Theory of ice premelting in porous media

Hendrik Hansen-Goos\textsuperscript{1} and J. S. Wettlaufer\textsuperscript{1,2,3}

\textsuperscript{1}Department of Geology and Geophysics, Yale University, New Haven, CT 06520, USA
\textsuperscript{2}Department of Physics, Yale University, New Haven, CT 06520, USA
\textsuperscript{3}Program in Applied Mathematics, Yale University, New Haven, CT 06520, USA

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Premelting describes the confluence of phenomena that are responsible for the stable existence of the liquid phase of matter in the solid region of its bulk phase diagram. Here we develop a theoretical description of the premelting of water ice contained in a porous matrix, made of a material with a melting temperature substantially larger than ice itself, to predict the amount of liquid water in the matrix at temperatures below its bulk freezing point. Our theory combines the interfacial premelting of ice in contact with the matrix, grain boundary melting in the ice, and impurity and curvature induced premelting, the latter occurring in regions which force the ice-liquid interface into a high curvature configuration. These regions are typically found at points where the matrix surface is concave, along contact lines of a grain boundary with the matrix, and in liquid veins. Both interfacial premelting and curvature induced premelting depend on the concentration of impurities in the liquid, which, due to the small segregation coefficient of impurities in ice are treated as homogeneously distributed in the premelted liquid. Our principal result is an equation for the fraction of liquid in the porous medium as a function of the undercooling, which embodies the combined effects of interfacial premelting, curvature induced melting, and impurities. The result is analyzed in detail and applied to a range of experimentally relevant settings.

I. INTRODUCTION

Due to advances in our understanding of the premelting of ice and other materials over the last fifteen years\textsuperscript{1}, and an increasing appreciation of the influence and extent of this phenomenon in technological, terrestrial and extraterrestrial settings, it is prudent to revisit the detailed thermodynamic basis of its manifestation in porous media where water is invariably found in adsorbed layers or in bulk. In 1992 Cahn, Dash, and Fu (CDF)\textsuperscript{2} developed the original premelting based theory for the liquid fraction $f_l$ of water in a porous medium at sub-freezing temperatures. Given experimental and theoretical advances in the interim, we extend their work in a number of respects. CDF considered a matrix of monodisperse spheres arranged either in a simple cubic packing or in a cubic close packing, regarded as lower and upper bounds for the experimental systems. We design our theory for the \textit{random} close packing of monodisperse spheres, which may be a more realistic description for most applications, e.g. frozen soils, where even a small degree of polydispersity prevents a periodic packing of the matrix particles. The assumed periodic packing of the matrix particles leads CDF to take the grain size of the interstitial ice to be on the order of the size of the matrix lattice unit cell. Given that the degree of polycrystallinity of the interstitial ice has a complicated dependence on the freezing rate [see e.g., Ref.\textsuperscript{3} and references therein] and the sample annealing history, the ice grain size may differ substantially from the matrix particle size. Whence, polycrystallinity of the interstitial ice enters into our theory as a parameter, namely the density of grain boundary surface relative to the interstitial volume. Calculations since CDF showed that the dispersion force contribution to interfacial melting could be repulsive or attractive\textsuperscript{1,4} and thus favor or suppress melting in a manner and strength that depends on the effective dielectric response of the ice/water/matrix interface. Moreover, it has been understood that the great inter-experimental variability in both the interfacial and surface melting of ice was likely due to the sensitivity of the phenomena to impurities which have a rich and complex influence on the melting behavior\textsuperscript{5}. Thus, our theory allows for both attractive and repulsive dispersion forces of arbitrary strength (depending on the matrix material) and includes the effect of charges at the interface, which give rise to an electrostatic force. Finally, most importantly, our work includes the effect of charged impurities in the premelted liquid on both interfacial and curvature induced premelting.

The article is structured as follows. In Section\textsuperscript{II} we lay out the tenets and details of our theory, discussing interfacial premelting and curvature induced premelting in the presence of charged impurities and combine them in order to describe premelting in a porous medium. The final result of the theoretical treatment is an implicit functional equation for the liquid fraction $f_l$, Eq. (21). The first part of Section\textsuperscript{III} is dedicated to the analysis of interfacial premelting as described by Eq. (4), and in the second part Eq. (21) for the liquid fraction $f_l$ is analyzed by extracting exact solutions for certain limiting cases and numerical solution in the general case. The porous media which we consider consist of fused quartz, gold, and silicon. We summarize in Section\textsuperscript{IV}.
II. THEORY

A. Interfacial melting and grain boundary melting: Planar interfaces

In order to calculate the equilibrium thickness $d$ of a premelted layer for a given undercooling $\Delta T = T_m - T$, where $T_m$ is the bulk melting temperature of ice and $T$ is the temperature of the system, we consider a three layer planar system. Layer I is given by the substrate, i.e., in the case of interfacial melting, the layer consists of the material which forms the porous medium, or of ice when grain boundary melting is considered. Layer II is the premelted quasi-liquid layer of thickness $d$ and finally layer III consists of ice. In what follows we consider the complete free energy of the system and then focus on the individual complications and contributions in turn to show how known asymptotic results emerge from the theory.

1. General theory in the presence of impurities

It is assumed that layer II contains a certain amount of impurities. The Gibbs free energy of the system per unit area is given by

$$G(T, P, N_s, N_i, N_l) = \mu_s(T, P) N_s + \mu_l(T, P) N_l + \mu_i(T, P) N_i + R_g T N_i \left[ \ln N_i - 1 \right] + T(d),$$

(1)

where $T$ and $P$ are temperature and pressure, $\mu_s$, $\mu_l$, $\mu_i$ are the chemical potentials per mole of the solid, the liquid, and the impurity species, respectively, $N_s, N_l, N_i$ are the numbers of mole per unit area of the solid, the liquid, and the impurities, respectively. The term $R_g T N_i \left[ \ln N_i - 1 \right]$, where $R_g$ is the gas constant, corresponds to the mixing entropy of the impurities assuming low impurity concentration so that the ideal gas result applies. Finally, $T(d)$ denotes the total effective interfacial free energy described in more detail below. Note that $d$ is directly proportional to $N_i$ through $N_i = \rho_l d$, where $\rho_l$ is the molar density of the liquid which is a constant in the calculation.

The solid (ice) and the liquid (water) parts of the system can exchange material and hence the condition for thermodynamic equilibrium is given by

$$\frac{\partial G}{\partial N_s} = \frac{\partial G}{\partial N_l}.$$  

(2)

Applying this criterion to the free energy given above, and using the fact that $\partial \mu_{s,l}/\partial T \equiv -s_{s,l}$ where $s_{s,l}$ is the specific entropy of the solid (liquid), we can approximate $\mu_l - \mu_s \approx q_m \frac{T_m}{d}$, where $q_m$ is the latent heat of melting per mole and arrive at

$$\rho_l q_m \frac{\Delta T}{T_m} \frac{T_m N_i}{d} = T'(d).$$

(3)

In the absence of interfacial forces we have $T'(d) = 0$, and the result reduces to $\Delta T = \frac{R_g T_m N_i}{q_m \rho_l} \rho_i$, where the molar density of impurities is $\rho_i = N_i/d$. This undercooling corresponds precisely to the classical colligative effect—the depression of the bulk freezing temperature due to dissolved impurities. Obviously, compared to the force free case, repulsive interfacial forces for which $T'(d) < 0$ result in a larger undercooling $\Delta T$ thereby enhancing the layer thickness. Conversely, attractive interfacial forces wherein $T'(d) > 0$ suppress the formation of a premelted liquid film. Indeed, for sufficiently large $\Delta T$, Eq. (3) may have no solution showing that the attractive interfacial forces can dominate the colligative effect of the impurities and the premelted liquid layer will vanish.

