁻³ and ~1,200 415 cm⁻³, respectively). The simulated s_{max} is in close agreement with estimates (~0.1%) 416 based on observational data over the eastern Mediterranean (Bougiatioti et al., 2016a; 417 Kalkavouras et al., 2016). CDNC over these seas is subject to high seasonal variation 418 ranging from ~400 cm⁻³ (~800 cm⁻³) over the Mediterranean (Yellow) Sea during JJA, 419 to over $1,000 \text{ cm}^{-3}$ (2,000 cm⁻³) during DJF due to the higher updraft velocities during 420 boreal winter (exceeding 1 m s⁻¹) compared to summer (below 0.2 m s⁻¹). Over the 421 northern coasts, the annual mean CDNC is significantly enhanced compared to the 422 oceans of the Southern Hemisphere due to the transport of pollutants from 423 industrialized areas in the Northern Hemisphere. Despite the high updraft velocities 424

calculated over the southern oceans throughout the year (up to 1 m s⁻¹), the lack of aerosol (typically below 100 cm⁻³) results in CDNC below 50 cm⁻³. Finally, the calculated CDNC decreases with altitude due to the decrease in aerosol concentration by dilution and atmospheric removal (Figure 4). The global mean CDNC is predicted to be 231 cm⁻³, 171 cm⁻³, 120 cm⁻³, 87 cm⁻³, and 60 cm⁻³ at 940 hPa, 900 hPa, 860 hPa, 820 hPa, and 770 hPa, respectively.

431

432 **3.2 Model Evaluation**

433 The predicted in-cloud CDNC are compared to observational data from continental, polluted marine and clean marine regions around the world (Karydis et 434 al., 2011a). The locations of observations (i.e., longitude, latitude, and altitude) and 435 time of year have been taken into account in sampling the model results. Given that 436 the observations span a decade, in contrast to the simulation which represents one 437 year, the month of each campaign has been used to account for the seasonal 438 variability of the CDNC. Thus, the implicit assumption is that inter-annual variability 439 can be neglected. It should also be mentioned that the observations typically do not 440 represent monthly means over 1.9° grid squares, as sampled from the model results, 441 442 so that the comparison is more qualitative than quantitative. Furthermore, the cloudaveraged CDNC for stratocumulus clouds, which are described by EMAC, is typically 443 444 well captured by the cloud droplet formation parameterization used in this study (Meskhidze et al., 2005; Fountoukis et al., 2007; Morales et al., 2011), while the 445 446 droplet collision and coalescence processes, which are neglected here, are becoming important only in the presence of clouds with substantial amount of drizzle. A 447 summary of the comparison results is presented in Table 1 and Figure 5. The mean 448 bias (MB), mean absolute gross error (MAGE), normalized mean bias (NMB), 449 450 normalized mean error (NME), and the root mean square error (RMSE) are used to assess the model performance (Table 2). 451

The model captures the low values (below 100 cm⁻³) observed over the remote Pacific, Atlantic and Indian Oceans and at the same time is capable of simulating the higher concentrations (>100 cm⁻³) observed over the eastern Pacific Ocean (Table 1). On the other hand, it falls short in reproducing the relatively high CDNC (>100 cm⁻³) observed during summer over the western Arctic Ocean and over the remote area west of Australia. Overall, the model tends to underestimate the CDNC over remote oceans with a MB = -33 cm⁻³ and NMB = -39% (Table 2). 459 Both the observed and simulated CDNC show significant increases over polluted marine regions close to the coasts (Table 1; Figure 5a). Compared to satellite 460 retrievals (Bennartz, 2007; Rausch et al., 2010), the model reproduces the CDNC over 461 the American and African coasts well, but it significantly overestimates CDNC along 462 the Asian coasts (Table 1). Compared to in situ observations, the model reproduces 463 the high CDNC along coastal areas in the Northern Hemisphere (e.g., the Yellow Sea, 464 Oregon, Florida, Canary Islands), but systematically overestimates CDNC over the 465 British coasts. Further, the model does not reproduce some of the high CDNC 466 observations over more remote areas (i.e., over the Azores and eastern Atlantic 467 Ocean). Overall, the model tends to overestimate the CDNC over polluted marine 468 areas with a MB = 127 cm^{-3} and NMB = 75% (Table 2). 469

The observed CDNC over continental regions is subject to high spatial variability, 470 with reported values ranging from $<100 \text{ cm}^{-3}$ over Alaska (Dong and Mace, 2003) to 471 >1,000 cm⁻³ over China (Zhao et al., 2006), England (Bower et al., 1999), and the 472 continental USA (Fountoukis et al., 2007). The model captures the observed 473 variability with low values over remote areas (e.g., over Alaska) and high values over 474 the industrialized parts of the Northern Hemisphere (i.e., East Asia, Europe, and 475 476 China). Overall, the model overestimates CDNC over continental regions (MB= 269 cm⁻³ and NMB=58%; Table 2). Over China, the simulated CDNC is within the 477 observed range with the exception of Hebei Province where it significantly 478 overestimates measured CDNC (Table 1). In Europe, the model reproduces the high 479 480 CDNC observed over Central Europe and England but it clearly overestimates the low CDNC values observed over Finland. Over North America, the model captures the 481 482 variability of the observed CDNC, predicting lower values over remote areas (e.g., Alaska) and higher values over the industrialized areas of USA (e.g., Ohio and 483 484 Michigan). It tends to overestimate the CDNC over the continental USA and underestimate the observed values over Alaska. 485

Globally, the calculated NMB is 56% and the NME is 82%, indicating that some of the discrepancy between the modelled and the observed CDNC is explained by uncertainties in the observations and the numerical simulations. Around 60% of the simulated CDNC are within a factor of 2 compared to the measurements (Figure 5a) and 40% of the simulated CDNC differ less than 30% from the measurements. Based on the typical properties of marine stratus clouds, a uniform increase in global CDNC by 30% (or 50%) leads to an increase in cloud albedo of 2.25% (or 3.75%) and a 493 perturbation of -1.1 W m^{-2} (or -1.7 W m^{-2}) in the global mean cloud radiative forcing 494 (Schwartz, 1996). However, the simulated CDNC presented here refers to the number 495 concentration of droplets nucleated in clouds and represents an upper limit with 496 respect to the comparison with observations, since collision and coalescence 497 processes, which are not taken into account here, can reduce the CDNC.

