1	Ice-Nucleating Particles Near Two Major Dust Source Regions
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Ice-Nucleating Particles Near Two Major Dust Source Regions

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33 Abstract

34 Mineral dust and sea spray aerosol represent important sources of ice_nucleating particles (INPs), the minor 35 fraction of aerosol particles able to trigger cloud ice crystal formation and, consequently, influence multiple climate-relevant cloud properties including lifetime, reflectivity, radiative properties and precipitation 36 37 initiation efficiency. Mineral dust is considered the dominant INP source in many parts of the world due to its ice nucleation efficiency and its sheer abundance, with global emission rates of up to 4700 Tg a⁻¹. 38 39 However, INPs emitted from the ocean surface in sea spray aerosol frequently dominate INP populations 40 in remote marine environments, including parts of the Southern Ocean where cloud-resolving model 41 simulations have demonstrated that cloud reflectivity isradiative properties are likely strongly controlled 42 by INPs. Here we report INP concentrations measured in aerosol and seawater samples during Air Quality 43 and Climate Change in the Arabian BAsin (AQABA), a shipborne campaign that spanned the Red Sea, Gulf of Aden, Arabian Sea, Arabian Gulf, and part of the Mediterranean. In aerosol samples collected 44 45 within a few hundred kilometers of the first and second ranked sources of dust globally, the Sahara and 46 Arabian Peninsula, INP concentrations ranged from 0.2 to 11 L⁻¹ at -20 °C with observed ice-active surface nucleation site densities (n_s) 1-3 orders of magnitude below levels predicted by mineral dust INP 47 parameterizations. Over half of the samples (at least 14 of 26) were collected during dust storms with 48 average dust mass concentrations between 150 and 490 µg m⁻³ (PM₁₀), as simulated by the Modern-Era 49 50 Retrospective analysis for Research and Application, version 2 (MERRA-2). The impacts of heat and peroxide treatments indicate that organics were responsible fordominated the observed ice nucleation (IN) 51 52 -activity at temperatures \geq -15 °C with proteinaceous (heat-labile) INPs frequently observed at higher 53 freezing temperatures > -10 °C. INP concentrations in seawater samples ranged between 3 and 46 mL⁻¹ at 54 -19 °C, demonstrating the relatively low INP source potential of seawater in the region as compared to 55 seawater from multiple other regions reported previously. Overall, our results demonstrate that despite 56 proximity to the Sahara and the Arabian Peninsula and the dominance of mineral dust in the aerosol 57 sampled, existing mineral dust parameterizations alone would not skillfully represent the near-surface n_s in 58 the observed temperature regime (-6 to -25 °C). The decreased n_s, and results demonstrating that organics dominated the observed IN activity > 15 °C, indicate that the IN active organic species are limited 59 60 compared to the mineral IN components of dust. Future efforts to develop or improve representations of 61 dust INPs at modest supercooling (≥ -15 °C) would benefit from a characterization of the specific organic 62 species associated with dust INPs. More generally, an improved understanding of the organic species associated with increased IN -activity and their variability across dust source regions would directly inform 63 64 efforts to determine whether n_s -based parameterizations are appropriate for faithful representation of dust

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65 INPs in this sensitive temperature regime, whether region-specific parameterizations are required, or 66 whether an alternative to the n_s approach is necessary.

1 Introduction 67

Ice-nucleating particles (INPs) modulate the temperature and relative humidity at which ice 68 69 particle formation occurs in the atmosphere and thus are a key factor that controls ice-phase 70 partitioning in clouds. As initiators of ice formation and related phase-partitioning processes, INPs 71 affect multiple cloud properties and exert a strong influence on cloud lifetime, reflectivity radiative 72 properties_and precipitation_initiation_efficiency (e.g. Lohmann and Feichter, 2005; Vergara-73 Temprado et al., 2018): (Brunner et al., 2021).

74 Globally, desert dust is likely the most abundant aerosol type by mass (Kinne et al., 2006; Kok et 75 al., 2021). Furthermore, multiple studies have demonstrated that mineral dust is the dominant ice-76 nucleating (IN) species in many parts of the world based on observations (Ardon-Dryer and Levin, 2014; Boose et al., 2016; DeMott et al., 2015a; Price et al., 2018) and modeling of global INP 77 distributions (Burrows et al., 2013; Hoose et al., 2010; Murray et al., 2012; Vergara-Temprado et 78 al., 2017). Annual global dust emission rate estimates range between 400 and 4700 Tg a⁻¹ (Huneeus 79 et al., 2011; Kok et al., 2021). Of the average global dust loading in the atmosphere (20-29 Tg), 80 81 North African source regions are estimated to contribute ~50% (11-15 Tg), and the Middle East and Central Asian source regions account for the bulk of the remainder, ~30% (7.7 Tg) (Kok et 82 al., 2021). Analysis of satellite products indicates that dust emissions rates are increasing over the 83 Middle East at a rate of 15% a⁻¹ (Klingmüller et al., 2016; Yu et al., 2018).

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85 While Hoose and Möhler (2012) showed that mineral dust INPs generally activate ice crystals at freezing temperatures < -15 °C, dust containing K-feldspar has been shown to nucleate ice at much 86 warmer temperatures, up to -4 °C (Atkinson et al., 2013; Harrison et al., 2016; Niedermeier et al., 87 2015; Wex et al., 2014; Whale et al., 2015; Zolles et al., 2015). K-feldspars represent up to ~24% 88 89 of Saharan and Asian dusts by mass (Nickovic et al., 2012). However, knowledge of the abundance and the available surface fraction of aerosolized K-feldspar would be necessary to evaluate the IN 90 efficiency of dust at temperatures > -15 °C (Kanji et al., 2017). 91

92 Though mineral dust is considered to be the dominant INP source in many regions, multiple modeling and observational studies suggest that marine INPs are frequently dominant by number 93

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94 in remote ocean regions in air masses with low concentrations of terrestrial aerosol (McCluskey et al., 2018b, 2018c; Vergara-Temprado et al., 2017; Wilson et al., 2015; DeMott et al., 2016). Using 95 96 a global aerosol model to simulate marine organic and K-feldspar INP populations, Vergara-Temprado et al. (2017) showed that the relative contribution of marine organic vs. dust INPs in 97 98 remote regions varies seasonally, and that marine organic INPs frequently outnumber K-feldspar INPs (up to 100% of the simulated days in the Southern Ocean during summer). Results from a 99 100 follow-on cloud-resolving model study showed that Southern Ocean cloud reflectivity is strongly modulated by INP concentrations, indicating that accurate estimates of the radiative energy budget 101 in the Southern Ocean likely require improved and reliable representation of both dust and marine 102 organic INPs (Vergara-Temprado et al., 2018). By generating isolated nascent sea spray aerosol 103 over a range of biological conditions, mesocosm studies have shown that marine INPs are 104 105 comprised of two classes: a dissolved organic carbon (DOC) type composed of IN-active molecules and a particulate organic carbon (POC) type linked to the death phase of phytoplankton 106 107 blooms (McCluskey et al., 2017, 2018b).

108 Parameterizations for both marine and mineral dust populations are commonly implemented in atmospheric models to estimate dust and marine INP concentrations. There are multiple existing 109 mineral dust INP parameterizations used to estimate their concentrations in aerosolized desert dust, 110 111 some based exclusively on laboratory measurements (e.g., Niemand et al., 2012; Ullrich et al., 112 2017), and others derived from a combination of laboratory and field measurements (DeMott et al., 2015). There are, additionally, multiple mineral-specific INP parameterizations including illite 113 114 (Broadley et al., 2012), kaolinite (Welti et al., 2012), quartz (Harrison et al., 2019) and K-feldspar 115 (Atkinson et al., 2013; hereafter, "A13"). The parameterizations by Ullrich et al. (2017; hereafter, "U17") and Niemand et al. (2012;, hereafter, "N12") were developed using dust samples from 116 117 multiple deserts, and both found little variability in the IN activity between dusts from locations as disparate as the Sahara and Asia. DeMott et al. (2015;, hereafter, "D15") found agreement 118 119 between their observations-based parameterization and N12, supporting the validity of laboratorybased parameterizations. Results in D15 also confirmed the conclusions of N12 and U17: that to 120 first order, dusts from distinct regions can be parameterized as a single particle type. The D15 121 parameterization has been considered to be representative of dust that has undergone atmospheric 122 photochemical and oxidative processes in transport (i.e., "aged" dust), because the 123

parameterization was derived from observations made far (1000s of kilometers) from the dustemissions sources (Boose et al., 2016).

By contrast, few studies report *in situ* INP measurements near (e.g., < 1 day of transport) a major 126 127 dust source, and the lack of observations near dust source regions inhibit the evaluation of the 128 ability of existing dust INP parameterizations to represent nascent dust populations (Boose et al., 129 2016; Gong et al., 2020; Price et al., 2018). INP observations are particularly lacking for the 130 sensitive temperature regime > -20 °C. Boose et al. (2016) found that D15 overpredicted INPs 131 observed during Saharan dust events at a location within 100s of km west of the Sahara (Izaña, Tenerife, Spain) by 2-3 orders of magnitude, suggesting that aging may lead to increased IN 132 133 efficiency in mineral dust and that D15 may be less representative of nascent dust. These conclusions were supported by Conen et al. (2015), who found that concentrations of INPs at -20 134 135 °C measured during Saharan dust events were one order of magnitude higher at Jungfraujoch in the Swiss Alps than in Izaña, where dust events had occurred 1-7 d prior to reaching Jungfraujoch. 136 Gong et al. (2020) measured INPs in a variety of atmospheric and seawater sample types at Cabo 137 138 Verde and determined mineral dust to be the dominant source of INPs observed in the atmosphere but found that INPs with freezing temperatures > -10 °C were likely biological. At altitudes 139 between 30 and 3500 m in the same region, Price et al. (2018) found that measured concentrations 140 of INPs ranged two orders of magnitude at a given temperature, and that the observed 141 142 concentrations related to the atmospheric dust loadings.

Recently, multiple studies have provided new, much-needed observations of ambient atmospheric 143 INPs in marine environments (DeMott et al., 2016; Hartmann et al., 2020; McCluskey et al., 2018c, 144 2018d; Yang et al., 2020) where data was historically lacking and, consequently, an impediment 145 146 to achieving predictive understanding of global INP distributions (Burrows et al., 2013). There are now two parameterizations available for the estimation of atmospheric concentrations of 147 148 marine INPs emitted from the ocean surface: Wilson et al. (2015), which estimates cumulative INPs from total organic carbon (TOC) concentrations in simulated sea spray aerosol (SSA), and 149 150 McCluskey et al. (2018), which estimates ice-active surface-nucleation-site density (n_s) from 151 aerosol surface area. Wilson et al. (2015) and McCluskey et al. (2018; hereafter "M18") derived 152 marine INP parameterizations from field measurements of INPs in Atlantic and Arctic Ocean sea 153 surface microlayer samples and pristine SSA samples over the North Atlantic Ocean, respectively.

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154 Here, we report observations of INPs measured in air masses influenced by both desert dust and marine aerosol (Edtbauer et al., 2020) in close proximity to the two greatest global dust aerosol 155 156 sources: the Sahara (#1) and the Arabian Peninsula (#2) (Kok et al., 2021). INP concentrations 157 were measured in 26 ambient aerosol samples collected during Air Quality and Climate Change 158 in the Arabian Basin (AQABA), a shipborne campaign which took place July – August 2017 on a 159 transect that spanned the -central and eastern parts of the Mediterranean, the Red Sea, the Gulf of 160 Aden, the Arabian Sea and the Arabian Gulf. The rest of this study will be structured as follows. 161 We present an overview of measurements and data sources in Sect. 2 Methods. In Results Sect. 162 3.1, an overview of INP concentrations observed is presented, followed by an assessment of 163 subsurface seawater (SSW) source potential (Sect. 3.2). Observed Observed ns were are compared 164 to dust and marine INP parameterizations in Sect. 3.3, followed by an analysis of the, and the 165 contributions of heat-labile (e.g., proteinaceous) and heat-stable organic compounds to observed 166 INP populations in aerosol (Sect. 3.4). The same analysis is applied to assess organic contributions 167 to observed INPs in a soil dust sample from a likely source region in Sect. 3.5. We discuss the findings, potential INP sources and compare with prior studies in the Discussion Sect. 5. Finally, 168 169 in Sect. 5 we offer strategies to address the challenges of evaluating dust-specific INP 170 parameterizations and recommend measurements needed to develop predictive understanding of 171 dust INPs at modest supercooling (T \geq 15°C). were assessed via heat and peroxide treatments. 172 Finally, the potential INP source strengths of subsurface seawater (SSW) were assessed and 173 compared with SSW INP measurements from prior studies of remote and coastal seawater.

174 2 Methods

175 2.1 Project Overview

The AQABA campaign was conducted from 25 June to 3 September 2017 onboard the RV *Kommandor Iona*. The research voyage was conducted in two transects: the first leg beginning in
La-Seyne-sur-Mer, France, heading through the Suez Canal, around the Arabian Peninsula and
ending in Kuwait, and second leg a return transect via the same route (Figs. S1-S2). The campaign
supported a large suite of on- and offline aerosol and gas-phase measurements (Bourtsoukidis et

al., 2019, 2020; Celik et al., 2020; Edtbauer et al., 2020; Eger et al., 2019; Friedrich et al., 2021;
Pfannerstill et al., 2019; Tadic et al., 2020; Wang et al., 2020).

183 2.2 Aerosol and Trace Gas Measurements

Aerosol size distributions were measured using an Optical Particle Spectrometer (OPC, Grimm 184 model 1.109) and a Fast Mobility Particle Spectrometer (FMPS, TSI model 3091). The OPC 185 measures particles in the size range $0.25 - 32 \mu m$, and the FMPS measures particles with sizes 186 187 between 5.6 nm and 560 nm with 6s and 1s time resolution, respectively. The inlet for the aerosol 188 instrumentation was located at the top of a measurement container at a horizontal distance of about 189 1525 m from the INP filter sampling unit (Figs. S3-S4). To avoid condensation in inlet lines, 190 aerosol samples were passed through a drying system, which reduced ambient relative humidity 191 (RH) to an average value of $\approx 40\%$ in the measurement container. Ambient RH ranged between 192 67 and 81% during INP sampling periods. OPC and FMPS data were averaged over 1-minute time 193 intervals. A filter flag based on aerosol measurements was derived to identify and eliminate stack 194 emissions and applied to all aerosol data. The filter flag was based on short term variation in 195 particle number concentration measured by a Condensation Particle Counter (CPC, TSI model 196 3787), black carbon concentrations (Aethalometer, Magee AE33), wind direction and speed. The 197 flag was set when the apparent wind direction was from the direction of the stack $(\pm 30^\circ)$ as seen from the aerosol inlet position (Fig. S3) and strong fluctuations of black carbon and/or particle 198 199 number concentrations were observed relative to background levels. Two additional measurements 200 provided aerosol data from which a filter flag intended to identify and eliminate stack emissions 201 was derived: particle number concentrations as measured by a Condensation Particle Counter 202 (CPC, TSI model 3787) and black carbon concentrations (Aethalometer, Magee AE33). The filter 203 flag, based on short term variation in particle number concentration, black carbon concentration, 204 wind direction and speed, was applied to all aerosol data so that samples contaminated by stack 205 emissions could be identified. Particle losses were estimated using the Particle Loss Calculator (von der Weiden et al., 2009), Losses were negligible (<1%) up to 3.5 µm and increased to 40% 206 207 <u>at 10 µm.</u>

Particle surface area concentrations were derived from the 1-min time-averaged FMPS and OPC
 measurements as follows. Geometric diameters were estimated from the measured mobility

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210 diameters (FMPS) and optical particle diameters (OPC). Aerosols were assumed dry at sampling 211 conditions following the drying system described above. To convert optical particle diameters into 212 geometric diameters, it was assumed that all coarse particles ($d_p > 3000$ nm) were composed of 213 sea salt and dust with a mass ratio of 25% to 75%, and <u>for using</u> the respective refractive indices 214 and shapes the measured optical particle diameters were converted into geometric diameters <u>(Sect.</u> 215 <u>S1.1)</u>. The sea salt:dust mass ratio was based on average dust and sea salt concentrations as 216 measured in particles < 10 µm (PM₁₀, see Sect. S2 for details).

217 Fine particle ($d_p < 700 \text{ nm}$) sizes were was converted from optical diameter (d_{opt}) into geometric 218 diameter (dgeo) using the optical properties calculated from the PM4-chemical composition of 219 particles $< 1 \mu m$ (PM₁) as measured by an Aerosol Mass Spectrometer (Aerodyne HR-ToF-AMS), 220 assuming spherical particles (Celik et al., 2020). For particles in the intermediate size range (700 221 -3000 nm), log-linear interpolation of optical and spherical properties was applied for conversion 222 of optical into geometric particle diameters (Sect. S1.2). The mobility diameters measured by the 223 FMPS were considered equivalent to the geometric diameter, assuming spherical particles. From 224 the resulting particle size distributions, particle surface area was calculated for each size bin. Total 225 particle surface concentrations were determined by integrating the surface area distribution for 226 particles up to 10 µm (dgeo). The overall uncertainty of derived particle surface area concentrations 227 is estimated to be 30%, including the uncertainty due to particle losses (see Sect. S3).