2. Contributions to the effective interfacial free energy $T(d)$

The interfacial energy has two contributions: $T(d) = T_{\text{dis}}(d) + T_{\text{elec}}(d)$, where $T_{\text{dis}}(d)$ is due to dispersion forces, acting between the three interfaces defining the system, and $T_{\text{elec}}(d)$ captures the effect of immobile charges at the interface. Although we compute the strength of the dispersion force interaction from the full frequency dependent theory, we take the small thickness limit in which only non-retarded forces contribute to $T_{\text{dis}}(d)$ which consequently can be written as

$$T_{\text{dis}}(d) = -\frac{A_H}{12\pi d^2},$$

(4)

where $A_H$ is the Hamaker constant for the given layered system $[1, 3]$. For symmetric systems, such as the ice-liquid-ice configuration, which occurs in the case of grain boundary melting, $A_H$ is positive and hence dispersion forces are attractive. At the interface between the ice and the porous matrix $A_H$ can assume both signs, depending on the material properties of the porous medium.

Immobile surface charges with a given surface charge density $q_s$ on both interfaces which bound the quasi-liquid layer are screened by the counterions in the liquid. Making use of the Debye-Hückel theory this leads to the contribution

$$T_{\text{elec}}(d) = \frac{2q_s^2 e^{-\kappa d}}{\kappa},$$

(5)

where $\kappa$ is the inverse Debye length for monovalent ions, with the electron charge $e$, Avogadro's number $N_A$, and Boltzmann's constant $k_B$. The Debye length describes the characteristic decay of the ion field adjacent to the charged surface due to the screening by counter ions. A repulsive force between two charged surfaces originates in the restriction.
of the entropy of the ions as the surfaces are brought closer. Whereas ions of charge opposite to those of the surface are attracted to it, they are repelled from each other, and the increased proximity induced by decreasing the film thickness increases the free energy. Note the relatively complex $d$-dependence of $F_{\text{elec}}(d)$ which is due to the fact that $\kappa^{-1}$ is also a function of $d$. Thus, the richness of the role of impurities in the premelting of ice is a consequence of this and the efficient segregation of ions into the film; since nearly all of the impurities remain in the liquid, as the temperature changes the amplitude and range of the repulsive electrostatic interaction varies. In principle, the situation could be even more complex when for example $q_s = q_s(d)$. Indeed, when $d$ is small the Debye-Hückel expression is less accurate and steric effects become more important. However, to introduce such complications here exceeds our understanding of the situation based on present experimental evidence.

Using these expressions for $I_d = F_{\text{dis}}(d) + F_{\text{elec}}(d)$ in Eq. (8) we obtain an implicit equation for the thickness $d$ of the premelted layer as a function of the undercooling $\Delta T$:

$$\rho m \frac{\Delta T}{T_m} = \frac{R_g T_m N_i}{d} - \frac{A_H}{6\pi \rho m} + \frac{q_m^2}{\epsilon_0} \left[ 1 - \frac{1}{c \sqrt{ho m d}} \right] e^{-c \sqrt{\rho m d}},$$

(7)

where the constant $c$ is

$$c = \sqrt{\frac{e^2 N_A}{\epsilon_0 k_BT_m}}.$$ 

(8)

This result is analyzed in detail in Section III A.

### B. Curvature induced melting: The Gibbs-Thomson Effect

In order to assess the effect of the ice-liquid interfacial curvature on the freezing temperature, we neglect the known anisotropy of the surface energy of ice and consider a spherical ice crystal of radius $r$ surrounded by undercooled water. As opposed to the planar case, the interfacial area is no longer a constant as it was in the treatment of Section III A. With this new complication we reconsider the Gibbs free energy $G$ of the total system. The same holds for the numbers of moles $N_s$, $N_l$, and $N_i$ which are now taken for the total system and not per unit surface. For ease of notation we use the same symbols for these new quantities. All other quantities are as defined above. Whence, we have

$$G(T, P, N_s, N_l, N_i) = \mu_s(T, P) N_s + \mu_l(T, P) N_l + \mu_i(T, P) N_i + R_g T N_l \left[ \ln \frac{N_l}{N_i} - 1 \right] + 4\pi r^2 \gamma_{sl},$$

(9)

where $r$ is related to $N_s$ through $N_s = \frac{4}{3} \pi r^3$ with the constant molar density $\rho_s$. We note here that we understand that $\gamma_{sl} = \gamma_{sl}(r, N_i, \phi)$, where $\phi$ is the angle that defines the surface normal. Thus, the surface energy, itself a Gibbs free energy conjugate to area, is known in general to depend on the size of the system, the adsorption of impurities and the crystallographic orientation but the simplification of treating it as a constant is justified both on the basis of experimental evidence for the first two dependencies and on the level of our analysis.

The condition for thermodynamic equilibrium is again

$$\frac{\partial G}{\partial N_l} = \frac{\partial G}{\partial N_i},$$

(10)

and hence a calculation analogous to that in the Section III A yields the result

$$q_m \frac{\Delta T}{T_m} - R_g T_m \frac{\rho_i}{\rho_l} = \frac{2\gamma_{sl}}{\rho_s r^2},$$

(11)

or, for the general case of a surface with principal radii of curvature $r_1$ and $r_2$,

$$q_m \frac{\Delta T}{T_m} - R_g T_m \frac{\rho_i}{\rho_l} = \gamma_{sl} \left( \frac{1}{r_1} + \frac{1}{r_2} \right).$$

(12)

This result is the well known Gibbs-Thomson effect, i.e. the freezing point depression due to convexity of the ice-liquid interface, calculated here for the case where impurities are present in the liquid water. The larger the undercooling $\Delta T$, the smaller the radii of curvature of an ice-liquid interface in equilibrium. At the bulk freezing point, $\Delta T \gg R_g T_m^2 \rho_i / q_m \rho_l$, the radius of curvature diverges and a planar interface is the stable configuration.

### C. Premelting in a porous medium

We use the random close packing (RCP) of hard spheres as a model for the porous matrix. Apart from choosing the matrix material and hence calculating the properties which enter the determination of the Hamaker constant (see Section III A), the only free parameter characterizing the porous matrix which contains the partially premelted ice is the radius $R$ of the hard spheres. The properties of RCP hard spheres which are required for our calculation are the sphere packing fraction $\eta$ and the average coordination number $C$. Recent estimates of these numbers are $\eta = 0.634$ and $C = 6$. It should be noted, however, that the basic concept of RCP is not well defined mathematically, so these numbers vary slightly according to the protocol and algorithm used to determine the exact configuration.

At a given undercooling $\Delta T$ we are interested in the fraction of the interstitial volume which is occupied by quasi-liquid water $f_l$ which we will more compactly refer to as the liquid fraction. The first contribution we consider comes from interfacial premelting at the surface of the porous matrix. In the limit where the premelted layer is much thinner than the radius of the matrix particles, $d \ll R$, this fraction is well approximated by

$$f_l^{\text{mat}}(\Delta T, \rho_i) = \frac{q_{sl, i}}{1 - \eta} \times \left[ (\Delta T, \rho_i; A_H^{\text{mat}}, q_{sl, i}) \right],$$

(13)
where \( a_{\text{mat}} \) is the surface area of the matrix particles per unit volume and \( d \) is obtained from Eq. (7) using the Hamaker constant \( A_H^{\text{mat}} \) for the ice-water-matrix layered system and the appropriate surface charge density \( q_s^{\text{mat}} \). The sphere number density of the matrix is \( n_s = \eta/(4\pi R^3/3) \) and thus we obtain \( a_{\text{mat}} = 4\pi R^2 n_s = 3\eta/R \) and

\[
f_l^{\text{mat}}(\Delta T, \rho_i) = \frac{3\eta}{1-\eta} \times d(\Delta T, \rho_i; A_H^{\text{mat}}, q_s^{\text{mat}}) R^{-1}. \tag{14}
\]

The second contribution to the liquid fraction is due to grain boundary melting. Its importance depends sensitively on the grain size of the interstitial ice, which is a function of such factors as (1) the rate of cooling, (2) the undercooling \( \Delta T \), and (3) the time allowed for annealing. Even were we to attempt a complete computation of the coarsening rate of the ice grains in the matrix, there is no unique way to specify the initial value problem and hence to arrive at a concrete grain size as a function of the properties of the porous matrix. Thus, we introduce as an additional parameter the grain boundary density \( \rho_{gb} \), which is the surface area of grain boundaries per unit volume of the interstitial ice. We denote \( \tilde{\rho}_{gb} = \rho_{gb} R \) as the related dimensionless density. The corresponding liquid fraction follows immediately

\[
f_l^{\text{gblayer}}(\Delta T, \rho_i) = \tilde{\rho}_{gb} \times d(\Delta T, \rho_i; A_H^{\text{gb}}, q_s^{\text{gb}}) R^{-1}, \tag{15}
\]

where \( A_H^{\text{gb}} \) is the Hamaker constant for the ice-liquid-ice layered system and \( q_s^{\text{gb}} \) is the surface charge density for the grain boundary. The thickness \( d \) of the quasi-liquid layer is again obtained from Eq. (7).

