The Urmia playa as a source of airborne dust and ice-nucleating particles – Part 2: Unraveling the relationship between soil dust composition and ice nucleation activity

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Received: 23 June 2022 – Discussion started: 26 July 2022
Revised: 2 October 2022 – Accepted: 20 October 2022 – Published: 23 November 2022

Abstract. Ice-nucleating particles (INPs) originating from deserts, semi-arid regions, and dried lakebeds may cause heterogeneous ice nucleation, impacting cloud properties. Recently, due to climate change and water scarcity, abandoned agricultural lands with little surficial crust and negligible vegetation cover have become an increasing source of atmospheric dust worldwide. Unlike deserts, these areas are rich in soluble salt and (bio-)organic compounds. Using soil samples from various sites of the Lake Urmia playa (LUP) in northwestern Iran and airborne dusts collected at nearby meteorological stations, we elucidate how minerals, soluble salts, and organic matter interact to determine the IN activity of saline soils and dust. X-ray powder diffraction shows that the mineralogical composition is dominated by K-feldspars (microcline), quartz, carbonates, and clay minerals. The samples were stripped stepwise of organic matter, carbonates, and soluble salts. After each removal step, the ice nucleation (IN) activity was quantified in terms of onset freezing temperatures ($T_{\text{het}}$) and heterogeneously frozen fractions ($F_{\text{het}}$) by emulsion freezing experiments using differential scanning calorimetry (DSC).

We examined the influence of soluble salts and pH on microcline and quartz in emulsion freezing experiments, comparing these with reference suspensions of microcline and quartz exposed to salt concentrations and pH levels characteristic of the LUP samples. These analyses, combined with correlations between $T_{\text{het}}$ and $F_{\text{het}}$, allow us to identify the components that contribute to or inhibit IN activity. The LUP dusts turn out to be very good INPs, with freezing onset temperatures around 248 K in immersion freezing experiments. Interestingly, their IN activity proves to be dominated by the relatively small share of (bio-)organic matter (1 %–5.3 %). After organic matter removal, the remaining IN activity ($T_{\text{het}} \approx 244$ K) can be traced back to the clay fraction, because $T_{\text{het}}$ and $F_{\text{het}}$ correlate positively with the clay mineral content but negatively with quartz and microcline. We attribute the inability of quartz and microcline to act as INPs to the basic pH of the LUP samples as well as to the presence of soluble salts. After additionally removing soluble salts and carbonates, the IN activity of the samples increased again significantly ($T_{\text{het}} \approx 249$ K), and the negative correlation with quartz and microcline turned into a slightly positive one. Removing carbonates and salts from the natural samples leads to an increase in $T_{\text{het}}$ and $F_{\text{het}}$ as well, indicating that their presence also suppresses the IN activity of the (bio-)organic INPs. Overall, this study demonstrates that mineral and organic INPs do not just add up to yield the IN activity of soil dust but that the freezing behavior is governed by inhibiting and promoting interactions between the components.

Published by Copernicus Publications on behalf of the European Geosciences Union.
1 Introduction

The ice content in clouds determines key cloud properties such as albedo, lifetime, and precipitation formation (e.g., Lohmann, 2006; Field and Heymsfield, 2015; Mülmenstädt et al., 2015). An accurate modeling of ice formation in clouds is therefore key to estimating the Earth’s radiation budget in response to climate change (e.g., IPCC, 2021; Lohmann and Neubauer, 2018). Ice formation through homogeneous nucleation occurs readily within the volume of cloud droplets at temperatures below about 237 K; yet, at higher temperatures, this process becomes too improbable to be of relevance, and seeds are required to initiate cloud glaciation. These seeds can be ice crystals falling from higher cloud layers to layers below through a seeder-feeder process (Purdy et al., 2005; Ramelli et al., 2021; Prosko et al., 2021), or they can be ice-nucleating particles (INPs), which contain sites that catalyze the formation of the ice phase (e.g., Kanji et al., 2017). The number of airborne particles that are able to induce ice at the highest temperatures is low but increases with decreasing temperature. Considerable effort has been undertaken to quantify the atmospheric INP population and its temporal fluctuation globally (e.g., Bigg, 1973; Kanji et al., 2017; Welti et al., 2020; Brunner et al., 2021). Measurements have been performed by means of online techniques on a single-particle level, e.g., in continuous flow diffusion chambers (DeMott et al., 2010; Tobo et al., 2013; Lacher et al., 2018, 2021; Welti et al., 2018, 2020; Brunner et al., 2021) or through filter sampling with subsequent analysis of the IN activity directly on the filters (Schroed et al., 2017; Price et al., 2018) or in drop-freezing assays (O’Sullivan et al., 2018; Testa et al., 2021; Chen et al., 2021; Welti et al., 2018, 2020). In most of these studies, direct physicochemical characterization of samples is limited. Usually, the types of INPs are indirectly inferred through air mass back trajectory analysis. In some of these studies, the sensitivity of IN activity to heat or H$_2$O$_2$ digestion is tested to infer the contribution of biogenic material to INPs (O’Sullivan et al., 2018; Testa et al., 2021). Some studies present an elemental characterization (Schroed et al., 2017; Welti et al., 2018; Price et al., 2018) or fluorescence measurements to infer the presence of biological INPs (Tobo et al., 2013).

Alternatively, samples have been surface collected and subsequently analyzed in drop-freezing assays (Conen et al., 2011; O’Sullivan et al., 2014; Tobo et al., 2014; Hill et al., 2016; Suski et al., 2018) in cloud chambers (Steinke et al., 2016, 2020) or in continuous flow diffusion chambers (Boose et al., 2016, 2019; Paramonov et al., 2018). Here, samples are sufficiently abundant to perform more detailed analyses and physicochemical characterization. One major finding from these studies is that (bio-)organic material present in soil dusts enables them to freeze water at higher temperatures than pure mineral dust does. INPs that are active at temperatures above 255–258 K have been ascribed to biogenic material due to their heat lability (O’Sullivan et al., 2018) and their sensitivity to H$_2$O$_2$ digestion, through the correlation of IN activity with the organic share of the samples, or a combination of these methods (Conen et al., 2011; Tobo et al., 2014; Hill et al., 2016; Testa et al., 2021).

Below 255 K, the role of (bio-)organic material as source of INPs is unclear. Paramonov et al. (2018) found a decrease of IN activity in the temperature range 233–243 K after removing heat-sensitive material from soil samples; yet, the decrease in IN activity was not proportional to the amount of organic matter present in the samples. The H$_2$O$_2$ treatment did not affect or it even enhanced IN activity of the investigated soil samples. Boose et al. (2019) found that the mineralogical composition determined the IN activity of most desert dust samples that they investigated at temperatures up to 242 K, with the exception of one carbonaceous sample, where most of the IN activity stemmed from organic matter.

To gain direct insight into the IN activity of aerosol constituents, freezing experiments with surrogate samples for different types of aerosols have been performed. Minerals and biological materials have attracted most attention, as they are considered the dominant INP types below and above 255–258 K, respectively. In laboratory studies, diverse types of organic and biogenic material proved to be IN active. Biological INPs include whole bacteria, pollen, and spores, but also fragments of plants and animals (e.g., Morris et al., 2013; Després et al., 2012; Kanji et al., 2017). Moreover, organic macromolecules of biological origin – such as proteinaceous material (e.g., Huang et al., 2021), humic and fulvic substances (Fornea et al., 2009; Wang and Knopf, 2011; Borduas-Dedekind et al., 2019), cellulose (Hiranuma et al., 2015a, 2019), polysaccharides (Steinke et al., 2020), and lignin (Bogler and Borduas-Dedekind, 2020; Steinke et al., 2020) – also proved to be IN active.

Among the minerals present in atmospheric dust (e.g., Murray et al., 2012; Kanji et al., 2017), feldspars, clay minerals, and quartz are considered most relevant as INPs. Other common minerals like calcite, dolomite, and micas proved to be hardly IN active (e.g., Kaufmann et al., 2016). Among the feldspars, K-feldspars showed higher IN activity than Na–Ca feldspars (e.g., Zolles et al., 2015; Kaufmann et al., 2016; Harrison et al., 2016, 2019; Kumar et al., 2019b; Welti et al., 2019); in particular, microcline remained IN active up to the highest temperatures observed for mineral particles (e.g., Atkinson et al., 2013; Kaufmann et al., 2016; Welti et al., 2019). Yet, the IN activity of feldspars also proved to be highly sensitive to the presence of solutes: while low concentrations of ammonia and ammonium ions enhanced it, alkali ions and acids deteriorated it (Kumar et al., 2018, 2019b; Whale et al., 2018; Perkins et al., 2020; Yun et al., 2020). Reference quartz samples showed high IN activity, but only after being freshly milled. This indicates that
IN-active sites on quartz depend on its surface structure and are most probably related to defects (Zolles et al., 2015; Kumar et al., 2019b). Clay minerals dominate the submicron fraction of atmospheric dust and can be transported over long distances and even between continents (e.g., Murray et al., 2012). All clay minerals that have been tested as INPs so far – that is, kaolinite, montmorillonite, and illite – proved to be IN active, though at lower temperatures than K-feldspars and quartz (e.g., Pinti et al., 2012; Hoose and Möhler, 2012; Hiranuma et al., 2015b; Kanji et al., 2017).