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499 **4. Mineral Dust Effect on CDNC**

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501 4.1 Total Impact of Mineral Dust on CDNC

To estimate the overall effect of mineral dust on CDNC a sensitivity run was 502 conducted switching off the mineral dust emissions. Figure 6 depicts the difference in 503 CDNC between the base case simulation and the sensitivity test. A positive change 504 corresponds to an increase of the CDNC due to the presence of dust. The predicted 505 CDNC is typically increased by the presence of dust aerosols over the main deserts 506 (Figure 6). Over the Sahara, CDNC increases less than 50 cm^{-3} (up to 20%). The 507 largest change is calculated downwind of the Patagonian ($\sim 150 \text{ cm}^{-3}$ or 70%) and 508 Atacama (~350 cm⁻³ or 40%) deserts. Over these deserts dust emissions increase the 509 aerosol concentration by more than $5,000 \text{ cm}^{-3}$ (Figure 6c). The effect of mineral dust 510 on CDNC close to Sahara varies significantly throughout the year due to the 511 seasonality of the mineral dust emissions. Over the sub-Sahelian region, CDNC 512 increases by up to 150 cm⁻³ during DJF, owing to the northeasterly trade winds (i.e., 513 Harmattan winds) which blow from the Sahara Desert over West Africa during 514 winter. Over the eastern Sahara and the Arabian deserts CDNC increases up to 150 515 cm⁻³ during spring (i.e., MAM) and autumn (i.e., SON) when the Sirocco winds are 516 most common. 517

In contrast to regions close to deserts, CDNC decreases over the polluted regions 518 of the Northern Hemisphere and especially over southern Europe (~100 or less than 519 10%) and northeastern Asia (up to 400 cm⁻³ or 20%). In these areas, dust particles 520 transported from the Sahara over Europe and from the Gobi and Taklimakan deserts 521 over Asia, are mixed with anthropogenic particles decreasing the total aerosol number 522 concentration (Figure 6c), due to coagulation, and affecting the aerosol-water vapor 523 interactions. As the insoluble fraction of aerosols increases due to the addition of 524 mineral dust (Figures 3b and 3c), the exponent x in Eq. 4 changes, resulting in a 525 decrease of the number of activated droplets. Furthermore, the relatively large, aged 526

dust particles over these areas activate early on in the cloud formation process, taking up much water per particle and thus reducing s_{max} (~15%), and consequently cloud droplet formation on the smaller anthropogenic particles (e.g., the activated fraction of the particles in the accumulation mode reduces by 20%). Beside microphysical effects, the presence of mineral dust can also affect cloud formation by altering the energy balance of the atmosphere, and thus turbulent motions and the updraft velocity.

Nevertheless, the calculated updraft velocity does not change significantly between 534 the two simulations (less than 5%), in part because the meteorology is dynamically 535 nudged to analysis data (Jeuken et al., 1996). CDNC also decreases over the oceans 536 downwind of deserts in the Northern Hemisphere, and even over the rainforests in the 537 538 Southern Hemisphere (~150 or 30%). Overall, the impact of mineral dust on CDNC is positive only in areas with low cloud cover (i.e., over the main deserts where cloud 539 540 cover is typically lower than 5%; Figure 2d). On the other hand, mineral dust negatively affects cloud droplet formation over areas with high cloud cover (e.g., over 541 542 Europe and Eastern Asia). Despite that CDNC increases over the deserts due to the presence of dust particles, the decrease of CDNC over the industrialized and forested 543 544 continental areas dominates the calculated global average change, i.e., the calculated global average CDNC decreases by 11% (or 26 cm⁻³). 545

546

547 **4.2 Impact of Mineral Dust Chemistry on CDNC**

To estimate the effects of thermodynamic mineral dust interactions with inorganic 548 anions on the predicted CDNC, a sensitivity run was conducted by switching off the 549 dust-aerosol chemistry. Karydis et al. (2016) have shown that dust can significantly 550 affect the partitioning of inorganic aerosol components and especially nitrate. 551 Analogous to Karydis et al. (2016), accounting for thermodynamic interactions of 552 mineral dust in our simulations results in an increase of the tropospheric burden of 553 nitrate, chloride, and sulfate aerosols by 44%, 9%, and 7%, respectively. On the other 554 hand, ammonium decreases by 41%. The dust presence itself also decreases by 14% 555 since it becomes significantly more soluble, mostly due to the condensation of nitric 556 557 acid on its surface, and is removed more efficiently through wet and dry deposition, the latter partially due to the increased sedimentation by dust particles that have a 558 larger water content. Therefore, the calculated change of CDNC (Figures 7a and 7b) 559

560 is the net result of counterbalancing effects. Due to the increase of the soluble fraction by considering mineral dust chemistry, the CDNC activated from dust particles 561 increases (Figure 7c), while the total number of dust particles and the CDNC from 562 insoluble particles decreases (Figure 7d). Taking as an example a grid cell over the 563 Sahara Desert, the model simulations indicate that by accounting for the mineral dust 564 chemistry, the soluble fraction of the dust containing particles increases by 0.07, 565 resulting in an increase of CDNC activated from soluble aerosol modes by 150 cm⁻³ 566 (Figure 7c). On the other hand, the aerosol number concentration decreases by 90 cm⁻ 567 ³ due to the more efficient atmospheric removal of the aged dust particles, resulting in 568 a decrease of the CDNC activated from the insoluble modes by 50 cm^{-3} (Figure 7d). 569 The net effect is that the total CDNC increases by 100 cm^{-3} (Figure 7a). 570