228 The water-soluble fraction of total suspended particles (TSPs) was monitored with hourly 229 resolution using a Monitor for AeRosols and Gases in Ambient Air, MARGA (Metrohm Applikon 230 model S2, Herisau, Switzerland). Sea salt concentrations were estimated by scaling measured 231 soluble Na⁺ concentrations by 3.27 following Manders et al. (2009) and were used as a proxy for 232 SSA number concentrations. Size-resolved single particle chemical composition measurements 233 have shown that sea salt represents 50-70% of SSA particles by number ($d_p > 0.5 \mu m$) (Collins et 234 al., 2014)-Hourly composition data was linearly interpolated for 4 samples where 1-3 hours (of 7-235 24 hours total sampling time) was missing (Fig. S53). The MARGA sampling line was

236 equipped with a PM₁₀ cyclone, but the sample was not dried as the instrument is not prone to

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(GOCART) model (Chin et al., 2002; Colarco et al., 2010). Dust emissions and deposition rates in MERRA-2 are estimated by summing the emissions and deposition rates across GOCART

259 simulated dust particles between 0.1 - 10 µm in size (dry diameter) (Gelaro et al., 2017). Dust 260 emissions are constrained by wind-driven erosion over the source locations, which are identified various satellite products and are jointly assimilated within GEOS 5 with meteorological

from the topographic depression map (Ginoux et al., 2001). Acrosol observations are derived from 261 262

263 observations (Buchard et al., 2017). MERRA 2 has been shown to successfully reproduce the

interannual variability of North-Atlantic dust transport. Additionally, the improved aerosol 264

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The NO measurements were used to prevent stack sampling during INP collection (see Sect. 2.4). 243

244 2.3 Dust Mass Concentrations from MERRA-2

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239 Nitric oxide (NO) concentrations were measured using a commercially available two-channel chemiluminescence monitor, CLD 790 SR (ECO Physics AG, Dürnten, Switzerland). During the 240 AQABA campaign, the CLD 790 SR, MARGA, FMPS, OPC, HR-ToF-AMS, CPC and 241

Aethalometer were operated within laboratory containers on the main deck of the research vessel.

Area averaged hSince dust concentrations were not measured during the campaign, hourly dust

surface mass concentrations along the cruise track were obtained from the (0.5 \times \star 0.625 °)

Modern-Era Retrospective analysis for Research and Application, version 2 (MERRA-2; Gelaro

et al., 2017) and were averaged over the region covered during each sampling period. (Buchard et

al., (-2017) showed a high degree of correlation between MERRA-2 and surface dust concentration

observations (r \ge 0.69), particularly during dust storms (r \ge 0.92). MERRA-2 surface dust mass

concentrations also correlated well with PM_{10} observed during AQABA (r ≥ 0.71) (Fig. S6).

MERRA-2 uses the GEOS-5 Earth system model (Molod et al., 2015; Rienecker et al., 2011) with 72 vertical layers between the surface and 0.01 hPa (~ 80 km) and the three-dimensional variational

data assimilation Gridpoint Statistical Interpolation analysis system (Kleist et al., 2009; Wu et al.,

2002, additional details in Sect. S4). It simulates 5 types of aerosols (dust, sea salt, sulfate and

black and organic carbon) using the Goddard Chemistry, Aerosol, Radiation, and Transport

to be consistent with the aerosol sizing instruments described above.

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condensation. Particle transmission losses to the MARGA were estimated using the PLC and found Formatted: Font: (Default) Times New Roman, 12 pt

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assimilation scheme in MERRA-2 was shown to have a positive impact on the representation of
 long-range dust transport from the Sahara compared to prior versions (Buchard et al., 2017).

267 2.4 INP Measurements in Aerosol Measurement of Ice Nucleating Particles

Ambient aerosol sampling for offline measurement of INPs was conducted from 5 Jul – 31 Aug
2017 on the *Kommandor Iona's* wheelhouse top (platform above the bridge), ~<u>1</u>25 m horizontally
from the online aerosol measurements inlet and ~<u>1</u>35 m from the ocean surface (Figs. S<u>3</u>-4).
Sampling locations along the cruise transect corresponding to each aerosol sample are shown in
Figs. S1-2.

Aerosol samples were collected over 3-28 hour periods on polycarbonate filters (47 mm diameter, 273 0.2 µm pore-size, Whatman® Nuclepore, Chicago, Illinois, USA) placed in open-face Nalgene® 274 275 Analytical Filter Units (Waltham, Massachusetts, USA). Sampling intervals and frequency were 276 chosen with the aim of collecting > 5000 L during dust events and > 10,000 L when OPC particle counts were relatively low (e.g., during sampling periods f040-44), as conditions allowed. Aerosol 277 sampling flow rates through the filter units were set to 10-13 Lpm-LPM using a MassStream[™] 278 mass flow controller (Bethlehem, PA, USA) connected inline with a rotary vane pump (Thomas 279 280 QR-0100, Gardner Denver ©, Monroe, LA, USA). To decrease exposure to stack emissions, the pump was automated to switch off when online measurements of NO exceeded one standard 281 282 deviation above the average background concentration for over 1 minute (~ 0.4 ± 0.8 ppb). 283 Comparing the stack contamination filter flag for aerosol measurements (Sect. 2.2) with INP 284 sampling periods additionally indicates no influence of stack emissions on INP filter samples. 285 Lacking a size-selective inlet for INP sampling, it is possible that aerosols $> 10 \,\mu\text{m}$ were present 286 in INP samples during dust events. Surface area may be underestimated for these samples due to the PM₁₀ cutoff for aerosol sizing (Sect. 2.2 and S3), but we do not expect this to affect our overall 287 288 conclusions as increased aerosol surface area would further reduce ns (see Results Sect. 3.3 and 289 Discussion Sect. 4). -

Prior to sampling, filters were cleaned by soaking in $10 \% \text{ peroxide (H}_2O_2)$ for 10 minutes followed by rinsing three times with deionized water, the last rinse further "polished" by passage through a 0.1 µm pore-size syringe filter (Puradisc, Whatman ®, Maidstone, U.K). Filters were pre-loaded into filter units in a laminar flow hood to further minimize contamination from handling. After

294 collection, each aerosol filter was placed in a 60 mm diameter sterile Petri dish (Life Science Products, Frederick, Colorado, USA) using pre-cleaned acetyl plastic forceps (Fine Science Tools, 295 296 Foster City, California, USA), sealed with Parafilm and stored frozen (-20 °C). Samples were shipped in a dry shipper via Cryoport® High Vol Shipper at -180 °C and upon arrival at the 297 298 laboratory were stored at -80 °C until processed, within 18 to 38 months of since collection. To release collected particles, filters were immersed in 5-8 mL ultrapure water (Cat. Number W4502, 299 300 Sigma-Aldrich®, St. Louis, MO, USA) and shaken by hand for 20 minutes just prior to measurement. Eight samples were additionally diluted 100-fold to measure INP concentrations 301 302 at lower freezing temperatures (Fig. S5).

303 INP concentrations were measured using the Scripps Institution of Oceanography SIO-Automated Ice Spectrometer (SIO-AIS), an immersion freezing droplet assay instrument that is described in 304 305 detail in Beall et al. (2017). Briefly, the aerosol sample suspensions and SSW samples were 306 distributed in 30 ×* 50-µL aliquots into clean 96-well polypropylene sample trays (OPTIMUM®) ULTRA Brand, Life Science Products). An equal number and volume of aliquots of ultrapure 307 308 water accompanied each sample in the tray as a control. Trays were then inserted into an aluminum 309 block that was cooled at -0.87 °C min⁻¹ until the samples are frozen. Cumulative INP number concentrations per temperature per volume liquid are calculated using the fraction (f) of unfrozen 310 311 wells per given temperature interval:

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$$n_{\text{INP,L}} = \frac{-ln(f)}{V_d}$$
 Eq. (1)

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where V_d is the volume of the sample in each well_(Vali, 1971). For aerosol filter samples, cumulative INP number concentrations are calculated using the ratio of the <u>ultrapure water</u> volume used for resuspension of the particles (V_{re}) to the volume of air sampled (V_A):

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$$n_{\text{INP}} = \frac{-ln(f) \cdot V_{re}}{V_d \cdot V_A}$$
 Eq. (2)

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321 Prior to calculating n_{INP} , the fraction of unfrozen wells (f) was adjusted for contamination in the 322 water used for suspension by subtracting the number of frozen ultrapure water wells per 323 temperature interval from both the total number of unfrozen wells and total wells of the sample. The $n_{\rm INP}$ was additionally adjusted for background INPs from filters and sampling handling 324 325 processes. Background $n_{\rm INP}$ INP concentrations were estimated using measured $n_{\rm INP}$ INP concentrations in aerosol sample field blanks, which had been momentarily placed placed in the 326 327 sampling apparatus $\sim 5s$ (without actuating the pump) before removal and unloading and storage 328 of the filter. Seven field blank samples were collected, one every ~ 7 days of the cruise (Fig. S7). 329 INP concentrations were measured in field blanks as described above, and the n_{INP} simulated using the mean air volume sampled (6680 L). Figure S $\frac{76}{10}$ shows the estimated $n_{INP} n_{INP}$ across the 7 330 field blanks, which ranged between 13.0×10^{-4} and 3.0×10^{-32} L⁻¹ at -20 °C. The freezing onset 331 332 temperatures detected in the field blanks ranged between -6 and -27 °C. To correct $n_{\rm INP}$ $n_{\rm TAPP}$ 333 measured in aerosol samples for background INPs from sample handling, a linear regression-of 334 the average based on the geometric mean $n_{\rm INP}$ to concentration measured in field blank 335 suspensions (mL⁻¹ water) was used to estimate background concentrations of INPs in samples at 336 all temperatures between -14.5 °C and -27 °C. The estimated background $n_{\rm INP}$ in the estimated background $n_{\rm INP}$ is the estimated $n_{\rm INP}$ is the estimated 337 was then subtracted from the INP concentration measured in filter sample suspension volumes in 338 this temperature range prior to calculating $n_{\rm INP}$ $n_{\rm INP}$. The $n_{\rm INP}$ $n_{\rm INP}$ -measured in one aerosol 339 sample (f033) fell within the estimated INP background levels.

For this study, the detection limit was 0.68 n_{INP} _INPs mL⁻¹ liquid or 0.001-0.0024 n_{INP} _INPs L⁻¹ air for the maximum and minimum and maximum air volume sampled, respectively. To extend the upper limit of detection (i.e., the point at which all droplets have frozen) dilutions of 1:10 and 1:100 were performed on 8 samples (Fig. S<u>8</u>5).

The ice-active surface site density, n_s , is a metric used to define the ice-nucleating capabilities of an aerosol species (i.e., an aerosol sample of all the same particle type) (Kanji et al., 2017) as follows:

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$$n_s = \frac{N_{ice}}{N_{tot} \times A \ (cm^2)}$$
 Eq. (3)

where N_{ice} is the number of frozen droplets, N_{tot} is the total number of particles in a monodisperse aerosol population, and A is the surface area per particle. The value of n_s can also be approximated for polydisperse aerosol samples containing multiple aerosol types:

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$$n_s = \frac{N_{ice}}{A_{tot} (cm^2)}$$
 Eq. (4)

353 where A_{tot} is the total surface area of the polydisperse aerosol-sample. The difference between the 354 n_s approximation (Eq. 4) and n_s (Eq. 3) is that many particle types are typically included in the n_s 355 approximation, and in an ambient aerosol measurement most of these are not IN-activeice 356 nucleating (see also Hiranuma et al., (2015) Sect. 2.4). Furthermore, the subset of INPs in the 357 sample are likely also of different types, which likely have different n_s in the strict sense (Eq. 3). Nevertheless, the n_s approximation is a useful metric for comparing the ice-nucleating ability of 358 different air masses and source regions and is often used for comparing data across studies of INPs 359 measured in ambient air. It is extremely challenging to separate measurements of INPs and surface 360 area by each particle type, and requires, for example, combining online measurements of single 361 362 particle chemistry, size distributions and INPs (Cornwell et al., 2019). All n_{INP} and n_s are reported 363 normalized to a standard temperature of 273.15 °K and pressure of 1013 hPa.

364 Heat and hydrogen peroxide_treatments were applied to a subset of samples (12 of 26) to test for 365 heat-labile biological (e.g., proteinaceous) and organic INP composition, respectively, following 366 the procedure described in (McCluskey et al., (2018b) and; Suski et al., (2018). The 12 samples 367 were selected based on sampling location with the aim of getting a representative measurement 368 from each region. For each heat-treated sample, a 2 mL aliquot of the original ultrapure water suspension was heated to 95 °C for 20 min in a water bath and re-tested to assess the reduction in 369 370 <u>INP concentrations for changes in n_{INP} </u>. For peroxide treatments, 1.6 mL of the original suspension 371 was combined with 0.8 mL of 30% H2O2-peroxide (Sigma Aldrich®, St. Louis, Missouri, USA) 372 to achieve a final concentration of 10%, then the mixture was immersed in water, and heated to 95 373 °C for 20 min while being illuminated with two 26-W UVB fluorescent bulbs to generate hydroxyl 374 radicals. To remove residual- H_2Q_2 peroxide and, to prevent otherwise significant freezing point 375 depression, the solution was cooled and catalase (Cat. Number IC10042910, MP Biomedicals, 376 Santa Ana, California, USA) was added. Since catalase is itself decomposed by H₂O₂, while 377 simultaneously catalyzing peroxide's disproportionation into water and oxygen, the enzyme was 378 added in several 20 µL aliquots, allowing several minutes between each, until no effervescence 379 resulted upon its addition. Figure S7 shows the estimated $n_{\rm INP}$ in a heat and peroxide-treated blank

380 sample. Fisher's Exact Test was applied to frozen and unfrozen well fractions between each 381 untreated sample and its corresponding treated sample to test for significant differences (p < 0.05). 382 Note that significant difference in frozen well fraction is insufficient as a sole indicator of sensitivity in peroxide treated samples because samples are diluted 2:3 (by 33%) compared to 383 384 untreated samples. -As $n_{\rm INP}$ -<u>INP concentrations</u>-can be corrected for the dilution by scaling (as opposed to frozen well fractions), the overlap in 95% binomial sampling confidence intervals 385 386 (Agresti and Coull, 1998) between the untreated and peroxide-treated sample is an additional indicator of sensitivity for a given data point in the peroxide-treated sample spectrum within ± 0.2 387 388 °C, the uncertainty in the SIO-AIS temperature measurement (Beall et al., 2021). A lack of overlap in the 95% binomial sampling confidence interval within ± 0.2 °C equates to a significance 389 390 threshold of p < 0.005 (Krzywinski and Altman, 2013). INP concentrations were additionally measured in in untreated, heat-treated, and peroxide-treated 391 392 subsamples from an archived suspension of the soil dust sample N12 SD for comparison with this 393 study (DeMott et al., 2018; hereafter referred to as "N12-SD"). N12 SD (sample CD1 in Megahed

394 <u>2007</u>) is sedimented airborne dust that was carried in boundary layer winds north and then east

395 from the desert in central Egypt during a sandstorm to a sampling point ~50 km north of Cairo,

396 collected on 18th Feb 2003. The sample was stored in a "clean bottle" (no temperature of storage

given) for further analysis (Megahed 2007). Briefly, the sample was generated during the recent

<u>laboratory intercomparison of INP measurements-(DeMott et al., (2018), collected on a 0.2 μm</u>

399 <u>Nuclepore polycarbonate membrane filter (Whatman ®, Chicago, Illinois, USA) and stored frozen</u>

400 <u>at -20 °C until processed</u>, as described in DeMott et al (2018).

402 <u>2.5 INP Measurements in SSW</u>

401

INP concentrations were additionally measured in 10 SSW samples. For seawater sampling, a
water intake vertical steel pipe was positioned on the starboard of the ship approximately 2 m
below the sea_surface level. The seawater was pumped into a 200 L stainless steel tank and
continuously exchanged at a rate of 3000 L h⁻¹. SSW samples for INP analysis were collected in
15 mL sterile centrifuge tubes (Falcon[™], ThermoFisher Scientific, Waltham, Massachusetts,
USA) and stored frozen at -20 °C until they could be shipped in a dry shipper via Cryoport[®] (-180
°C) and ultimately stored at -80 °C as for aerosol samples <u>until processed as described above (Sect.</u>