We now deal with curvature induced premelting. Important contributions to the liquid water fraction come from regions where the ice-liquid interface has a large curvature. In our idealized matrix these are the points where two matrix spheres are in contact. Following CDF there is a liquid pocket forming at the contact point with a volume \( 2\pi r^2 R \) to lowest order in \( r \), which is the radius of curvature of the ice-liquid interface (Fig. 1). Taking into account that the second principal radius of curvature is much larger than \( r \), we can use Eq. (12) to relate \( r \) to the undercooling \( \Delta T \):

\[
r(\Delta T, \rho_i) = \frac{\gamma_{st}}{\rho_s} \left( \frac{q_m \Delta T}{T_m} - R_g T_m \rho_i \rho_i \right)^{-1}. \tag{16}
\]

Using the number density \( n_s \) of the sphere matrix and the matrix coordination number \( C \) we can express the corresponding fraction of liquid water with respect to the total interstitial volume:

\[
f_l^{\text{contact}}(\Delta T, \rho_i) = \frac{C/2 \times n_s \times 2\pi r(\Delta T, \rho_i)^2 R}{1-\eta} = \frac{3}{2} \times \frac{\eta}{1-\eta} \times \frac{C}{2} \times \left[ \frac{r(\Delta T, \rho_i)}{R} \right]^2, \tag{17}
\]

with \( r(\Delta T, \rho_i) \) from Eq. (16).

A further curvature induced contribution to the liquid fraction brought out by CDF comes from the lines where a grain boundary and a matrix sphere meet. In the limit that the matrix particles are small on the scale of the linear scale of the ice grains, but that \( R \) is much larger than \( r(\Delta T, \rho_i) \) the corresponding liquid fraction can be calculated using elementary geometry (see Appendix A)

\[
f_l^{\text{gbmat}}(\Delta T, \rho_i) = 3 \left( 2 - \frac{\pi^2}{8} \right) \times \frac{\eta}{1-\eta} \times \tilde{\rho}_{gb} \times \left[ \frac{r(\Delta T, \rho_i)}{R} \right]^2. \tag{18}
\]

Finally, curvature induced premelting gives rise to liquid veins along the lines where several grain boundaries meet. With the simplifying assumption that the ice grains are arranged in a cubic packing, the total length of veins per interstitial volume, or the density of veins \( \rho_{\text{veins}} \), is obtained from \( \rho_{gb} \) as \( \rho_{\text{veins}} = \rho_{gb}^2 / 3 \). Given that the cross-sectional area of a vein in cubic packing is \( (4 - \pi) r^2 \), we can calculate the corresponding contribution to the liquid fraction as

\[
f_l^{\text{veins}}(\Delta T, \rho_i) = \rho_{\text{veins}} \times (4 - \pi) \times \left[ r(\Delta T, \rho_i) \right]^2 = \frac{1}{3} \left( 4 - \pi \right) \times \tilde{\rho}_{gb}^2 \times \left[ \frac{r(\Delta T, \rho_i)}{R} \right]^2. \tag{19}
\]

Therefore, the total liquid fraction is obtained as the sum of the five individual contributions which we have derived above; \( f_l^{\text{mat}}, f_l^{\text{gblayer}}, f_l^{\text{contact}}, f_l^{\text{gbmat}}, \) and \( f_l^{\text{veins}} \).

D. Implicit equation for the total liquid fraction

Thus far we have taken impurities into account in terms of a molar density \( \rho_i \) or, in the case of the pre-
melted layer, a surface density of impurities \( N_i = \rho_i d \). Laboratory probes using scattering or related approaches involve relatively small samples with good thermal control and a well defined (fixed for a given run) dopant level \( \rho_0 \). The temperature changes in these systems are quasi-static so that the sample is at coexistence. In the natural environment there are of course temperature gradients. Therefore, for applicability of our work there, the typical broad temperature gradients found for example in natural soils imply that a local region within will be nearly isothermal and thus isosolene. Clearly, there will be many environmental circumstances where this may be violated, but because the first test of such a theory should be of high precision we focus mainly on the simplest experimental setting and exploit the fact that the impurities are almost exclusively in the liquid fraction of the interstitial water it follows that

\[
\rho_i = \frac{\rho_0}{d},
\]

thereby properly accounting for the increase in the impurity concentration in the premelted liquid as the liquid fraction decreases. Due to the fact that the regions of premelted water are all interconnected, \( \rho_i \) is homogeneous over the entire sample.

Having now considered the essential physical phenomena we can formulate an equation for the total liquid fraction \( f_i^{\text{tot}} \) in the porous medium as a function of the undercooling \( \Delta T \) and the impurity concentration \( \rho_i \) in the completely melted sample. Combining all the contributions discussed in Section IIIC we obtain

\[
f_i^{\text{tot}} = f_i^{\text{mat}}(\Delta T, \rho_0/f_i^{\text{tot}}) + f_i^{\text{layer}}(\Delta T, \rho_0/f_i^{\text{tot}}) + f_i^{\text{contact}}(\Delta T, \rho_0/f_i^{\text{mat}}) + f_i^{\text{globmat}}(\Delta T, \rho_0/f_i^{\text{mat}}) + f_i^{\text{veins}}(\Delta T, \rho_0/f_i^{\text{tot}})
\]

which is an implicit function for \( f_i^{\text{tot}} = f_i^{\text{tot}}(\Delta T, \rho_0) \) and in the general case has no analytical solution. Moreover, the underlying equation (7) for the thickness \( d \) of the premelted layer, which is required for \( f_i^{\text{mat}} \) and \( f_i^{\text{layer}} \), is itself an implicit function of \( d \). This calls for a careful stepwise analysis first of Eq. (4) and subsequently of Eq. (21), which we perform in Sections IIIA and IIIB.

### III. ANALYSIS AND APPLICATION

#### A. Thickness of the premelted layer

In Section IIIA we derived Eq. (2) for the thickness \( d \) of the premelted layer as a function of the undercooling \( \Delta T \), the impurity concentration \( N_i \), the dispersion forces as embodied in the Hamaker constant \( A_H \), and the surface density \( q_s \) of immobile charges on the solid-liquid interfaces. This expression cannot be solved analytically in the general case. Hence, presently we analyze Eq. (4) analytically, to demonstrate that we can obtain known distinguished limits and facilitate intuition, and numerically, to provide specific ranges of results for particular physical systems of interest. We begin by rewriting this equation in terms of the reduced, or dimensionless, undercooling \( t = \Delta T/T_m \) as follows

\[
t = \frac{bN_i}{d} - \frac{a}{d^3} + \frac{1}{c\sqrt{N_i}}e^{-c\sqrt{N_i}d},
\]

where the constants \( b = R_g T_m/(\rho_q m) \), \( a = A_H/(6\pi \rho_q m) \), and \( q = q_s/(\epsilon_0 \rho_q m) \). For attractive dispersion forces which suppress premelting we have \( a > 0 \), and for repulsive dispersion forces which promote premelting we have \( a < 0 \). All other quantities in Eq. (22) are positive.