Only few studies have tried to relate the IN activity of natural samples with their mineralogical composition (O’Sullivan et al., 2014; Kaufmann et al., 2016; Boose et al., 2016; Paramonov et al., 2018). Kaufmann et al. (2016) were able to explain the freezing behavior of most of their surface-collected dust samples by their mineralogical composition. Boose et al. (2016, 2019) found a correlation between the IN activity of dust samples sourced from different deserts and their quartz and feldspar content. Yet, some of the samples were milled prior to performing freezing experiments, which might have enhanced the IN activity of quartz (Zolles et al., 2015; Kumar et al., 2019a). Paramonov et al. (2018) found a good correlation of IN activity with total feldspar and K-feldspar content of surface-collected dust. A relationship between mineralogical composition and IN activity was also observed by O’Sullivan et al. (2014) for temperatures below 255 K.

Different parameterizations to quantify the immersion freezing of dust particles have been proposed. Niemand et al. (2012) developed a parameterization of ice-active site density per surface area from cloud chamber measurements of desert dust samples. The parameterization proposed by DeMott et al. (2015) is based on laboratory studies and atmospheric measurements of dust representative of Saharan and Asian desert sources and links the prediction of INP number concentrations to particle number concentrations of sizes larger than 0.5 µm. These parameterizations premise that, to a first order, mineral dust may be assumed to be one particle type. Alternatively, a specific mineral type is taken to be representative for the mineral dust fraction like K-feldspars (Atkinson et al., 2013) or illite (Hiranuma et al., 2015b). These parameterizations have the advantage of being simple; yet, they neglect the complexity of soil dust and potential aging effects in the atmosphere.

Soil types that expand due to human influence are dried lakebeds, abandoned agricultural land, and saline soils on the margin of drying lakes (Abduwuaili et al., 2010; Goodman et al., 2019; Sweeney et al., 2016; Varga et al., 2014; Perez and Gill, 2009; Washington et al., 2006; Prospero et al., 2002). Such soils contribute significantly to aerosols locally, regionally, and potentially globally, as exemplified by the enormous amounts of dust that are deflated annually from the Aral Sea in Kazakhstan and Uzbekistan and Ebinur Lake in northwestern China (Abduwuaili et al., 2010). In this study, we focus on desiccation of Lake Urmia (LU), one of the biggest salt lakes in the world, in the northwest of Iran. Increasing soil salinity, which negatively affects plant growth and crop production, led to soil erosion and land degradation (Gorji et al., 2020; Jiang et al., 2019; Scudiero et al., 2015) and resulted in the remains of vast, barren lands composed of fine textured, lacustrine sediments (Hamzehpour et al., 2018; Shadkam et al., 2016; Farokhnia and Morid, 2014). These saline areas with limited vegetation cover in most parts might be highly vulnerable to wind erosion due to their unique physicochemical characteristics. Recent studies have indeed demonstrated the increase in the intensity of dust storms over LUP and nearby cities (Ahmady-Birgani et al., 2020; Boroughani et al., 2019; Sotoudeheian et al., 2016).

In the companion paper of this work (Hamzehpour et al., 2022), we identified highly erodible playa surfaces along the western Lake Urmia playa (LUP) and characterized four surface-collected soil samples together with airborne dust samples from nearby meteorological stations with respect to physicochemical properties, mineralogical composition, and IN activity. Emulsion freezing experiments in a differential scanning calorimeter (DSC) revealed that some of the lower concentrated suspensions of the dust samples (2 wt %) exhibited a higher heterogeneously frozen fraction (\(f_{\text{inh}}\)) of droplets than the higher concentrated samples (5 wt %). Such a reciprocal proportionality points to interactions between sample constituents that inhibit IN activity. Moreover, we found a negative correlation of IN activity with K-feldspar and quartz content, which is opposite to findings from previous studies (O’Sullivan et al., 2014; Boose et al., 2016, 2019; Paramonov et al., 2018). To elucidate the potential contribution or inhibition of (bio-)organic matter, minerals, and soluble salts to the IN activity, we performed treatments to remove carbonates, soluble salts, and organic matter from the samples. To attribute the observed change in IN activity to specific interactions between soluble salts and minerals, we performed reference freezing experiments with quartz and microcline suspensions, with salt concentration and pH in the typical range of the LUP samples. The insights gained from these treatments are the subject of this paper.

## 2 Material and methods

### 2.1 Sampling locations

The sampling procedure has been described in detail in Part 1 of this work (Hamzehpour et al., 2022). Here, we recapitulate the relevant information for the present study. Four soil samples were collected from the top 5 cm surface layer of highly wind-erodible areas from the north to the south of the Lake Urmia playa (LUP). Together with the soil samples, airborne dust samples were collected at meteorological stations near the soil sampling areas. Dust samples were collected using high-volume samplers manufactured by Graseby–Andersen (Smyrna, Georgia, USA) on a 20.3 cm × 25.4 cm glass micro-fiber filter (Whatman Inc.,...
USA) at flow rates of 1.13–1.41 m$^3$ min$^{-1}$ for 24 h. Detailed discussion of the dust sources in the region are presented in the companion paper. Here, we just give a summary of the sampling locations in Table 1.

### 2.2 Physicochemical analysis

A detailed characterization of the samples has been given and discussed in Part 1 of this work (Hamzehpour et al., 2022). Here, we summarize the applied methods and replicate the findings with relevance to this study.

The soil samples were passed through a 2 mm sieve and, along with the dust samples, their physicochemical properties were determined, as summarized in Table 2. Soil electrical conductivity (EC) and acidity (pH) were measured in a 1 : 2.5 soil to water suspension using a Jenway conductivity meter (model 4510) and VWR Symphony SB70P pH meter, respectively (Rhoades, 1996). In addition, soil organic carbon (OC) was measured using a wet oxidation technique (Nelson and Sommers, 1996) and converted to organic matter (OM) by multiplying by a factor of 2. Soil total carbonates were determined through back titration of the remaining HCl (Sparks et al., 2020).

### 2.3 Quantitative mineralogy

The mineralogical composition of the bulk dust samples and surface-collected soil samples was investigated with X-ray diffraction (XRD) analysis (Bish and Plötze, 2010) followed by quantitative Rietveld analysis. More details are given in Part 1 (Hamzehpour et al., 2022). We replicate the results in Table 3.

### 2.4 Thermogravimetric analysis

Thermogravimetric analysis (TGA) and differential scanning calorimetry (DSC) of soil and dust samples smaller than 63 µm were measured on a STA 449 F5 Jupiter from Netzsch coupled to a QMS 403 D Aëolos mass spectrometer. The gas flow was set to 60 + 20 mL Ar min$^{-1}$ and 20 mL O$_2$ min$^{-1}$ to obtain an atmospheric mixture. A soil or dust sample (10–20 mg) was placed in an Al$_2$O$_3$ crucible and heated up with a heating rate of 10°C min$^{-1}$ from 40 to 1000°C. In TGA thermograms, the region 120–190°C is a transition between endothermic water loss and exothermic organic matter oxidation. The data were analyzed and plotted with the Proteus software from Netzsch.

### 2.5 Ice nucleation activity of treated soil and dust samples

A statistical design with six treatments, each with one replication, was used to study the effects of soluble salt removal (SR), carbonates removal (CR), and organic matter removal (OMR) on the IN activity of soil and dust samples, as illustrated in Fig. 1. Untreated samples (natural samples) are used as the reference. Apart from the single treatments, combined treatments of salt and carbonate removal (SR + CR), salt and organic matter removal (SR + OMR), and carbonate, salt, and organic matter removal (CR + SR + OMR) were performed. The combined organic matter and carbonates removal (OMR + CR) was excluded from further analysis, because the measurement of electrical conductivity of the thus-treated samples revealed a concomitant major loss of salts, making them similar to the combined CR + SR + OMR-treated samples.

To remove stepwise-soluble salts, carbonates, and organic matter from the soil and dust samples, we followed the procedure described by Jackson (1985), Kittrick and Hope (1963), and Dane and Topp (2020), as outlined below.

#### 2.5.1 Removal of soluble salts

In order to remove soluble salts, samples were washed with distilled water. The samples were suspended in distilled water and shaken for 1 min, followed by centrifugation with 6000–7000 rpm for 5 min. A ratio of 1 : 2.5 soil to water was found to be efficient enough for removing salts after examining different ratios in test experiments. The procedure was repeated until the leached-salt concentration dropped below 10 mM. After drying in an oven at 60°C, the samples were stored in 20 mL glass vials for further analysis.

#### 2.5.2 Removal of carbonates

Carbonates were removed from the samples by using a 1 M sodium acetate buffer solution with adjusted pH of 5. Careful pH monitoring is important, because for pH > 5, the dis-
solution of carbonates remains incomplete, while pH < 5 can lead to the dissolution of iron hydroxides. A total of 40 mL of sodium acetate buffer solution was added to 1000 mg of each sample in a 50 mL Erlenmeyer flask and then kept at 60°C in a water bath until CO₂ bubbles were no longer visible. Then, the suspension was centrifuged and the supernatant decanted. In order to prevent organic matter decomposition of the samples, the temperature should not exceed 60°C and stored in 20 mL glass vials for further analysis.