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4), within 18 to 38 months since collection. Storage duration was not correlated with INP
ncentration changes in frozen marine and coastal precipitation samples (Beall et al., 2020). Heat
d hydrogen peroxide treatments as described above were applied to five <u>5 SSW samples from</u>
Arabian Sea and the Gulf of Aden. The focus on these regions was motivated by the detection
marine aerosol originating from the upwelling region in Somalia reported in (Edtbauer et al.,
020; see Sect. 3.3).of these. To assess the contribution of submicron INPs to total measured
Ps, 2 mL of SSW was filtered through a 0.2 µm sterile syringe-filter (Acrodisc® Pall®, Port
ashington, New York, USA) and re-tested.
P concentrations in SSW collected at the Ellen Browning Scripps Memorial Pier at Scripps
stitution of Oceanography (SIO: 32,8662 N, 117,2544 W) were assessed in 17 samples for
mparison with SSW collected during AOABA. Samples were collected between 31 Jan and 7
av 2016 in 15-30 mL sterile centrifuge tubes (Falcon™, ThermoFisher Scientific, Waltham,
assachusetts, USA) at depths of 1-3m and processed immediately using the SIO-AIS as
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Scribed above. SFLEXPART Back Trajectories r mass 72 hour back trajectories for each sample were simulated using the FLEXible PARTicle rpersion model (FLEXPART) in backward mode (Stohl et al., 1998). NOAA Climate Forecast
Scribed above. SFLEXPART Back Trajectories r mass 72 hour back trajectories for each sample were simulated using the FLEXible PARTicle r persion model (FLEXPART) in backward mode (Stohl et al., 1998). NOAA Climate Forecast stem (CFS) short duration (t < 6 h) forecasts (Saha et al., 2014) were used as three-dimensional
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Scribed above. SFLEXPART Back Trajectories SFLEXPART Back Trajectories for each sample were simulated using the FLEXible PARTicle persion model (FLEXPART) in backward mode (Stohl et al., 1998). NOAA Climate Forecast stem (CFS) short duration (t < 6 h) forecasts (Saha et al., 2014) were used as three-dimensional reing datasets. Particle releases from 35 m above sea level (ASL) followed the vessel track using seel position information from the European Common Automatic Weather Station (EUCAWS; p://eumetnet.eu/; last access Sept. 2021).
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Scribed above. S S FLEXPART Back Trajectories File Contract trajectories for each sample were simulated using the FLEXible PARTicle persion model (FLEXPART) in backward mode (Stohl et al., 1998). NOAA Climate Forecast stem (CFS) short duration (t < 6 h) forecasts (Saha et al., 2014) were used as three dimensional coing datasets. Particle releases from 35 m above sea level (ASL) followed the vessel track using seel position information from the European Common Automatic Weather Station (EUCAWS; p://eumetnet.eu/; last access Sept. 2021).
Seribed above. SELEXPART Back Trajectories F HEXPART Back Trajectories for each sample were simulated using the FLEXible PARTicle persion model (FLEXPART) in backward mode (Stohl et al., 1998). NOAA Climate Forecast stem (CFS) short duration (t < 6 h) forecasts (Saha et al., 2014) were used as three dimensional reing datasets. Particle releases from 35 m above sea level (ASL) followed the vessel track using seel position information from the European Common Automatic Weather Station (EUCAWS; p://eumetnet.eu/; last access Sept. 2021).
Seribed above. SFLEXPART Back Trajectories r mass 72 hour back trajectories for each sample were simulated using the FLEXible PARTicle persion model (FLEXPART) in backward mode (Stohl et al., 1998). NOAA Climate Forecast stem (CFS) short duration (t < 6 h) forecasts (Saha et al., 2014) were used as three dimensional
seribed above. FIEXPART Back Trajectories r mass 72 hour back trajectories for each sample were simulated using the FLEXible PARTicle persion model (FLEXPART) in backward mode (Stohl et al., 1998). NOAA Climate Forecast stem (CFS) short duration (t < 6 h) forecasts (Saha et al., 2014) were used as three-dimensional reing datasets. Particle releases from 35 m above sea level (ASL) followed the vessel track using ssel position information from the European Common Automatic Weather Station (EUCAWS; <u>p://eumetnet.eu/;</u> last access Sept. 2021). Results and Discussion - Characteristics of INPs Observed During AQABA
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437	Mediterranean Sea, the Red Sea, the Gulf of Aden, Arabian Sea, Gulf of Oman, and Arabian Gulf
438	spanned up to 23° orders of magnitude at -15 °C (Fig. 1, Table 1), between 5×10^{-3} and 5×10^{-1} L ⁻
439	¹ . <u>This range agrees within an order of magnitude with that of (Prodi et al., (1983) who measured</u>
440	<u><i>n</i>_{INP}</u> in the Mediterranean, Red Sea, Gulf of Aden and Indian Ocean nearly 4 decades prior to the
441	present study (4 × 10^{-2} to 2 L ⁻¹ at -16 °C). Average ambient dust concentrations during each
442	sampling period ranged from 2-490 μg m $^{-3}$ (PM $_{10}$ Table 1). There is no agreed-upon standard for
443	definition of extreme dust events in the literature, though the 24-hr average WHO or US EPA
444	health standards for average PM_{10} are commonly used (Gandham et al., 2020; Khaniabadi et al.,
445	2017). Using the US EPA health standard for PM_{10} as a threshold for extreme events (150 $\mu\text{g}~\text{m}^-$
446	³), 14 of the 26 samples were collected during dust events. This is conservative given the equivalent
447	WHO guideline for PM_{10} is 50 μg m $^{\text{-3}}$ (WHO, 2005), in which case 22 of the 26 sampling periods
448	would be classified as dust events. Prior studies have reported comparable PM_{10} levels during dust
449	events in the region (Gandham et al., 2020; Krasnov et al., 2016; Shahsavani et al., 2012).
450	Figs \$0.510 show the extent of k means clustered ELEYPAPT back trajectories below the
450	altitude of 1500 m (see Sect. 55 for details). This threshold was applied to aliminate most of the
451	free tropospheric parts of the back-trajectories and was selected based on the MERRA-2 monthly
452	average planetary boundary laver (PBL) heights during the campaign period, which were 200,700
455	m over the ocean and up to 1700 m over land ELEXPAPT 72 hour Aair mass back trajectories
455	show that source regions included the Mediterranean Nile Delta Sinai Peninsula (f006-f008
455	f038) Northeast Egypt (f009-f010) Iran (f023-f024) and Southern and Eastern Europe (f040
450	f042_f044) (S9-S16) Mmany of samples collected during extreme dust events (f013_f014_f016_
458	f018 and f020) were influenced by emissions from North Africa and the Arabian Peninsula (Figs
459	S98-S108 S12-14) Other source regions included the Mediterranean Nile Delta Singi Peninsula
460	(f006, f008) Northeast Equat (f009, f010) Iran (f024) and Southern and Eastern Europe (f040.
461	$\frac{1000}{1000}$, Normeast Egypt (1009 1010), main (1024), and Southern and Eastern Europe (1040, $\frac{1042}{1000}$).
462	μ^2 cm ⁻³ with the exception of f024 for which aerosol surface area range was > 600 μ^2 cm ⁻³
463	(Table 1).
100	

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464 465



468Figure 1. Map of the sample locations for 26 aerosol samples collected on the RV Kommandor469Iona during Air Quality and climate change in the Arabian BAsin (AQABA; see also Figs. S1-S2470and S9-S10). Measured n_{INP} INP concentrations spanned three-2 orders of magnitude at -15 °C,471from 5×10^{-3} to $5 \times 10^{-1}0.5$ L⁻¹. Marker sizes indicate abundance of INPs. Marker colors indicate472the average ambient dust mass concentration during the sampling period from hourly MERRA-2473reanalysis data.

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Table 1. Summary of aerosol samples collected during AQABA. "—" indicates where data are
missing; "NaN" indicates values below detection limit. Locations are given at the transect
midpoint during each sampling period.

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Sl-	Start 1-4-4	Store datafina	T - 44 J-	T		Sample	Average Aerosol	Aerosol Surface Area	Average Dust	Average Seasalt
ID Sample	(UTC)	(UTC)	(° N)	(° E)	$n_{\rm INP}$ 15 °C (L ⁻¹)	Volume (Loir)	Area	[min,	tion	Formatted
						(L all)	(PM ₁₀ , µm ² cm ⁻³)	(PM ₁₀ , μm ² cm ⁻³)	(PM ₁₀ , μg m ⁻³)	(PM ₁₀ , µg m ⁻³)
f006	05-Jul 05:46	05-Jul 11:37	26.224	35.025	0.0146	3370	290	[199, 375]	170	-
f007	05-Jul 16:40	05-Jul 19:51	26.291	34.933	0.0475	2588	260	[222, 289]	70	-
f008	06-Jul 07:09	06-Jul 14:08	25.225	35.775	0.1161	5225	177	[106, 259]	100	-
f009	07-Jul 05:50	07-Jul 15:07	25.011	35.947	0.0838	6940	352	[253,416]	110	-
f010	08-Jul 16:33	09-Jul 05:59	23.623	36.931	0.0592	8073	219	[163, 287]	50	-
f013	14-Jul 12:26	14-Jul 16:13	18.687	39.672	0.0585	2283	264	[176, 352]	490	10
f014	15-Jul 05:10	15-Jul 11:49	16.552	40.834	0.0348	4000	271	[204, 343]	300	5
f016	18-Jul 07:04	18-Jul 14:52	11.939	45.334	0.0534	4690	265	[158, 391]	430	-
f018	22-Jul 10:20	22-Jul 18:44	20.941	59.474	0.0166	5025	212	[171, 238]	340	-
f019	23-Jul 04:48	23-Jul 13:34	21.410	59.691	0.0145	5270	218	[190, 240]	240	-
f020	25-Jul 17:15	26-Jul 04:02	23.976	58.809	0.0184	6511	-	-	390	5
f023	04-Aug 04:05	04-Aug 11:56	28.084	50.284	0.0112	4720	835	[756, 965]	150	4
f024	05-Aug 05:57	05-Aug 13:53	25.432	53.853	0.0371	5221	357	[206, 827]	90	-
f025	07-Aug 09:26	07-Aug 16:46	23.814	59.186	0.0129	4410	55	[46, 72]	220	12
f030	13-Aug 07:08	14-Aug 11:06	15.970	54.705	0.0132	15111	28	[16, 144]	160	-
f031	14-Aug 15:03	15-Aug 09:03	14.003	52.357	0.0121	12972	25	[19, 105]	230	-
f032	15-Aug 09:42	15-Aug 15:07	13.354	49.432	0.0059	3260	96	[82, 147]	80	6
f033	16-Aug 09:30	16-Aug 13:17	12.208	45.706	NaN	2280	73	[51, 135]	130	2

I	f034	16-Aug 13:27	17-Aug 07:04	12.177	45.429	0.0206	8464	168	[51, 372]	150	1
	f035	17-Aug 07:30	17-Aug 14:55	13.308	42.974	0.0365	4460	340	[244, 409]	210	2
	f036	18-Aug 06:36	18-Aug 15:03	16.290	41.038	0.0057	6634	208	[160, 428]	280	2
	f037	19-Aug 07:05	20-Aug 07:04	18.699	39.609	0.0326	18806	240	[175, 331]	190	7
	f038	21-Aug 07:22	21-Aug 16:01	24.112	36.554	0.0422	6700	256	[202, 295]	150	-
	f040	26-Aug 16:02	27-Aug 07:04	33.803	24.814	0.0314	9030	90	[58, 142]	< 10	3
	f042	28-Aug 07:51	28-Aug 16:02	35.310	17.965	0.0279	6396	163	[131, 222]	< 10	2
	f044	31-Aug 08:30	31-Aug 20:16	39.569	13.380	0.4572	11296	211	[148, 255]	< 10	-

Sample ID	Start datetime (UTC)	Stop datetime (UTC)	Latitude	Longitud e		Sample Volume (L.air)	Aerosol Surface Area (µm² em³)	Average Dust Concentra tion (µg m ⁻ ³)	Average Seasalt Concentrat ion (µg m ⁻³)	
					0.0146				Formatted:	Font: (Default) Times New Roman, 8 pt
f006unt	05 Jul 05:46	05-Jul 11:37	26.224	35.025	0.0475	3370	290	170	Formatted:	Font: (Default) Times New Roman, 8 pt
f007unt	05-Jul-16:40	05-Jul 19:51	26.291	34.933	0.1161	2588	260	70	Formatted:	Font: (Default) Times New Roman, 8 pt
f008unt	06 Jul 07:09	06-Jul 14:08	25.225	35.775	0.0838	5225	180	100	Formatted:	Font: (Default) Times New Roman, 8 pt
f009unt	07-Jul 05:50	07-Jul 15:07	25.011	35.947	0.0592	6940	350	110	Formatted:	Font: (Default) Times New Roman, 8 pt
f010unt	08-Jul 16:33	09-Jul 05:59	23.623	36.931	0.0585	8073	220	50	Formatted:	Font: (Default) Times New Roman, 8 pt
f013unt	14-Jul 12:26	14-Jul 16:13	18.687	39.672	0.0348	2283	260	490	Formatted:	Font: (Default) Times New Roman, 8 pt
f014unt	15 Jul 05:10	15 Jul 11:49	16.552	40.834	0.0534	4000	270	300	Formatted:	Font: (Default) Times New Roman, 8 pt
f016unt	18-Jul 07:04	18-Jul 14:52	11.939	4 5.334	0.0166	4690	260	430	Formatted:	Font: (Default) Times New Roman, 8 pt
f018unt	22-Jul 10:20	22-Jul 18:44	20.941	59.474	0.0145	5025	210	340	Formatted:	Font: (Default) Times New Roman, 8 pt
f019unt	23-Jul 04:48	23-Jul 13:34	21.410	59.691	0.0184	5270	220	230	Formatted:	Font: (Default) Times New Roman, 8 pt
f020unt	25 Jul 17:15	26 Jul 04:02	23.976	58.809	0.0112	6511	-	390	Formatted:	Font: (Default) Times New Roman, 8 pt
	04 Aug	04 Aug								
f023unt	04:05	11:56	28.084	50.284	0.0371	4 720	830	150	Formatted:	Font: (Default) Times New Roman, 8 pt
	05-Aug	05-Aug								
f024unt	05:57	13:53	25.432	53.853	0.0129	5221	360	90	Formatted:	Font: (Default) Times New Roman, 8 pt
	07-Aug	07-Aug								
f025unt	09:26	16:46	23.81 4	59.186	0.0132	4410	50	220	Formatted:	Font: (Default) Times New Roman, 8 pt
	13-Aug	14-Aug								
f030unt	07:08	11:06	15.970	54.705	0.0121	15111	30	160	Formatted:	Font: (Default) Times New Roman, 8 pt
	14 Aug	15 Aug								
f031unt	15:03	09:03	14.003	52.357	0.0059	12972	30	220	Formatted:	Font: (Default) Times New Roman, 8 pt

	15-Aug	15-Aug							
f032unt	09:42	15:07	13.35 4	4 9.432	NaN	3260	100	80	Formatted: Font: (Default) Times New Roman, 8 pt
	16 Aug	16 Aug							
f033unt	09:30	13:17	12.208	45.706	0.0206	2280	90	130	Formatted: Font: (Default) Times New Roman, 8 pt
	16-Aug	17-Aug							
f034unt	13:27	07:04	12.177	4 5.429	0.0365	8464	170	150	Formatted: Font: (Default) Times New Roman, 8 pt
	17-Aug	17-Aug							
f035unt	07:30	14:55	13.308	4 2.974	0.0057	4460	340	210	Formatted: Font: (Default) Times New Roman, 8 pt
	18-Aug	18-Aug							
f036unt	06:36	15:03	16.290	4 1.038	0.0326	6634	210	280	Formatted: Font: (Default) Times New Roman, 8 pt
	19 Aug	20 Aug							
f037unt	07:05	07:04	18.699	39.609	0.0422	18806	240	190	Formatted: Font: (Default) Times New Roman, 8 pt
	21-Aug	21-Aug							
f038unt	07:22	16:01	24.112	36.554	0.0314	6700	260	140	Formatted: Font: (Default) Times New Roman, 8 pt
	26-Aug	27-Aug							
f040unt	16:02	07:04	33.803	24.814	0.0279	9030	90	<10	Formatted: Font: (Default) Times New Roman, 8 pt
	28-Aug	28-Aug							
f042unt	07:51	16:02	35.310	17.965	<u>0.4572</u>	6396	160	10	Formatted: Font: (Default) Times New Roman, 8 pt
	31-Aug	31 Aug			n				
f044unt	08:30	20:16	39.569	13.380	$\frac{(L^{-1})}{(L^{-1})}$	11296	210	40	Formatted: Font: (Default) Times New Roman, 8 pt

478 <u>3.2 Seawater Source Potential</u>

479 The n_{INP} values in 10 SSW samples collected during AQABA were used to characterize the INP 480 source potential of SSA generated by bubble bursting (Wang et al., 2017). Results from prior 481 studies have demonstrated that jet droplets are a more efficient transfer vehicle than film drops of INPs into SSA particles (Mitts et al., 2021; Wang et al., 2017). We measured the n_{INP} in SSW to 482 test whether the seawater source strength was comparable to that of prior studies, or whether the 483 SSW was possibly enriched with INPs due to biological activity or even dust deposition (Cornwell 484 et al., 2020). 485 486 Figure S17 shows how the $n_{\rm INP}$ measured at -19 °C in 10 seawater samples varied by the sample

collection location. Concentrations ranged between 1 and 50 n_{INP} mL⁻¹ and were highest between the Gulf of Oman and the Gulf of Aden. This region exhibited relatively high chlorophyll *a* during the cruise, with levels between 1 and 30 mg m⁻³ (Fig. S18). In Fig. 2, n_{INP} were compared with SSW from the Ellen Browning Scripps Memorial Pier in coastal Southern California (SIO Pier), Formatted: Font: Not Italic

l	491	Cabo	Verde	in f	he No	ortheast	Atlantic.	the	Southern	Ocean	(McCluskey	et al.	. 2017).	and t	the
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- 492 Northwest Atlantic (Schnell, 1977). AQABA $n_{\rm INP}$ were most comparable with Gong et al.'s
- 493 (2020) observations in Cabo Verde. The lack of any unusually high INP spectra suggests that INP
- 494 enrichment due to dust deposition (Cornwell et al. 2020) was absent or infrequent. It is possible
- 495 that storage of SSW samples (Sect. 2.5) could have decreased measured n_{INP} , though we expect







499

Figure 2. Measured n_{INP} in 10 SSW samples collected during AQABA. Also shown are the 500 501 composite INP spectrum of 14 coastal SSW samples collected on São Vincente Island, Cabo Verde 502 (Gong et al., 2020), 17 coastal SSW samples collected at the Ellen Browning Scripps Pier (green shading), and 12 SSW samples collected in the Southern Ocean (McCluskey et al., 2018d). 503 Schnell's (1977) SSW measurements are represented as a composite spectrum of 24 samples (blue 504 shaded region) and 5 additional spectra (blue markers) from samples that exhibited higher freezing 505 506 temperatures. All spectra presented are uncorrected for freezing point depression. 507 Offline treatments for testing heat lability, organic composition, and size were applied to 5 of the

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	10 approximation communical (Mintheada Vece, 2.5) Heat and 0.2 une taltering a transference as a gradest that a longe	
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500	TO seawater samples threthous bee, 2.37. freat and 0.2 and filtering treatments suggest that a farge	

509 fraction of the seawater INPs were heat sensitive and larger than 0.2 µm (Fig. S19). These results

510 are indicative of the POC type of marine INP defined in McCluskey et al. (2018a), though this

511 result should be interpreted with caution as storage could potentially have increased sensitivity to

512 filtering treatments.

