

# A Mathematical Model of Microbially-Induced Convection in Sea Ice

Noa Kraitzman<sup>1</sup>, Jean-David Grattepanche<sup>2</sup>, Robert Sanders<sup>2</sup>, and Isaac Klapper<sup>3</sup>

<sup>1</sup>Department of Mathematics, Macquarie University, New South Wales 2109, Australia

<sup>2</sup>Department of Biology, Temple University, Philadelphia, PA 19122, USA

<sup>3</sup>Department of Mathematics, Temple University, Philadelphia, PA 19122, USA

Corresponding author: Isaac Klapper, [klapper@temple.edu](mailto:klapper@temple.edu)

**Abstract.** Through its role as an interface between ocean and atmosphere, sea ice is important both physically and biologically. We propose here that the resident microbial community can influence the structure of sea ice, particularly near its ocean interface, by effectively lowering the local freezing point via an osmolytic mechanism. This lowered freezing point can enhance fluid flow, linking a bottom, convective ice layer with the underlying ocean, resulting in improved nutrient uptake and byproduct removal. A mathematical model based on a previously suggested abiotic one dimensional simplification of mushy ice fluid dynamics is used to illustrate, and supporting measurements of freezing point depression by lab grown sea ice-associated organisms are provided.

## 10 1 Introduction

Sea ice in the polar regions is a fundamental cryospheric habitat that serves as an important component of Earth's climate system by regulating heat exchange between the atmosphere and ocean and also by reflecting sunlight (Ledley, 1991; Ledley; Eicken and Lemke, 2001; Boeke and Taylor, 2018). It also supports diverse microscopic assemblages, including bacteria, algae, and various invertebrates, all essential for maintaining ecological balance (Arrigo and Thomas, 2004; Thomas and Dieckmann, 15 2008; Arrigo, 2014). During sea ice formation and growth, the progressive development of brine-filled channels establishes distinct physicochemical microenvironments that facilitate the colonization of specialized microbial communities, which can propagate to impact large-scale systems. This can have macroscale implications: in the polar oceans, sea ice serves as a critical environment influencing ecology and biogeochemical cycles (Comeau et al.; Arrigo, 2014; Swadling et al., 2023), functioning as a refuge for larger organisms like seals and penguins, while providing habitat for communities of prokaryotes and eukaryotes 20 on ice surfaces and within brine channels (Garrison et al., 1986; Bluhm et al., 2017; Kohlbach et al., 2018). Indeed, it appears that the structure and activity of sea ice microbial communities is considerably more complex than once thought (Audh et al., 2023).

Sea ice microbial communities serve as vital food resources for copepods and krill (Bluhm et al., 2017; Kohlbach et al., 2018) and influence fundamental ecological and atmospheric processes, including dimethylsulfide emissions, carbon dioxide

25 uptake, methane release, and halogen chemistry (Fadeev et al., 2021; Steiner et al., 2021), influencing global biogeochemical  
cycles (Lannuzel et al., 2020). Laboratory studies have demonstrated that microbial communities can modify their habitat  
chemically (Zhou et al., 2014) but also physically, through alteration of sea-ice microstructure by pore clogging and depression  
of brine freezing points (Krembs et al., 2011). As sea ice continues to thin and transition toward predominantly first-year ice,  
30 these communities are adapting by initiating colonization earlier in the season and accumulating greater biomass over shortened  
periods, possibly facilitated by enhanced light penetration, with their ecological importance increasing as sea ice loss continues  
due to their adaptability to seasonal fluctuations in the carbonate system (Torstensson et al., 2021). While these mechanisms  
are known, their precise dynamics and quantification at large scales remain poorly understood and are generally absent from  
large-scale modeling (Vancoppenolle et al., 2013). Given the rapid transformation of polar regions, exploring the potential for a  
reciprocal relationship between large-scale marine biogeochemical processes and their resident microbial communities seems  
35 ~~worthwhile~~timely. In that spirit, we propose here the possibility that sea ice microbes can not only adapt to changing sea ice  
conditions but also modify the ice itself to maintain nutrient supplies.