2.5.3 Removal of organic matter

In order to remove the organic matter from the samples, 5 mL of 30% (wt/wt) H₂O₂ was added to a known amount of the samples in a tall, 500 mL beaker and stirred until the reaction slowed. Then, the suspension was placed in a water bath at 65–70°C until the reaction ceased and the solution became clear. H₂O₂ solution was added in 5 mL portions until no more reaction was evident. Thereafter, the samples were centrifuged for 15 min in order to remove the supernatant. In case of sample dispersion, a few droplets of 0.5 M MgCl₂ were added to the sample. Finally, the samples were dried in the oven at 60°C and stored in 20 mL glass vials for further analysis. Based on the study by Daily et al. (2022), samples needed to be heated above 90°C to damage the IN activity of the minerals. Therefore, we do not expect a negative effect on the IN activity of the mineral components due to the heating required to remove organic matter; yet, we cannot exclude it.

2.6 Reference experiments with the minerals microcline and quartz

In order to investigate the role of salt type and concentration and pH on the IN activity of microcline and quartz, we performed experiments with a microcline sample from North Macedonia (the same as has been used in Klumpp et al., 2022) and a quartz sample from Sigma-Aldrich (the same as has been used in Kumar et al., 2019a), with 2 wt% and 1 wt% concentrations, respectively. NaCl, CaCl₂, and MgCl₂ (all Sigma-Aldrich) were chosen as soluble salts, since they

Table 1. The location of the collected dust samples and their corresponding soil samples.

<table>
<thead>
<tr>
<th>Dust sample</th>
<th>Location (wrt to LU)</th>
<th>Longitude (E)</th>
<th>Latitude (N)</th>
<th>Corresponding soil sample (s)</th>
<th>Land type</th>
<th>Longitude (E)</th>
<th>Latitude (N)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Salmas (Sa)</td>
<td>North</td>
<td>44°49′10″</td>
<td>38°12′38″</td>
<td>Soil Sa</td>
<td>AAL</td>
<td>45°01′17″</td>
<td>38°13′27″</td>
</tr>
<tr>
<td>Jabal (Jab)</td>
<td>Northwest</td>
<td>45°01′34″</td>
<td>37°51′44″</td>
<td>Soil Jab</td>
<td>Sa-Sh</td>
<td>45°03′22″</td>
<td>37°50′36″</td>
</tr>
<tr>
<td>Merang (Mer)</td>
<td>West</td>
<td>45°09′14″</td>
<td>37°47′35″</td>
<td>Soil Mer</td>
<td>Sa-SC</td>
<td>45°07′47″</td>
<td>37°49′24″</td>
</tr>
<tr>
<td>Miandoab (MD)</td>
<td>South</td>
<td>45°58′04″</td>
<td>37°03′58″</td>
<td>Soil MD</td>
<td>SC-CF</td>
<td>45°46′44″</td>
<td>37°06′49″</td>
</tr>
</tbody>
</table>

AAL: abandoned agricultural lands; Sa-Sh: sand sheets; Sa-SC: sandy salt crusts; SC-CF: salt crusts-clay flats.

Table 2. Physicochemical properties of the studied natural dust samples and surface-collected soil samples.

<table>
<thead>
<tr>
<th>Sample</th>
<th>EC (dS m⁻¹)</th>
<th>TC (%)</th>
<th>OM (%)</th>
<th>pH</th>
<th>Particle size classes (&lt; 2 mm)</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Clay (≤ 2 µm)</td>
<td>Silt (2–50 µm)</td>
</tr>
<tr>
<td>Soil Sa</td>
<td>0.9</td>
<td>21.1</td>
<td>5.3</td>
<td>8.2</td>
<td>15.8</td>
<td>20.2</td>
</tr>
<tr>
<td>Dust Sa</td>
<td>27.5</td>
<td>14.4</td>
<td>2.6</td>
<td>7.5</td>
<td>5.7</td>
<td>63.2</td>
</tr>
<tr>
<td>Soil Jab</td>
<td>35.3</td>
<td>28.5</td>
<td>1.0</td>
<td>8.6</td>
<td>1.8</td>
<td>6.9</td>
</tr>
<tr>
<td>Dust Jab</td>
<td>43.9</td>
<td>43.3</td>
<td>1.3</td>
<td>8.3</td>
<td>1.1</td>
<td>2</td>
</tr>
<tr>
<td>Soil Mer</td>
<td>40</td>
<td>19.1</td>
<td>3.3</td>
<td>8.2</td>
<td>9.3</td>
<td>59.2</td>
</tr>
<tr>
<td>Dust Mer</td>
<td>19.9</td>
<td>17.1</td>
<td>2.9</td>
<td>8.2</td>
<td>3.0</td>
<td>10.2</td>
</tr>
<tr>
<td>Soil MD</td>
<td>16.3</td>
<td>13.1</td>
<td>1.2</td>
<td>8.1</td>
<td>3.0</td>
<td>13.1</td>
</tr>
<tr>
<td>Dust MD</td>
<td>4.9</td>
<td>13.5</td>
<td>1.7</td>
<td>8.0</td>
<td>3.9</td>
<td>15.6</td>
</tr>
</tbody>
</table>

EC: electrical conductivity; TC: total carbonates; OM: organic matter.
are the dominant salts in the sediments of LUP. Concentrations of 0.01, 0.1, and 0.4 M were chosen to cover the salinity range of the LUP samples. To investigate the combined effect of soluble salts and pH on the IN activity of quartz and microcline, each salt solution was adjusted to pH 5 (acidic soils without carbonates), pH 7 (neutral), and pH 8 (representative of carbonate soils) by adding small portions of dilute NaOH or HCl solutions. Figure 2 illustrates the combinations of salt type, salt concentration, and pH used to perform the immersion freezing experiments with the reference microcline and quartz samples.

2.7 Freezing experiments

To characterize the average ice nucleation efficiency of both natural and treated soil and dust samples, emulation freezing experiments with a differential scanning calorimeter (DSC) Q10 from TA instruments were performed. The < 63 μm fraction was suspended in water (molecular bioreagent water, Sigma-Aldrich) at concentrations of 2 wt% and 5 wt% and was sonicated for 5 min in order to minimize particle aggregation. Emulsions were prepared by combining the suspensions with a mixture of mineral oil and lanolin (both Sigma-Aldrich) at a ratio of 1:4 and by stirring the mixture with a rotor–stator homogenizer (Polytron PT 1300D with a PT-DA 1307/2EC dispersing aggregate) for 40 s at 7000 rpm. For DSC measurements, 5 to 10 mg of emulation were placed in an aluminum pan, and the pan was hermetically sealed. Thermograms were registered at a rate of 1 K min⁻¹ in the temperature ranges of freezing and melting and evaluated in terms of the onset temperatures of the heterogeneous ($T_{het}$) and homogeneous freezing peaks ($T_{hom}$), the heterogeneously frozen fraction ($F_{het}$), and the melting temperature $T_{melt}$, as has been explained in more detail in Kumar et al. (2018) and Klumpp et al. (2022). The stability of the emulsions was tested for some samples by subjecting them to three freezing cycles, with the first and third cycles performed with a cooling rate of 10 K min⁻¹ as control cycles, following the procedure introduced by Marcolli et al. (2007). Before each experiment, emulsions were freshly prepared. Experiments were repeated at least once with a freshly prepared suspension. The average precision in $T_{het}$ is ±0.2 K. Uncertainties in $F_{het}$ are, on average, ±0.05 but may be much larger when heterogeneous freezing signals are weak or overlap (forming a shoulder) with the homogeneous freezing signal.

The same procedure was used to investigate the IN activity of the reference minerals (microcline with 2 wt% and quartz with 1 wt% concentrations) in combination with different solute compositions, as described in Sect. 2.6.
Figure 2. Statistical design of solution composition to investigate the role of salt type, salt concentration, and pH on the IN activity of microcline and quartz samples. The experiments were performed for suspension concentrations of 2 wt % for microcline and 1 wt % for quartz in solutions of the three salts NaCl, CaCl₂, and MgCl₂, with each salt at the three indicated concentrations and with each concentration at three different pH values.

3 Results and discussion

3.1 Treated natural soil and dust samples

To investigate the impact of soluble salts, carbonates, and organic matter on the IN activity in the immersion mode of the soil and dust samples Sa, Jab, Mer, and MD, we removed salts (SR treatment), carbonates (CR treatments), and organic matter (OMR treatments) from the samples in a systematic way, as illustrated in Fig. 1. DSC thermograms of the treated samples at suspension concentrations of 2 wt % and 5 wt % were recorded and are displayed together with the thermograms of the untreated samples in Fig. 3 (dust samples) and Fig. A1 in Appendix A (soil samples). T₁₀₀ and F₁₀₀ retrieved from the natural and treated samples are summarized in Table 4. In Figs. 4 and 5, the effects of different treatments – given as \( \Delta T_{\text{het}}(\text{treated-natural}) \) (\( T_{\text{het}} \) of treated sample minus \( T_{\text{het}} \) of natural sample) and \( F_{\text{het}}(\text{treated/natural}) \) (\( F_{\text{het}} \) of treated sample divided by \( F_{\text{het}} \) of natural sample) – of each investigated dust and soil sample at 5 wt % concentrations are presented. Triangle plots of \( \Delta T_{\text{het}}(\text{treated-natural}) \) and \( F_{\text{het}}(\text{treated/natural}) \) for 2 wt % concentrations are shown in Figs. A2 and A3 in Appendix A. Finally, Fig. 6 summarizes the effects of all treatments.