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513	Understanding of storage impacts on INPs measured in SSW is lacking. However, Beall et al.,
514	(2020) showed that average INP concentration changes for untreated coastal precipitation samples
515	due to frozen storage were within $2 \times $ of n_{INP} measured in fresh samples, with changes at the upper
516	or lower end of the 95% CI exceeding 10× for some freezing temperatures. If SSW samples are
517	similarly sensitive to storage, we would expect INP concentration changes to be within $2 \times $ on
518	average, but up to $> 10 \times$ for any particular untreated sample. Beall et al. (2020) also reported
519	similar changes INPs < 0.45 µm with a greater tendency toward losses, which indicates that storage
520	may have caused increased sensitivity to the filter treatments applied to stored samples.
521	3.3 Ice-active Surface Site Densities in Aerosol

In Fig. ure 32, approximated ice nucleation active surface site densities (n_s) in aerosol samples are 522 compared with multiple population-specific observations and parameterizations for dust and 523 524 marine INPs. The AQABA measurements are also compared with observations from dust-laden 525 air over the Tropical Atlantic (Price et al., 2018). Overall, observations nearly bridge the full 526 regime between the M18 parameterization for marine INPs (hereafter "M18"; McCluskey et al., 527 2018c), and multiple dust INP parameterizations based on laboratory studies of surface dust. At higher temperatures, between -5 and -12 °C, most observations show agreement with the 528 composite spectrum of n_s observed in a range of marine and coastal environments from DeMott et 529 al. (2016) and (Yang et al., (2020) Yang et al. (2019), and/or the Atkinson et al. (2013) A13 K-530 531 feldspar parameterization. Between -10 and -20 °C, several samples agree with the M18 marine INP parameterization within an order of magnitude, whereas two to three n_s spectra approach the 532 U17 and N12 laboratory-derived dust INP parameterizations within an order of magnitude 533 534 (Niemand et al., 2012; Ullrich et al., 2017), depending on temperature. Multiple samples (~8) 535 additionally agreed with Price et al.'s (2018) observations of INPs between 30-3500 m above the 536 dust-ladeny Tropical Atlantic, and most agree with the Gong et al. (2020) surface-level observations, measured at Cabo Verde in the same region as Price et al. (2018) (Cabo Verde). 537

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540 **Figure 32.** Ice-active surface-nucleation-site densities (n_s) as a function of temperature for 25 of \bullet 541 26 aerosol samples collected during AQABA. Gong et al. (2020) and Price et al. (2018) measured 542 INPs in dust-dominant air masses in the tropical east Atlantic, with minor contributions from SSA, 543 while the DeMott et al. (2016) measurements were collected across a range of locations and conditions within the marine boundary layer comprising air masses mostly dominated by relatively 544 pristine marine SSA. INP concentrations measured in sample f033 were below the detection limits 545 546 imposed by field blanks (see Sect. 2.4, Fig. S76). Error bars represent 95% binomial sampling confidence intervals (Agresti and Coull, 1998). Sample f020 is not shown due to missing aerosol 547 surface area data during the sampling period. For the 8 samples on which a dilution was performed 548 (Fig. S8), ns for both the raw undiluted and diluted sample are shown. *DeMott et al. (2016) data 549 550 shown have been updated with additional data from Yang et al. (2020).

551

Considering the frequency of dust events encountered (dust concentration > 150 μ g m³, see Table 552 553 1), and the high probability that dust was the dominant aerosol source during most sampling 554 periods, it is striking that most n_s spectra observed are 1-3 orders of magnitude lower than the values predicted by dust parameterizations. As noted in Gong et al. (2020), some deviations could 555 556 be expected due to the difference between approximated n_s based on total particle surface area in 557 ambient measurements and true n_s based on surface area of a homogeneous aerosol population (see 558 Methods Sect. 2.4). The FLEXPART back trajectories show that air masses for multiple samples 559 originated from densely populated regions such as Southern and Eastern Europe (f040, f042, f044, Fig S9). The back trajectories also show that for samples f006 f008, f010, and f038, air masses 560 were influenced by the populous region around the Nile River Delta. Agricultural soil dusts 561 562 represent a potential constituent of the INPs observed from these regions. A range of n, has been 563 reported in studies of agricultural soil dusts, the lower end of which agrees with the n_{*} observed in 564 the present study between 8 and 25 °C (Steinke et al., 2016; Tobo et al., 2014; O'Sullivan et al., 2014). 565

Given the marine environment in which sampling <u>occurredtook place</u>, a significant amount of sea spray aerosol (SSA) was also detected in many of the sampled airmasses, <u>using sea salt as a proxy</u> (Table 1), and likely present in others for which no composition data were available. Edtbauer et al. (2020) reported the detection of high levels of dimethyl sulfide (DMS, up to 800 ppt) in the Formatted: Superscript

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570 Gulf of Aden associated with a local phytoplankton bloom during AOABA (as evidenced by 571 visible bioluminescence around the ship at night) as well as high levels of dimethyl sulfone 572 (DMSO₂) and other marine biogenic volatile organic carbons (VOCs) from the Somalian upwelling region. As mentioned above, the n_s for most samples between -6 and -18 °C agree with 573 n_s derived from observations across various locations within the marine boundary layer (Fig. 32). 574 However, considering that SSA is associated with 1000 times fewer IN sites per unit surface area 575 576 than dust (i.e. $1000 \times \text{lower } n_s$) (McCluskey et al., 2018c), the characteristically low IN activity of untreated SSW (even in light of the modest changes expected from storage, Sect. 3.2), and the 577 frequency of dust events during AQABA, our findings suggest it is unlikely that the observed INPs 578 579 originated from SSA. and the high relative abundance of dust compared to sea salt concentrations (Table 1), it is unlikely that the observed INPs originated from SSA. In general, detection of marine 580 581 INPs in ambient aerosol is challenging due to their low relative abundance and decreased 582 efficiency compared to dust (DeMott et al., 2016; McCluskey et al., 2018c). Thus, while SSA 583 contributed to the measured aerosol surface area (Table 1), it is unlikely that the INPs observed in 584 this study were marine in origin, or at least that this is indiscernible in the present study or based 585 on present parameterizations of these populations. Heterogeneous aerosol composition in the sampled air masses likely contributed to some of the 586

587 low ns spectra observed due to the contribution of non-INPs to the measured aerosol surface area (see description of n_s approximation in Sect. 2.4). However, the difference between n_s observed 588 during the most extreme dust events, i.e., when the aerosol population was likely approaching 589 homogeneity in composition, and the ns predicted from N12 and U17 was still greater than 2 orders 590 591 of magnitude. Figures 43(a) and (b) show overlap in n_{INP} and n_s observed in samples collected in 592 low dust and high dust conditions, indicating that the INP populations observed during AQABA exhibited similar efficiencies-IN activity despite variation in total aerosol composition and dust 593 loading. No correlation was found between $n_{\rm JNP}$ and aerosol surface area (Fig. S207), PM₁₀ or dust 594 595 concentration. This result is in contrast to Price et al. (2018) who found the variability in $n_{\rm INP}$ to 596 be largely determined by variability in dust loading or aerosol surface area. (Price et al., (2018) 597 reported higher maximum aerosol surface area concentrations of ~1500 μ m cm⁻³ from three samples collected in an exceptionally optically thick layer, compared to the maximum of 965 µm 598 599 cm⁻³ in the present study (Table 1). Yet overall, the aerosol surface area concentrations compare very well with those observed by (Price et al., 2018), indicative of comparable dustiness in the two 600

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601 studies. Excluding the three case mentioned above samples. Price et al. (2018) collected in an 602 exceptionally optically thick dust layer, the average aerosol surface area was 227 ± 68 μ m² cm⁻³ 603 vs. 226 ± 26 μ m² cm⁻³ for the present study. Furthermore, the sample with the highest *n_s* at -15 °C 604 (f04<u>4</u>0) was collected when dust concentrations were lowest (< 102 μ g m⁻³) (Fig. <u>43</u>, Table 1). 605 This is also in direct contrast to Price et al. (2018), who found that the highest *n_s* observed 606 corresponded to the highest dust loading.

607 Gong et al. (2020) also observed n. lower by more than 2 orders of magnitude compared to N12 and U17 despite the large fraction of supermieron INPs (77-83% depending on temperature), and 608 609 that the supermieron particles were mainly mineral dust. The large differences between parameterized n. for dust, and n. observed in both Gong et al. (2020) and the present study between 610 -12 and -25 °C demonstrate that existing na-based parameterizations may not faithfully represent 611 m, at moderate freezing temperatures, despite proximity to major source regions. Whereas DeMott 612 613 et al. (2015a) found that for temperatures < -20 °C, mineral dust particles from Saharan and Asian deserts may be parameterized as a common particle type, our findings suggest that characteristic 614 n, parameterizations for dust from different source regions may be 615 20 °C or needed alternatively, that this temperature regime requires an alternative to an n_{*} based parameterizations. 616 617 Gong et al. (2019a) demonstrated that predicting minp from surface area size distributions alone may not be feasible in environments where the acrosol and/or INP composition are unknown and 618 619 proposed a probability density function PDF-based approach to predicting INPs at a given freezing 620 temperature.







Figure 43. INP concentrations (n_{INP}) (**a**) and ice <u>nucleatione-active surface</u> site densities (n_s) (**b**) as a function of temperature for 26 aerosol samples collected during AQABA. Markers are colored by the average ambient dust concentration for the respective sampling period. <u>Error bars represent</u> <u>95% binomial sampling confidence intervals (Agresti and Coull, 1998).</u> The n_s measured in samples collected during low dust conditions are equal to or greater than (up to 100×) the n_s

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628 measured during dust events between -9 and -18 °C. INP concentrations measured in sample f033 629 were below the detection limits imposed by field blanks (see Sect. 2.4, Fig. S₇₆). Sample f020 is 630 not shown in (b) due to missing aerosol surface area data during the sampling period. For the 8 631 samples on which a dilution was performed (Fig. S8), n_5 for both the raw undiluted and diluted

632 <u>sample are shown.</u>

633 3.4 Characterization of INPs in Aerosol

Offline treatments for testing heat lability and organic composition of INPs were performed on 12 634 635 samples via heat and $\frac{H_2O_2}{Peroxide}$ treatments, respectively (Fig. 54). Prior studies have shown 636 that the IN-active component of various types of mineral dusts are insensitive to heat treatments (Conen et al., 2011; Hara et al., 2016; Hill et al., 2016; O'Sullivan et al., 2014). The IN activity of 637 638 K-feldspar, the dominant ice-nucleatingIN component of mineral dust, was additionally found to be insensitive to digestion with peroxide (O'Sullivan et al., 2014). A small number of studies 639 reported degradation of IN activity with peroxide treatment and/or heat treatment in Arizona Test 640 641 Dust (ATD), that they attributed to organic material (Perkins et al., 2020; Yadav et al., 2019). 642 Thus, we assume here that any degradation of IN activity due to heat and peroxide treatment are 643 duecorrespond to loss of heat-labile (e.g. proteinaceous) and heat-stable organic INPs, respectively. 644

645 Fisher's Exact Test was applied to frozen and unfrozen well fractions for each untreated sample 646 and its corresponding treated sample to test for significant differences (p < 0.05). Sensitivity to 647 peroxide in most samples (i.e., INP degradation) demonstrate the consistent presence of stable organic INPs at temperatures \geq -15 °C. The lack of peroxide sensitivity at temperatures $<\frac{below}{below}$ -648 15 °C indicates dominance by mineral dust INPs at lower temperatures. Heat sensitivity in five 649 650 samples suggests that biological INPs contributed to their warmest freezing INPs. Gong et al. (2020) similarly found heat-sensitivity in INPs at temperatures > -10 °C. Four of the 12 samples 651 exhibited heat sensitivity at relatively moderate temperatures -11 to -18 °C, including the two 652 653 samples collected in the Mediterranean Sea. One sample (f010) exhibited increased $n_{\rm INP}$ INP 654 concentrations in freezing temperatures < below -18 °C after heat and peroxide treatments. That 655 the response to both heat and peroxide were nearly identical (Fig. 54) suggests that compounds 656 may have been released from the surface during heating, uncovering a more IN active surface

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underneath (heating was common to both procedures). The increased n_{INP} post heat and peroxide

treatment is an unexpected result given previous studies on treated soil dust measurements (Conen

659 et al., 2011; Hill et al., 2016; O'Sullivan et al., 2014; Tobo et al., 2014). <u>However, increases in IN</u>

660 <u>activity after heat treatment have been reported previously for airborne Saharan desert dust and</u>

661 <u>aerosol collected during Saharan dust intrusions (Boose et al., 2019; Conen et al., 2022) as well as</u>

662 SSA and precipitation (Martin et al., 2019; McCluskey et al., 2018a) and should be further

663 <u>investigated in future studies.</u>, though an<u>An</u> increase in IN activity after <u>peroxide treatment has</u>

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also been reported in a Himalayan dust sample (Paramonov et al., 2018). -peroxide treatment has

665 also been reported in a Himalayan dust sample (Paramonov et al., 2018).





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Figure 54. INPs in aerosol samples treated with heat and H2O2-peroxide (Methods Sec. 2.4) to test 668 for INP heat-lability and organic composition. Markers of untreated spectra are colored by the 669 670 average dust concentration during the sampling period. Markers of heat-treated and H2O2peroxide-671 treated samples are filled to indicate significant INP concentration difference from untreated 672 samples according to Fisher's Exact Test (p < 0.05). Sensitivity to $\frac{H_2O_2}{peroxide}$ is evident for all samples \geq -15 °C, indicative of stable organic INPs. Heat-lability is also evident at high to 673 moderate temperatures in multiple samples, demonstrating that biological (e.g., proteinaceous) 674 675 INPs also contributed to INPs observed during AQABA.

Given the frequency of dust storms events and generally high concentrations of dust during most
sampling periods, it is surprising that most samples exhibit peroxide sensitivity. Aridisols and
entisols are the dominant soil types in North Africa and the Arabian Peninsula (Nortcliff, 2012).
Both types are associated with the lowest levels of organic carbon, commonly used as a proxy for

total soil organic matter, compared to other soil types (3 and 9 g kg⁻¹, respectively) (Yost and
Hartemink, 2019).

682 <u>3.5 Characterization of INPs in a Soil Dust Sample</u>

683 INP measurements of soil dusts in this region are scarce and have only been reported for a single. 684 surface dust soil sample, sample "SD", collected 50 km north of Cairo (Niemand et al., 2012), FLEXPART back-trajectories indicate this source region for several samples (f006-10, f038), 685 though it should be noted that dust sources cannot be confirmed in this study lacking aerosol and 686 soil dust minerology. For comparison with this study, we measured INPs in untreated, heat-treated, 687 and peroxide-treated subsamples of an archived-aliquot suspension of the N12-SD sample 688 (Methods Sect. 2.4; DeMott et al., 2018) -described in-. Niemand et al. (2012), Sample N12-SD 689 exhibits sensitivity to both heat and peroxide at temperatures > -16 °C, indicating biological 690 composition of INPs at high freezing temperatures. Multiple AQABA samples influenced by 691 692 desert air mass sources show similar sensitivities at higher temperatures: f006, f007, f019, and 693 f020. Several others exhibit only peroxide-sensitivity in this temperature range. Overall, the heat 694 and peroxide sensitivities in the N12-SD sample indicate that desert dusts may contribute 695 biological and/or organic INPs at moderate to high-freezing temperatures, such as those observed in AQABA samples. Gong et al.'s (2020) results showing heat-sensitivity in INPs at temperatures 696 > -10 °C further demonstrate the contribution of biological INPs at high temperatures in dust-697 698 ladeny air masses near North- Africa.

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Figure 65. Measured concentrations of INPs in an aerosolized soil dust sample<u>"N12-SD"</u>, collected 50 km north of Cairo, Egypt_(Niemand et al., 2012), that was treated with heat and peroxide to test for INP heat-lability and organic composition, same as in Fig. 54 above (Methods Sec. 2.4). A 1:20 dilution of the sample is shown (triangles) and markers of heat-treated and H₂O₂peroxide-treated samples are filled to indicate significant INP concentration differences from untreated samples according to Fisher's Exact Test (p < 0.05). Sensitivity to peroxide and heat treatments indicates biological INPs between -6 and -16 °C.

708 <u>4 Discussion</u>

709 Considering the high freezing temperatures observed, evidence of organic composition, and 710 FLEXPART back trajectories showing that aerosol sources included populous regions and at least 711 one agriculturally active region (the Nile River Delta; Figs. S9-S16), it is possible that agricultural 712 soil dusts contributed to some of the relatively higher n_s , $n_{\rm INP}$, and heat and peroxide sensitivity observed during AQABA. A range of ns has been reported in studies of agricultural soil dusts, the 713 714 lower end of which agrees with the n_s observed in the present study between -8 and -25 °C (Fig. 715 3; Steinke et al., 2016; Tobo et al., 2014; O'Sullivan et al., 2014). Samples from air masses 716 influenced by the Nile River Delta or Southern Europe (f007-8, f010, f038, f042, f044) show a 717 higher fraction of heat-sensitive INPs (Fig. 54). Heat-sensitivity is indicative of biological INPs, 718 which have been associated with agricultural soil dusts in prior studies (Hill et al., 2016; O'Sullivan et al., 2014). Hill et al. (2016) and O'Sullivan et al. (2014) showed peroxide sensitivity 719 720 in agricultural soil dusts at temperatures > -18 to -15 °C, respectively, a range which aligns with 721 the peroxide sensitivity exhibited in the present study. A range of n_s has been reported in studies 722 of agricultural soil dusts, the lower end of which agrees with the n_s-observed in the present study between 8 and -25 °C (Steinke et al., 2016; Tobo et al., 2014; O'Sullivan et al., 2014). Agricultural 723 724 soil dusts are relatively rich in organic and biological material (Conen et al., 2011, 2016; Ellerbrock 725 et al., 2005; Kögel-Knabner et al., 2008; O'Sullivan et al., 2014) and contribute up to 20-25% of 726 the global dust load (Ginoux et al., 2012). Furthermore, they are associated with IN activities 727 higher than that of mineral dust (Conen et al., 2011; Fornea et al., 2009; Isono and Ikebe, 1960; 728 O'Sullivan et al., 2014; Steinke et al., 2016; Tobo et al., 2014). High onset temperatures, up to -6 729 °C, are the norm (Conen et al., 2011; Garcia et al., 2012; Hill et al., 2016; O'Sullivan et al., 2014),

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and the high activity of agricultural soil particles has been attributed to internally mixed organic
 matter (O'Sullivan et al., 2014; Tobo et al., 2014).