### 1.1 Sea Ice

Sea ice itself would seem to be a difficult habitat, and while clear differences in community composition ~~were not always~~  
~~observed between sea ice~~have not always been observed between sea ice and the surrounding water (Garrison et al., 1986;  
40 Dawson et al., 2023), metabolomics show differences suggesting mechanisms to cope with the changes in temperature and  
salinity between these two environments (Dawson et al., 2023). We focus here on interaction between the ice and its microbial  
community (the structure of sea ice is notable for its length scale dependence through a variety of physical mechanisms) at  
small scales, centimeters and below, where individual ice crystals, fundamental to sea ice's formation, are typically on the  
millimeter size. This scale reveals sea ice as a composite material composed of two main components: solid ice ~~;~~ and liquid  
45 brine. Because of the connection to transport properties, composite details influence to a significant extent macro-properties  
including, notably here, biological productivity.

Sea ice is formed during colder seasons when air temperature can reach temperatures well below 0° C. On the other hand, the  
temperature at the sea ice-ocean interface remains approximately -1.8° C throughout the year and so, when the air temperatures  
drop sufficiently, ice formation can occur. As freezing occurs, impurities are excluded, leading to the formation of relatively  
50 high salinity brine inclusions in almost pure-water ice. Inclusion sizes become larger as the warmer (at least in colder seasons)  
ice-ocean interface is approached for thermodynamic reasons (discussed below in Section 1.2) (Kraitzman et al., 2022). Near  
this interface, in a layer in which the ice temperature rises above a threshold temperature (Golden et al., 1998), brine inclusions  
interconnect to the point of forming a permeable material through which fluid can, in principle, flow and thus bring fresh  
seawater and accompanying nutrients (and microbes) (Tedesco et al., 2010).

55 The presence of an interconnected network does not by itself result in fluid flow though; there is also a need for a driving  
force capable of overcoming viscous drag. During periods of growth, ~~flows in~~flow within sea ice can be driven by density  
inversions in large part due to brine concentration (Worster, 1992, 1997; Worster et al., 2000; Feltham et al., 2006; Worster and  
Jones, 2015). That is, freezing of seawater results in relatively dense brine, through the ~~salinity~~ concentration process described

above, which can result in convective, Rayleigh-Taylor instabilities (Chandrasekhar, 1961) in which, roughly, the gravitational potential energy of a density inversion can be more easily released by convective versus diffusive transport. Convective flow requires interconnected channels, and the larger those channels, the easier convective transport becomes since viscous drag is sensitive to channel size. This draining process, however, relies on freezing of “new”, salty seawater by the advancing ice front though, and cannot sustain flow when freezing is not occurring. Hence the question arises of how microbial communities can sustain themselves within the ice, even in the permeable layer, when sufficient freezing is not occurring. We argue here that the microbes themselves can drive a convective flow.

## 1.2 Effective Salinity

Locally, sea ice temperature and brine inclusion concentration are often approximated as being close to thermodynamic equilibrium, since the local heat diffusion time scale is generally short compared to time scales of other relevant diffusive and advective transport processes of interest. That is, local brine inclusion salinity (we will use the term salinity in a specific sense related to effect of freezing temperature, discussed below) must be such that the local brine freezing point matches the local temperature, quantified through an empirical relation of the form  $\mathcal{L}(T) = S_{\text{brine}}$ , called a liquidus relation (see Appendix A3 for details). Here  $T$  ( $^{\circ}\text{C}$ ) is the local temperature and  $S_{\text{brine}}$  (ppt) is the local brine salinity. Note also that total, or bulk, salinity  $S_{\text{bulk}}$  is conserved, locally, on short time scales, and that  $S_{\text{brine}} = S_{\text{bulk}}/\phi_{\text{brine}}$  (where  $\phi_{\text{brine}}$  is local brine volume fraction) so that the liquidus relation can be written in the form  $\widehat{\mathcal{L}}(T, \phi_{\text{brine}}) = S_{\text{bulk}}$ . As  $\widehat{\mathcal{L}}$  is monotone increasing in  $\phi_{\text{brine}}$ , then, given  $T$  and  $S_{\text{bulk}}$ , this relation can be inverted to solve for  $\phi_{\text{brine}}$ .