1. Maximal undercooling and minimal layer thickness

The simplest case occurs when both dispersion and electrostatic forces are absent, i.e., \( a = 0 \) and \( q = 0 \). The thickness \( d \) of the liquid film is then simply given by

\[
d = bN_i t^{-1},
\]

representing the colligative effect of the dissolved impurities which, due to the small segregation coefficient of ice, are strongly enriched in the liquid layer. The situation immediately becomes more complex if we take dispersion forces into account. Let us first consider materials that have \( a > 0 \), which means that dispersion forces suppress the formation of a premelted liquid film. Assuming \( q = 0 \) for the moment, Eq. (22) possesses a solution as long as the undercooling \( t \) does not exceed a critical value \( t_{\text{max}} = 2(6N_i)^{3/2} \), corresponding to a minimum film thickness \( d_{\text{min}} = \sqrt{36\pi/4N_i} \). If the temperature is reduced such that \( t > t_{\text{min}} \) the attractive dispersion forces dominate the colligative effect and the liquid film collapses to zero thickness.

For the most general form of Eq. (22), the maximal undercooling \( t_{\text{max}} \) and corresponding minimal layer thickness \( d_{\text{min}} \) at which the quasi-liquid layer collapses cannot be expressed analytically. We obtain the global maximum \( (d_{\text{min}}, t_{\text{max}}) \) by extremizing the right hand side of Eq. (22) with respect to \( d \). Two limiting cases can be distinguished. For large impurity concentrations \( N_i \) the contribution due to surface charges becomes negligible...
and hence we recover the result

\[ d_{\text{min}}(N_i \to \infty) \to \frac{3a}{b N_i}, \]
\[ t_{\text{max}}(N_i \to \infty) \to \frac{2 (b N_i)^{3/2}}{3 \sqrt{3a}}, \] (24)

for the case \( q = 0 \) which we derived above. On the other hand, it can be shown that in the limiting case \( N_i \to 0 \) the values \( d_{\text{min}} \) and \( t_{\text{max}} \) follow from the surface charge dominated limit \((b = 0 \text{ and } a = 0)\) which has

\[ d_{\text{min}}(N_i \to 0) \to \frac{2.62}{c^2 N_i}, \quad t_{\text{max}}(N_i \to 0) \to 0.0757 \times q. \] (25)

In Fig. 2 we plot the minimal thickness \( d_{\text{min}} \) of the quasi-liquid layer corresponding to the maximal undercooling \( t_{\text{max}} \) as a function of the impurity concentration \( N_i \) for different values of the surface charge \( q_s \). We use a Hamaker constant \( A_H = 3.0 \times 10^{-23} \text{ J} \) which is the result obtained by Wilen et al. [4] for the ice/liquid/fused quartz layered system. The analytical results for \( N_i \to 0 \) and \( N_i \to \infty \) are characterized by the \( N_i \)-dependencies of the asymptotic behavior. Note that the minimal thickness in the dispersion force dominated limit \((q_s = 0)\) is below 0.1 nm and therefore equal to the dimension of a single molecule. Hence, the notion of a quasi-liquid layer can no longer be applied and for the ice/liquid/fused quartz system layer collapse can only be observed experimentally at low impurity concentrations and it is always triggered by surface charges. This is seen directly from Eq. (22) where the contribution of the surface charge for sufficiently small \( d \) becomes negative, indicating attractive forces, which suppress liquid layer formation. In order to observe a collapse of the quasi-liquid layer driven by dispersion forces, the latter have to be more strongly attractive. We have included in Fig. 2 the case of the ice-liquid-gold system which has a Hamaker constant \( A_H = 1.573 \times 10^{-21} \text{ J} \), and hence dispersion forces are indeed strong enough to yield a liquid-layer collapse at thicknesses of the order of those calculated for the surface charge dominated case. The contrast between the two scenarios will be studied further in the second part of this section (Figs. 4 and 5). In Fig. 3 the corresponding results for the maximal undercooling \( \Delta T_{\text{max}} = T_{\text{m}} t_{\text{max}} \) are plotted. As we expect from Eqs. (22) and (23), \( \Delta T_{\text{max}} \) increases with surface charge \( q_s \) resulting in a \( q_s \)-dependent offset for \( N_i \to 0 \). Moreover, for all \( q_s \), \( \Delta T_{\text{max}} \) increases with impurity concentration \( N_i \). A large Hamaker constant, for example of the ice/liquid/gold system, results in a low \( \Delta T_{\text{max}} \) as shown by Eq. (24), where \( \Delta T_{\text{max}} \propto A_H^{1/2} \). There are interesting experimental consequences of Fig. 3. Obviously, measuring \( \Delta T_{\text{max}} \) for just one given impurity concentration \( N_i \) is not sufficient to determine the surface charge \( q_s \) at the interface. This follows from the fact that any two curves intersect, and hence measurements for at least two different impurity concentrations are required.

The collapse is caused by the combined effect of immobile surface charges (dominating as \( N_i \to 0 \)) and dispersion forces (dominating as \( N_i \to \infty \)). For the ice/liquid/fused quartz system the collapse caused by dispersion forces occurs at thicknesses on the scale of the size of a single molecule where the notion of a quasi-liquid layer no longer applies. In order to observe collapse induced by dispersion forces a larger Hamaker constant is required such as provided by the ice/liquid/gold system.

2. Layer thickness for a given undercooling

When \( q = 0 \), Eq. (22) can still be solved analytically but because the resulting expression is lengthy and not particularly instructive we present the solution only in the small \( t \) limit of moderate undercooling,

\[ d \approx \frac{b N_i}{t} - \frac{a t}{b^2 N_i^2} - O(t^2). \] (26)

This result clearly illustrates the fact that the presence of attractive dispersion forces \((a > 0)\) acting across the system suppresses the formation of a quasi-liquid layer, whereas repulsive dispersion forces \((a < 0)\) tend to increase the thickness of the premelted layer.

When surface charges on the interfaces of the quasi-liquid layer with the substrate/ice are involved, Eq. (22) becomes transcendental and has no analytical solution. In order to get some insight in the effect of surface charges, we consider several special cases. First, let us assume that for moderate undercooling the system is dominated by surface charges. This scenario corresponds to setting \( a = 0, b = 0 \) in Eq. (24) and considering only the leading contributions as \( d \) approaches infinity. The solution is then

\[ d = \frac{[\ln(q_s)]^2}{c^2 N_i}. \] (27)
surface charges enhance the formation of a quasi-liquid

This result reveals an interesting consequence of the presence of charges at the interface. Under appropriate conditions the layer thickness $d$ decreases when the impurity concentration increases. This observation is contrary to the naive expectation based on the colligative effect and originates in the fact that for larger $N_i$ the electrostatic repulsion is screened for effectively.

For repulsive dispersion forces ($a < 0$) the solution of Eq. (22) can be obtained asymptotically in the limit of large undercooling $t \to \infty$, whereas for attractive dispersion forces ($a > 0$) the premelted layer collapses to zero thickness for $t$ larger than some finite value. The repulsive case is

$$d(t, N_i; a, q) = \frac{(-a)^{1/3}}{t^{1/3}} + \frac{bN_i}{3t} \frac{(-a)^{1/6}q}{3c\sqrt{N_i}t^{4/3}} + \mathcal{O}\left(\frac{1}{t^{4/3}}\right)$$

and seems to contradict Eq. (27) for two reasons. First, the negative sign of the term proportional to $q$ in Eq. (27) indicates that immobile surface charges impede the formation of the premelted layer, whereas in Eq. (27) immobile surface charges alone are sufficient to stabilize a premelted layer. Second, while Eq. (27) implies that $d$ decreases when the impurity concentration $N_i$ is increased, the opposite conclusion follows from Eq. (28) where, due to the term $\propto q$, the inhibition due to surfaces charges is reduced when $N_i$ increases.

The apparent contradiction can be reconciled by the following observations. A direct consequence of Eq. (22) is that immobile surface charges do not contribute to the film thickness $d$ when the latter equals the Debye length, namely $d_0 = \frac{1}{e\sqrt{2\pi N_i}}$, in which case the surface charge term vanishes. The film thickness $d_0$ translates into an undercooling $t_0 = b^2 \sqrt{N_i} - a\sqrt{\beta} c N_i$. For an undercooling $t < t_0$ surface charges enhance the formation of a quasi-liquid layer while for $t > t_0$ surface charges suppress layer formation. Considering that Eq. (27) applies to the $t \to 0$ case while Eq. (28) to the $t \to \infty$ case, this explains the essential difference between Eqs. (27) and (28) in terms of whether they support or suppress the formation of a liquid film.