732 Organic material can condense or adsorb onto aerosols during photochemical and oxidative processes, representing another potential source of organic INPs during AQABA (Dall'Osto et al., 733 2010; Hinz et al., 2005; Krueger et al., 2004). Could aging explain the organics and decreased n_s 734 735 observed? Though dust aerosol was collected within 1 day's transport from source regions throughout this study, we cannot rule out the possibility of aging impacts, lacking single particle 736 chemistry measurements (e.g., Sullivan et al., 2007). In addition to field observations of n_{INP} INP 737 738 concentrations demonstrating that aging increased the IN efficiency of desert dust INPs (see 739 Introduction; Boose et al., 2016; Conen et al. 2015), prior studies of the effects of aging on mineral 740 dust INPs have yielded mixed and sometimes contradictory results, indicating that the impact of 741 aging on IN properties likely depends on multiple factors including the ice nucleation pathway, 742 the type of aging process, surface morphology, and mineralogy (Perkins et al., 2020 and references therein). Multiple studies have investigated the effects of various aging processes on Arizona Test 743 Dust (ATD) as a proxy for diverse natural dust samples. These included exposure to sulfuric acid, 744 745 nitric acid vapor, and solution-phase processes (Cziczo et al., 2009; Eastwood et al., 2009; Knopf and Koop, 2006; Salam et al., 2007; Sullivan et al., 2010b, 2010a). Perkins et al. (2020) 746 747 demonstrated the INP lability in ATD through multiple solution-phase aging processes (e.g., 748 incubation in water, exposure to acid or salt), with up to $1000 \times \frac{1}{1000}$ reductions in INP abundance at freezing temperatures > 10 °C. This result contrasts with the increase in IN activity attributed to 749 aging reported in Boose et al. (2016) and Conen et al. (2015). Perkins et al. (2020) additionally 750 751 reported that the lability of IN activity in ATD is temperature dependent, with large reductions 752 evident at freezing temperature > 10 °C, yet little to no change at temperatures < below -15 °C. By 753 contrast, most of the n_s spectra in AQABA samples were $10 - 1000 \times$ lower than established dust 754 parameterizations even at temperatures < -below -15 °C. In summary, it has proven difficult to 755 determine any consistent impact of atmospheric processing on the IN activity of dust in model 756 systems such as ATD (Perkins et al., 2020), and few studies have investigated impacts of aging on 757 ambient desert dust, especially at modest supercooling (Boose et al., 2016). Furthermore, the use of ATD as a proxy for natural dust in INP studies has been questioned due to the complex ice-758 nucleatingIN-properties of natural dust, including mineral composition and defect sites at the 759

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particle surface, the latter of which is likely affected by the mechanical processing and millinginvolved in ATD production (e.g., Perkins et al., 2020 and references therein).

762 Gong et al. (2020) also observed n. lower by more than 2 orders of magnitude compared to N12 and U17 despite the large fraction of supermicron INPs (77-83% depending on temperature), and 763 that the supermicron particles were mainly mineral dust. The large differences between 764 765 parameterized n_x for dust, and n_x observed in both Gong et al. (2020) and the present study between -12 and -25 °C demonstrate that existing n_g based parameterizations may not faithfully represent 766 na at moderate freezing temperatures, despite proximity to major source regions. Whereas DeMott 767 et al. (2015a) found that for temperatures < 20 °C, mineral dust particles from Saharan and Asian 768 769 deserts may be parameterized as a common particle type, our findings suggest that characteristic 770 $n_{\rm r}$ parameterizations for dust from different source regions may be needed > 20 °C, or, 771 alternatively, that this temperature regime requires an alternative to an n,-based parameterizations. 772 Gong et al. (2019a) demonstrated that predicting n_{INP} from surface area size distributions alone may not be feasible in environments where the aerosol and/or INP composition are unknown and 773 774 proposed a probability density function PDF-based approach to predicting INPs at a given freezing 775 temperature. 776 Gong et al. (2020) also observed n_s lower by more than 2 orders of magnitude compared to N12 777 and U17 despite the large fraction of supermicron INPs (77-83% depending on temperature), and 778 that the supermicron particles were mainly mineral dust. The cause of the decreased n_s observed 779 here and in Gong et al. (2020) compared to dust n_s parameterizations remains elusive. Both studies 780 were conducted in air masses dominated by dust near major sources. In contrast, Price et al. (2018) 781 found agreement near the region of the Gong et al. (2020) study._One obvious difference is that Price et al. (2018) conducted measurements at higher altitudes, between 30 and 3500 m. A prior 782 783 study that compared $n_{\rm INP}$ in dust-ladeny air masses at the surface with $n_{\rm INP}$ collected between 0.5 784 and 3 km above sea level found that median $n_{\rm INP}$ increased by up to 10× above the surface and

785 correlated to dust loading (Schrod et al., 2017). -<u>The differences between Price et al. (2018) and</u>

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786 the two surface-based studies draws attention to the need for vertical profiles of $n_s > -25$ °C in **787** dust-laden air masses.

788 The decreased n_s compared to Price et al. (2018) is also unlikely to be related to differences in INP measurement. In all three studies, cold stage or droplet assay measurements of immersion mode 789 INPs were used in resuspensions of aerosol collected on filter samples. Recent studies that 790 791 intercompared instruments designed for measurement of immersion mode INPs showed excellent agreement (i.e., within measurement uncertainty) in measurements of standardized dust and 792 biological samples (DeMott et al., 2018) and when co-sampling ambient aerosol (DeMott et al., 793 794 2017). Moreover, the DeMott et al. (2018) intercomparison study demonstrated good agreement 795 in multiple natural dust samples between the various measurement methods used to derive D15, 796 N12 and U17 and the droplet assay methods applied in Gong et al. (2020), Price et al. (2018), 797 and the present study.

798Storage protocol represents another difference between (Price et al., (2018) and the two surface-799based studies. (Gong et al., (2020) and the present study stored samples frozen prior to analysis,800whereas (Price et al., (2018) processed samples immediately after collection. An understanding of801storage impacts on INPs collected on filters is lacking (Wex et al., 2019), but we note that the802discrepancies in n_s between the two surface-based studies and (Price et al., (2018) exceed the range803of INP concentration changes reported in untreated INP precipitation samples stored frozen (Beall804et al., 2020).

805 Thus, the large differences between parameterized n_s for dust, and n_s observed in both Gong et al. (2020) and the present study between -12 and -25 °C indicate that existing n_s -based 806 807 parameterizations may not faithfully represent n_s at moderate freezing temperatures, despite 808 proximity to major source regions. Whereas DeMott et al. (2015) found that for temperatures < -20 809 °C, mineral dust particles from Saharan and Asian deserts may be parameterized as a common 810 particle type, our findings suggest that characteristic ns parameterizations for dust from different 811 source regions may be needed > -20 $^{\circ}$ C, or, alternatively, that this temperature regime requires an alternative to an n_s -based parameterizations. Gong et al. (2019a) demonstrated that predicting $n_{\rm INP}$ 812 813 from surface area size distributions alone may not be feasible in environments where the aerosol and/or INP composition are unknown and proposed a probability density function (PDF)-based 814

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815 <u>approach to predicting INPs at a given freezing temperature.</u>

In light of the evidence from this study that INPs were primarily influenced by organics associated 816 817 with dust, especially at higher temperatures, and the lack of relationship between dust loading, n_s, and n_{INP}, we offer the following points for consideration.__Prior studies of aerosolized dust 818 819 demonstrated that it is frequently enriched in organic matter (6-20×) compared to soil dust \neq and 820 that wind erosion selectively removes the chemically-enriched, fine portion of the soil higher -of 821 in plant nutrients, organic matter and metals_(Aryal et al., 2012; Delany and Zenchelsky, 1976; 822 Van Pelt and Zobeck, 2007). Furthermore, a recent study that measured airborne concentrations of 823 prokaryotic cells over the Red Sea characterized the region as a "global hot spot" with average concentrations of 155,000 (± 65,000) cells m⁻³, 19× higher than that over the subtropical and 824 tropical open oceans (Mayol et al., 2014; Yahya et al., 2019). Yahya et al. (2019) demonstrated 825 that the microbial loading was very likely related to the high concentrations of dust, as 99.9% of 826 the cells were attached to dust particles. Organic and biological species have been shown to 827 828 dominate IN activity at temperatures > ~-15 °C in many studies (e.g., Kanji et al., 2017; Ladino et al., 2019; O'Sullivan et al., 2018, Kanji et al., 2017, and references therein). Thus, a faithful 829 830 representation of dust INPs may require two parameterizations: one for the IN activity dominated 831 by minerals < ~-15 °C such as D15, U17 and N12, and another for the dust-associated organics > ~-15 °C. As IN-active organics are limited compared to the IN-active mineral component of dust, 832 we could expect an increase in n_s slope between warm and cold regimes. The apparent decreased 833 834 n_s observed in this study between -18 and -12 °C could potentially be related to a plateau in n_s 835 through the transition between the mineral and organic "modes" (see untreated samples in Figs. 5-836 Fig. 65). This study underscores the need to characterize the IN-active organic species associated with dust from major source regions and to investigate the extent to which biological and/or 837 organic particles contribute to INP populations in dust-laden air masses at high to moderate 838 839 freezing temperatures $\geq -15 \,^{\circ}C$.

840 3.2 Seawater Source Potential

- 841 The *n_{INP}* values in 10 SSW samples collected during AQABA were used to characterize the INP+
- 842 source potential of SSA generated by bubble bursting (Wang et al., 2017). Results from prior
- 843 studies have demonstrated that jet droplets are a more efficient transfer vehicle than film drops of

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844	INPs into SSA particles (Mitts et al., 2021; Wang et al., 2017). While it is unlikely that many of
845	the INPs detected in aerosol samples were marine in origin (see Sec. 3.1), we measured the INP
846	concentrations in SSW to test whether the seawater source strength was comparable to that of prior
847	studies, or were possibly enriched with INPs due to biological activity or even dust deposition
848	(Cornwell et al., 2020).

- Figure S10 shows how the INP concentrations measured at -19 °C in 10 seawater samples varied
 by the sample collection location. Concentrations ranged between 1 and 50 INPs mL⁻¹ and were
 highest between the Gulf of Oman and the Gulf of Aden. This region exhibited relatively high
 chlorophyll *a* during the cruise, with levels between 1 and 30 mg m⁻³ (Fig. S11). In Fig. 6, INP
 concentrations were compared with SSW from the Ellen Browning Scripps Memorial Pier in
- 854 coastal Southern California (SIO Pier), Cabo Verde in the Northeast Atlantic, the Southern
- 855 Ocean (McCluskey et al., 2017), and the Northwest Atlantic (Schnell, 1977). AQABA INP
- 856 concentrations were most comparable with Gong et al.'s (2020) observations in Cabo Verde. The



857 lack of any unusually high INP spectra suggests that INP enrichment due to dust deposition

858 (Cornwell et al. 2020) was absent or infrequent.

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872Heat resilience but peroxide sensitivity in sample s001 additionally indicates the presence of873non-proteinaceous organic INPs, such as the dissolved organic carbon (DOC) type defined in874McCluskey et al. (2018a). Considering the characteristically low IN activity of the SSW, the875lower n_s -of SSA compared to mineral dust (McCluskey et al., 2018b), and the frequency of dust876events during AQABA, our findings suggest that dust was highly likely to be the dominant INP877class observed in this study.

878

879 <u>54</u> Conclusions

Observations from the two-month AQABA campaign in the Mediterranean, Red Sea, Arabian Sea and Arabian Gulf are among the first INP measurements made in close proximity to the two largest dust sources globally: the Sahara and the Arabian Peninsula (Kok et al., 2021). <u>Observed *p*INP INP</u> concentrations-measured in 26 aerosol samples spanned two-2 or more orders of magnitude (5×10^{-3} to 5×10^{-1} L⁻¹0.002 to 0.5 L⁻⁴-at -15 °C).

885 In summary, INPs observed during AQABA were very likely dominated by mineral dust with 886 some additional contributions possibly from densely-populated and/or agricultural regions 887 including the Nile River Delta region and Southern or Eastern Europe. Despite proximity to major 888 dust sources and a high frequency of dust events with MERRA-2 simulated mass concentrations up to 490 μ g m⁻³ (PM₁₀), the observed n_s for most samples was lower by 1-3 orders of magnitude 889 890 compared to n_s predicted by dust parametrizations N12 and U17 at T < -12 °C-(Niemand et al., 891 2012; Ullrich et al., 2017). Observed n_s in 48% of the samples were equivalent within uncertainty to that of Many INPs measured in AQABA showed agreement with the A13 parameterization for 892 893 K-feldspar (Atkinson et al., 2013), an ice-active component of desert dust. Observed ns agreed -894 with with many observations within the marine boundary layer (DeMott et al., 2016; Yang et al., 895 2020), and with the Price et al.'s (2018) measurements of $n_{\rm INP}$ INP concentrations in dust-laden air masses over the Tropical Atlantic, within measurement uncertainty. Peroxide sensitivity was 896 evident in all samples tested (12 of 26), at temperatures \geq -15 °C, demonstrating a consistent 897 contribution of organic material to warm-temperature INPs. Heat-sensitivity further suggested the 898 presence of biological (e.g., proteinaceous) INPs in a subset of samples, particularly at high 899 freezing temperatures <u>>-10 °C</u>. While the dominant mineral dusts in the region are associated with 900

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901	the lowest concentrations of soil organic carbon globally (e.g., Yost and Hartemink, 2019 and
902	references therein), aerosolized fine dust is known to be enriched in organic matter (Aryal et al.,
903	2012; Delany and Zenchelsky, 1976; Van Pelt and Zobeck, 2007) and is additionally associated
904	with high microbial loading in the Red Sea (Yahya et al., 2019). A soil dust sample from North
905	Africa (originally from N12) exhibited heat and peroxide sensitivity between -5 and -16 °C, further
906	demonstrating that the IN activity of mineral dust could be associated with organic and/or
907	biological material. Contrary to Price et al. (2018), who measured INP in the dust-laden Tropical
908	Atlantic, no correlation was found between dust loading and nINP or ns. Results from this study and
909	Gong et al. (2020) indicate that the existing n_s parameterizations alone do not skillfully represent
910	mineral dust associated INPs at modest supercooling near major dust sources.
911	The source strengths of Red Sea, Mediterranean, Arabian Sea, and Arabian Gulf bulk seawater
912	were also evaluated. The maximum source potential was observed in the Arabian Sea (50 INP
913	mL ⁻¹ at -19 °C).) Overall, the The observed n_{INP} range for SSW samples were equivalent to agreed
914	well with the those of Gong et al. (2020) SSW measurements at Cabo Verde within the 95%
915	binomial sampling confidence intervals (Agresti and Coull, 1998).
916	Considering that desert dust parameterizations overpredicted the <i>n_s</i> values observed during
917	AQABA, despite proximity to major global emissions sources, this study demonstrates the need
918	to evaluate the fidelity of dust INP parameterizations in nascent versus aged dust populations.
919	The discrepancies underscore the challenges of evaluating dust-specific INP parameterizations:
920	limited observations at modest supercooling, few assured methods for distinguishing between
921	different INP sources in ambient aerosol, a dearth of characteristic soil dust samples from major
922	dust sources, and limited knowledge of the specific composition and characteristics of dust INPs
923	at temperatures > -15 °C. Vertical profiles of n_s in dust-laden air masses are also needed to
924	determine whether n_s is consistently lower at the surface and examine the variability of n_s with
925	altitude, Potential storage impacts on INPs collected on filters are an additional factor worthy of
926	future investigation, though storage alone does not likely explain the relatively decreased n_s
927	compared to parameterizations observed in this study, as U17 and N12 were both derived from
928	stored dust samples.

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930 In addition to providing observations at high to moderate freezing temperatures, future studies could apply the methods developed in Gong et al. (2020) to estimate the contribution of marine 931 932 INPs to the aerosol sampled by assuming equivalent distributions of sea salt and INPs between seawater and air. Furthermore, given the combination of marine, dust, and anthropogenically-933 934 influenced air masses encountered, and the evidence of organic and biological INPs at modest supercooling in this study and Gong et al. (2020), future studies could benefit from advances in 935 936 on-line Light-Induced Fluorescence (LIF) measurement techniques. Whereas the interpretation of fluorescence data from most LIF-based instruments has been limited by the lack of spectroscopic 937 information, newer instruments support real-time spectrally-resolved size and fluorescence 938 measurement information for single particles (Fennelly et al., 2018; Huffman et al., 2020; 939 940 Könemann et al., 2019). This information could be used to potentially "tag" different classes of 941 organics and biological aerosols, enabling investigations of relationships between n_s , $n_{\rm INP}$ and 942 organic signatures in, e.g., mineral dusts and agricultural soil dusts. Finally, the decreased n_s 943 observed in this study further motivate comprehensive aerosol-ice nucleation studies, which aim to achieve closure between measured and predicted ambient $n_{\rm INP}$ <u>tNP concentrations</u> by 944 simultaneously characterizing ambient INPs and ice nucleation relevant properties of the total 945 946 aerosol population, such as composition and aerosol chemical mixing state (Sullivan et al., 2007).

947 Data Availability: The data set supporting this manuscript is hosted by the UCSD Library
948 Digital Collections (<u>https://doi.org/10.6075/J0X0676P</u>) (Beall et al., 2021).

949 Author contributions:

- CMB, TCH, PJD, MOA, CP, JL, JS, FD, BW, HH, MDS, and KAP designed the study. CMB
- 951 performed the INP measurements, FLEXPART modeling and analysis with support from TCH,
- 952 PJD, MOA, MDS, MP and KAP. TCH, PJD and MOA contributed significantly to the writing,
- 953 preparation of figures and analysis. TK, MI, RP and HH supported the field collection of aerosol
- 954 for INP analysis and TK additionally provided aerosol number concentration data. JS and MP

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956	composition measurements and analysis. All authors contributed to the writing of the article.	
957	Competing interests:	
958	The authors declare they have no conflict of interest.	
959		
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975	References	
976	Agresti, A. and Coull, B. A.: Approximate Is Better than "Exact" for Interval Estimation of Binomial	
977	Proportions, Am. Stat., 52(2), 119, doi:10.2307/2685469, 1998.	
978	Ardon-Dryer, K. and Levin, Z.: Ground-based measurements of immersion freezing in the eastern	

provided aerosol water-soluble composition data. FD oversaw the aerosol sizing and AMS

955

- 979 Mediterranean, Atmos. Chem. Phys., 14(10), 5217–5231, doi:10.5194/acp-14-5217-2014, 2014.
- 980 Aryal, R., Kandel, D., Acharya, D., Chong, M. N. and Beecham, S.: Unusual Sydney dust storm and its
- 981 mineralogical and organic characteristics, Environ. Chem., 9(6), 537–546 [online] Available from:

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982 https://doi.org/10.1071/EN12131, 2012.