3.1.1 Salt-removed (SR) samples

DSC thermograms of the samples after SR treatment are shown in Figs. 3 and A1 in Appendix A in a dark orange color. For almost all samples, removing soluble salts increased the IN activity for both suspension concentrations (see Figs. 4, 6a, and A2 in Appendix A): \( T_{\text{het}} \) increased by \( \Delta T_{\text{het}}(\text{SR-natural}) = 1.0 \text{ K} \) from a mean value of 247.8 to 248.8 K (Table 4). In all samples, with the exception of Soil Sa, at both 5 wt % and 2 wt % concentrations (Figs. 5 and A3 in Appendix A), \( F_{\text{het}} \) increased after SR treatment (see Fig. 6b). The highest \( F_{\text{het}}(\text{SR/natural}) \) value (1.69) was observed for Dust Jab at 5 wt % concentration (\( F_{\text{het}} = 0.78 \) after SR versus 0.46 in the natural sample), which is the sample with the highest electrical conductivity (EC) of 43.8 dS m⁻¹ (Table 2). The smallest difference in IN activity before and after SR treatment was observed for Soil Sa, the sample with the lowest soil EC (0.85 dS m⁻¹). Therefore, removing salts from this sample had almost no or even a negative effect on both \( T_{\text{het}}(\Delta T_{\text{het}}(\text{SR-natural}) = -0.16 \text{ K} \) as the average of 5 wt % and 2 wt % samples) and \( F_{\text{het}}(F_{\text{het}}(\text{SR/natural}) = 1.0) \).

In Part 1 of the manuscript (Hamzehpour et al., 2022), we showed that there was a significantly negative correlation between EC as an indicator of the presence of soluble salts and \( T_{\text{het}} \) and \( F_{\text{het}} \) of the 5 wt % samples (−0.68 and −0.75, respectively), implying that high soluble salt content had a negative influence on IN activity. The increase in \( T_{\text{het}} \) and \( F_{\text{het}} \) after SR treatments confirms the inhibiting effect of soluble salts on the IN activity of the investigated soil and dust samples at these concentrations.

3.1.2 Carbonate-removed (CR) samples

DSC thermograms of the samples after carbonate removal are shown in Figs. 3 and A1 in Appendix A in a black color. Similar to SR treatment, in almost all of the samples, the CR treatment increased the IN activity for both concentrations (Figs. 4, 6a, and A2 in Appendix A A). \( T_{\text{het}} \) increased, on average, by 1.2 K after carbonate removal (Table 4). The highest \( \Delta T_{\text{het}}(\text{CR-natural}) \) was measured for Soil Sa (4.1 K for 2 wt % concentration), while the lowest one occurred in Soil MD, with a slightly negative value of −0.2 K at 2 wt % concentration (Fig. A1 in Appendix A). This sample also exhibits one of the lowest total carbonate contents (TC from back titration (Table 2): 13.1 %; from mineralogical composition (Table 3): 13.1 %; from TGA-MS (Table 5): 11.95 %).

The CR treatment led to \( F_{\text{het}}(\text{CR/natural}) \) > 1 for all samples (Figs. 5, 6b, and A3 in Appendix A). The highest \( F_{\text{het}}(\text{CR/natural}) \) was observed in Soil Jab at 2 wt % concentration (\( F_{\text{het}}(\text{CR/natural}) = 1.78) \), which exhibits one of the highest TC contents (28.5 %), and the lowest value belonged to...
Figure 3. DSC thermograms of treated dust samples with 5 wt % (a, c, e, g) and 2 wt % (b, d, f, h) concentrations in comparison with the untreated samples. SR: salt removed; CR: carbonate removed; OMR: organic matter removed; SR + CR: salt and carbonate removed; SR + OMR: salt and organic matter removed; SR + CR + OMR: salt, carbonate, and organics removed. $T_{\text{het}}$ in Kelvin and $F_{\text{het}}$ in fraction are displayed directly on the curves.
Dust MD at 2 wt % (r_{het(CR/natural)} = 0.99), which has one of the lowest TC contents (13.5 % in Table 2).

In Part 1 (Hamzehpour et al., 2022) of the manuscript, we showed that \( T_{het} \) of the natural samples are negatively correlated to TC and pH. The increase in \( T_{het} \) after carbon removal confirms this negative correlation.

### 3.1.3 Salt- and carbonate-removed samples (SR + CR)

DSC thermograms of the samples after combined salt and carbonate removal are shown in Figs. 3 and A1 in Appendix A in a blue color. \( T_{het(SR+CR)} \) was, on average, 249.2 K, i.e., 1.4 K higher than for the natural samples (Table 4). Thus, the increase in \( T_{het(SR+CR)} \) after the combined treatment is higher than that of each single treatment (Fig. 6a). Soil Sa exhibited the highest increase in \( T_{het} \) after combined salt and carbonate removal, with \( \Delta T_{het(SR+CR/natural)} = 3.8 \) K for the 5 wt % sample and 3.5 K for 2 wt % (Fig. A1 in Appendix A), although its EC is negligible (0.9 dS m\(^{-1}\)) and TC (21.1 %) is average (Table 2). Comparison with \( T_{het} \) after carbonate removal reveals that this treatment alone also led to a strong increase in \( T_{het} \) by \( \Delta T_{het(CR/natural)} = 1.9 \) K for the 5 wt % and even 4.1 K for the 2 wt % sample. Thus, we identify a strong interference of carbonates with the IN-active species in this sample.

\( F_{het} \) increased due to salt and carbonate removal for all samples (Fig. 6b). The highest \( r_{het(SR+CR/natural)} \) was observed in Dust Jab (Fig. 4), with a value of 1.9 K for the 5 wt % sample. Simultaneously, \( T_{het} \) increased by \( \Delta T_{het(SR+CR/natural)} = 2.2 \) K for the 5 wt % sample and even by 2.8 K for the 2 wt % sample. Indeed, Dust Jab exhibits the highest EC (43.9 dS m\(^{-1}\)) and TC values (43 %) of all samples and the second highest pH value (8.3). This again confirms that the presence of both carbonates and soluble salts reduce IN activity.

### 3.1.4 Organic-matter-removed samples (OMR)

DSC thermograms taken after organic matter removal, given in a red color in Figs. 3 and A1 in Appendix A, show that this treatment reduced the freezing onset temperatures of most samples (blue-colored region of the triangles in Figs. 4, 5, A2, and A3 in Appendix A). For the 5 wt % concentration, \( \Delta T_{het(OMR/natural)} \) was negative for all samples, with an average decrease of 1.5 K. For the 2 wt % samples, \( \Delta T_{het(OMR/natural)} \) ranged between −2.5 and 1 K, with an average of \( \Delta T_{het(OMR/natural)} = −0.7 \) K. Soil Sa, which exhibited the strongest shift to lower temperatures after organics removal (\( \Delta T_{het(OMR/natural)} = −2.9 \) K), had the highest organic matter content among all studied samples (5.28 %). This demonstrates that freezing occurring at the highest temperatures strongly depends on the presence of organic matter and confirms the positive correlation (correlation coefficients \( cc = 0.79 \) for 5 wt % and \( cc = 0.44 \) for 2 wt % samples) between organic matter and the freezing onset temperatures that had been reported in Part 1 (Hamzehpour et al., 2022). The dependence of \( T_{het} \) on the presence of organic matter is also in agreement with various studies, which showed that the IN activity of collected soil dust samples decreases when the samples are treated with heat or hydrogen peroxide to remove organics from soil dust samples (Conen et al., 2011; Tobo et al., 2014; Hill et al., 2016).

Although the correlation coefficient between \( F_{het} \) and OM was significantly positive (0.77 for 5 wt % and 0.74 for 2 wt % concentration, Hamzehpour et al., 2022), \( F_{het} \) did not decrease after organics removal with \( r_{het(OMR/natural)} = 1.06 \) for 5 wt % and 1.04 for 2 wt % samples. For both concentrations, almost half of the samples showed an increase in \( r_{het(OMR/natural)} \) (Figs. 5 and 6b). In Soil and Dust Mer, \( F_{het} \) increased due to OMR treatments (Fig. A1 in Appendix A) despite their rather high OM content (3.32 % and 2.89 %, respectively). Therefore, it seems that the total content of organic matter is not the determining factor for the heterogeneously frozen fraction of the natural samples. Moreover, organic molecules may also have a hampering effect on the IN activity of the minerals, e.g., through adsorption on the mineral surfaces and blocking of IN-active sites (Kleber et al., 2021).