In order to isolate the dependence of $d$ on the impurity concentration $N_i$ we consider Eq. (22) in the case where dispersion forces can be neglected ($a = 0$) and the immobile interfacial charge density is small (i.e. $q \to 0$). To lowest order $d$ depends linearly on $q$, and hence we obtain

$$d(t, N_i; a = 0, q \to 0) = \frac{bN_i}{t} + \frac{bq}{t^2} \left( N_i - \frac{\sqrt{7}}{c\sqrt{b}} \right) e^{-cN_i \sqrt{b}/t} + \mathcal{O}(q^2).$$

From this result the sign of $\partial d/\partial N_i$ can be shown to equal the sign of

$$S \equiv \alpha^2 (2 - \alpha) + \epsilon^\alpha / \beta,$$

where $\alpha = cN_i \sqrt{b}/t$ and $\beta = q/(bc^2 N_i^2)$. We find $S > 0$ for all $\alpha > 0$ as long as $\beta < \beta_c = \epsilon^4/32 \approx 1.71$. Therefore if $N_i > \sqrt{q/(\beta_c b^2)}$ the layer thickness $d$ increases with $N_i$ for any given undercooling $t$. For lower impurity concentrations ($\beta > \beta_c$) there is a finite interval for $\alpha$ which contains $\alpha_c = 4$, where $S < 0$. Therefore, at sufficiently low impurity concentration there exists an interval of intermediate undercooling $t$ where $d$ decreases when $N_i$ increases. This result implies that for large undercooling $t$ the layer thickness $d$ must always increase with $N_i$ which was precisely a conclusion obtained from Eq. (28). However, the result cannot be applied to the charge dominated case, Eq. (27), because the corresponding limit $b \to 0$ renders Eq. (27) trivial. It is possible though to obtain $S < 0$ in the related limit $b \to 0$, $t \to 0$, if we assume that $b/t$ approaches a large constant such that $\alpha > 2$ holds.

Attractive dispersion forces The non-monotonic dependence of $d$ on $N_i$ is illustrated in Fig. 4 where the full numerical solution of Eq. (22) is plotted for the case of the ice-liquid-fused quartz layered system. At low undercooling $\Delta T$ the system displays characteristic colligative behavior as described by Eq. (28). The effect of immobile surface charges vanishes in the limit $t \to 0$ as indicated by Eq. (29) and the layer thickness $d$ increases when the impurity density $N_i$ increases. The layer thickness for small $\Delta T$ is obtained by requiring that the impurity concentration $\rho_i = N_i/d$ equals the impurity concentration which would be necessary in a bulk system to see a freezing point depression equal to the undercooling. When $\Delta T$ is increased, surfaces charges start to increase the layer thickness relative to the system without surface charges ($q_i = 0$). Interestingly, the resulting layer thickness for the system with the lower impurity density ($N_i = 0.5 \, \text{mmol/m}^2$) becomes even larger than for the system with more impurities ($N_i = 2.0 \, \text{mmol/m}^2$). This illustrates the result from the above analysis displaying

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{fig3.png}
\caption{The maximal undercooling $\Delta T_{\text{max}}$ as a function of the impurity concentration $N_i$ for the same parameters as used for the calculation of $d_{\text{min}}$ in Fig. 4. The zero slope asymptote for $N_i \to 0$ is observed as expected from Eq. (25). The large Hamaker constant of the ice-liquid-gold system leads to low values of $\Delta T_{\text{max}}$.}
\end{figure}
the existence of an interval of intermediate undercooling where the layer thickness decreases when the impurity density increases. Recall that the reason for this is that at low impurity concentrations the electrostatic force is less effectively screened and hence the repulsion due to surface charges has a stronger effect, leading to a larger layer thickness than for higher impurity concentrations. When the system is undercooled further the collapse of the quasi-liquid layer can be observed in the figure. Here, because of the weakness of dispersion forces, the collapse is a consequence of the electrostatic force becoming attractive at low layer thicknesses. Collapse for the cases without surface charges cannot be observed in Fig. 4 as it occurs at thicknesses below 0.1 nm (c.f., Fig. 3). In order to see layer collapse at larger layer thicknesses a larger Hamaker constant is required, as provided by the ice-liquid-gold system. To see this, we plot solutions of Eq. (22) for this system in Fig. 5 for the same parameters as considered in Fig. 4 for the ice-liquid-fused quartz system. Layer collapse is now observed at thicknesses above 0.1 nm even for the configurations without surface charges. For the configurations with surface charges we observe that for \( N_i = 0.5 \) \( \mu \text{mol}/\text{m}^2 \) layer collapse is not altered by the stronger dispersion forces while for \( N_i = 2.0 \) \( \mu \text{mol}/\text{m}^2 \) the stronger attractive dispersion forces lead to a layer collapse at a larger layer thickness compared to the corresponding configuration considered in Fig. 4. Thus, in this case stronger dispersion forces drive layer collapse.

**Repulsive dispersion forces** The result of the analysis following Eq. (29) can be tested on the ice/liquid/fused silicon system which has a Hamaker constant of \( A_H = -1.66 \times 10^{-21} \) J, the negative sign implying repulsive dispersion forces. As shown in Fig. 6 the system displays the expected colligative behavior for low undercooling and an enhanced layer thickness for intermediate undercooling where the effect of surface charges enhances the colligative effect, which alone would yield a straight line, \( d \propto N^{-1} \). For intermediate undercooling the figure reveals a transition from a regime of low impurity density where the layer thickness decreases when \( N_i \) increases, to a regime of high impurity density where the colligative effect dominates and accordingly an increase of \( d \) is observed when \( N_i \) increases. From the figure the locus of the transition can be estimated to be \( \Delta T_c \approx 10 - 20 \) K, \( d_c \approx 1 \) nm, and \( N_{i,c} \approx 2.5 \) \( \mu \text{mol}/\text{m}^2 \). This is in good agreement with the analytical result obtained from Eq. (29). \( \Delta T_c = 9.6 \) K, \( d_c = 1.25 \) nm, and \( N_{i,c} = 3.97 \mu \text{mol}/\text{m}^2 \). It should be noted that the analytical result is exact only in the limit \( q_s \to 0 \) but the complete solution of Eq. (22) shows that it applies qualitatively and semi-quantitatively for surface charge densities as high as \( q_s = 0.5 \) C/m². For the ice/liquid/silicon system no layer collapse is observed. This results from the fact that dispersion forces, which dominate at low layer thicknesses, are repulsive and stabilize the quasi-liquid layer. However, at impurity densities \( N_i \) smaller than those considered in Fig. 6 the Debye-Hückel expression for \( F_{\text{elc}} \) generates an attractive force in the ice/liquid/silicon system with \( q_s = 0.5 \) C/m² at suf-
cients small layer thicknesses which results in a discon-
tinuous drop in the layer thickness at a certain under-
cooling. The layer, however, does not collapse to zero
thickness but rather is stabilized at a smaller thickness
due to the repulsive dispersion forces which dominate for
very thin quasi-liquid layers. In this article we do not
investigate this phenomenon of layer discontinuity which
occurs in a regime where both the $d$-dependence of $q_s$
the peculiarities of the Debye-Hückel result, and other
effects, have to be accounted for.

**B. Volume fraction of premelted water**

Having analyzed premelting in planar layered systems,
we now return to the implicit equation (Eq. 21) for the
volume fraction of premelted liquid at a given undercool-
ing $t$ that we introduced in Section [11]. In addition to
being implicit for $f_l$ the equation uses the thickness of the
premelted layer, which itself is obtained from an equation
which does not have an analytical solution (Eq. 22). Ac-
cordingly, we do not possess an analytical expression for
$f_l$ in the general case.