Over longer time scales, though, local bulk salinity can change due to diffusive or advective transport, and, particularly in the presence of biological activity, due to chemical sources or sinks. We argue here that sea ice microorganisms can use such production to in fact manipulate ice permeability and, ultimately, drive advective flow. Some sea ice micro-algae including diatoms, e.g., *Rhizosolenia*, *Corethron*, and haptophyta, e.g., *Phaeocystis*, and also heterotrophic species such as nanoflagellates and ciliates (Garrison et al., 1986; Caron and Gast, 2010; Kohlbach et al., 2018; Hop et al., 2020; Dawson et al., 2023) ~~produced~~ are known to produce metabolites in different distributions and concentrations in relation to change in environmental conditions including temperature and salinity (Dawson et al., 2020). For example, dimethylsulfoniopropionate (DMSP) and dimethylsulfide (DMS), which show antifreeze properties (Uhlir et al., 2019; Sheehan and Petrou, 2020), tend to increase inside the cells of micro-algae associated with sea ice compared to seawater: seawater at approximately 0.19 nmol metabolite C per total  $\mu\text{mol C}$  and sea ice at approximately 1.81 nmol metabolite C per total  $\mu\text{mol C}$  (Dawson et al., 2023). Other metabolites, such as proline and glycine betaine, are also produced in larger concentrations in ice conditions, and are known to increase freeze tolerance in other species (e.g., yeast for proline (Morita et al., 2003), *Arabidopsis* for glycine betaine (Xing and Rajashekar, 2001)).

We explore here the possibility that at least some of these various substances may, among other things, effectively act as external antifreeze agents in the manner of osmolytes. To do so, we define an effective brine salinity as a combination  $S_{\text{brine}} + Y_{\text{sal}}O_{\text{brine}}$  and replace  $S_{\text{brine}}$  in the liquidus relation by this ~~effective~~ effective brine salinity, see Appendix A for details. Here  $O_{\text{brine}}$  is a brine (microbially-originated) osmolyte concentration and  $Y_{\text{sal}}$  is a yield coefficient, converting osmolyte

concentration to effective salinity concentration. Osmolyte is measured in normalized units so that one unit of osmolyte has the same density effect as one unit of salt. The parameter  $Y_{\text{sal}}$  takes account of the relative salinity effect of osmolyte in these normalized units. For modeling purposes, we don't distinguish a particular choice of microbially produced osmolyte, or osmolytes, and so ~~don't do not~~ try to estimate a particular value for  $Y_{\text{sal}}$ . ~~We don't see~~ There did not seem to be a lot of sensitivity to choice of  $Y_{\text{sal}}$  in computations though, possibly because of two competing effects: increasing/decreasing  $Y_{\text{sal}}$  tends to increase/decrease brine volume fraction which ~~increase~~ increases/decreases permeability, but at the same time decreasing/increasing added density (from the extra osmolyte) which decreases/increases the density driven fluid driving potential.

## 100 2 Freezing Point Measurements Materials and Methods

To test the impact of microalgae on ice formation temperature, we measured freezing points (e.g., Fujino et al. (1974)) using the diatom *Chaetoceros neogracilis* (W-126Chaetoceros, 97% similarity to EU090012) from the Antarctic Protist Culture Collection, USA. This organism has been reported in sea ice and in seawater in the Arctic Ocean during pack ice melting (Gérikas Ribeiro et al., 2020) and often dominates phytoplankton community (Katsuki et al., 2009; Crawford et al., 2018). *C. neogracilis* is a solitary rectangular cell, measuring 4-10  $\mu\text{m}$  (Scott and Marchant, 2005; Katsuki et al., 2009), and was isolated from the Ross Sea in 1999 on board the RVIB Nathaniel B Palmer. It has been maintained since in the laboratory in a non-axenic culture kept in 36 ~~PSU~~ ppt Instant Ocean® with f/2 + Si media (Guillard and Ryther, 1962) in an incubator at 4° C under illumination at 20  $\mu\text{mol photons m}^{-2} \text{ s}^{-1}$  14:10 light:dark cycle. Cell abundance was approximately  $7 \cdot 10^8 \text{ cell L}^{-1}$  during the experiment.