- 983 Atkinson, J. D., Murray, B. J., Woodhouse, M. T., Whale, T. F., Baustian, K. J., Carslaw, K. S., Dobbie,
- 984 S., O'Sullivan, D. and Malkin, T. L.: The importance of feldspar for ice nucleation by mineral dust in
- 985 mixed-phase clouds, Nature, 498(7454), 355–358, doi:10.1038/nature12278, 2013.
- 986 Beall, C. M., Stokes, M. D., Hill, T. C., DeMott, P. J., DeWald, J. T. and Prather, K. A.: Automation and
- 987 Heat Transfer Characterization of Immersion Mode Spectroscopy for Analysis of Ice Nucleating
- 988 Particles, Atmos. Meas. Tech., (February), 1–25, doi:10.5194/amt-2016-412, 2017.
- 989 Beall, C. M., Lucero, D., Hill, T. C., DeMott, P. J., Stokes, M. D. and Prather, K. A.: Best practices for
- 990 precipitation sample storage for offline studies of ice nucleation in marine and coastal environments,
- 991 Atmos. Meas. Tech., 13(12), 6473–6486, doi:10.5194/amt-13-6473-2020, 2020.
- 992 Beall, C. M., Michaud, J. M., Fish, M. A., Dinasquet, J., Cornwell, G. C., Stokes, M. D., Burkart, M. D.,
- 993 Hill, T. C., Demott, P. J. and Prather, K. A.: Cultivable halotolerant ice-nucleating bacteria and fungi in
- 994 coastal precipitation, Atmos. Chem. Phys., 21(11), 9031–9045, doi:10.5194/acp-21-9031-2021, 2021.
- 995 Boose, Y., Sierau, B., Isabel García, M., Rodríguez, S., Alastuey, A., Linke, C., Schnaiter, M.,
- 996 Kupiszewski, P., Kanji, Z. A. and Lohmann, U.: Ice nucleating particles in the Saharan Air Layer, Atmos.
- 997 Chem. Phys., 16(14), 9067–9087, doi:10.5194/acp-16-9067-2016, 2016.
- 998 Boose, Y., Baloh, P., Plötze, M., Ofner, J., Grothe, H., Sierau, B., Lohmann, U. and Kanji, Z. A.:
- 999 Heterogeneous ice nucleation on dust particles sourced from nine deserts worldwide -- Part 2: Deposition
 1000 nucleation and condensation freezing, Atmos. Chem. Phys., 19(2), 1059–1076, doi:10.5194/acp-19-10591001 2019, 2019.
- 1002 Bourtsoukidis, E., Ernle, L., Crowley, J. N., Lelieveld, J., Paris, J.-D., Pozzer, A., Walter, D. and
- 1003 Williams, J.: Non-methane hydrocarbon ($chem{C_2}-chem{C_8}$) sources and sinks around the
- 1004 Arabian Peninsula, Atmos. Chem. Phys., 19(10), 7209–7232, doi:10.5194/acp-19-7209-2019, 2019.
- 1005 Bourtsoukidis, E., Pozzer, A., Sattler, T., Matthaios, V. N., Ernle, L., Edtbauer, A., Fischer, H.,
- 1006 Könemann, T., Osipov, S., Paris, J.-D., Pfannerstill, E. Y., Stönner, C., Tadic, I., Walter, D., Wang, N.,
- 1007 Lelieveld, J. and Williams, J.: The Red Sea Deep Water is a potent source of atmospheric ethane and
- 1008 propane, Nat. Commun., 11(1), 447, doi:10.1038/s41467-020-14375-0, 2020.
- 1009 Broadley, S. L., Murray, B. J., Herbert, R. J., Atkinson, J. D., Dobbie, S., Malkin, T. L., Condliffe, E. and
- 1010 Neve, L.: Immersion mode heterogeneous ice nucleation by an illite rich powder representative of
- 1011 atmospheric mineral dust, Atmos. Chem. Phys., 12(1), 287–307, doi:10.5194/acp-12-287-2012, 2012.

- 1012 Brunner, C., Brem, B. T., Collaud Coen, M., Conen, F., Hervo, M., Henne, S., Steinbacher, M., Gysel-
- 1013 Beer, M. and Kanji, Z. A.: The contribution of Saharan dust to the ice-nucleating particle concentrations
- 1014 at the High Altitude Station Jungfraujoch (3580\,m\,a.s.l.), Switzerland, Atmos. Chem. Phys., 21(23),
- 1015 18029–18053, doi:10.5194/acp-21-18029-2021, 2021.
- 1016 Buchard, V., Randles, C. A., da Silva, A. M., Darmenov, A., Colarco, P. R., Govindaraju, R., Ferrare, R.,
- 1017 Hair, J., Beyersdorf, A. J., Ziemba, L. D. and Yu, H.: The MERRA-2 Aerosol Reanalysis, 1980 Onward.
- 1018 Part II: Evaluation and Case Studies, J. Clim., 30(17), 6851–6872, doi:10.1175/JCLI-D-16-0613.1, 2017.
- Burrows, S. M., Hoose, C., Pöschl, U. and Lawrence, M. G.: Ice nuclei in marine air: biogenic particles or
 dust?, Atmos. Chem. Phys., 13(1), 245–267, doi:10.5194/acp-13-245-2013, 2013.
- 1021 Celik, S., Drewnick, F., Fachinger, F., Brooks, J., Darbyshire, E., Coe, H., Paris, J.-D., Eger, P. G.,
- 1022 Schuladen, J., Tadic, I., Friedrich, N., Dienhart, D., Hottmann, B., Fischer, H., Crowley, J. N., Harder, H.
- 1023 and Borrmann, S.: Influence of vessel characteristics and atmospheric processes on the gas and particle
- 1024 phase of ship emission plumes: in situ measurements in the Mediterranean Sea and around the Arabian
- 1025
 Peninsula, Atmos. Chem. Phys., 20(8), 4713–4734, doi:10.5194/acp-20-4713-2020, 2020.
- 1026 Collins, D. B., Zhao, D. F., Ruppel, M. J., Laskina, O., Grandquist, J. R., Modini, R. L., Stokes, M. D.,
- 1027 Russell, L. M., Bertram, T. H., Grassian, V. H., Deane, G. B. and Prather, K. A.: Direct aerosol chemical
- 1028 composition measurements to evaluate the physicochemical differences between controlled sea spray
- aerosol generation schemes, Atmos. Meas. Tech., 7(11), 3667–3683, doi:10.5194/amt-7-3667-2014,
 2014.
- 1031 Conen, F., Morris, C. E., Leifeld, J., Yakutin, M. V and Alewell, C.: Biological residues define the ice
 1032 nucleation properties of soil dust, Atmos. Chem. Phys., 11(18), 9643–9648, doi:10.5194/acp-11-96431033 2011, 2011.
- 1034 Conen, F., Rodríguez, S., Hüglin, C., Henne, S., Herrmann, E., Bukowiecki, N. and Alewell, C.:
- Atmospheric ice nuclei at the high-altitude observatory Jungfraujoch, Switzerland, Tellus, Ser. B Chem.
 Phys. Meteorol., 67(1), 1–10, doi:10.3402/tellusb.v67.25014, 2015.
- 1037 Conen, F., Einbock, A., Mignani, C. and Hüglin, C.: Measurement report: Ice-nucleating particles active
- 1039 3444, doi:10.5194/acp-22-3433-2022, 2022.
- 1040 Cornwell, G. C., McCluskey, C. S., Levin, E. J. T., Suski, K. J., DeMott, P. J., Kreidenweis, S. M. and
- 1041 Prather, K. A.: Direct Online Mass Spectrometry Measurements of Ice Nucleating Particles at a California
- 1042 Coastal Site, J. Geophys. Res. Atmos., 124(22), 12157–12172, doi:doi:10.1029/2019JD030466, 2019.

- 1043 Cornwell, G. C., Sultana, C. M., Prank, M., Cochran, R. E., Hill, T. C. J., Schill, G. P., DeMott, P. J.,
- 1044 Mahowald, N. and Prather, K. A.: Ejection of Dust From the Ocean as a Potential Source of Marine Ice
- 1045 Nucleating Particles, J. Geophys. Res. Atmos., 125(24), e2020JD033073,
- 1046 doi:https://doi.org/10.1029/2020JD033073, 2020.
- 1047 Cziczo, D. J., Froyd, K. D., Gallavardin, S. J., Moehler, O., Benz, S., Saathoff, H. and Murphy, D. M.:
- **1048** Deactivation of ice nuclei due to atmospherically relevant surface coatings, Environ. Res. Lett., 4(4),
- 1049 44013, doi:10.1088/1748-9326/4/4/044013, 2009.
- 1050 Dall'Osto, M., Harrison, R. M., Highwood, E. J., O'Dowd, C., Ceburnis, D., Querol, X. and Achterberg,
- E. P.: Variation of the mixing state of Saharan dust particles with atmospheric transport, Atmos. Environ.,
 44(26), 3135–3146, doi:https://doi.org/10.1016/j.atmosenv.2010.05.030, 2010.
- 1053 Delany, A. C. and Zenchelsky, S.: THE ORGANIC COMPONENT OF WIND-EROSION-
- 1054 GENERATED SOIL-DERIVED AEROSOL, Soil Sci., 121(3) [online] Available from:
- 1055 https://journals.lww.com/soilsci/Fulltext/1976/03000/THE_ORGANIC_COMPONENT_OF_WIND_ER
 1056 OSION_GENERATED.2.aspx, 1976.
- 1057 Demott, P. J., Prenni, A. J., Mcmeeking, G. R., Sullivan, R. C., Petters, M. D., Tobo, Y., Niemand, M.,
- 1058 Möhler, O., Snider, J. R., Wang, Z. and Kreiden: Integrating laboratory and field data to quantify the
- 1059 immersion freezing ice nucleation activity of mineral dust particles, 393–409, doi:10.5194/acp-15-3931060 2015, 2015.
- 1061 DeMott, P. J., Hill, T. C. J., McCluskey, C. S., Prather, K. A., Collins, D. B., Sullivan, R. C., Ruppel, M.
- 1062 J., Mason, R. H., Irish, V. E., Lee, T., Hwang, C. Y., Rhee, T. S., Snider, J. R., McMeeking, G. R.,
- 1063 Dhaniyala, S., Lewis, E. R., Wentzell, J. J. B., Abbatt, J., Lee, C., Sultana, C. M., Ault, A. P., Axson, J.
- 1064 L., Diaz Martinez, M., Venero, I., Santos-Figueroa, G., Stokes, M. D., Deane, G. B., Mayol-Bracero, O.
- 1065 L., Grassian, V. H., Bertram, T. H., Bertram, A. K., Moffett, B. F. and Franc, G. D.: Sea spray aerosol as
- 1066 a unique source of ice nucleating particles, Proc. Natl. Acad. Sci., 113(21), 5797–5803,
- 1067 doi:10.1073/pnas.1514034112, 2016.
- 1068 DeMott, P. J., Hill, T. C. J., Petters, M. D., Bertram, A. K., Tobo, Y., Mason, R. H., Suski, K. J.,
- 1069 Mccluskey, C. S., Levin, E. J. T., Schill, G. P., Boose, Y., Rauker, A. M., Miller, A. J., Zaragoza, J.,
- 1070 Rocci, K., Rothfuss, N. E., Taylor, H. P., Hader, J. D., Chou, C., Huffman, J. A., Pöschl, U., Prenni, A. J.
- 1071 and Kreidenweis, S. M.: Comparative measurements of ambient atmospheric concentrations of ice
- 1072 nucleating particles using multiple immersion freezing methods and a continuous flow diffusion chamber,
- 1073 Atmos. Chem. Phys., 17(18), 11227–11245, doi:10.5194/acp-17-11227-2017, 2017.

- 1074 DeMott, P. J., Möhler, O., Cziczo, D. J., Hiranuma, N., Petters, M. D., Petters, S. S., Belosi, F.,
- 1075 Bingemer, H. G., Brooks, S. D., Budke, C., Burkert-Kohn, M., Collier, K. N., Danielczok, A., Eppers, O.,
- 1076 Felgitsch, L., Garimella, S., Grothe, H., Herenz, P., Hill, T. C. J., Höhler, K., Kanji, Z. A., Kiselev, A.,
- 1077 Koop, T., Kristensen, T. B., Krüger, K., Kulkarni, G., Levin, E. J. T., Murray, B. J., Nicosia, A.,
- 1078 O'Sullivan, D., Peckhaus, A., Polen, M. J., Price, H. C., Reicher, N., Rothenberg, D. A., Rudich, Y.,
- 1079 Santachiara, G., Schiebel, T., Schrod, J., Seifried, T. M., Stratmann, F., Sullivan, R. C., Suski, K. J.,
- 1080 Szakáll, M., Taylor, H. P., Ullrich, R., Vergara-Temprado, J., Wagner, R., Whale, T. F., Weber, D., Welti,
- 1081 A., Wilson, T. W., Wolf, M. J. and Zenker, J.: The Fifth International Workshop on Ice Nucleation phase
- 1082 2 (FIN-02): laboratory intercomparison of ice nucleation measurements, Atmos. Meas. Tech., 11(11),
- 1083 6231-6257, doi:10.5194/amt-11-6231-2018, 2018.
- 1084 Eastwood, M. L., Cremel, S., Wheeler, M., Murray, B. J., Girard, E. and Bertram, A. K.: Effects of
- 1085 sulfuric acid and ammonium sulfate coatings on the ice nucleation properties of kaolinite particles,
- 1086 Geophys. Res. Lett., 36(2), doi:https://doi.org/10.1029/2008GL035997, 2009.
- 1087 Edtbauer, A., Stönner, C., Pfannerstill, E. Y., Berasategui, M., Walter, D., Crowley, J. N., Lelieveld, J.
- and Williams, J.: A new marine biogenic emission: methane sulfonamide (MSAM), dimethyl sulfide
- 1089 (DMS), and dimethyl sulfone (\chem{DMSO_{2}}) measured in air over the Arabian Sea, Atmos. Chem.
- 1090 Phys., 20(10), 6081–6094, doi:10.5194/acp-20-6081-2020, 2020.
- 1091 Eger, P. G., Friedrich, N., Schuladen, J., Shenolikar, J., Fischer, H., Tadic, I., Harder, H., Martinez, M.,
- 1092 Rohloff, R., Tauer, S., Drewnick, F., Fachinger, F., Brooks, J., Darbyshire, E., Sciare, J., Pikridas, M.,
- 1093 Lelieveld, J. and Crowley, J. N.: Shipborne measurements of CINO\$_{2}\$ in the Mediterranean Sea and
- around the Arabian Peninsula during summer, Atmos. Chem. Phys., 19(19), 12121–12140,
- 1095 doi:10.5194/acp-19-12121-2019, 2019.
- 1096 Fennelly, M. J., Sewell, G., Prentice, M. B., O'Connor, D. J. and Sodeau, J. R.: Review: The Use of Real-
- 1097 Time Fluorescence Instrumentation to Monitor Ambient Primary Biological Aerosol Particles (PBAP),
 1098 Atmosphere (Basel)., 9(1), doi:10.3390/atmos9010001, 2018.
- 1099 Friedrich, N., Eger, P., Shenolikar, J., Sobanski, N., Schuladen, J., Dienhart, D., Hottmann, B., Tadic, I.,
- 1100 Fischer, H., Martinez, M., Rohloff, R., Tauer, S., Harder, H., Pfannerstill, E. Y., Wang, N., Williams, J.,
- 1101 Brooks, J., Drewnick, F., Su, H., Li, G., Cheng, Y., Lelieveld, J. and Crowley, J. N.: Reactive nitrogen
- around the Arabian Peninsula and in the Mediterranean Sea during the 2017 AQABA ship campaign,
- 1103 Atmos. Chem. Phys., 21(10), 7473–7498, doi:10.5194/acp-21-7473-2021, 2021.
- 1104 Gandham, H., Dasari, H. P., Langodan, S., Karumuri, R. K. and Hoteit, I.: Major Changes in Extreme

- 1105 Dust Events Dynamics Over the Arabian Peninsula During 2003–2017 Driven by Atmospheric
- 1106 Conditions, J. Geophys. Res. Atmos., 125(24), e2020JD032931,
- 1107 doi:https://doi.org/10.1029/2020JD032931, 2020.
- 1108 Gelaro, R., McCarty, W., Suárez, M. J., Todling, R., Molod, A., Takacs, L., Randles, C. A., Darmenov,
- 1109 A., Bosilovich, M. G., Reichle, R., Wargan, K., Coy, L., Cullather, R., Draper, C., Akella, S., Buchard,
- 1110 V., Conaty, A., da Silva, A. M., Gu, W., Kim, G.-K., Koster, R., Lucchesi, R., Merkova, D., Nielsen, J.
- 1111 E., Partyka, G., Pawson, S., Putman, W., Rienecker, M., Schubert, S. D., Sienkiewicz, M. and Zhao, B.:
- 1112 The Modern-Era Retrospective Analysis for Research and Applications, Version 2 (MERRA-2), J. Clim.,
- 1113 30(14), 5419–5454, doi:10.1175/JCLI-D-16-0758.1, 2017.
- 1114 Gong, X., Wex, H., Müller, T., Wiedensohler, A., Höhler, K., Kandler, K., Ma, N., Dietel, B., Schiebel,
- 1115 T., Möhler, O. and Stratmann, F.: Characterization of aerosol properties at Cyprus, focusing on cloud
- 1116 condensation nuclei and ice-nucleating particles, Atmos. Chem. Phys., 19(16), 10883–10900,
- 1117 doi:10.5194/acp-19-10883-2019, 2019a.
- 1118 Gong, X., Wex, H., van Pinxteren, M., Triesch, N., Fomba, K. W., Lubitz, J., Stolle, C., Robinson, T.-B.,
- Müller, T., Herrmann, H. and Stratmann, F.: Ice nucleating particles measured in air, cloud and seawater
 at the Cape Verde Atmospheric Observatory (CVAO), , doi:10.1594/PANGAEA.906946, 2019b.
- 1121 Gong, X., Wex, H., van Pinxteren, M., Triesch, N., Fomba, K. W., Lubitz, J., Stolle, C., Robinson, T.-B.,
- 1122 Müller, T., Herrmann, H. and Stratmann, F.: Characterization of aerosol particles at Cabo Verde close to
- 1123 sea level and at the cloud level -- Part 2: Ice-nucleating particles in air, cloud and seawater, Atmos. Chem.
- 1124 Phys., 20(3), 1451–1468, doi:10.5194/acp-20-1451-2020, 2020.
- 1125 Hara, K., Maki, T., Kakikawa, M., Kobayashi, F. and Matsuki, A.: Effects of different temperature
- treatments on biological ice nuclei in snow samples, Atmos. Environ., 140, 415–419,
- 1127 doi:10.1016/j.atmosenv.2016.06.011, 2016.
- 1128 Harrison, A. D., Whale, T. F., Carpenter, M. A., Holden, M. A., Neve, L., O'Sullivan, D., Vergara
- 1129 Temprado, J. and Murray, B. J.: Not all feldspars are equal: a survey of ice nucleating properties across
- the feldspar group of minerals, Atmos. Chem. Phys., 16(17), 10927–10940, doi:10.5194/acp-16-109272016, 2016.
- 1132 Harrison, A. D., Lever, K., Sanchez-Marroquin, A., Holden, M. A., Whale, T. F., Tarn, M. D., McQuaid,
- 1133 J. B. and Murray, B. J.: The ice-nucleating ability of quartz immersed in water and its atmospheric
- importance compared to K-feldspar, Atmos. Chem. Phys., 19(17), 11343–11361, doi:10.5194/acp-19-
- **1135** 11343-2019, 2019.