*Low surface charge* However, where immobile surface
charges can be neglected ($q_s = 0$), Eq. 21 becomes more
tractable. In the case where the matrix material has a
negative Hamaker constant ($A_H^{\text{mat}} < 0$), i.e., dispersion
forces are repulsive and support the formation of a pre-
melted layer, Eq. 21 becomes

$$f_l = \frac{A}{(t - c_0/f_l)^{1/3}} + \frac{B}{(t - c_0/f_l)^2}, \quad (31)$$

where

$$A = \frac{1}{R} \times \frac{3\eta}{1 - \eta} \times (\frac{-A_H^{\text{mat}}}{6\pi q_m \rho_l})^{1/3} \quad \text{and}$$

$$B = \frac{\xi^2}{R^2} \times \frac{3}{2} \times \frac{\eta}{1 - \eta} \times \frac{C}{2} + 3 \left(2 - \frac{\pi^2}{8}\right) \times \eta \times \bar{\rho}_{gb} \times \frac{1}{3} (4 - \pi) \times \bar{\rho}_{gb}^2,$$  \quad (33)

with $\xi = \gamma_{sl}/\rho_s q_m$ and where $c_0 = R_d T_m \rho_0 / q_m \rho_l$ is, up
to a constant factor, the density of impurities $\rho_0$ in the
fully melted system.

Obviously, Eq. 31 requires that $t > c_0/f_l$ must hold
and thus, because $0 \leq f_l \leq 1$, when $t < c_0$ Eq. 31
is not applicable. The case $t < c_0$ corresponds to an
undercooling which is smaller than the depression of the
bulk freezing point due to the colligative effect originating
from the presence of the impurities. Therefore, we must
assume a liquid fraction $f_l = 1$ when $t < c_0$.

While Eq. 31 does not possess a general analytical
solution, it can be solved in the limit of low impurity
concentration $c_0$ to yield

$$f_l = \frac{A}{t^{1/3}} + \frac{B}{t^2} + g_1(\tilde{t}) \times \frac{c_0}{3t} + g_2(\tilde{t}) \times \frac{c_0^2}{9 A H^{5/3}} + O(c_0^3), \quad (34)$$

with $\tilde{t} = \frac{A}{t} t^{5/3}$ where

$$g_1(\tilde{t}) = 1 + \frac{5}{t + 1} \quad \text{and}$$

$$g_2(\tilde{t}) = 1 + \frac{14 \tilde{t}^2 - 12 \tilde{t} - 1}{(t + 1)^3}. \quad (35)$$

Note that the term $\propto c_0$ is sufficient to obtain the cor-
correct leading contribution due to impurities in both lim-
iting cases $t \to 0$ and $t \to \infty$. The factor $\propto c_0^3$ becomes
a constant in the limit $t \to 0$ where $g_2$ vanishes. The
coefficient belonging to $c_0^3$ vanishes as $t^{-7/3}$ in the limit
$t \to \infty$ and is thus subdominant to $\frac{B}{t}$.

In the case where dispersion forces are attractive ($A_H^{\text{mat}} > 0$) the formation of a premelted layer requires
an impurity concentration which is higher than the one
which would be required to prevent the bulk liquid from
freezing. Hence, in this case all the liquid water in the
matrix pores must originate from liquid veins which form
due to curvature induced premelting. In contrast to the
premelted layer with $A_H^{\text{mat}} > 0$ these veins have formed
when $t > c_0/f_l$, a temperature for which the premelted
layers are still collapsed and do not contribute to the
liquid fraction. Thus, premelted layers do not actually
form because at higher temperatures, beginning with the

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**FIG. 6:** Thickness $d$ of the quasi-liquid layer for the
ice/liquid/silicon system ($A_H = -1.66 \times 10^{-21}$ J) as a func-
tion of the undercooling $\Delta T$. As dispersion forces in this
system are repulsive they stabilize thin quasi-liquid layers and
hence no layer collapse is observed. The figure illustrates the
analytical result obtained from Eq. (29) which states that
for intermediate undercooling $\Delta T$ and $N_l$ below a transition
value the layer thickness decreases when the impurity concen-
tration increases. For large $N_l$ the colligative effect dominates
and $d$ increases when $N_l$ is increased. The transition value
$N_{lc}$ as well as the locus $\Delta T_c$ and $d_c$ where the interval with
$\partial d / \partial N_l < 0$ appears as derived from the figure are in good
agreement with the analytical result (values given in the fig-
ure). Note the sigmoidal transition to small but finite film
thickness for the lowest value of $N_i$. 

\begin{align*}
q_s &= 0.5 \text{ C/m}^2 \\
N_i &= 1.25 \text{ mmol/m}^2 \\
N_i &= 2.5 \text{ mmol/m}^2 \\
N_i &= 5.0 \text{ mmol/m}^2 \\
N_{lc} &= 3.97 \text{ mmol/m}^2 \\
\Delta T_c &= 9.6 \text{ K} \\
\end{align*}

\begin{align*}
\delta_c &= 1.25 \text{ nm} \\
\end{align*}
veins, the sample completely liquifies. Therefore, the corresponding equation for \( f_l \) is

\[
   f_l = \frac{B}{(t - c_0/f_l)^2},
\]

which has a simple analytical solution. Again, we examine the leading order contributions in the limiting cases \( t \to 0 \) and \( t \to \infty \) to find

\[
   f_l \to 0 \quad \frac{B}{t^2} + \frac{2c_0}{t} - \frac{\rho_0}{B} + \mathcal{O}(t) \quad \text{and} \quad \frac{c_0}{t} + \frac{\sqrt{Bc_0}}{t^{3/2}} + \frac{B}{2t^2} + \mathcal{O}\left(\frac{1}{t^{5/2}}\right),
\]

where the latter relation is an extension of Eq. (4) which is undefined in the limit \( t \to \infty \) and \( A \to 0 \).

\textbf{Non-zero surface charge} \quad \text{In the general case with immobile surface charges (} \( q_s \neq 0 \) \text{)} we solve Eq. (21) numerically. In Fig. 7 we show results for a matrix of fused quartz with a grain size of \( R = 500 \, \mu m \), which may serve as a model for frozen soils. The surface charge density is chosen to be \( q_s = 0.5 \, C/m^2 \) which has the correct order of magnitude for mineral/water interfaces at high pH [10]. The ice which fills the voids of the porous matrix is taken to be polycrystalline with a grain boundary density \( \rho_{gb} = 1 \, \mu m^{-1} \). A comparison of Eqs. (14) and (15) shows that for the given \( \rho_{gb} \) the surface area provided by grain boundaries is by a factor 100 larger than the surface area of the porous matrix. Therefore, we expect the impurity concentration in the sample to be very low when grain boundaries are melted and, according to Eq. (21), the maximal undercooling \( \Delta T_{\text{max}} \) for which quasi-liquid interfacial water at the grain boundaries is stable should be low. An upper limit for \( \Delta T_{\text{max}} \) can be calculated by assuming that all the impurities in the sample are located at the premelted grain boundaries. For the values of \( \rho_0 \) from Fig. 7 we use \( N_t = \rho_0/\rho_{gb} \) to obtain a range \( N_t = 0.01 - 0.1 \, \mu mol/m^2 \) and, with the aid of Eq. (21), this corresponds to a range in \( \Delta T_{\text{max}} \) of 0.01 - 0.26 K. The range of \( \Delta T_{\text{max}} \) uses the Hamaker constant \( \Delta H = 3.3 \times 10^{-22} \, J \) computed from complete dispersion force theory [3] in the limit of thin layers with the data for the dielectric functions from Elbaum and Schick [11]. Including a typical surface charge density of \( q_s = 0.01 \, C/m^2 \) for the grain boundary [12] lowers these values to \( \Delta T_{\text{max}} = 0.01 - 0.20 \, K \). Hence, the grain boundaries are always collapsed to zero film thickness for the range of \( \Delta T \) considered in the figure. A prominent feature of \( f_l = f_l^{\text{tot}} \) with \( \rho_0 = 0.01 \, mol/m^3 \) in the fused quartz matrix is the discontinuity at \( \Delta T = 22.6 \, K \) where \( f_l \) drops by almost an order of magnitude. This is the signature of the collapse of the premelted layer at the ice/fused quartz interface described in detail in Section 11A. In the strongly undercooled, layer collapsed system, the only contribution to \( f_l \) comes from curvature induced melting; \( f_l^{\text{curv}} = f_l^{\text{contact}} + f_l^{\text{mat}} + f_l^{\text{veins}} \) (cf. Eq. (21)), which discontinuously increases at the undercooling where the premelted layer collapses. This is because, due to the efficient rejection of impurities by the ice lattice, the impurity concentration in the remaining liquid increases discontinuously when the liquid layers collapse. At low undercooling curvature melting becomes increasingly important exceeding the contribution from interfacial premelting \( f_l^{\text{layer}} = f_l^{\text{mat}} \). However, at larger undercooling, close to the location of layer collapse, curvature induced melting yields only a minute contribution to \( f_l \). The importance of curvature induced melting at low undercooling is reflected in Eq. (33) where the curvature term \( B/t^2 \) dominates for \( t \to 0 \). In the opposite limit of large undercooling the analytical result Eq. (39) applies. The leading term \( c_0/t \) originates from the collicative effect and curvature induced melting enters only through the subdominant term. Hence, for large undercooling the principal contribution to the liquid fraction is the collicative effect. However, liquid water is located in regions of high curvature: contact of matrix spheres, contact lines of grain boundaries and matrix spheres and liquid veins. Figure 7 also shows \( f_l \) in the same system with a higher impurity concentration (\( \rho_0 = 0.1 \, mol/m^3 \)). Here the collapse of the liquid layer is moved to a lower temperature (\( \Delta T \approx 110 \, K \)) so that no discontinuity is observed in the range of undercooling considered in the figure.