110 For the measurements, we used a simplified thermal calorimetric apparatus, see Figure 1. A volume of 25 mL of liquid was placed in a ~~50mL~~ 50 mL glass culture tube inside a 150mL Nalgene plastic bottle, Fig. 1. To create a cold bath, the plastic bottle was placed in dry ice (-70° C). Dry ice was used because its temperature is well below that of the freezing point of the culture liquid as well as to reduce the duration of the experiment and avoid any metabolic interference or death of the microalgae due to the freezing, rendering more complicated the interpretation of the results. To ensure the homogeneity of the cooling process, a magnetic stir bar was introduced into the ~~50mL~~ 50 mL glass culture tube and the liquid was stirred at medium speed. Temperature was recorded each minute with a Traceable® Excursion-Trac™ datalogging thermometer with stainless-steel probe, see Fig. 1. Using this apparatus, the freezing point of seawater (Instant Ocean) plus f/2 media (nutrients for phytoplankton) and a diatom culture of *Chaetoceros* sp., suspended in seawater plus media, were assessed. Salinity of the culture medium itself ( $\sim 26.5 \text{ mg/L}$ ) was negligible compared to that of the seawater ( $\sim 11.1 \text{ g/L}$ ). To understand the impact of osmolytes produced by the diatoms on the freezing point, cultures were centrifuged at 7500 rpm for 10 minutes at 4° C and the resulting liquid supernatant was used to measure the impact of osmolytes.

125 Each experiment was run for 30-40 minutes, until solidified ice was observed. To determine the freezing point, temperature was plotted over time. After a phase of cooling and then supercooling, the liquid changed state and ice nucleation began. The resulting ice formation released heat until the freezing point (plateau) was attained, at which an approximate equilibrium between crystal formation and melting occurs. The temperature was stable during this phase.



**Figure 1.** Freezing point measurement apparatus. (Left) the thermal calorimetric apparatus constituted of an isolated bucket on a magnetic stir plate. The bucket contains dry ice in which a chamber is placed. (Right) the chamber is composed of a 50 mL glass culture tube inside a polycarbonate 150 mL bottle.

### 3 Mathematical Model

Mathematical descriptions of phase behavior in abiotic sea ice have been based, typically, on reaction-advection-diffusion equations for transport of bulk heat and salt, coupled to a liquidus relation (as described in Section 1.2) maintaining a local equilibrium between solid and brine phases (e.g., Worster (1992, 1997)) (e.g., Worster, 1992, 1997). This equilibrium also sets local solid and brine phase volumes, which couple into a fluid dynamics description, typically modeled as a porous media (Darcy) flow. The entire system is driven by a temperature gradient vertically through the ice sheet column, and can become quite complex. We employ a significant simplification of full sea ice dynamics, following the references Vancoppenolle et al. (2010); Griewank and Notz (2013); Turner et al. (2013); Griewank and Notz (2015), in which three dimensional mushy layer fluid dynamics are replaced by a one-dimensional (1D), parameterized reduction. That is, mushy ice velocity field  $\mathbf{u}(\mathbf{x}, t)$  is replaced by a 1D, non-negative upwards velocity  $U(z, t)\hat{\mathbf{z}}$ , driven by a modeled Rayleigh-Taylor instability mechanism. When the computed local Rayleigh number  $Ra(z, t)$  exceeds a critical Rayleigh number  $Ra_{\text{crit}}$ , material is removed and flushed into the ocean at rate proportional to  $Ra - Ra_{\text{crit}}$ , with  $U(z, t)$  computed so as to conserve mass, see Appendix A for more details. The principle novelty here arises from the addition of microbes and microbially-produced osmolyte, equations (A3) and (A5), and the possibility of microbially-induced fluid flow. Note that a 1D sea ice system with microbes and limiting nutrient included, but without osmolyte and resulting induced convective flow, was considered in Vancoppenolle et al. (2010).

Two types of microbial communities are separately considered, though it can be expected that both types may occur simultaneously in actuality. First, we consider a sessile community (i.e., a biofilm community) that is fixed to the ice, and, second,

we consider a mobile community (i.e., a planktonic community) that advects with the local fluid velocity. In the second case, there is the possibility of transient occupation in which microbes from an ocean reservoir only pass through the ice without any significant residence time, so we also consider growth and decay processes. In the sessile case, though, transience is not an issue so for simplicity we do not include growth/decay in the microbial model. That is, in the biofilm case, we suppose that growth and decay roughly balance.

For both types of communities, we include a mechanism for inhibition of activity in the upper part of the ice-sheet/sea ice column—namely via high levels of effective salinity. (Recall that brine salinity must be high enough so that the freezing point matches temperature, so is high where temperature is low.) Other possible inhibitors include nutrient limitation or temperature effects, but we do not aim to explore their relative impacts here and thus focus on a single mechanism. Note that inhibition is not strictly necessary to examine the hypothesis of microbially induced convection and is, in fact, somewhat cosmetic – we parameterize it so that it is significant only in regions above the convective zones.