- 1136 Hartmann, M., Adachi, K., Eppers, O., Haas, C., Herber, A., Holzinger, R., Hünerbein, A., Jäkel, E.,
- 1137 Jentzsch, C., van Pinxteren, M., Wex, H., Willmes, S. and Stratmann, F.: Wintertime Airborne
- 1138 Measurements of Ice Nucleating Particles in the High Arctic: A Hint to a Marine, Biogenic Source for Ice
- 1139 Nucleating Particles, Geophys. Res. Lett., 47(13), e2020GL087770,
- 1140 doi:https://doi.org/10.1029/2020GL087770, 2020.
- 1141 Hill, T. C. J., DeMott, P. J., Tobo, Y., Fröhlich-Nowoisky, J., Moffett, B. F., Franc, G. D. and
- 1142 Kreidenweis, S. M.: Sources of organic ice nucleating particles in soils, Atmos. Chem. Phys., 16(11),
- 1143 7195–7211, doi:10.5194/acp-16-7195-2016, 2016.
- 1144 Hinz, K.-P., Trimborn, A., Weingartner, E., Henning, S., Baltensperger, U. and Spengler, B.: Aerosol
- single particle composition at the Jungfraujoch, J. Aerosol Sci., 36(1), 123–145,
- 1146 doi:https://doi.org/10.1016/j.jaerosci.2004.08.001, 2005.
- 1147 Hiranuma, N., Augustin-Bauditz, S., Bingemer, H., Budke, C., Curtius, J., Danielczok, A., Diehl, K.,
- 1148 Dreischmeier, K., Ebert, M., Frank, F., Hoffmann, N., Kandler, K., Kiselev, A., Koop, T., Leisner, T.,
- 1149 Möhler, O., Nillius, B., Peckhaus, A., Rose, D., Weinbruch, S., Wex, H., Boose, Y., DeMott, P. J., Hader,
- 1150 J. D., Hill, T. C. J., Kanji, Z. A., Kulkarni, G., Levin, E. J. T., McCluskey, C. S., Murakami, M., Murray,
- 1151 B. J., Niedermeier, D., Petters, M. D., O'Sullivan, D., Saito, A., Schill, G. P., Tajiri, T., Tolbert, M. A.,
- 1152 Welti, A., Whale, T. F., Wright, T. P. and Yamashita, K.: A comprehensive laboratory study on the
- 1153 immersion freezing behavior of illite NX particles: a comparison of 17 ice nucleation measurement
- techniques, Atmos. Chem. Phys., 15(5), 2489–2518, doi:10.5194/acp-15-2489-2015, 2015.
- Hoose, C. and Möhler, O.: Heterogeneous ice nucleation on atmospheric aerosols: A review of resultsfrom laboratory experiments., 2012.
- 1157 Hoose, C., Kristjánsson, J. E., Chen, J.-P. and Hazra, A.: A Classical-Theory-Based Parameterization of
- 1158 Heterogeneous Ice Nucleation by Mineral Dust, Soot, and Biological Particles in a Global Climate Model,
- 1159 J. Atmos. Sci., 67(8), 2483–2503, doi:10.1175/2010JAS3425.1, 2010.
- 1160 Huffman, J. A., Perring, A. E., Savage, N. J., Clot, B., Crouzy, B., Tummon, F., Shoshanim, O., Damit,
- 1161 B., Schneider, J., Sivaprakasam, V., Zawadowicz, M. A., Crawford, I., Gallagher, M., Topping, D.,
- 1162 Doughty, D. C., Hill, S. C. and Pan, Y.: Real-time sensing of bioaerosols: Review and current
- 1163 perspectives, Aerosol Sci. Technol., 54(5), 465–495, doi:10.1080/02786826.2019.1664724, 2020.
- 1164 Huneeus, N., Schulz, M., Balkanski, Y., Griesfeller, J., Prospero, J., Kinne, S., Bauer, S., Boucher, O.,
- 1165 Chin, M., Dentener, F., Diehl, T., Easter, R., Fillmore, D., Ghan, S., Ginoux, P., Grini, A., Horowitz, L.,
- 1166 Koch, D., Krol, M. C., Landing, W., Liu, X., Mahowald, N., Miller, R., Morcrette, J.-J., Myhre, G.,

- Penner, J., Perlwitz, J., Stier, P., Takemura, T. and Zender, C. S.: Global dust model intercomparison in
 AeroCom phase I, Atmos. Chem. Phys., 11(15), 7781–7816, doi:10.5194/acp-11-7781-2011, 2011.
- 1169 Kanji, Z. A., Ladino, L. A., Wex, H., Boose, Y., Burkert-Kohn, M., Cziczo, D. J. and Krämer, M.:
- 1170 Overview of Ice Nucleating Particles, Meteorol. Monogr., 58, 1.1-1.33, doi:10.1175/amsmonographs-d-1171 16-0006.1, 2017.
- 1172 Khaniabadi, Y. O., Daryanoosh, S. M., Amrane, A., Polosa, R., Hopke, P. K., Goudarzi, G., Mohammadi,
- 1173 M. J., Sicard, P. and Armin, H.: Impact of Middle Eastern Dust storms on human health, Atmos. Pollut.
- 1174 Res., 8(4), 606–613, doi:https://doi.org/10.1016/j.apr.2016.11.005, 2017.
- 1175 Kinne, S., Schulz, M., Textor, C., Guibert, S., Balkanski, Y., Bauer, S. E., Berntsen, T., Berglen, T. F.,
- 1176 Boucher, O., Chin, M., Collins, W., Dentener, F., Diehl, T., Easter, R., Feichter, J., Fillmore, D., Ghan,
- 1177 S., Ginoux, P., Gong, S., Grini, A., Hendricks, J., Herzog, M., Horowitz, L., Isaksen, I., Iversen, T.,
- 1178 Kirkevåg, A., Kloster, S., Koch, D., Kristjansson, J. E., Krol, M., Lauer, A., Lamarque, J. F., Lesins, G.,
- 1179 Liu, X., Lohmann, U., Montanaro, V., Myhre, G., Penner, J., Pitari, G., Reddy, S., Seland, O., Stier, P.,
- 1180 Takemura, T. and Tie, X.: An AeroCom initial assessment optical properties in aerosol component
- 1181 modules of global models, Atmos. Chem. Phys., 6(7), 1815–1834, doi:10.5194/acp-6-1815-2006, 2006.
- 1182 Kleist, D. T., Parrish, D. F., Derber, J. C., Treadon, R., Wu, W.-S. and Lord, S.: Introduction of the GSI
- 1183 into the NCEP Global Data Assimilation System, Weather Forecast., 24(6), 1691–1705,
- 1184 doi:10.1175/2009WAF2222201.1, 2009.
- Klingmüller, K., Pozzer, A., Metzger, S., Stenchikov, G. L. and Lelieveld, J.: Aerosol optical depth trend
 over the Middle East, Atmos. Chem. Phys., 16(8), 5063–5073, doi:10.5194/acp-16-5063-2016, 2016.
- 1187 Knopf, D. A. and Koop, T.: Heterogeneous nucleation of ice on surrogates of mineral dust, J. Geophys.
- 1188 Res. Atmos., 111(D12), doi:https://doi.org/10.1029/2005JD006894, 2006.
- 1189 Kok, J. F., Adebiyi, A. A., Albani, S., Balkanski, Y., Checa-Garcia, R., Chin, M., Colarco, P. R.,
- 1190 Hamilton, D. S., Huang, Y., Ito, A., Klose, M., Li, L., Mahowald, N. M., Miller, R. L., Obiso, V., Pérez
- 1191 Garc\'\ia-Pando, C., Rocha-Lima, A. and Wan, J. S.: Contribution of the world's main dust source regions
- to the global cycle of desert dust, Atmos. Chem. Phys., 21(10), 8169–8193, doi:10.5194/acp-21-81692021, 2021.
- 1194 Könemann, T., Savage, N., Klimach, T., Walter, D., Fröhlich-Nowoisky, J., Su, H., Pöschl, U., Huffman,
- 1195 J. A. and Pöhlker, C.: Spectral Intensity Bioaerosol Sensor (SIBS): an instrument for spectrally resolved
- fluorescence detection of single particles in real time, Atmos. Meas. Tech., 12(2), 1337–1363,
- 1197 doi:10.5194/amt-12-1337-2019, 2019.

- Krasnov, H., Katra, I. and Friger, M.: Increase in dust storm related PM10 concentrations: A time series
 analysis of 2001–2015, Environ. Pollut., 213, 36–42, doi:https://doi.org/10.1016/j.envpol.2015.10.021,
 2016.
- 1201 Krueger, B. J., Grassian, V. H., Cowin, J. P. and Laskin, A.: Heterogeneous chemistry of individual
- 1202 mineral dust particles from different dust source regions: the importance of particle mineralogy, Atmos.
- 1203 Environ., 38(36), 6253–6261, doi:https://doi.org/10.1016/j.atmosenv.2004.07.010, 2004.
- 1204 Krzywinski, M. and Altman, N.: Error bars, Nat. Methods, 10(10), 921–922, doi:10.1038/nmeth.2659,
 1205 2013.
- 1206 Ladino, L. A., Raga, G. B., Alvarez-Ospina, H., Andino-Enr\'\iquez, M. A., Rosas, I., Mart\'\inez, L.,
- 1207 Salinas, E., Miranda, J., Raml'\irez-D\'\iaz, Z., Figueroa, B., Chou, C., Bertram, A. K., Quintana, E. T.,
- 1208 Maldonado, L. A., Garc\'\ia-Reynoso, A., Si, M. and Irish, V. E.: Ice-nucleating particles in a coastal
- tropical site, Atmos. Chem. Phys., 19(9), 6147–6165, doi:10.5194/acp-19-6147-2019, 2019.
- Lohmann, U. and Feichter, J.: Global indirect aerosol effects: a review, Atmos. Chem. Phys., 5(3), 715–
 737, doi:10.5194/acp-5-715-2005, 2005.
- 1212 Manders, A. M. ., Schapp, M., Jozwicka, M., van Arkel, F., Weijers, E. . and Matthijsen, J.: The
- 1213 contribution of sea salt to PM 10 and PM in the Netherlands, [online] Available from:
- 1214 http://www.pbl.nl/sites/default/files/cms/publicaties/500099004.pdf, 2009.
- 1215 Martin, A. C., Cornwell, G., Beall, C. M., Cannon, F., Reilly, S., Schaap, B., Lucero, D., Creamean, J.,
- 1216 Ralph, F. M., Mix, H. T. and Prather, K.: Contrasting local and long-range-transported warm ice-
- 1217 nucleating particles during an atmospheric river in coastal California, USA, Atmos. Chem. Phys., 19(7),
- 1218 4193–4210, doi:10.5194/acp-19-4193-2019, 2019.
- 1219 Mayol, E., Jiménez, M. A., Herndl, G. J., Duarte, C. M. and Arrieta, J. M.: Resolving the abundance and
- 1220 air-sea fluxes of airborne microorganisms in the North Atlantic Ocean, Front. Microbiol., 5, 557,
- doi:10.3389/fmicb.2014.00557, 2014.
- 1222 McCluskey, C. S., Hill, T. C. J., Malfatti, F., Sultana, C. M., Lee, C., Santander, M. V, Beall, C. M.,
- 1223 Moore, K. A., Cornwell, G. C., Collins, D. B., Prather, K. A., Jayarathne, T., Stone, E. A., Azam, F.,
- 1224 Kreidenweis, S. M. and DeMott, P. J.: A Dynamic Link between Ice Nucleating Particles Released in
- 1225 Nascent Sea Spray Aerosol and Oceanic Biological Activity during Two Mesocosm Experiments, J.
- 1226 Atmos. Sci., 74(1), 151–166, doi:10.1175/JAS-D-16-0087.1, 2017.
- 1227 McCluskey, C. S., Hill, T. C. J., Sultana, C. M., Laskina, O., Trueblood, J., Santander, M. V., Beall, C.

- 1228 M., Michaud, J. M., Kreidenweis, S. M., Prather, K. A., Grassian, V., DeMott, P. J., McCluskey, C. S.,
- 1229 Hill, T. C. J., Sultana, C. M., Laskina, O., Trueblood, J., Santander, M. V., Beall, C. M., Michaud, J. M.,
- 1230 Kreidenweis, S. M., Prather, K. A., Grassian, V. and DeMott, P. J.: A mesocosm double feature: Insights
- 1231 into the chemical make-up of marine ice nucleating particles, J. Atmos. Sci., JAS-D-17-0155.1,
- 1232 doi:10.1175/JAS-D-17-0155.1, 2018a.
- 1233 McCluskey, C. S., Hill, T. C. J., Sultana, C. M., Laskina, O., Trueblood, J., Santander, M. V, Beall, C.
- 1234 M., Michaud, J. M., Kreidenweis, S. M., Prather, K. A., Grassian, V. and DeMott, P. J.: A Mesocosm
- 1235 Double Feature: Insights into the Chemical Makeup of Marine Ice Nucleating Particles, J. Atmos. Sci.,
- 1236 75(7), 2405–2423, doi:10.1175/JAS-D-17-0155.1, 2018b.
- 1237 McCluskey, C. S., Ovadnevaite, J., Rinaldi, M., Atkinson, J., Belosi, F., Ceburnis, D., Marullo, S., Hill,
- 1238 T. C. J., Lohmann, U., Kanji, Z. A., O'Dowd, C., Kreidenweis, S. M. and DeMott, P. J.: Marine and
- 1239 Terrestrial Organic Ice-Nucleating Particles in Pristine Marine to Continentally Influenced Northeast
- 1240 Atlantic Air Masses, J. Geophys. Res. Atmos., 123(11), 6196–6212, doi:10.1029/2017JD028033, 2018c.
- 1241 McCluskey, C. S., Hill, T. C. J., Humphries, R. S., Rauker, A. M., Moreau, S., Strutton, P. G., Chambers,
- 1242 S. D., Williams, A. G. and McRobert, I.: Observations of Ice Nucleating Particles Over Southern Ocean
- 1243 Waters, Geophys. Res. Lett., 989–997, doi:10.1029/2018GL079981, 2018d.

Megahed, K., 2007: The impact of mineral dust aerosol particles on cloud formation. Ph.D. dissertation,
 Rheinische Friedrich-Wilhelms-University Bonn, 175 pp. [Available online at https://bonndoc.ulb.uni bonn.de/xmlui/handle/20.500.11811/3083]

- 1247 Mitts, B., Wang, X., Lucero, D., Beall, C., Deane, G., DeMott, P. and Prather, K.: Importance of
- 1248 Supermicron Ice Nucleating Particles in Nascent Sea Spray, Geophys. Res. Lett., n/a(n/a),
- 1249 e2020GL089633, doi:https://doi.org/10.1029/2020GL089633, 2021.
- 1250 Molod, A., Takacs, L., Suarez, M. and Bacmeister, J.: Development of the GEOS-5 atmospheric general
- 1251 circulation model: evolution from MERRA to MERRA2, Geosci. Model Dev., 8(5), 1339–1356,
- 1252 doi:10.5194/gmd-8-1339-2015, 2015.
- Murray, B. J., O'Sullivan, D., Atkinson, J. D. and Webb, M. E.: Ice nucleation by particles immersed in
 supercooled cloud droplets., Chem. Soc. Rev., 41(19), 6519–54, doi:10.1039/c2cs35200a, 2012.
- 1255 Nickovic, S., Vukovic, A., Vujadinovic, M., Djurdjevic, V. and Pejanovic, G.: Technical Note: High-
- 1256 resolution mineralogical database of dust-productive soils for atmospheric dust modeling, Atmos. Chem.
- 1257 Phys., 12(2), 845-855, doi:10.5194/acp-12-845-2012, 2012.