In Fig. 8 many of the same conditions as in Fig. 7 are considered (\( R = 500 \, \mu m, \, q_s = 0.5 \, C/m^2, \, \rho_0 = 0.1 \, mol/m^3 \)) the principal difference being that the pore ice is single-crystalline (\( \rho_{gb} = 0 \)). We find that \( f_l^{\text{tot}} \) in the fused quartz matrix is qualitatively the same as for
the polycrystalline ice (see Fig. 7) in that it does not show liquid layer collapse on the scale of the plot for $\rho_0 = 0.1$ mol/m$^3$. When replacing the matrix with gold, which has stronger attractive dispersion forces at the ice-gold interface, we see a collapse of the premelted layer at $\Delta T \approx 7$ K. This is accompanied by, as in the case of layer collapse for the fused quartz matrix at $\rho_0 = 0.01$ mol/m$^3$ (Fig. 6), a discontinuous increase of the liquid fraction $f_i^{tot}$. As for the fused quartz matrix, for sufficiently low undercooling that the premelted layer has collapsed, the only contribution to $f_i^{tot}$ comes from curvature induced melting: $f_i^{tot} \approx f_i^{curv}$. While, on the scale of the plot $f_i^{tot}$ does not display a discontinuity at the point of layer collapse, there is a discontinuity $\Delta f_i$ which can be seen by expanding the full result in the limit of large impurity concentration $c_0$

$$\Delta f_i = \frac{1}{2} \times f_i^{layer} \times \left( \frac{B}{c_0 t_{coll}} \right)^{1/2} + O \left( \frac{1}{c_0^{3/2}} \right), \quad (40)$$

where $t_{coll}$ is the undercooling at layer collapse and $f_i^{layer}$ the liquid fraction from interfacial melting at a temperature slightly above collapse $t = t_{coll} - \delta t$, with $\delta t > 0$. Clearly, $\Delta f_i \rightarrow 0$ for $c_0 \rightarrow \infty$.

An important result seen in Fig. 8 is that although the Hamaker constants for gold and fused quartz differ by almost two orders of magnitude, the results for $f_i$ are almost identical over a large range of $\Delta T$; the deviation increases from 0.02 % to 0.4 % at layer collapse and is 6 % for $\Delta T = 60$ K. This is because dispersion forces are only relevant for thin premelted layers and hence at temperatures slightly above layer collapse. However, when layer collapse occurs, according to Eq. (40) the high impurity concentration, combined with single-crystallinity (i.e. small $B$), insures that the Debye length and the discontinuity of $f_i$ are small. Therefore, the deviation between the results for the two different matrix materials remains small. The conclusions drawn from Fig. 8 are important to note for experimental determination of $f_i$ in different materials because they imply that even though the underlying microscopic behavior of two samples may fundamentally differ (interfacial premelting vs. collapsed premelted layer) this may not be resolved in the integrated/homogenized quantity $f_i$. Spatially resolved measurements may be required to detect characteristic transitions such as the collapse of interfacially premelted layers.

**Effect of polycrystallinity** Finally, we study the influence of the degree of polycrystallinity of the interstitial ice for the case where the porous matrix consists of silicon, which leads to dispersion forces that are repulsive ($A_H < 0$) and prevent the interfacially premelted layer from collapsing. As in the previous examples the grain size of the matrix is $R = 500 \mu$m and as in Fig. 7 the grain boundaries are collapsed. Thus, Fig. 9 shows the curves for $f_i = f_i^{tot}$, the interfacial melting ($f_i^{layer}$) and curvature induced melting ($f_i^{curv}$) contributions for single-crystalline ($\rho_{gb} = 0$) and polycrystalline ice ($\rho_{gb} = 1 \mu$m$^{-1}$). As expected, the polycrystallinity leads to a higher liquid fraction $f_i$. In particular, for small undercooling $\Delta T$, where curvature induced melting is enhanced and hence liquid veins greatly contribute, $f_i$ consists almost entirely of $f_i^{curv}$. The values of $f_i^{curv}$ for the single crystal ice, for which contributions derive only from the contact points between matrix spheres, are always by at least an order of magnitude lower than for the polycrystalline ice case. This is not as obvious as it may first appear. Due to the higher liquid fraction in polycrystalline ice the impurity concentration $\rho_i$ is smaller than for the single crystal case. Thus, polycrystalline ice has a smaller ice-water radius of curvature (see Eq. (10)). Consequently, the volume of liquid contained for example at the vicinity of layer collapse. The discontinuity for fused quartz is not visible in the figure because it occurs at large undercooling ($\Delta T \approx 103$ K).
pronounced at small undercooling. In this regime a polycrystalline sample always has a higher liquid fraction, $(\rho_{gb} = 1 \text{ \mu m}^{-1})$. Cooling $\Delta T$ is assured. Will provide irrelevant and misleading predictions unless the microscopic phenomena in a theory that is intended to predict the effective medium behavior of a porous medium, will provide irrelevant and misleading predictions unless the proper asymptotic limits of a complete theory are assured.