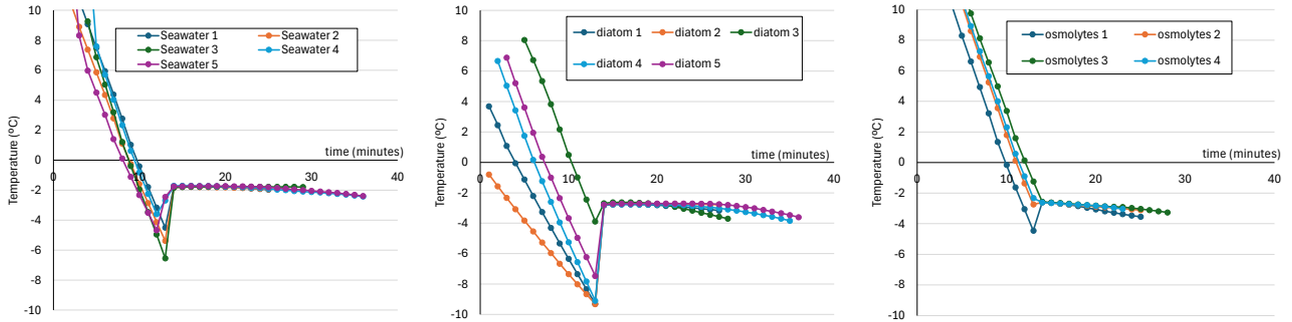
A fixed thickness  $L$  is supposed for the ice sheet-column (in computations below,  $L = 1$  m). Ice thickness is fixed in time so as to better isolate the effects of the model microbial community on ice thickness and structure, without the additional complications of growth/decay of the ice itself. In physical terms, we are thus assuming that heat flux in/out of the ice-ocean interface is balanced without any latent heat sink or source. Including interface latent heat contributions, i.e., ice sheet-column growth/decay, changes a fixed boundary problem to a free boundary one and effectively requires knowledge of heat transport characteristics on the ocean side of the boundary which are subject to ~~(additional)~~ additional significant effects of ocean transport. A growing ice sheet-column also introduces abiotically induced flow resulting from excessively dense brine, again potentially obscuring effects of a local microbial community.

The 1 m thick ice domain consists of the transverse (through the ice sheet-column) profiles of conserved quantities bulk enthalpy mass density  $H_{\text{bulk}}(\mathbf{x}, t)$ , bulk salt volume density  $S_{\text{bulk}}(\mathbf{x}, t)$ , bulk osmolyte volume density  $O_{\text{bulk}}(\mathbf{x}, t)$ , limiting nutrient density  $C_{\text{bulk}}(\mathbf{x}, t)$ , and bulk microbial volume density  $B_{\text{bulk}}(\mathbf{x}, t)$ . From these quantities, temperature  $T(\mathbf{x}, t)$ , brine volume fraction  $\phi_b(\mathbf{x}, t)$ , salt brine density  $S_{\text{brine}}(\mathbf{x}, t)$ , osmolyte brine density  $O_{\text{brine}}(\mathbf{x}, t)$ , limiting nutrient brine density  $C_{\text{brine}}(\mathbf{x}, t)$  and microbial brine density  $B_{\text{brine}}(\mathbf{x}, t)$  can be computed, see Appendix A for details. Boundary temperatures are set at  $T = -20^\circ \text{C}$  at the air-ice interface, and  $T = -2^\circ \text{C}$  at the ice-ocean interface.

## 4 Results

### 4.1 Freezing Point Measurements

Experiments were conducted to measure effects of cultured diatoms on seawater freezing temperature, see Figure 2. *Chaetoceros* abundance in the sample was approximately  $7 \cdot 10^8$  cell/L, as compared to reported measurements ranging (at least) from  $10^4$  to  $10^9$  cell/L from algal cells in sea ice (Arrigo, 2017), with diatoms dominating in the bottom layer (van Leeuwe et al., 2018). Seawater solution used to culture diatoms froze at  $-1.75 \pm 0.03^\circ \text{C}$  (Fig. 2, left), while seawater solution with the diatom culture froze at  $-2.73 \pm 0.05^\circ \text{C}$  (Fig. 2, middle). That is, the microalgae culture decreased the freezing temperature by approximately  $1^\circ \text{C}$ . Supernatant liquid, for which the algal cells are centrifuged out, froze at  $-2.70 \pm 0.12^\circ \text{C}$ , see Fig. 2, right,