Overall, the presence of reactive dust components results in an increase of CDNC 571 over the deserts that are close to anthropogenic sources, e.g., up to 100 cm^{-3} (or 20%) 572 over the Sahara and up to 200 cm⁻³ (or 30%) over the Arabian Peninsula. In these 573 areas, the CCN activity of mineral dust (initially hydrophilic) is enhanced by the 574 acquired hygroscopicity from the anthropogenic (including biomass burning) aerosol 575 compounds (mainly nitrate). Even though the chemically aged dust particles activate 576 577 into droplets more efficiently than insoluble ones, their reduced number concentration dominates the calculated effect on CDNC over the relatively pristine remote desert 578 regions, e.g., CDNC decreases up to 200 cm⁻³ (or 20%) downwind of the Taklimakan, 579 250 cm⁻³ (or 30%) around the Atacama, and up to 100 cm⁻³ (or 40%) over the 580 Patagonian deserts. Even over the rainforests, HNO₃ from NO_X, emitted by biomass 581 burning, thermodynamically interacts with the coarse soil particles from the upwind 582 deserts, resulting in an increase of CDNC by around 50 cm⁻³. CDNC is also slightly 583 increased over Europe and eastern Asia (up to 150 cm⁻³ or about 10%) where HNO₃ 584 from anthropogenic NO_x sources interacts with mineral dust from the surrounding 585 deserts. While the global average CDNC does not change much by taking into 586 account thermodynamic and chemical interactions of mineral dust with inorganic air 587 pollutants, CDNC spatial distributions change substantially. 588

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4.3 Impact of Water Adsorption by Mineral Dust on CDNC

To estimate the effects of water adsorption onto the surface of insoluble dust 591 particles on CDNC, a sensitivity run was conducted by switching off the FHH 592 adsorption calculations. In this sensitivity simulation, the soluble modes follow the k-593

Köhler theory while insoluble modes do not participate in cloud droplet formation 594 calculations. Figure 8 depicts the difference in CDNC between the base case 595 simulation and this sensitivity test. A positive change corresponds to an increase of 596 the CDNC from water adsorption on mineral dust. The calculations show that CDNC 597 is increased by applying FHH theory over several arid areas where the insoluble dust 598 concentration is high (Figure 8), since κ -Köhler theory does not take into account the 599 contribution of insoluble particles to cloud droplet formation. CDNC is increased in 600 the vicinity of the Saharan, Arabian and Thar deserts ($\sim 100 \text{ cm}^{-3}$ or about +20%) 601 where the insoluble fraction of mineral dust is larger due to the small anthropogenic 602 emission influence that makes the particles hygroscopic. On the other hand, CDNC 603 decreases over the polluted regions of the Northern Hemisphere and especially over 604 Europe (~100 cm⁻³ or about -10%) and Asia (up to ~400 cm⁻³ or -20%). Over these 605 areas, the added hydrophilicity by the soluble coatings on the surface of the aged dust 606 particles increases their water uptake during activation. Therefore, the aged dust 607 particles relatively strongly compete for water vapor, reducing the $s_{\rm max}$ (~15%) and 608 thus cloud droplet formation from the smaller anthropogenic particles. Over the 609 tropical rainforests CDNC decreases by ~150 cm⁻³ (about -30%). Overall, the use of 610 611 the UAF results in a decrease of the global average CDNC by $\sim 10\%$ (or about -23 cm^{-3}). 612

613

614 **5 Additional Sensitivity Tests**

Three additional sensitivity simulations were conducted to investigate the CDNC dependency on i) the chemical composition of the emitted dust aerosols, ii) the hydrophilicity of mineral dust, and iii) the strength of the dust aerosol emissions. Figure 9 depicts the absolute annual mean changes in CDNC compared to the reference simulation for each of the sensitivity tests. A positive change corresponds to an increase of the CDNC relative to the reference.

621

622 5.1 Sensitivity to the emitted dust aerosol composition

The first sensitivity test assumes a globally uniform chemical composition of mineral dust (Sposito, 1989), in contrast to the reference simulation where the mineral dust composition depends on the soil characteristics of each desert (Karydis et al., 2016). While the emitted mineral dust load remains the same in the sensitivity

simulation, the different mineral dust composition results in significant changes in the 627 calculated tropospheric burdens of dust components (Karydis et al., 2016). In 628 particular, the fraction of the mineral components relative to the total dust in the 629 sensitivity simulation is lower over most of the deserts compared to the reference. 630 This reduction of the chemically reactive mineral components in the sensitivity 631 simulation results in a slowdown of the mineral dust aging and hence in an increase of 632 its concentration due to the reduced atmospheric removal. Conversely, the CCN 633 activity of dust particles is higher in the reference simulation since the chemical aging 634 635 is more efficient compared to the sensitivity simulation. These counterbalancing effects result in negligible changes of CDNC worldwide (less than 10%). 636

637

638 **5.2 Sensitivity to the hydrophilicity of dust**

The second sensitivity test assumes increased hydrophilicity of mineral dust 639 aerosols by using a 10% lower B_{FHH} parameter (B_{FHH} =1.1). The B_{FHH} parameter 640 directly affects the CCN activity of dust particles by changing the equilibrium 641 supersaturation (Eq. 3) and the "CCN spectrum" (Eq. 4) through the exponent x. 642 Kumar et al. (2011b) tested the CCN activity of aerosols dry generated from clays, 643 644 calcite, quartz, and desert soil samples from Northern Africa, East Asia/China, and Northern America. They found that B_{FHH} , which strongly affects the equilibrium 645 646 curve, varied from 1.12 to 1.30 (i.e., ±10% from 1.2 which is the value used in our base case simulation). Therefore, the sensitivity test presented here can represent the 647 potential impacts on the results due the simplification of using a globally uniform set 648 of FHH parameters to describe the hydrophilicity of mineral dust independently of its 649 source and composition. The higher hydrophilicity of mineral dust in the sensitivity 650 simulation results in increased CDNC over areas close to deserts by up to 30% (e.g., 651 100 cm⁻³ over Sahara and 200 cm⁻³ over Gobi and Taklimakan). A notable increase is 652 also calculated over eastern China and northern India (up to 150 cm⁻³ or 10%) where 653 mineral dust is mixed with anthropogenic compounds. These results indicate that 654 changes in the hydrophilicity of the freshly emitted dust, due to the variability of its 655 composition with source region, can have an important impact on the calculated 656 CDNC. Remote from the main deserts (e.g., over central Europe), the change in 657 CDNC is negligible since the contribution of mineral dust particles on cloud droplet 658 formation is low. Overall, the calculated global average CDNC increases in the 659 sensitivity simulation by about 5% (or $\sim 12 \text{ cm}^{-3}$). 660

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662 **5.3 Sensitivity to the emitted dust aerosol load**