IV. SUMMARY AND CONCLUSION

In this work we construct a theory for the premelting of ice in a porous matrix (Section III), which is modeled as a random close pack of monodisperse spheres. The theory includes premelting at the matrix-ice interface, grain boundary premelting, as well as curvature and impurity induced premelting. The latter includes premelting in liquid veins, at contact lines of the matrix with a grain boundary, and at the contact points between two matrix spheres. The matrix is characterized by the sphere radius $R$ and the matrix material, which enters the theory via the Hamaker constant $A_H$ for the ice/liquid/matrix layered system. In addition to dispersion forces at the ice-matrix interface, which, depending on the matrix material, either stabilize or destabilize the premelted quasi-liquid layer, we include the effect of electrostatic forces at the interface due to adsorbed surface charges at the interface along the lines of our previous work Ref. 5. The theory includes the effect of charged impurities in the premelted water both on the thickness of the premelted layers (grain boundaries, ice-matrix interface) as well as on the curvature induced melting. The main idea of the present approach is to make use of the small segregation coefficient of ice which leads to an enrichment of impurities in the premelted liquid (making the impurity concentration a function of the fraction of premelted liquid) and of the fact that regions of premelted liquid in a sample are all interconnected and therefore the impurity concentration is homogeneous. This approach allowed us to derive an implicit equation for the liquid fraction $f_l$ as a function of the undercooling $\Delta T$, Eq. (21), which we analyze for general parameters by considering limiting cases where analytical solutions are possible (Section III). To understand each contribution we began by performing a careful study of interfacial melting, analyzing an implicit equation for the thickness $d$ of the premelted layer as a function of $\Delta T$, Eq. (4), for general parameters. In particular, we addressed the discontinuous collapse of the premelted layer under certain conditions and the non-monotonic dependence of $d$ on the impurity concentration (cf. Ref. 5). However, in the general case both Eq. (7) and Eq. (21), the latter of which makes use of the former, have to be solved simultaneously through a numerical scheme. In so doing we chose physically relevant parameters of interest to a wide community of scientists. For the porous matrix we use $R = 500 \mu m$ and considered two materials (fused quartz and gold) that yield attractive dispersion forces at the ice/matrix interface, thereby inhibiting the formation of a premelted quasi-liquid layer, and one material (silicon) which enhances the premelted layer. The surface charge at the interface is taken to be $q_s = 0.5 \text{ C/m}^2$. The impurity concentrations lie in the range of $0.01 - 0.1 \text{ mol/m}^3$ defined with respect to the interstitial volume, and we consider both single- and polycrystalline ice. We calculate $f_l$ for $\Delta T$ in the range of $0.5 - 60 \text{ K}$ and obtain liquid fractions which range between $10^{-6}$ and $10^{-3}$ of the interstitial volume.
For the parameters under consideration the grain boundaries are always of zero thickness (i.e., collapsed), and hence only the collapse of the quasi-liquid layer at the ice-matrix interface is observed in $f_l$ (see Fig. 7 for the case of the fused quartz matrix). When the temperature of the sample is decreased below the point of layer collapse, only curvature induced premelting contributes to $f_l$. While dispersion forces are too weak in the fused quartz system to significantly influence the results, in the case of the gold matrix dispersion forces are sufficiently strong to drive layer collapse, as opposed to the surface charge triggered layer collapse in the fused quartz system (see Fig. 8 for a comparison of the two cases). An important conclusion from Fig. 8 is that for high impurity concentration and/or low degree of polycrystallinity the characteristic discontinuity of $f_l$ upon layer collapse is minute and most likely not resolved in experiments. Thus, transitions that would be dramatic and easily detected at planar interfaces will be much more challenging to extract from measurements of the integrated/homogenized quantity $f_l$. Finally, we study the influence of the degree of polycrystallinity on $f_l$ for the case of the silicon matrix where, due to the repulsive dispersion forces, layer collapse does not occur (Fig. 9). The results reveal an interesting consequence of the non-monotonic dependence of $d$ on the impurity concentration. Although the actual impurity concentration is higher in the single-crystalline sample than in the polycrystalline sample at all temperatures, the thickness of the premelted layer can be both larger and smaller in either case, depending on the temperature range. This peculiarity is obscured by the curvature induced contribution to $f_l$ providing further evidence that additional measurements are required $f_l$ to capture the physics controlling the sample microstructure.

We view our approach as having provided a thorough theory of premelting in a porous medium which can be applied to a plethora of experimental situations and material systems. For the sake of brevity and clarity, we presented examples of specific materials systems; gold, fused quartz, and silicon. However, we chose these materials because their dielectric properties, as reflected in their Hamaker constants, span all the range of materials as diverse as sapphire, polystyrene, and polyvinylchloride [2]. We did not explore the effect of varying the matrix sphere radius $R$, its roughness or its surface charge density $q_s$. The value $R = 500 \, \mu m$ was chosen with the example of frozen soils in mind, while $q_s = 0.5 \, \text{C/m}^2$ provides a realistic order of magnitude as inferred from experimental data for mineral-water interfaces at large pH [10]. However, not much is known experimentally about $q_s$ in the ice/liquid/matrix systems and the simplifying assumptions of symmetry (same charge at the ice-liquid and liquid-matrix interfaces) as well as independence of $q_s$ on the impurity concentration and layer thickness are not yet established experimentally on planar systems. Nevertheless, given the importance of the impurity concentration dependent screening length for regulating the strength and range of the electrostatic force, the precise value of $q_s$ may be considered secondary.

An important question for future research concerns the dynamical evolution of the porous matrix system in response to changes in the control parameters, in particular the sample undercooling $\Delta T$. In order to guarantee a homogeneous impurity concentration throughout a sample, $\Delta T$ must be changed on a time scale that is slow compared to the typical impurity diffusion time. An even more ambitious question concerns impurity diffusion during layer collapse. The situation where the premelted quasi-liquid layer between the ice and the matrix collapses, which forces the impurities to migrate to remaining liquid regions with curvature induced melting constitutes a challenging task for further investigations.

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Appendix A: Melting volume at a grain boundary–matrix contact line

We consider a matrix sphere that is embedded at the interface between two ice grains. The grains are assumed to have a planar grain boundary and to be much larger than the matrix sphere. Let the distance between the sphere center and the grain boundary be $z < R$. Curvature induced melting leads to a circular “vein” along the line where the grain boundary and the sphere intersect (Fig. 10a). Because $r$ is assumed to be much smaller than

![FIG. 10: a) Geometry of a matrix sphere intersecting with a grain boundary which is located at distance $z$ from the sphere center. b) Zoom on the curvature melting induced liquid “vein”. As $r \ll R$ the matrix surface can be considered as flat in the vicinity of the liquid vein.](image-url)
we can consider the matrix surface as being planar in the vicinity of the vein as shown in Fig. 10b. The angle between the grain boundary and the matrix sphere is

\[ \alpha = \arccos \left( \frac{z}{R} \right). \]  

(A1)

Using elementary geometry we obtain the surface areas

\[ A = \cot \left( \frac{\alpha}{2} \right) r^2 - \left( \frac{\pi}{2} - \frac{\alpha}{2} \right) r^2, \]  

(A2)

\[ A' = \tan \left( \frac{\alpha}{2} \right) r^2 - \frac{\alpha}{2} r^2, \]  

(A3)

and the total surface area

\[ A_{\text{tot}} = A + A' = \left( \frac{2}{\sin \alpha} - \frac{\pi}{2} \right) r^2. \]  

(A4)

Accordingly, the volume of premelted water for the configuration in Fig. 10 is \( 2\pi h A_{\text{tot}} \). Taking all possible distances of the matrix sphere to the grain boundary into account yields an average volume \( \bar{v} \) of premelted liquid as

\[
\bar{v} = \frac{1}{R} \int_0^R dz 2\pi h A_{\text{tot}} \\
= \frac{1}{R} \int_0^R dz 2\pi R \sin \alpha \left( \frac{2}{\sin \alpha} - \frac{\pi}{2} \right) r^2 \\
= 2\pi r^2 \int_0^R dz \left( 2 - \frac{\alpha}{2} \sqrt{1 - \frac{z^2}{R^2}} \right) \\
= 2\pi r^2 R \left( 2 - \frac{\pi^2}{8} \right). 
\]  

(A5)

Now consider a random close packing with packing fraction \( \eta \) (the porous medium), which contains ice with two different crystallographic orientations being separated by a planar grain boundary. The volume fraction \( \eta \) is also the surface fraction of the grain boundary which is cut out in the form of circles with radii between 0 and \( R \) due to the presence of the matrix spheres. Given that the average surface area per intersecting sphere is

\[ \bar{a} = \frac{1}{R} \int_0^R dz \pi (R^2 - z^2) = \frac{2}{3} \pi R^2 \]  

(A6)

a sample of volume \( V_s = A_s x_s \) with a grain boundary perpendicular to the \( x \)-axis contains \( \eta A_s / \bar{a} \) intersection circles of matrix spheres with the grain boundary, if \( A_s \) is sufficiently large. If the sample contains the total grain boundary surface \( \rho'_\text{gb} V_s \) this corresponds to a net surface \( A_s = \rho'_\text{gb} V_s / (1 - \eta) \) and hence a density of intersection circles

\[ n_c = \frac{\eta \rho'_\text{gb}}{(1 - \eta) \bar{a}} = \frac{\eta \rho'_\text{gb}}{\bar{a}}, \]  

(A7)

where \( \rho'_\text{gb} \) is the grain boundary density with respect to the interstitial volume. The volume of premelted liquid along the intersection circles (per unit volume of the sample) is \( \bar{v} n_c \). Or, relative to the interstitial volume:

\[ \bar{v} n_c = \frac{3}{1 - \eta} \left( 2 - \frac{\pi^2}{8} \right) \times \frac{\eta}{1 - \eta} \times \rho'_\text{gb} \times \left( \frac{r}{R} \right)^2. \]  

(A8)

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