The combined removal of soluble salts and organic matter shows similar trends in IN activity as the removal of OMR alone, though these are less pronounced. For the 5 wt % samples, \( T_{het} \) decreased, on average, by −1.3 K after combined SR + OMR treatment compared with −1.5 K for the OMR

<table>
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<tr>
<th>Conc. (wt %)</th>
<th>Natural</th>
<th>SR</th>
<th>CR</th>
<th>OMR</th>
<th>SR + CR</th>
<th>SR + OMR</th>
<th>SR + CR + OMR</th>
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<tr>
<td>( T_{het}(K) )</td>
<td>2</td>
<td>247.4 ± 1.4</td>
<td>248.6 ± 0.9</td>
<td>249.0 ± 1.7</td>
<td>249.1 ± 1.4</td>
<td>247.4 ± 1.3</td>
<td>247.7 ± 0.83</td>
</tr>
<tr>
<td>5</td>
<td>248.2 ± 1.3</td>
<td>248.9 ± 1.1</td>
<td>249.0 ± 1.6</td>
<td>246.6 ± 1.1</td>
<td>249.3 ± 2.2</td>
<td>246.9 ± 1.3</td>
<td>248.6 ± 0.21</td>
</tr>
<tr>
<td>Mean</td>
<td>247.8</td>
<td>248.8</td>
<td>249.0</td>
<td>246.7</td>
<td>249.2</td>
<td>247.1</td>
<td>248.1</td>
</tr>
<tr>
<td>( F_{het}(%) )</td>
<td>2</td>
<td>0.62 ± 0.11</td>
<td>0.74 ± 0.05</td>
<td>0.77 ± 0.07</td>
<td>0.64 ± 0.06</td>
<td>0.79 ± 0.04</td>
<td>0.70 ± 0.09</td>
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<tr>
<td>5</td>
<td>0.63 ± 0.12</td>
<td>0.77 ± 0.06</td>
<td>0.78 ± 0.06</td>
<td>0.64 ± 0.1</td>
<td>0.80 ± 0.05</td>
<td>0.64 ± 0.1</td>
<td>0.74 ± 0.06</td>
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<tr>
<td>Mean</td>
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<td>0.755</td>
<td>0.775</td>
<td>0.640</td>
<td>0.795</td>
<td>0.670</td>
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Figure 4. Effects of treatments in terms of $\Delta T_{\text{het}}$ (treatment/natural) for dust and soil samples at 5 wt% concentration. SR: salt removed; CR: carbonate removed; OMR: organic matter removed; SR + CR: salt and carbonate removed; SR + OMR: salt and organic matter removed; SR + CR + OMR: salt-, carbonate-, and organics-removed treatments.

Figure 5. Effects of treatments in terms of $r_{\text{het}}$ (treatment/natural) for soil and dust samples at 5 wt% concentration. SR: salt removed; CR: carbonate removed; OMR: organic matter removed; SR + CR: salt and carbonate removed; SR + OMR: salt and organic matter removed; SR + CR + OMR: salt-, carbonate-, and organics-removed treatments.
The correlation of $T_{\text{het}}$ with organic matter content and the decrease of $T_{\text{het}}$ after $\text{H}_2\text{O}_2$ digestion are strong evidence that the minor fraction of organics present in the soil and dust samples dominates freezing onset temperatures. To gain further insight into the chemical composition of the organic matter present in the samples, we performed TGA-MS coupled with DSC. The thermograms show a stepwise decrease of mass (Fig. 7) that we can attribute to $\text{H}_2\text{O}$ or $\text{CO}_2$ loss, as registered by the concurrently measured mass spectra. Furthermore, we can attribute mass loss to exothermic and endothermic processes based on the sign of the heat flow registered by DSC (Table 5, Fig. A4 in Appendix A).

Figure 6. (a) Effect of treatments on $T_{\text{het}}$. Points above 0 (shown as dashed lines) represent the increase in $T_{\text{het}}$ after the treatment. (b) Relative change in $F_{\text{het}}$ after the treatments. Points above 1 show an increase in $F_{\text{het}}$ of the treated samples compared to the natural samples.
Overall, OC determined through TGA-MS is similar and correlates well ($cc = 0.93$) with OM determined through wet oxidation, as displayed in Table 2. According to Table 5, the ratio of EO-OS and TS-OC varies between samples. Soil Sa and Soil Mer had the highest total organic carbon (TOC) content among samples; however, the easily oxidizable fraction was highest in Soil Sa (1.42 %), while the thermally more stable fraction was largest in Soil Mer (1.91 %). Overall, IN activity of the soil and dust samples correlates better with the easily oxidizable organic fraction ($cc = 0.47$ with $T_{het}$ and 0.61 with $F_{het}$ as the average of 2 wt % and 5 wt % samples) than with the thermally more stable organics ($cc = 0.22$ with $T_{het}$ and 0.28 with $F_{het}$). Therefore, carbohydrates such as cellulose and proteinaceous molecules are the more likely contributors to the IN activity of the investigated dust and soil samples than the thermally more stable organics like lignin. Interestingly, the positive correlation between EO-OC increases after carbonate removal to $cc = 0.83$ with $T_{het}$ and 0.65 with $F_{het}$, pointing to an inhibiting effect of carbonates on the IN activity of these organics. Such an inhibiting effect seems to be especially strong for Soil Sa, with $\Delta T_{het}((CR)\_natural) = 1.9$ K and $\Delta T_{het}((SR+CR)\_natural) = 3.8$ K for the 5 wt % and $\Delta T_{het}((CR)\_natural) = 4.1$ K and $\Delta T_{het}((SR+CR)\_natural) = 3.5$ K for the 2 wt % concentration. This sample exhibits a strong decrease of $T_{het}$ after OMR treatment, which indicates that its onset freezing temperature is determined by the organic fraction. Thus, as the IN activity of the organic material is reduced in the presence of carbonates, there seem to be interactions between these species.

3.1.5 Salt-, carbonate-, and organic-matter-removed samples (SR + CR + OMR)

DSC thermograms of the samples after SR + CR + OMR treatments are shown in Figs. 3 and A1 in Appendix A in a purple color. In Figs. 4, 5, A2, and A3 in Appendix A, the centers of the triangles represent this treatment. With a value of 248.1 K, the mean $T_{het}((SR+CR)+OMR)$ is only 0.35 K higher (Table 4) than the mean $T_{het}(natural)$ (247.8 K) but lower than $T_{het}$ after SR, CR, and SR + CR treatments. Yet, for the individual samples, the trends are diverse: for three samples (Soil Sa, Soil Mer, and Soil MD), $\Delta T_{het}((SR+CR)+OMR)$ is negative at both concentrations (2 wt % and 5 wt %); for Dust MD, it is negative for 2 wt % and slightly positive for 5 wt % suspensions; for the other samples, it is positive at both concentrations. The highest decrease compared to the natural sample was in Soil Sa (5 wt %), with a 2.4 K reduction (Figs. 4 and A1 in Appendix A). The highest increase in $T_{het}$ was for the 2 wt % sample of Dust Jab, with $\Delta T_{het}((SR+CR)+OMR) = 3.2$ K.

In most samples, $F_{het}$ increased after combined SR + CR + OMR treatments. Yet, for some samples with a negative $\Delta T_{het}((SR+CR)+OMR)$, the decrease in $T_{het}$ was accompanied by a slight decrease in $F_{het}$ (Figs. 3, 6 and A1 in Appendix A). The strongest increase was for the 2 wt % sample of Soil Jab, with $r_{het}((SR+CR)+OMR) = 1.62$, and for the 5 wt % sample of Dust Jab, with $r_{het}((SR+CR)+OMR) = 1.61$. This strong increase is in
Tables and captions

Table 5. Weight loss in percent, quantified with TGA-MS coupled with DSC thermograms in terms of different volatile and oxidizable fractions.

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<tbody>
<tr>
<td></td>
<td>H2O</td>
<td>EO-OC</td>
<td>TS-OC</td>
<td>TOC</td>
<td>CaCO3</td>
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<tr>
<td>Soil Sa</td>
<td>1.56</td>
<td>1.42</td>
<td>1.16</td>
<td>2.58</td>
<td>18.95</td>
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<tr>
<td>Soil Jab</td>
<td>0.43</td>
<td>0.17</td>
<td>0.51</td>
<td>0.68</td>
<td>26.07</td>
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<tr>
<td>Soil Mer</td>
<td>0.9</td>
<td>0.44</td>
<td>1.91</td>
<td>2.35</td>
<td>12.52</td>
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<tr>
<td>Soil MD</td>
<td>0.89</td>
<td>0.14</td>
<td>0.44</td>
<td>0.58</td>
<td>11.95</td>
</tr>
<tr>
<td>Dust Sa</td>
<td>1.12</td>
<td>0.61</td>
<td>0.89</td>
<td>1.5</td>
<td>10.64</td>
</tr>
<tr>
<td>Dust Jab</td>
<td>0.72</td>
<td>0.39</td>
<td>0.92</td>
<td>1.31</td>
<td>64.91</td>
</tr>
<tr>
<td>Dust Mer</td>
<td>0.78</td>
<td>0.41</td>
<td>1.47</td>
<td>1.88</td>
<td>16.34</td>
</tr>
<tr>
<td>Dust MD</td>
<td>1.19</td>
<td>0.27</td>
<td>0.73</td>
<td>1</td>
<td>10.45</td>
</tr>
</tbody>
</table>


accordance with their high EC and TC values together with their rather low organic matter content. These samples exhibit both rather high microcline and quartz contents, such that the positive effects of removing salt and carbonates could have compensated the negative effect of organics removal, resulting in a net increase in heterogeneously frozen fractions.