The final sensitivity test assumes 50% lower emissions of mineral dust compared 663 to the reference simulation. The lower tropospheric dust load in the sensitivity 664 simulation (49%) results in a 10-30% (up to 150 cm⁻³) decrease of CDNC over the 665 main deserts. On the other hand, CDNC increases over the anthropogenic (e.g., East 666 Asia) and biomass burning (e.g., central Africa) regions by 5-10% (up to 150 cm⁻³). 667 The opposing responses of CDNC to mineral dust emissions result from the fact that 668 669 the tropospheric load of the other aerosol species does not change significantly between the two simulations since the chemical and thermodynamic interactions of 670 mineral cations with air pollution remain predominant, even after the 50% dust 671 emission reduction, i.e., the nitrate abundance rather than that of dust is rate limiting. 672 Therefore, the presence of inorganic anions (e.g., NO₃) in the aerosol phase remains 673 almost unchanged between the two simulations, which results in a decrease of the 674 insoluble fraction of the aerosol, given that mineral dust concentrations are 675 significantly lower in the reference simulation, leading to higher CCN activity. Over 676 the Taklimakan desert the insoluble fraction of the aerosol changes by less than 10%, 677 678 and therefore, the change in aerosol number concentration (~40%) due to the mineral dust emission change dominates the effect on CDNC, which is calculated to be about 679 100 cm⁻³ (or ~20%) lower in the sensitivity simulation. On the other hand, over 680 Southeast Asia, the aerosol number concentration changes less than 10% as the 681 insoluble fraction of the aerosols decreases by 40%. The significant decrease of ε_i in 682 Eq. (3) affects the calculated critical supersaturation of the particle as well as the 683 exponent x in Eq. (4) resulting in an increase of CDNC by about 150 cm⁻³ (or ~10%). 684 Overall, the impact of halving mineral dust emissions on the calculated global average 685 CDNC is remarkably small ($\sim 3\%$ or 6 cm⁻³). 686

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6 Summary and Conclusions

This study assesses the impact of mineral dust on global cloud droplet number concentrations by using an interactive aerosol-chemistry-cloud-climate model (EMAC). The "unified dust activation framework" (UAF) has been implemented into the EMAC model to account for the effects of dust particles through both the hydrophilicity from adsorption and the acquired hygroscopicity from pollution solutes (chemical aging) on CCN activity calculations. The calculation of cloud droplet

formation from soluble particles is carried out by using the κ-Köhler theory, while 695 that of insoluble particles is based on the FHH multilayer adsorption isotherm 696 approach. For atmospheric particles that contain a substantial fraction of both soluble 697 (e.g., nitrate) and insoluble material (e.g., mineral dust), cloud formation is calculated 698 using the UAF, which determines the maximum equilibrium water vapor 699 supersaturation over an aerosol consisting of an insoluble core with a soluble coating. 700 701 Furthermore, the model setup includes thermodynamic interactions between mineral dust anions (i.e., Na^+ , Ca^{2+} , K^+ , Mg^{2+}) and inorganic cations (i.e., NO_3^- , Cl^- , SO_4^{2-}). 702

The simulated CDNC at 940 hPa, i.e., at cloud base, is relatively high over the 703 industrialized areas of Europe, Asia and North America (exceeding 1,000 cm⁻³) and 704 over the biomass burning regions in the tropics (300-700 cm⁻³). Relatively high 705 CDNC is also calculated over the main deserts (100-1,000 cm⁻³) where the CCN 706 activity of pristine mineral dust is enhanced by chemical and thermodynamic 707 interactions with soluble compounds from anthropogenic (including biomass burning) 708 and natural sources. Low CDNC (around 50 cm⁻³) is calculated over the remote 709 oceans while CDNC is much higher (up to 1.000 cm⁻³) over more polluted marine 710 regions near the coast. In view of CDNCs from in situ and satellite observations, we 711 712 conclude that the model tends to underestimate CDNC over clean marine areas and overestimates CDNC over polluted regions. In the current application, CDNC 713 714 represents an upper limit with respect to the comparison with observations since we have not accounted for droplet depletion through collision and coalescence processes. 715 716 However, the model tendency to overestimate the high values of CDNC has small impact on the overall cloud radiative forcing since the sensitivity of cloud albedo to 717 718 CDNC over polluted areas is low (Seinfeld and Pandis, 2006).

To estimate the effects of mineral dust and its variable chemical composition on 719 CDNC, three main sensitivity simulations have been conducted. In the first, mineral 720 dust emissions were switched off. This reveals that despite the large tropospheric load 721 of mineral dust aerosols (35 Tg in the base case simulation) the dust presence 722 decreases the calculated global average CDNC by only 11%. This is the net result of 723 substantial positive and negative, partly compensating effects. Over polluted regions 724 (e.g., Europe), dust particles, mostly transported from the Sahara, are mixed with 725 pollution aerosols resulting in a significant reduction of the CCN activity of the 726 anthropogenic particles and hence cloud droplet formation. On the other hand, the 727 activation of freshly emitted dust particles through water adsorption results in an 728

increase of CDNC over the main deserts. However, on a global scale this is not equivalent with the calculated decrease over the polluted regions. While such sensitivity tests do not relate to real-world changes, they help understand the role of mineral dust in the climate system, and especially the importance of including these processes into climate models, being hitherto neglected.

A second simulation has been performed by switching off the mineral dust 734 chemistry to estimate the impact of interactions between inorganic and mineral 735 cations on the predicted CDNC. We find that the tropospheric burden of inorganic 736 737 anions (mainly nitrate) increases, resulting in a slight increase of CCN activity and cloud droplet formation efficiency in areas that are influenced by biomass burning and 738 industrial emissions. Furthermore, including crustal cation chemistry and 739 thermodynamics significantly affects the aging of mineral dust and its solubility, 740 especially due to the uptake of nitric acid, so that dust is removed more efficiently 741 through wet and dry deposition. This results in a decrease of CDNC over the remote 742 743 deserts (e.g., Taklimakan). Irrespective of the regional differences, the global average CDNC does not change significantly by considering mineral dust chemistry and 744 thermodynamics. 745