**Figure 2.** Change in temperature (°C) over time (minutes). (Left) artificial seawater growth media (36 PSU Instant Ocean® with  $f/2 + Si$  media) without diatoms, (middle) culture of diatoms in artificial seawater, (right) supernatant of the diatom in artificial seawater culture, Each panel shows the temperature time series for four to five replicates. Note the initial drop in temperature to a supercooled state, followed by an abrupt increase to the freezing temperature. Data is recorded at one minute intervals.

again lowering by approximately  $1^\circ\text{C}$  the freezing point compared to that of seawater solution alone, suggesting that algal cells, and any other material that centrifuges out, do not contribute substantially to the drop in freezing temperature. Using formula A6, an equivalent solution without supernatant would require an increase of approximately an additional 18 ppt in salinity. That is, the equivalent salinity would be approximately 150% that of seawater solution. While we don't know what dissolved compounds are in the supernatant, and the note that culture conditions differ from those of a sea ice environment, this suggests that there may be a significant contribution to lowering of freezing point from extracellular osmolytes.

## 4.2 Model: Biofilm Community

Biofilms, suspected to be commonly present in sea ice (Krembs et al., 2002; Roukaerts et al., 2021), are communities of organisms anchored in place by self-secreted extracellular matrices of polymers and other substances (Hall-Stoodley et al., 2004). In environmental settings they are often observed in a pseudo-steady state in which inputs and outputs roughly balance; we approximate sea ice biofilms to be in such a state here, dropping equations (A4)-(A5) and setting  $B_{\text{bulk}}(\mathbf{x}, t) = 1$  in equation (A3). The osmolyte production function in (A3) is defined to be

$$P(S_{\text{brine}}, O_{\text{brine}}, O_{\text{bulk}}, B_{\text{bulk}}) = Y_{\text{osmo}} r_O e^{-(S_{\text{brine}} + Y_{\text{sal}} O_{\text{brine}})/\lambda} B_{\text{bulk}} - \gamma_{\text{osmo}} O_{\text{bulk}} \quad (1)$$

where  $\lambda$  is a salinity inhibition coefficient,  $\gamma_{\text{osmo}}$  is a decay coefficient, and  $Y_{\text{osmo}}$  is a yield coefficient. Production is proportional to bulk microbial concentration with a decay proportional to bulk osmolyte concentration. The exponential encodes production inhibition at high effective salinity measured relative to  $\lambda$ . Production rate coefficient  $r_O$  is set at  $r_O = 0.1 \text{ hr}^{-1}$  for the computation discussed below. See Appendix C for other parameter values.

See Figure 3, left column, for results from a representative simulation. To set initial conditions, we simulate “inert” ice, that is, ice without biological activity ( $B(z, t) = 0$  and  $O_{\text{bulk}}(z, t) = 0$ ) until an approximate steady state is reached, and use the resulting values of  $S_{\text{bulk}}$  and  $H_{\text{bulk}}$  as initial-initial ( $t = 0$ ) conditions for the biofilm computation, together with  $B(z, 0) = 1$ .

After a transient period due to the abrupt introduction of biological activity, the simulation settles on an approximately periodic behavior with bursts of convective activity in an approximately 10 cm thick layer at the bottom, and an additional tapering flow up to an additional (approximately) 10 cm, Fig. 3 top left. Note also the slow build up of osmolyte in a layer near the top of the convective region, Fig. 3 middle left, where convection is slow due to low ice permeability; this layer eventually discharges as well (not shown). Thickness of the microbially-induced convective layer is limited by low permeability above this layer, which, effectively, does not allow convective draining to proceed fast enough to avoid salinity inhibition (recall the exponential inhibition term in (1)). Non-biologically sourced ~~salt concentrations~~ salinities reach levels similar to osmolyte concentrations, Fig. 3 bottom left (although levels are higher at the bottom where seawater enters the ice).