Overall, the negative correlation between $T_{het}$ and microcline in natural samples (cc of $-0.46$ and $-0.28$ for 2 wt% and 5 wt% samples, respectively) reverted to a slightly positive one after SR + CR + OMR treatment (cc of 0.30 and 0.32 for 5 wt% and 2 wt% samples, respectively). For quartz, the negative correlation with $T_{het}$ also reverted to a positive one for the 5 wt% samples (cc = 0.24) but remained negative for the 2 wt% samples. Moreover, the significantly positive correlation between $T_{het}$ and the total clay mineral content (cc = 0.79) reversed to negative (cc = $-0.22$) for the 5 wt% samples and remained only slightly positive for the 2 wt% samples (cc = 0.17) but with a much lower value than for the natural samples (cc = 0.65). Interestingly, for OMR treatment alone, the correlation between $T_{het}$ and clay minerals was almost as high as for the natural sample (cc of 0.62 and 0.71 for 5 wt% and 2 wt%, respectively), indicating that only the combined SR + CR + OMR treatment destroys the positive correlation. This again shows how intricate the interactions between the different soil and dust components are and that the observed IN activity is the product of all these interactions.

3.2 Effect of salt, salt concentration, and pH on IN activity of microcline and quartz

To substantiate further the impeding effect of soluble salts and carbonates on the IN activity of microcline and quartz in the natural samples, we performed freezing experiments with 2 wt% suspensions of microcline and 1 wt% suspensions of quartz reference samples in salt solutions of NaCl, CaCl$_2$, and MgCl$_2$ to mimic the saline environment in LUP. Moreover, we also varied pH and analyzed the thermograms with respect to $T_{het}$, $F_{het}$, and the melting point depression, as summarized in Figs. 8 and 9.

Salt concentrations of 0.01, 0.1, and 0.4 M were selected, as they comprise the salinity range of the soil and dust samples. The lowest concentration is representative for samples with EC lower than 1 dS m$^{-1}$ (Soil Sa and samples after salt removal), 0.1 M for samples with EC values of around 10 dS m$^{-1}$, and 0.4 M for samples with the highest salinity of around 40 dS m$^{-1}$ (Soil and Dust Jab); pH values were chosen to be 5 to represent the acidic conditions in soils in the absence of carbonates (samples after carbonate removal), pH 7 as the typical value for slightly carbonaceous conditions, and pH 8 to represent the basic environment in calcareous soils with a high content of carbonates (i.e., all samples before carbonate removal).

Results for $F_{het}$ and $T_{het}$ of the microcline and quartz samples, depending on the solution composition, are presented in Fig. 8 together with the reference value for microcline and quartz in pure water. Based on these experiments, the IN activity of both quartz and microcline decreases with increasing concentrations of all three alkali halides. With increasing salt concentration, a decrease in freezing onset temperatures as a function of water activity, as described by the “water-activity criterion” (Koop et al., 2000; Zobrist et al., 2008), is expected and can be derived from the melting point depression as $T_{mel}(a_w) = T_{mel}(a_w + \Delta a_{w,het})$. Figure 9 shows that the melting points of the salt solutions decrease to slightly below 271 K for the highest investigated salt concentration of 0.4 M. Table 6 compares the measured melting point depression $T_{mel}$ (DSC) with the ones based on water activities calculated with AIOMFAC, $T_{mel}(a_w)$ (Zuend et al., 2008, 2011). From these data, the freezing point depression $\Delta T_{het}(a_w)$ that is expected in the absence of specific interactions between the IN-active surface and the solute (Klumpp et al., 2022), which are given in Fig. 8 as blue dashed lines, can be derived. Comparison of the calculated freezing point depression with the actual reduction in onset freezing temperature shown in Fig. 8 shows a larger freezing...
Figure 8. (a) Heterogeneous freezing temperatures ($T_{het}$) and (b) $F_{het}$ of microcline (2 wt %) and quartz (1 wt %) in aqueous solutions of the salts NaCl (triangles), CaCl$_2$ (squares), and MgCl$_2$ (circles) at the salt concentrations 0.01, 0.1, and 0.4 M, and for pH 5 (green), pH 7 (blue), and pH 8 (red). Yellow diamonds represent $T_{het}$ and $F_{het}$ values for pure microcline (2 wt %) and quartz (1 wt %) in Milli-Q water. The blue dashed line in panel (a) represents the freezing temperature according to the water activity criterion; in panel (b), it marks the frozen fractions of the minerals in pure water.

A decrease of freezing temperatures in microcline samples with increasing salt concentrations that exceeds the prediction based on the water-activity criterion is in agreement with findings from ice nucleation studies with K-feldspars in the presence of monovalent cations (Kumar et al., 2018; Whale et al., 2018; Yun et al., 2020). Figure 8 shows now that divalent cations have a similar effect as monovalent ions. These findings are supported by Whale (2022), who recently showed that MgCl$_2$ has a similar effect as monovalent cations on ice nucleation by feldspar. For the investigated concentration and pH range, the cation concentration is more influential for $T_{het}$ of microcline than the nature of the cation. At pH 8, the freezing onset temperatures are only slightly reduced compared with neutral conditions. Yet, this decrease may amplify with increasing pH. For quartz, $T_{het}$ is less influenced by the solution concentration than in the case of microcline, but it is more sensitive to the nature of the solute, with MgCl$_2$ inducing a stronger shift to colder temperatures than NaCl and CaCl$_2$. Moreover, for the investigated pH range, the dependence of $T_{het}$ on pH is stronger for quartz than for microcline.
The decrease in IN activity of quartz with increasing pH is in accordance with findings of Kumar et al. (2019a) and is explicable by the increasing quartz dissolution with increasing pH, which destroys nucleation sites. Figure 8b shows that increasing the solute concentration also affects $F_{\text{het}}$ of microcline. Here, the valence of the cation seems to play a role, as the decrease in $F_{\text{het}}$ is stronger in the presence of the divalent Ca$^{2+}$ and Mg$^{2+}$ ions than for Na$^{+}$. In the case of Na$^{+}$, concentrations up to 0.1 M even lead to an increase in $F_{\text{het}}$, which is in accordance with the enhanced IN activity observed by Perkins et al. (2020) in dilute NaCl solutions. Similar trends are also present for quartz, though they are less pronounced.

Freezing onset temperatures of the samples after the removal of salts, carbonates, and organic matter ($T_{\text{het(SR+CR+OMR)}}$) are on average 248.6 K for the 5 wt % samples and 247.7 K for the 2 wt % samples compared with ($T_{\text{het(OMR)}}$) of 246.7 K for both the 5 wt % and 2 wt % samples after organics removal alone. The reference freezing measurements with quartz and microcline show that, based on the pH and salt content of the LUP samples, the higher freezing temperatures of SR + CR + OMR-treated samples compared with OMR-treated samples is in accordance with the recovered IN activity of microcline and quartz after soluble-salt and carbonate removal. Comparison with the effect of SR + OMR removal shows that the recovery in $T_{\text{het}}$ is mainly due to carbonate removal and indicates a high sensitivity of IN-active sites toward pH. The generally higher $F_{\text{het}}$ of the SR + CR + OMR samples (0.72 as the average of 5 wt % and 2 wt % samples) compared with $F_{\text{het}}$ of OMR samples (0.65) is in a range that can also be ascribed to the recovered IN activity of microcline and quartz after removal of soluble salts and carbonates. These findings, together with the slightly positive correlation of $T_{\text{het}}$ with quartz and microcline content of the samples after SR + CR + OMR treatments, can be considered as sufficient evidence that these minerals influence the heterogeneously frozen fraction and might even dominate freezing onset temperatures in the SR + CR + OMR-treated samples. Yet, the data are less conclusive regarding the respective contributions of either quartz or microcline to the IN activity of the LUP samples. Identifying quartz as a contributor to the IN activity of the SR + CR + OMR-treated samples would be among the first evidence that atmospheric quartz surfaces are sufficiently defectuous to act as a relevant atmospheric ice nucleator. The higher quartz (14.3 %–32 %) than microcline (3.5 %–6.9 %) content would speak for quartz as the dominating INP. Specifically, the quartz concentration in the suspensions varies from 0.3 wt %–0.6 wt % in the 2 wt % samples and from 0.7 wt %–1.6 wt % in the 5 wt % samples. The reference quartz sample (SA) at a suspension concentration of 1 wt % shows $F_{\text{het}} = 0.68$ and even 0.79 for the freshly milled sample (Kumar et al., 2019a). In comparison, the microcline concentration in the suspensions varies from 0.07 wt % to 0.14 wt % for the 2 wt % samples and from 0.18 wt % to 0.35 wt % for the 5 wt % samples, while Kumar et al. (2018) found $F_{\text{het}} = 0.21$ for a 0.2 wt % microcline suspension. The freezing onset temperatures of SR + CR +
Table 6. Water activities ($a_w$) of the salt solutions calculated with AIOMFAC (Zundel et al., 2008, 2010); melting temperatures, $T_{\text{melt}}(a_w)$, calculated as a function of $a_w$ with the parameterization by Koop et al. (2000) in comparison with the measured melting temperatures $T_{\text{melt}}$ (DSC) (average of quartz and microcline samples at all pH values), and corresponding freezing point depression as a function of $a_w$ derived with $T_{\text{het}}(a_w) = T_{\text{melt}}(a_w + \Delta a_{w,\text{het}})$, with $\Delta a_{w,\text{het}} = 0.2$ (Zobrist et al., 2008).