746 In the third simulation, the FHH calculations have been switched off to estimate the effects of water adsorption onto the surface of insoluble dust particles on the 747 748 predicted CDNC. The CDNC in the reference simulation is found to be higher over arid areas due to the adsorption activation of the freshly emitted insoluble dust 749 particles. On the other hand, CDNC is lower over polluted regions (e.g., over Europe) 750 since the aged dust particles experience significant water uptake during their 751 activation reducing the s_{max} and the activation of the smaller anthropogenic particles. 752 Overall, the use of the UAF results in a decrease of the global average CDNC by 753 ~10%. This result shows that for the modeling of cloud droplet formation, adsorption 754 activation of insoluble aerosols can be more important than mineral dust chemistry 755 and thermodynamics. However, taking into account the adsorption activation of 756 insoluble aerosols without mineral dust chemistry can result in a significant 757 overestimation of CDNC, mainly over the remote deserts. Conversely, considering 758 759 mineral dust chemistry and thermodynamics without UAF can result in significant overestimation of CDNC over polluted areas. 760

761 Finally, three additional sensitivity simulations have been conducted to investigate the sensitivity of the results to the physicochemical properties of the emitted mineral 762 dust (chemical composition, hydrophilicity and emission strength). The results 763 indicate that the calculated CDNC can be regionally sensitive to the mineral dust 764 hydrophilicity and emission load. Nevertheless, by assuming drastic differences in the 765 dust source and the dust hydrophilicity, we find only small (~5%) changes in the 766 global average CDNC. Larger CDNC changes are calculated over the main deserts 767 (up to 30%) and over highly polluted areas (up to 10%). Further, we find that the 768 769 global average CDNC is not sensitive to the chemical composition of mineral dust.

This study demonstrates that a comprehensive treatment of the CCN activity of 770 mineral dust aerosols and their chemical and thermodynamic interactions with 771 inorganic species by CCMs is important to realistically account for aerosol-chemistry-772 cloud-climate interactions. Neglecting the adsorption activation of freshly emitted 773 dust can result in significant biases over areas close to deserts. In addition, neglecting 774 the mineral dust chemistry and thermodynamics results in an underestimation of the 775 coating of dust by hygroscopic salts during atmospheric aging. The realistic 776 representation of soluble coating on dust is crucial since it affects its efficiency to 777 778 grow by water uptake, which significantly influences the local supersaturation and thus cloud droplet formation over anthropogenically polluted regions. In this first 779 780 study we apply the UAF diagnostically, while in future applications, e.g., to simulate climate responses, we plan prognostic calculations where effects on precipitation 781 782 formation and dynamical responses will also be accounted for.

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Table 1. Comparison of simulated and observed (Karydis et. al., 2011, and references)
therein) cloud droplet number concentrations.

Location	Lat.	Long.	Alt.	Time	Observation	Simulation	
S. Pacific Ocean			PBL	Annual	40	23	
S. Pacific Ocean	20S-35S	135W-175W	PBL	Annual	82	26	
Eastern Pacific Ocean	29N-32N	120W-123W	450-850m	July	49-279	133	
N. Pacific Ocean	41N	131W	<1500m	April	21-74	51	
N. Pacific Ocean			PBL	Annual	64	59	
W. of Canary Islands	32N	25W	PBL	July	17	115	
N. Atlantic Ocean			PBL	Annual	89	112	
S. Atlantic Ocean			PBL	Annual	67	51	
S. Indian Ocean			PBL	Annual	42	29	
West Australia (remote)	30S-40S	88E-103E	PBL	Annual	107	22	
Beaufort Sea (Western Arctic Ocean)	72N-78N	154W-159W	202-1017m	June	178-365	25	
Beaufort Sea (Western Arctic Ocean)	70.5N-73N	145N-147N	300-3000m	June	20-225	28	
Beaufort Sea (Western Arctic Ocean)	65N-75N	130W-170W	400-4600m	April	48-77	39	
Northeast Alaska coast	69N-71N	150W-158W	400-4000m	October	10-30	23	
Yellow Sea (Eastern coast of China)	28N-31N	127E-131E	PBL	Annual	30-1000	764	
SE Asia coast	10N-40N	105E-150E	PBL	Annual	186 (100-250)	522	
NE Asia coast			PBL	Annual	129	768	
N. America coast (Pacific)			PBL	Annual	96	91	
N. America coast (Pacific)	15N-35N	115W-140W	PBL	Annual	159 (150-300)	190	
S. America coast (Pacific)			PBL	Annual	77	75	
S. America coast (Pacific)	8S-28S	70W-90W	PBL	Annual	182 (100-300)	186	
N. Africa coast (Atlantic)			PBL	Annual	95	123	
S. Africa coast (Atlantic)			PBL	Annual	95	107	
S. Africa coast (Atlantic)	5S-25S	10W-15E	PBL	Annual	153 (130-300)	189	
Eastern N. Atlantic Ocean	50N-55N	25W-30W	800-2200m	April	65-300	39	
NW coast of Santa Maria, Azores	37N	25W	550-1000m	June	150 (74-192)	83	
Canary Islands Vicinity	28N	16.5W	PBL	June-July	51-256	174	
Canary Islands Vicinity	28N	16.5W	PBL	June-July	90-300	174	
Atlantic Ocean (W. of Morocco)	34N	11W	PBL	July	77	114	
Coast of Oregon	45.5N	124.5W	PBL	August	25-210	124	
Key West, FL	24.5N	82W	PBL	July	268-560	318	
Bay of Fundy, Nova Scotia, Canada	44N	66W	20-290m	August	61 (59-97)	246	
Cornwall Coast (SW UK)	50N	5.5W	450-800m	February	130	602	
British Isles, UK	55N	2.5W	Surface	April	172	287	
British Isles, UK	51N	6W	Surface	October	119	71	
British Isles, UK	53N	9.5W	Surface	December	96	318	
SE coast of England	51.5N-52N	1.5E-2.5E	380-750m	September	151-249	1019	
Indian Ocean (SW of India)	10S-10N	65E-75E	50-550m	February-March	100-500	520	