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- 1258 Niedermeier, D., Augustin-Bauditz, S., Hartmann, S., Wex, H., Ignatius, K. and Stratmann, F.: Can we
- 1259 define an asymptotic value for the ice active surface site density for heterogeneous ice nucleation?, J.
- 1260 Geophys. Res. Atmos., 120(10), 5036–5046, doi:https://doi.org/10.1002/2014JD022814, 2015.
- 1261 Niemand, M., Möhler, O., Vogel, B., Vogel, H., Hoose, C., Connolly, P., Klein, H., Bingemer, H.,
- 1262 Demott, P., Skrotzki, J. and Leisner, T.: A particle-surface-area-based parameterization of immersion
- 1263 freezing on desert dust particles, J. Atmos. Sci., 69(10), 3077–3092, doi:10.1175/JAS-D-11-0249.1, 2012.
- 1264 Nortcliff, S.: World Soil Resources and Food Security. Edited by R. Lal and BA Stewart. Boca Raton, Fl,
- 1265 USA: CRC Press (2012), pp. 574,£82.00. ISBN-13: 978-1439844502., Exp. Agric., 48(2), 305–306,
- **1266** 2012.
- 1267 O'Sullivan, D., Murray, B. J., Malkin, T. L., Whale, T. F., Umo, N. S., Atkinson, J. D., Price, H. C.,
- Baustian, K. J., Browse, J. and Webb, M. E.: Ice nucleation by fertile soil dusts: relative importance of
 mineral and biogenic components, Atmos. Chem. Phys., 14(4), 1853–1867, doi:10.5194/acp-14-18532014, 2014.
- 1271 O'Sullivan, D., Adams, M. P., Tarn, M. D., Harrison, A. D., Vergara-Temprado, J., Porter, G. C. E.,
- 1272 Holden, M. A., Sanchez-Marroquin, A., Carotenuto, F., Whale, T. F., McQuaid, J. B., Walshaw, R.,
- 1273 Hedges, D. H. P., Burke, I. T., Cui, Z. and Murray, B. J.: Contributions of biogenic material to the
- 1274 atmospheric ice-nucleating particle population in North Western Europe, Sci. Rep., 8(1), 13821,
- doi:10.1038/s41598-018-31981-7, 2018.
- Paramonov, M., David, R. O., Kretzschmar, R. and Kanji, Z. A.: A laboratory investigation of the ice
 nucleation efficiency of three types of mineral and soil dust, Atmos. Chem. Phys., 18(22), 16515–16536,
 doi:10.5194/acp-18-16515-2018, 2018.
- 1279 Van Pelt, R. S. and Zobeck, T. M.: Chemical Constituents of Fugitive Dust, Environ. Monit. Assess.,
 1280 130(1), 3–16, doi:10.1007/s10661-006-9446-8, 2007.
- Perkins, R. J., Gillette, S. M., Hill, T. C. J. and DeMott, P. J.: The Labile Nature of Ice Nucleation by
 Arizona Test Dust, ACS Earth Sp. Chem., 4(1), 133–141, doi:10.1021/acsearthspacechem.9b00304,
- **1283** 2020.
- 1284 Pfannerstill, E. Y., Wang, N., Edtbauer, A., Bourtsoukidis, E., Crowley, J. N., Dienhart, D., Eger, P. G.,
- 1285 Ernle, L., Fischer, H., Hottmann, B., Paris, J.-D., Stönner, C., Tadic, I., Walter, D., Lelieveld, J. and
- 1286 Williams, J.: Shipborne measurements of total OH reactivity around the Arabian Peninsula and its role in
- 1287 ozone chemistry, Atmos. Chem. Phys., 19(17), 11501–11523, doi:10.5194/acp-19-11501-2019, 2019.

- 1288 Price, H. C., Baustian, K. J., McQuaid, J. B., Blyth, A., Bower, K. N., Choularton, T., Cotton, R. J., Cui,
- 1289 Z., Field, P. R., Gallagher, M., Hawker, R., Merrington, A., Miltenberger, A., Neely III, R. R., Parker, S.
- 1290 T., Rosenberg, P. D., Taylor, J. W., Trembath, J., Vergara-Temprado, J., Whale, T. F., Wilson, T. W.,
- 1291 Young, G. and Murray, B. J.: Atmospheric Ice-Nucleating Particles in the Dusty Tropical Atlantic, J.
- 1292 Geophys. Res. Atmos., 123(4), 2175–2193, doi:https://doi.org/10.1002/2017JD027560, 2018.
- 1293 Prodi, F., Santachiara, G. and Oliosi, F.: Characterization of aerosols in marine environments
- 1294 (Mediterranean, Red Sea, and Indian Ocean), J. Geophys. Res. Ocean., 88(C15), 10957–10968,
- 1295 doi:https://doi.org/10.1029/JC088iC15p10957, 1983.
- 1296 Rienecker, M. M., Suarez, M. J., Gelaro, R., Todling, R., Bacmeister, J., Liu, E., Bosilovich, M. G.,
- 1297 Schubert, S. D., Takacs, L., Kim, G.-K., Bloom, S., Chen, J., Collins, D., Conaty, A., da Silva, A., Gu,
- 1298 W., Joiner, J., Koster, R. D., Lucchesi, R., Molod, A., Owens, T., Pawson, S., Pegion, P., Redder, C. R.,
- 1299 Reichle, R., Robertson, F. R., Ruddick, A. G., Sienkiewicz, M. and Woollen, J.: MERRA: NASA's
- 1300 Modern-Era Retrospective Analysis for Research and Applications, J. Clim., 24(14), 3624–3648,
- doi:10.1175/JCLI-D-11-00015.1, 2011.
- Salam, A., Lohmann, U. and Lesins, G.: Ice nucleation of ammonia gas exposed montmorillonite mineral
 dust particles, Atmos. Chem. Phys., 7(14), 3923–3931, doi:10.5194/acp-7-3923-2007, 2007.
- 1304 Schnell, R. C.: Ice Nuclei in Seawater, Fog Water and Marine Air off the Coast of Nova Scotia: Summer
- 1305 1975, J. Atmos. Sci., 34(8), 1299–1305, doi:10.1175/1520-0469(1977)034<1299:INISFW>2.0.CO;2,
 1306 1977.
- 1307 Schrod, J., Weber, D., Drücke, J., Keleshis, C., Pikridas, M., Ebert, M., Cvetković, B., Nickovic, S.,
- 1308 Marinou, E., Baars, H., Ansmann, A., Vrekoussis, M., Mihalopoulos, N., Sciare, J., Curtius, J. and
- 1309 Bingemer, H. G.: Ice nucleating particles over the Eastern Mediterranean measured by unmanned aircraft
- 1310 systems, Atmos. Chem. Phys., 17(7), 4817–4835, doi:10.5194/acp-17-4817-2017, 2017.
- 1311 Shahsavani, A., Naddafi, K., Jafarzade Haghighifard, N., Mesdaghinia, A., Yunesian, M., Nabizadeh, R.,
- 1312 Arahami, M., Sowlat, M. H., Yarahmadi, M., Saki, H., Alimohamadi, M., Nazmara, S., Motevalian, S. A.
- 1313 and Goudarzi, G.: The evaluation of PM10, PM2.5, and PM1 concentrations during the Middle Eastern
- 1314 Dust (MED) events in Ahvaz, Iran, from april through september 2010, J. Arid Environ., 77, 72–83,
- 1315 doi:https://doi.org/10.1016/j.jaridenv.2011.09.007, 2012.
- 1316 Sullivan, R. C., Guazzotti, S. A., Sodeman, D. A. and Prather, K. A.: Direct observations of the
- 1317 atmospheric processing of Asian mineral dust, Atmos. Chem. Phys., 7(5), 1213–1236, doi:10.5194/acp-71318 1213-2007, 2007.

- 1319 Sullivan, R. C., Miñambres, L., DeMott, P. J., Prenni, A. J., Carrico, C. M., Levin, E. J. T. and
- 1320 Kreidenweis, S. M.: Chemical processing does not always impair heterogeneous ice nucleation of mineral
- 1321 dust particles, Geophys. Res. Lett., 37(24), doi:https://doi.org/10.1029/2010GL045540, 2010a.
- 1322 Sullivan, R. C., Petters, M. D., DeMott, P. J., Kreidenweis, S. M., Wex, H., Niedermeier, D., Hartmann,
- 1323 S., Clauss, T., Stratmann, F., Reitz, P., Schneider, J. and Sierau, B.: Irreversible loss of ice nucleation
- 1324 active sites in mineral dust particles caused by sulphuric acid condensation, Atmos. Chem. Phys., 10(23),
- 1325 11471–11487, doi:10.5194/acp-10-11471-2010, 2010b.
- 1326 Suski, K. J., Hill, T. C. J., Levin, E. J. T., Miller, A., DeMott, P. J. and Kreidenweis, S. M.: Agricultural
- harvesting emissions of ice-nucleating particles, Atmos. Chem. Phys., 18(18), 13755–13771,
- 1328 doi:10.5194/acp-18-13755-2018, 2018.
- 1329 Tadic, I., Crowley, J. N., Dienhart, D., Eger, P., Harder, H., Hottmann, B., Martinez, M., Parchatka, U.,
- 1330 Paris, J.-D., Pozzer, A., Rohloff, R., Schuladen, J., Shenolikar, J., Tauer, S., Lelieveld, J. and Fischer, H.:
- 1331 Net ozone production and its relationship to nitrogen oxides and volatile organic compounds in the
- 1332 marine boundary layer around the Arabian Peninsula, Atmos. Chem. Phys., 20(11), 6769–6787,
- 1333 doi:10.5194/acp-20-6769-2020, 2020.
- 1334 Tobo, Y., DeMott, P. J., Hill, T. C. J., Prenni, A. J., Swoboda-Colberg, N. G., Franc, G. D. and
- Kreidenweis, S. M.: Organic matter matters for ice nuclei of agricultural soil origin, Atmos. Chem. Phys.,
 14(16), 8521–8531, doi:10.5194/acp-14-8521-2014, 2014.
- 1337 Ullrich, R., Hoose, C., Möhler, O., Niemand, M., Wagner, R., Höhler, K., Hiranuma, N., Saathoff, H. and
- 1338 Leisner, T.: A new ice nucleation active site parameterization for desert dust and soot, J. Atmos. Sci.,
- 1339 74(3), 699–717, doi:10.1175/JAS-D-16-0074.1, 2017.
- $\label{eq:alpha} \textbf{1340} \qquad \textbf{Vali, G.: Quantitative Evaluation of Experimental Results an the Heterogeneous Freezing Nucleation of } \textbf{Vali, G.: Quantitative Evaluation of Experimental Results and the Heterogeneous Freezing Nucleation of } \textbf{Vali, G.: Quantitative Evaluation of Experimental Results and the Heterogeneous Freezing Nucleation of } \textbf{Vali, G.: Quantitative Evaluation of Experimental Results and the Heterogeneous Freezing Nucleation of } \textbf{Vali, G.: Quantitative Evaluation of Experimental Results and the Heterogeneous Freezing Nucleation of } \textbf{Vali, G.: Quantitative Evaluation of Experimental Results and the Heterogeneous Freezing Nucleation of } \textbf{Vali, G.: Quantitative Evaluation of Experimental Results and the Heterogeneous Freezing Nucleation of } \textbf{Vali, G.: Quantitative Evaluation of Experimental Results and the Heterogeneous Freezing Nucleation of } \textbf{Vali, G.: Quantitative Evaluation of Experimental Results and the Heterogeneous Freezing Nucleation of } \textbf{Vali, G.: Quantitative Evaluation of Experimental Results and the Heterogeneous Freezing Nucleation of } \textbf{Vali, G.: Quantitative Evaluation of Experimental Results and the Heterogeneous Freezing Nucleation of } \textbf{Vali, G.: Quantitative Evaluation of } \textbf{Vali, G.: Quantitative Evaluative Evaluative Evaluative Evaluative Evaluative Evaluative Evaluative E$
- 1341 Supercooled Liquids, J. Atmos. Sci., 28(3), 402–409, doi:10.1175/1520-
- 1342 0469(1971)028<0402:QEOERA>2.0.CO;2, 1971.
- 1343 Vergara-Temprado, J., Murray, B. J., Wilson, T. W., O'Sullivan, D., Browse, J., Pringle, K. J., Ardon-
- 1344 Dryer, K., Bertram, A. K., Burrows, S. M., Ceburnis, D., Demott, P. J., Mason, R. H., O'Dowd, C. D.,
- 1345 Rinaldi, M. and Carslaw, K. S.: Contribution of feldspar and marine organic aerosols to global ice
- 1346 nucleating particle concentrations, Atmos. Chem. Phys., 17(5), 3637–3658, doi:10.5194/acp-17-3637-
- 1347 2017, 2017.
- 1348 Vergara-Temprado, J., Miltenberger, A. K., Furtado, K., Grosvenor, D. P., Shipway, B. J., Hill, A. A.,
- 1349 Wilkinson, J. M., Field, P. R., Murray, B. J. and Carslaw, K. S.: Strong control of Southern Ocean cloud

- 1350 reflectivity by ice-nucleating particles, Proc. Natl. Acad. Sci., 115(11), 2687 LP 2692,
- 1351 doi:10.1073/pnas.1721627115, 2018.
- 1352 Wang, N., Edtbauer, A., Stönner, C., Pozzer, A., Bourtsoukidis, E., Ernle, L., Dienhart, D., Hottmann, B.,
- 1353 Fischer, H., Schuladen, J., Crowley, J. N., Paris, J.-D., Lelieveld, J. and Williams, J.: Measurements of
- 1354 carbonyl compounds around the Arabian Peninsula: overview and model comparison, Atmos. Chem.
- 1355 Phys., 20(18), 10807–10829, doi:10.5194/acp-20-10807-2020, 2020.
- 1356 Wang, X., Deane, G. B., Moore, K. A., Ryder, O. S., Stokes, M. D., Beall, C. M., Collins, D. B.,
- 1357 Santander, M. V, Burrows, S. M., Sultana, C. M. and Prather, K. A.: The role of jet and film drops in
- controlling the mixing state of submicron sea spray aerosol particles, Proc. Natl. Acad. Sci., 114(27),
- 1359 6978–6983, doi:10.1073/pnas.1702420114, 2017.
- 1360 von der Weiden, S.-L., Drewnick, F. and Borrmann, S.: Particle Loss Calculator a new software tool for
- the assessment of the performance of aerosol inlet systems, Atmos. Meas. Tech., 2(2), 479–494,
- 1362 doi:10.5194/amt-2-479-2009, 2009.
- Welti, A., Lüönd, F., Kanji, Z. A., Stetzer, O. and Lohmann, U.: Time dependence of immersion freezing:
 an experimental study on size selected kaolinite particles, Atmos. Chem. Phys., 12(20), 9893–9907,
- 1365 doi:10.5194/acp-12-9893-2012, 2012.
- 1366 Wex, H., DeMott, P. J., Tobo, Y., Hartmann, S., Rösch, M., Clauss, T., Tomsche, L., Niedermeier, D. and
- 1367Stratmann, F.: Kaolinite particles as ice nuclei: learning from the use of different kaolinite samples and
- 1368 different coatings, Atmos. Chem. Phys., 14(11), 5529–5546, doi:10.5194/acp-14-5529-2014, 2014.
- 1369 Wex, H., Huang, L., Zhang, W., Hung, H., Traversi, R., Becagli, S., Sheesley, R. J., Moffett, C. E.,
- 1370 Barrett, T. E., Bossi, R., Skov, H., Hünerbein, A., Lubitz, J., Löffler, M., Linke, O., Hartmann, M.,
- 1371 Herenz, P. and Stratmann, F.: Annual variability of ice-nucleating particle concentrations at different
- 1372 Arctic locations, Atmos. Chem. Phys., 19(7), 5293–5311, doi:10.5194/acp-19-5293-2019, 2019.
- 1373 Whale, T. F., Murray, B. J., O'Sullivan, D., Wilson, T. W., Umo, N. S., Baustian, K. J., Atkinson, J. D.,
- 1374 Workneh, D. A. and Morris, G. J.: A technique for quantifying heterogeneous ice nucleation in microlitre
- 1375 supercooled water droplets, Atmos. Meas. Tech., 8(6), 2437–2447, doi:10.5194/amt-8-2437-2015, 2015.
- 1376 Wilson, T. W., Ladino, L. a., Alpert, P. a., Breckels, M. N., Brooks, I. M., Browse, J., Burrows, S. M.,
- 1377 Carslaw, K. S., Huffman, J. A., Judd, C., Kilthau, W. P., Mason, R. H., McFiggans, G., Miller, L. a.,
- 1378 Nájera, J. J., Polishchuk, E., Rae, S., Schiller, C. L., Si, M., Temprado, J. V., Whale, T. F., Wong, J. P. S.,
- 1379 Wurl, O., Yakobi-Hancock, J. D., Abbatt, J. P. D., Aller, J. Y., Bertram, A. K., Knopf, D. a. and Murray,
- 1380 B. J.: A marine biogenic source of atmospheric ice-nucleating particles, Nature, 525(7568), 234–238,

- 1381 doi:10.1038/nature14986, 2015.
- 1382 Wu, W.-S., Purser, R. J. and Parrish, D. F.: Three-Dimensional Variational Analysis with Spatially
- 1383 Inhomogeneous Covariances, Mon. Weather Rev., 130(12), 2905–2916, doi:10.1175/1520-
- 1384 0493(2002)130<2905:TDVAWS>2.0.CO;2, 2002.
- 1385 Yadav, S., Venezia, R. E., Paerl, R. W. and Petters, M. D.: Characterization of Ice-Nucleating Particles
- 1386 Over Northern India, J. Geophys. Res. Atmos., 124(19), 10467–10482,
- 1387 doi:https://doi.org/10.1029/2019JD030702, 2019.
- 1388 Yahya, R. Z., Arrieta, J. M., Cusack, M. and Duarte, C. M.: Airborne Prokaryote and Virus Abundance
- 1389 Over the Red Sea, Front. Microbiol., 10, 1112, doi:10.3389/fmicb.2019.01112, 2019.
- 1390 Yang, J., Wang, Z., Heymsfield, A. J., DeMott, P. J., Twohy, C. H., Suski, K. J. and Toohey, D. W.: High
- 1391 ice concentration observed in tropical maritime stratiform mixed-phase clouds with top temperatures
- 1392 warmer than -8 °C, Atmos. Res., 233, 104719, doi:https://doi.org/10.1016/j.atmosres.2019.104719, 2020.
- 1393 Yost, J. L. and Hartemink, A. E.: Chapter Four Soil organic carbon in sandy soils: A review, vol. 158,
- edited by D. L. Sparks, pp. 217–310, Academic Press., 2019.
- 1395 Yu, Y., Kalashnikova, O. V, Garay, M. J., Lee, H. and Notaro, M.: Identification and Characterization of
- 1396 Dust Source Regions Across North Africa and the Middle East Using MISR Satellite Observations,
- 1397 Geophys. Res. Lett., 45(13), 6690–6701, doi:https://doi.org/10.1029/2018GL078324, 2018.
- 1398 Zolles, T., Burkart, J., Häusler, T., Pummer, B., Hitzenberger, R. and Grothe, H.: Identification of Ice
- 1399 Nucleation Active Sites on Feldspar Dust Particles, J. Phys. Chem. A, 119(11), 2692–2700,
- 1400 doi:10.1021/jp509839x, 2015.

1401