<table>
<thead>
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<th>NaCl</th>
<th>CaCl$_2$</th>
<th>MgCl$_2$</th>
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<tr>
<td>$a_w$</td>
<td>0.9997</td>
<td>0.9966</td>
<td>0.9869</td>
</tr>
<tr>
<td>$T_{\text{melt}}(a_w)$ (K)</td>
<td>273.08</td>
<td>272.8</td>
<td>271.8</td>
</tr>
<tr>
<td>$T_{\text{melt}}$ (DSC) (K)</td>
<td>273.18</td>
<td>272.6</td>
<td>271.3</td>
</tr>
<tr>
<td>$\Delta T_{\text{het}}(a_w)$ (K)</td>
<td>0.04</td>
<td>0.45</td>
<td>1.76</td>
</tr>
</tbody>
</table>

OMR-treated samples are about 3–4 K lower than typical values of microcline emulsions in pure water (251–252 K; Kaufmann et al., 2016; Kumar et al., 2018; Klumpp et al., 2022) but are in the range observed for emulsion freezing experiments with quartz (247–250 K; Kumar et al., 2019a). Thus, the higher quartz concentration and the better agreement of the freezing onset temperatures of the LUP samples with the one of quartz than microcline would speak for quartz as the more relevant INP in the LUP samples. Yet, the evidence is not sufficient to exclude the opposite.

3.3 General discussion and conclusion

The investigated soil and dust samples proved to be complex mixtures of IN-promoting and -inhibiting agents. Through selective removal of some components and correlations between IN activity and mineralogical composition and physical-chemical properties, we have been able to identify the following main actors:

- The relatively small share of organic matter (1 %–5.3 %) provides the INPs that freeze at the highest temperatures in most samples with $T_{\text{het}} = 248.2$ K as the average of 5 wt % and 2 wt % concentrations. We inferred this finding from the reduction in $T_{\text{het}}$ after organic matter removal ($\Delta T_{\text{het}}(\text{OMR-natural}) = -1.1$ K) and from the positive correlation between organic matter content and $T_{\text{het}}$ (cc = 0.68). Conversely, $F_{\text{het}}$ is, on average, the same before and after OMR treatments, which indicates that organic INPs are substituted by mineral INPs present in the samples, which nucleate ice with a similar efficiency but at lower temperatures.

The (bio-)organic INPs seem to be carbohydrates, such as cellulose and/or proteinaceous molecules, rather than aromatic compounds, such as lignins, as evidenced by the better correlation of $T_{\text{het}}$ of the natural samples with the easily oxidizable fraction (250–350°C) than with the thermally more stable fraction (350–550°C) in TGA-MS. This finding is in accordance with studies by Hill et al. (2016) and Paramonov et al. (2018). Moreover, the organic matter seems to be associated with the clay mineral fraction, as evidenced by the strong correlation between total clay mineral content and OM content from wet oxidation (cc = 0.953). Indeed, (bio-)organic molecules are known to adsorb on clay mineral edges and basal surfaces (Kleber et al., 2021).

In most samples, the presence of soluble salts and carbonates seems to hamper the IN activity of the (bio-)organic INPs. This can be seen best for Soil Sa, in which (bio-)organic INPs dominate $T_{\text{het}}$, as evidenced by the strong reduction in the onset freezing temperature after organic matter removal. The freezing onset of this sample shifts to clearly higher temperatures after salt and carbonate removal, showing that these components hampered the IN activity of the organic INPs, either through direct interaction with the solid mineral or via dissolved ions or a combination of both. Some samples show an increase in $F_{\text{het}}$ after organic matter removal, pointing again to a direct interaction between organic matter and minerals, this time in form of an inhibiting effect of organic matter on the IN activity of the minerals. Overall, the IN activity of the organics seems to be influenced by inorganic species and vice versa. Further work is required to investigate to what extent such interactions enhance or inhibit the IN activity of organics and minerals.

- Clay mineral content of the investigated samples ranges from 14 % to 38 % and consists of kaolinite (2.2 %–8.4 %), smectites (0%–10.6 %), polygorskite (3.8 %–12.6 %), and chlorite (3.9 %–16.8 %). This mineral class has proven to be IN active, with onset freezing temperatures of $T_{\text{het}} = 239–242$ K for kaolinite and $T_{\text{het}} = 239–247$ K for montmorillonite in emulsion freezing experiments (Pinti et al., 2012; Kumar et al., 2019b). The other clay minerals present in the LUP samples have not yet been tested in IN experiments, but due to their structural similarity with kaolinite and montmorillonite, they most likely also contribute to IN activity. The significant correlation of total clay mineral content with $T_{\text{het}}$ (cc = 0.72 as the average of 5 wt % and 2 wt % samples) and $F_{\text{het}}$(cc = 0.76) suggests a large contribution of this mineral class to the IN activity of the LUP samples,
though this may also just be a co-correlation paralleling the high correlation between organic matter and IN activity.

After organic matter removal, the correlation of total clay minerals with $T_{\text{het}}$ decreased only slightly ($cc = 0.66$), although $T_{\text{het}}$ exhibited a clear downward shift from a mean value of 247.8 to 246.7 K. Although the mean $F_{\text{het}}$ remained almost the same, its correlation with the clay mineral content decreased drastically but remained positive ($cc = 0.36$). As the correlation of IN activity with all other minerals is even lower, clay minerals indeed seem to be the most dominant IN class after organic matter removal. This is supported by the heterogeneous freezing range (236–248 K) of the LUP samples after OMR removal, which is in general agreement with the freezing temperatures of clay minerals, although it is not specific for them, as other mineral types are IN active within the same temperature range.

- The K-feldspar (microcline) content of the investigated samples ranges from 3.5 %–6.9 %. Microcline has been shown to be a highly efficient INP, with freezing onset temperatures $T_{\text{het}} = 251–252$ K in emulsion freezing experiments (Kumar et al., 2018; Klumpp et al., 2022). Yet, it seems to be irrelevant as an INP in the natural LUP samples, since their microcline content is clearly negatively correlated, with both $T_{\text{het}}$ ($cc = -0.48$ as the average of 5 wt % and 2 wt % samples) and $F_{\text{het}}$ ($cc = -0.51$). Moreover, $T_{\text{het}}$ of most natural samples is too low for microcline suspended in pure water and further decreases after organic matter removal. The irrelevance of K-feldspar as an INP in the investigated samples is opposite to findings from previous freezing experiments with desert dust samples, which concluded that K-feldspars are relevant contributors to the observed IN activity of those samples (O’Sullivan et al., 2014; Boose et al., 2016, 2019; Paramonov et al., 2018).

Only after removal of soluble salts, carbonates, and organic matter, a positive correlation of microcline with $T_{\text{het}}$ ($cc = 0.31$ as the average of 5 wt % and 2 wt % samples) emerged, while the correlation with $F_{\text{het}}$ remained negative ($cc = -0.31$). Moreover, the onset freezing temperatures of the thus-treated samples (mean $T_{\text{het}} = 248.1$ K) remained below the typical freezing onset temperatures of microcline.

Reference emulsion freezing experiments with a microcline suspension in salt solutions covering the concentration and pH range of the LUP samples revealed that both the high salt concentration and pH of the natural samples hampers the IN activity of microcline. The increase in $T_{\text{het}}$ between the OMR- and the SR + CR + OMR-treated samples ($\Delta T_{\text{het}}(\text{SR+CR+OMR}–\text{OMR}) = 1.4$ K as the average of the 5 wt % and 2 wt % samples) suggests that mineral components within the samples can recover their IN activity after salt and carbonate removal. If this mineral component were microcline, the recovery would not be complete, as the onset freezing temperature remained below the values from emulsion freezing experiments with microcline reference samples in pure water.

In addition to the sensitivity of microcline towards soluble salts and pH, its minor concentration in the LUP samples relative to the larger contributions of other IN-active species might have obscured its contribution to IN activity. In other studies, the K-feldspar concentration and its contribution to the total of IN-active species showed more variation between samples. In Paramonov et al. (2018), the K-feldspar fraction varied from 1.9 % to 9.3 %; in O’Sullivan et al. (2014), it varied from 2.1 %–11 %. The microcline concentration in the samples investigated by Boose et al. (2016) ranged between 0 wt % and 3.9 wt %, with the exception of one sample with a microcline content of 30 wt %, which might have been decisive for the positive correlation between IN activity and microcline content.