Table 1. Continued

Location	Lat.	Long.	Alt.	Time	Observation	Simulation	
Qinghai Province (Western China)	34N-37N	98E-103E	PBL	Annual	30-700	585	
Beijing, China	37N-41N	113E-120E	PBL	Annual	30-1100	1185	
NE China (East of Beijing)	39N-40N	117.5E-118.5E	1719-1931m	April-May	200-800	813	
Hebei Province (Central Eastern China)	35N-40N	112E-119E	PBL	Annual	30-400	1150	
Cumbria, N. England	54.5N	2.5W	Surface	March-April	100-2000	743	
Cumbria, N. England	54.5N	2.5W	Surface	May	482-549	840	
Koblenz, Germany	50N	7.5E	901-914hPa	May	675-900	1258	
Koblenz, Germany	50N	7.5E	945hPa	October	965	1039	
Northern Finland	68N	24E	342-572m	Annual	154 (30-610)	332	
Kuopio, Finland	62.5N	27.5E	306m	August-November	138	1142	
Northern Finland	68N	24E	342-572m	October-November	55-470	336	
Cabauw, Netherland	51N	4.5E	PBL	May	180-360	946	
Jungfraujoch, Switzerland	46.5N	7.5E	Surface	July-August	112-416	176	
Barrow, AK	71.5N	156.5W	389-830m	August	56	47	
Barrow, AK	71.5N	156.5W	431-736m	May	222	26	
Barrow, AK	71.5N	156.5W	297-591m	June	121	31	
Barrow, AK	71.5N	156.5W	393-762m	July	54	29	
Barrow, AK	71.5N	156.5W	1059-1608m	September	81	23	
Southern Great Plains, OK	36.5N	97.5W	795-1450m	Winter	265-281	341	
Southern Great Plains, OK	36.5N	97.5W	343-1241m	Winter	244	341	
Southern Great Plains, OK	36.5N	97.5W	985-1885m	Spring	200-219	384	
Southern Great Plains, OK	36.5N	97.5W	671-1475m	Spring	203	537	
Southern Great Plains, OK	36.5N	97.5W	1280-2200m	Summer	128-159	393	
Southern Great Plains, OK	36.5N	97.5W	756-1751m	Summer	131	603	
Southern Great Plains, OK	36.5N	97.5W	1030-1770m	Autumn	217-249	505	
Southern Great Plains, OK	36.5N	97.5W	404-1183m	Autumn	276	642	
Southern Great Plains, OK	36.5N	97.5W	900-800hPa	March	200 (100-320)	563	
Southern Great Plains, OK	36.5N	97.5W	300-600m	April	650	1159	
Southern Great Plains, OK	36.5N	97.5W	700-1200m	September-October	457	740	
Cleveland, OH; Detroit, MI	40N-42.5N	80.5W-85W	300-1000m	August	320-1300	817	
Central Ontario, Canada	50N	85W	<2500m	October	147 (119-173)	201	
Central Ontario	50N	85W	2000-2100m	Summer	350-360	143	
Central Ontario	50N	85W	1300m	Winter	190	112	
Upper NY State	44N	75W	1500m	Autumn	240	583	
State College, Pennsylvania	41N	78W	1000-1600m	October	388	551	
Mount Gibbes, NC	35.5N	82W	Surface	Annual	238-754	392	
Cape Kennedy, FL	28.5N	80.5W	600-2800m	August	250-330	134	

Table 2. Statistical evaluation of EMAC CDNC against 75 observational datasets

 worldwide, derived from in situ measurements and satellite retrievals.

	Number of	Mean Observed	Mean Simulated	MAGE	MB	NME	NMB	RMSE
Site Type	datasets	(cm^{-3})	(cm^{-3})	(cm ⁻³)	(cm^{-3})	(%)	(%)	(cm ⁻³)
Clean marine	14	86	53	51	-33	60	-39	81
Polluted marine	24	169	296	159	127	94	75	263
Continental	37	339	536	269	198	80	58	358
Total	75	237	369	193	132	82	56	295

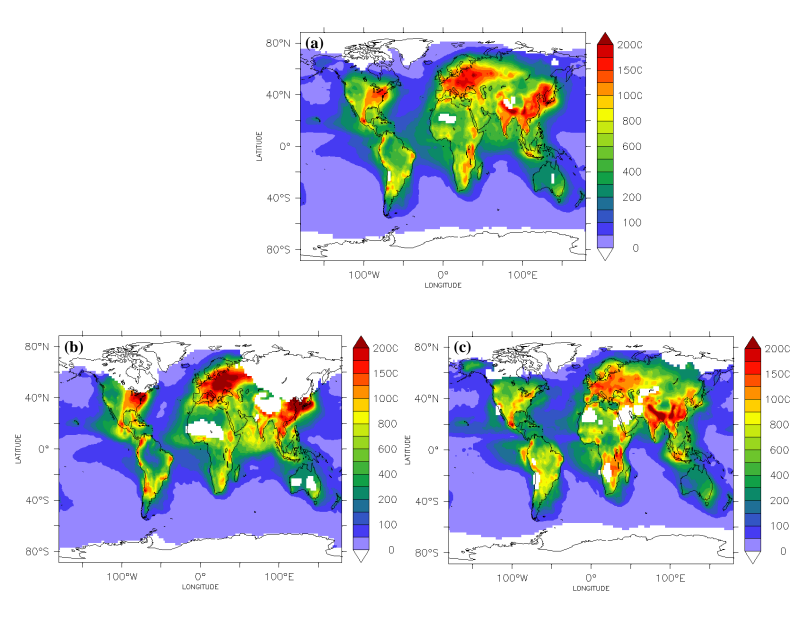


Figure 1: Predicted in-cloud (a) annual, (b) DJF, and (c) JJA mean cloud droplet number concentrations (cm⁻³) at the lowest cloud-forming level (940 hPa). White
color represents areas that are cloud-free or covered by ice clouds.