- Another IN-active mineral present in the LUP samples is quartz, with concentrations ranging from 14 % to 32 %. As the IN activity of quartz has been shown to depend on defects that can be introduced through milling (Zolles et al., 2015; Kumar et al., 2019a), it is unclear to what extent natural quartz samples are IN active. To avoid a high bias in IN activity through milling, we just sieved the soil samples to isolate the < 63 μm particle fraction. In emulsion freezing experiments, freezing onset temperatures of quartz ranged from 247 to 250 K (Kumar et al., 2019a), which coincides with the $T_{\text{het}}$ range of the untreated LUP samples. Yet, the correlations of quartz content with $T_{\text{het}}$ ($cc = -0.37$ as the average of the 5 wt % and 2 wt % samples) and $F_{\text{het}}$ ($cc = -0.39$) are both negative. This is opposite to findings by Boose et al. (2016, 2019), who found a positive correlation between quartz content and the IN activity of desert dust samples; yet, some of their samples had been milled prior to performing freezing experiments.

After removal of soluble salts, carbonates, and organic matter, the correlation of quartz content with $T_{\text{het}}$ ($cc = 0.24$) and $F_{\text{het}}$ ($cc = 0.18$) turned positive for the 5 wt % samples but remained negative for the 2 wt % samples, with correlations of $cc = -0.36$ for $T_{\text{het}}$ and $cc = -0.26$ for $F_{\text{het}}$. Reference emulsion freezing experiments with a quartz suspension with salt concentration and pH in the range of the LUP samples showed that, similar to microcline, the IN activity of quartz is also reduced in this environment. Thus, the recovery of quartz INPs provides a second explanation for the higher average IN activity of SR + CR + OMR-treated samples compared with OMR-treated samples. The higher quartz
than microcline content in the LUP samples together with the better agreement of freezing onset temperatures of reference quartz samples with \( T_{\text{het}}(\text{SR} + \text{CR} + \text{OMR}) \) suggest that quartz contributes more to the IN activity of the LUP samples than microcline; yet, the opposite assumption that microcline is the INP responsible for the increase in IN activity after SR + CR removal from the OMR sample cannot be excluded. Thus, the LUP dataset is not able to answer the question of whether natural quartz is a relevant contributor to IN activity in the atmosphere.

- Soluble salts may be present as crystalline phases and/or amorphous brine. As they are soluble, we can exclude them as INPs, but they may interfere with the IN activity of other INPs. Seven out of the eight LUP samples contain halite (crystallized NaCl) at concentrations from 1.3 % to 5.7 %, which completely dissolves in 5 wt % and 2 wt % aqueous suspensions. The low correlation between halite content and electrical conductivity (\( cc = 0.21 \)) indicates the presence of amorphous salts – likely chlorides, sulfates, carbonates, and bicarbonates of \( K^+ \), \( Ca^{2+} \), and \( Mg^{2+} \), as these are the most abundant ions in LUP sediments (Hamzehpour et al., 2022; Sharif et al., 2018). Dissolved salts are known to decrease freezing temperatures as a function of water activity according to the water-activity criterion (Koop et al., 2000; Zo- brist et al., 2008). Yet, reference freezing experiments with quartz and microcline evidenced a decrease in \( T_{\text{het}} \) and \( F_{\text{het}} \) that exceeds the expected effect of water activity for both minerals. The negative correlation of electrical conductivity with \( T_{\text{het}} \) (\( cc = -0.63 \) as the average of 5 wt % and 2 wt % samples) and \( F_{\text{het}} \) (\( cc = -0.60 \)) further supports the hampering effect that the high salinity has on the IN activity of the LUP samples. As the impact of soluble salts on IN activity depends on the suspension concentration, a higher dilution should reduce it. This can be seen for \( F_{\text{het}} \) of the untreated Dust Jab, which is the sample with the highest electrical conductivity (43.9 dS m\(^{-1}\)); while \( F_{\text{het}} \) of the 2 wt % sample is 0.62, the one of the 5 wt % sample is only 0.49. At an even lower concentration, the effect of soluble salts should decrease even more and might become negligible for immersion freezing in cloud droplets.

- Based on the XRD analysis, total carbonate concentrations vary between 13.1 % and 28.5 % in the LUP samples, with contributions of calcite (9.7 %–13.5 %), dolomite (1.9 %–4.5 %), aragonite (0 %–13.4 %), and magnesite (0 %–4.9 %). As calcite and dolomite did not show any IN activity in emulsion freezing experiments (Kaufmann et al., 2016), we consider the IN activity of all carbonate minerals to be negligible. The irrelevance of carbonates as INPs is further supported by the negative correlation of total carbonates with \( T_{\text{het}} \) (\( cc = -0.23 \) as the average of 5 wt % and 2 wt % samples) and \( F_{\text{het}} \) (\( cc = -0.40 \)). Rather, this negative correlation together with the increase in \( T_{\text{het}} \) (\( \Delta T_{\text{het}}(\text{SR} + \text{CR} + \text{OMR}) = 1.2 \% \)) and \( F_{\text{het}} \) (\( RT_{\text{het}}(\text{SR} + \text{OMR}) = 1.2 \)) after carbonate removal suggest that carbonates inhibit the INPs present in the samples. This hampering effect could occur by occlusion of active sites through direct interaction with the minerals or through release of the ions \( Ca^{2+} \), \( Mg^{2+} \), and \( CO_3^{2-} \) to the solution. Release of \( CO_3^{2-} \) is responsible for the basic pH of the suspensions, which may influence the IN activity of INPs through deprotonation of IN-active sites or by increasing the solubility of minerals – e.g., as is the case for quartz (Kumar et al., 2019b). Reference measurements showed that the IN activity of quartz and microcline is indeed already reduced at pH 8 compared to neutral conditions. Yet, the anticorrelation of IN activity with pH is less pronounced than the one with total carbonates – that is, the correlation of pH with \( T_{\text{het}} \) of the natural samples is only slightly negative (\( cc = -0.25 \) as the average of 5 wt % and 2 wt % samples) but slightly positive with \( T_{\text{het}} \) (\( cc = 0.19 \)). As the freezing onset of the natural samples is dominated by INPs originating from organic matter, the positive correlation of \( T_{\text{het}} \) with pH compared to the negative one with carbonates makes it unlikely that the inhibition of (bio-)organic INPs originates from a pH effect. Rather, it might be due to direct interactions with the minerals, either through occlusion of (bio-)organic INPs from the solution through a cementing effect, by their adsorption on the mineral surface, or through interactions with dissolved \( Ca^{2+} \), which can form inner-sphere and/or outer-sphere complexes with organic matter (Rowley et al., 2018). Further studies are required to elucidate which interactions are most relevant for the reduced IN activity.

In summary, the analysis of the effects that the different treatments exert on the IN activity of the LUP samples revealed complex interactions between organic matter and mineral particles in soils. This study was able to elucidate some of these interactions; yet, more research is required to disentangle how the mixing state of organic and inorganic soil components influence their ability to nucleate ice. While research over the past years established feldspars, quartz, and clay minerals as the most relevant mineral INPs in dust, less is known about the composition of the (bio-)organic matter that enhances the IN activity of fertile soils above the one of mineral dust. More research is required to elucidate the interactions between organic matter and mineral particles to improve our understanding of the origin of high IN activity in fertile soils.
Figure A1. DSC thermograms of treated soil samples with 5 wt % (a, c, e, g) and 2 wt % (b, d, f, h) concentrations in comparison with the untreated samples. SR: salt removed; CR: carbonate removed; OMR: organic matter removed; SR + CR: salt and carbonate removed; SR + OMR: salt and organic matter removed; SR + CR + OMR: salt, carbonate, and organics removed. \( T_{\text{het}} \) in Kelvin and \( f_{\text{het}} \) in fraction are displayed directly on the curves.
Figure A2. Effects of treatments in terms of $\Delta T_{\text{het}}$ (treatment-natural) for dust and soil samples at 2 wt % concentration. SR: salt removed; CR: carbonate removed; OMR: organic matter removed; SR + CR: salt and carbonate removed; SR + OMR: salt and organic matter removed; SR + CR + OMR: salt-, carbonate-, and organic-removed treatments.

Figure A3. Effects of treatments in terms of $rF_{\text{het}}$ (treatment-natural) for dust and soil samples at 2 wt % concentration. SR: salt removed; CR: carbonate removed; OMR: organic matter removed; SR + CR: salt and carbonate removed; SR + OMR: salt and organic matter removed; SR + CR + OMR: salt-, carbonate-, and organic-removed treatments.
Figure A4. TG (green) and DSC (blue) curves of soil and dust samples with corresponding mass data plotted against temperature. Light blue: total ion current of mass 44 (CO₂).
Author contributions. NH conducted the experiments and field works. KK provided equipment and facilities for the laboratory experiments. NH, CM, and TP contributed to the planning and interpretation of the experiments. NH and CM prepared the manuscript, with contributions from TP and KK. DT conducted the TGA analysis.

Competing interests. The contact author has declared that none of the authors has any competing interests.

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Acknowledgements. We thank Michael Plötze for the XRD measurements; we thank Ulrich Krieger and Uwe Weers for their support in the laboratory.

Financial support. Nikou Hamzehpour has been supported by visiting professors grant at ETH. Kristian Klumpp has been supported by the Schweizerischer Nationalfonds zur Förderung der Wissenschaftlichen Forschung (SNF; grant no. 200021_175716).

Review statement. This paper was edited by Xavier Querol and reviewed by two anonymous referees.

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