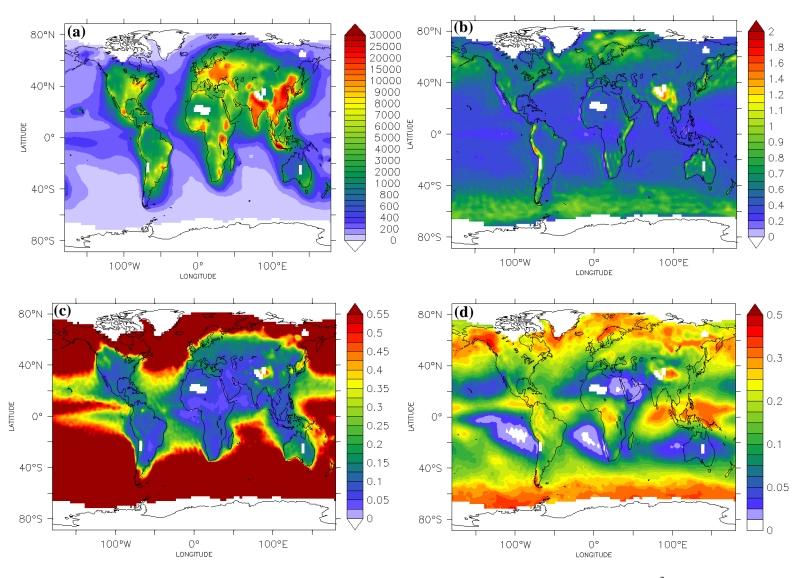


Figure 2: Predicted in-cloud annual mean (**a**) aerosol number concentration (cm⁻³), (**b**) cloud updraft velocity (m s⁻¹), (**c**) maximum supersaturation (%) at the lowest cloud-forming level (940 hPa), and (**d**) predicted annual mean low-level cloud cover. White areas correspond to regions where liquid cloud droplets do not form.

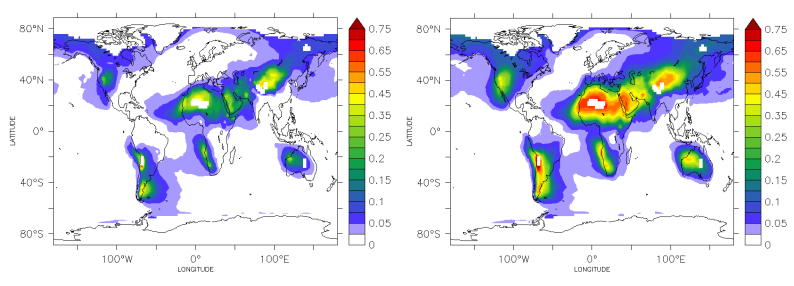


Figure 3: Predicted annual mean insoluble fraction of aerosols in the (a) accumulation and (b) coarse modes at the lowest cloud-forming level (940 hPa). White areas correspond to regions where liquid cloud droplets do not form.

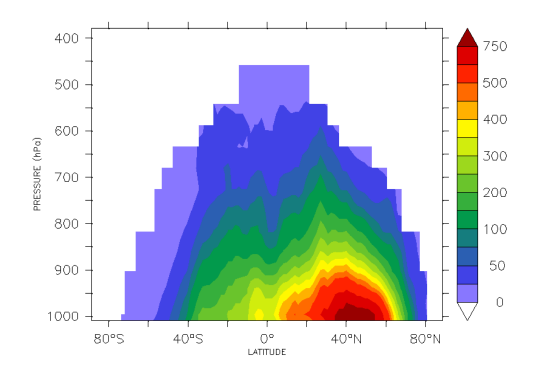


Figure 4: Predicted in-cloud zonal annual mean cloud droplet number concentration (cm⁻³). White areas correspond to regions where liquid cloud droplets do not form.

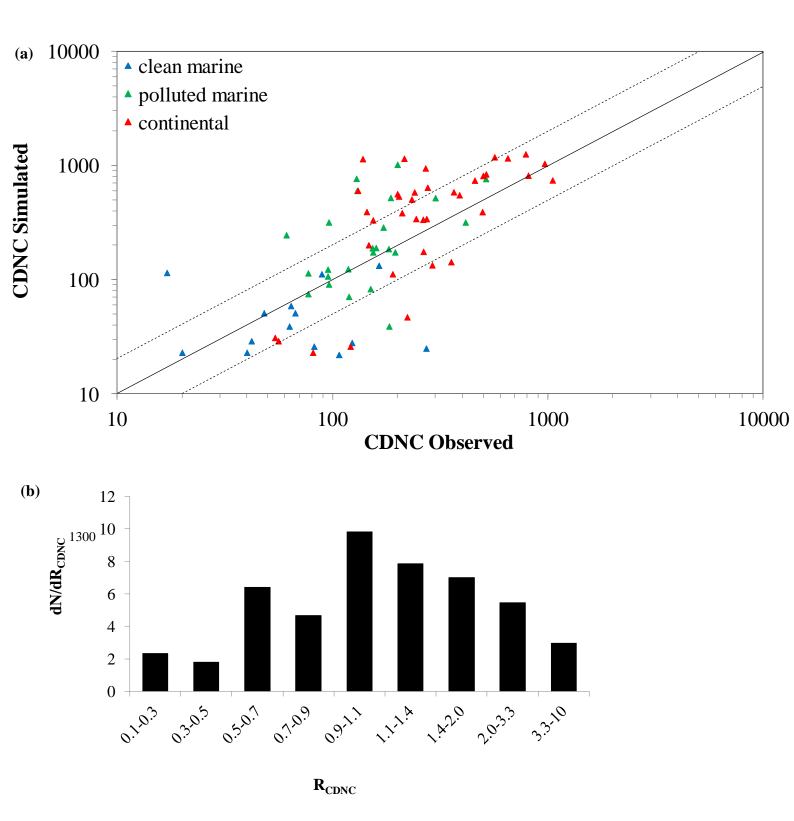


Figure 5: (a) Scatterplot comparing model simulated cloud droplet number concentrations (cm⁻³) against 75 observational datasets worldwide, derived from in situ measurements and satellite retrievals, also shown are the 1:1, 2:1, 1:2 lines, and (b) the probability distribution of the ratio of the simulated CDNC to the observed CDNC (R_{CDNC}), where N is the number of occurrences in each R_{CDNC} (inset plot).

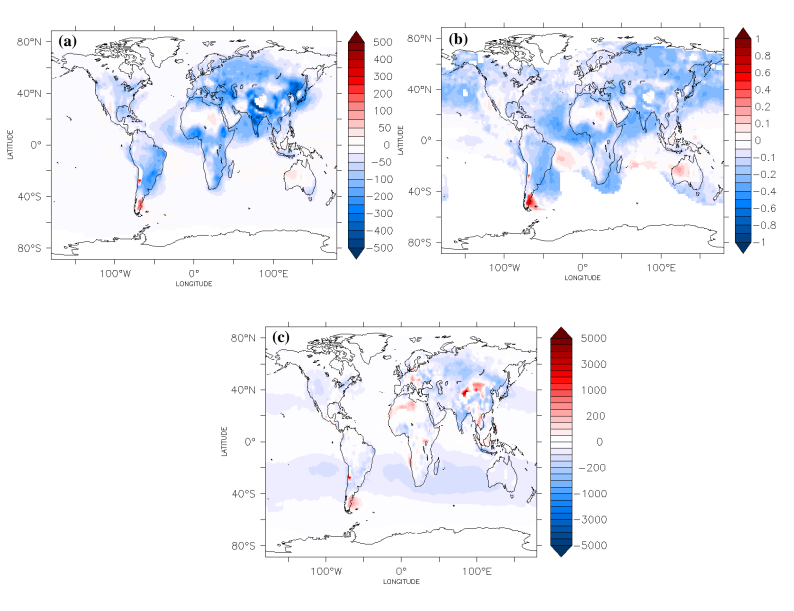


Figure 6: (a) Absolute (in cm⁻³) and (b) fractional annual change of the predicted CDNC, and (c) absolute (in cm⁻³) change of the predicted aerosol number concentration (at the lowest cloud-forming level, 940 hPa) by switching on/off the mineral dust emissions. A positive change corresponds to an increase from the presence of dust.

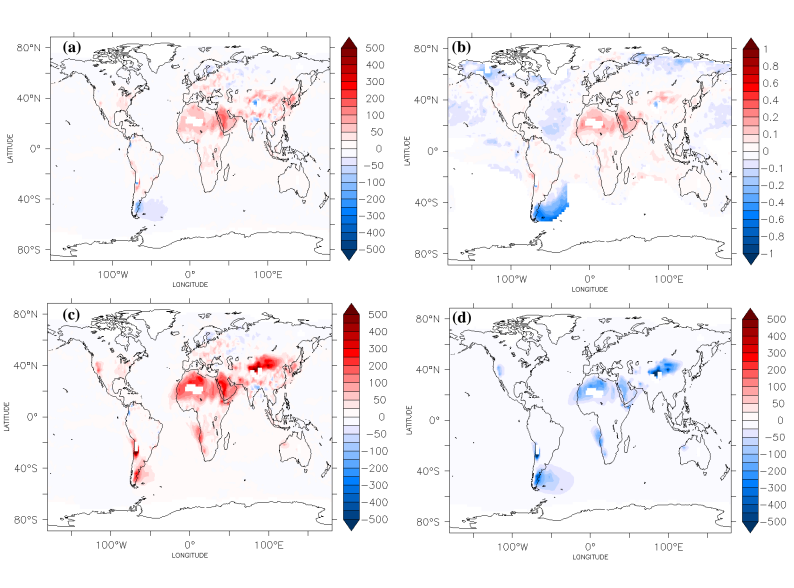


Figure 7: (a) Absolute (in cm⁻³) and (**b**) fractional annual average change of the predicted total CDNC, and absolute (in cm⁻³) annual average change of the CDNC from (**c**) soluble, and (**d**) insoluble particle modes, by switching on/off the mineral dust chemistry. Concentrations reported at the lowest cloud-forming level (940 hPa). A positive change corresponds to an increase from dust–chemistry interactions.

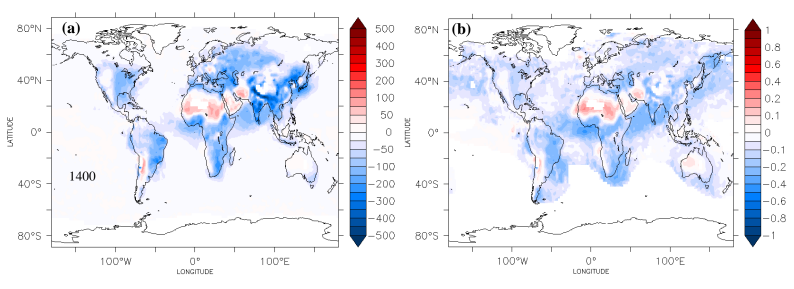


Figure 8: (a) Absolute (in cm⁻³) and (b) fractional annual average change of the predicted CDNC (at the lowest cloud-forming level, 940 hPa) by switching on/off the FHH adsorption activation physics. A positive change corresponds to an increase from water adsorption on mineral dust.

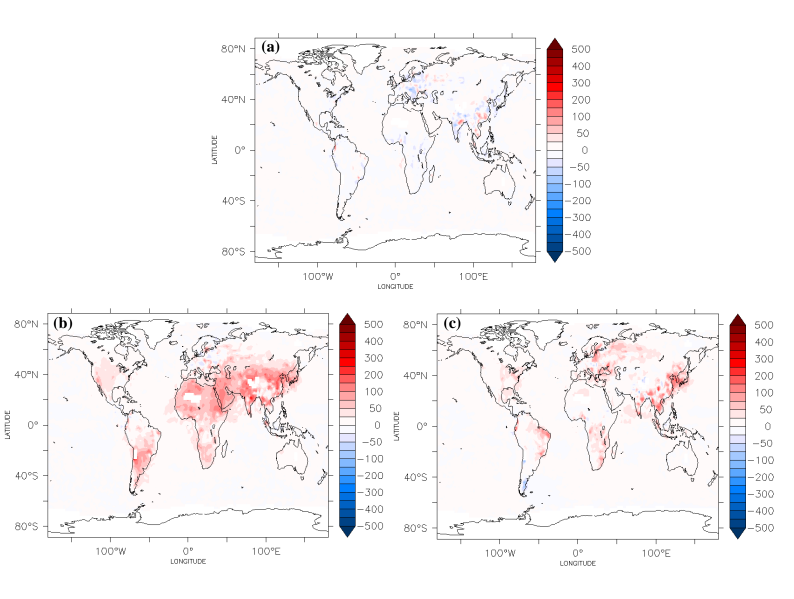


Figure 9: Absolute changes (in cm⁻³) of the predicted annual average CDNC by (**a**) assuming a globally uniform chemical composition of mineral dust, (**b**) increasing the B_{FHH} hydrophilicity parameter of dust by 10%, and (**c**) reducing mineral dust emissions by 50%. A positive change corresponds to an increase relative to the reference simulation.