### 4.3 Model: Planktonic Community

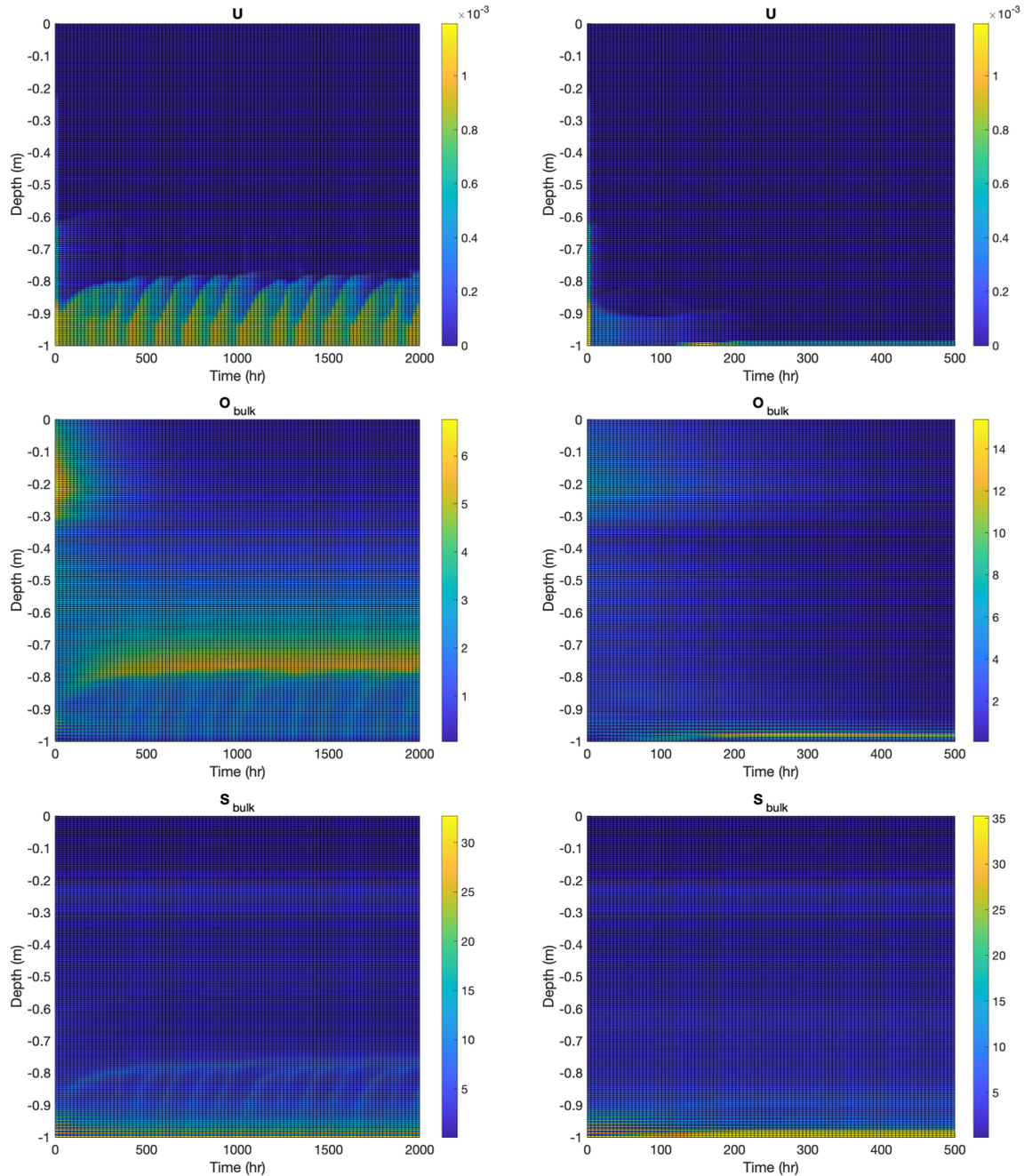
Alternatively, we consider the possibility of a microbial community in a planktonic state, i.e., not ice-attached, and allow this population to advect passively with any present convective flow. The ice ~~sheet-column~~ domain is appended with a sea compartment, near but below the ice, from which organisms can advect upward into the ice when the velocity  $U$  is non-zero, and into which organisms are ejected from ice regions where  $Ra$  exceeds  $Ra_{crit}$ . In addition, we include a nutrient supply, necessary (in the planktonic model) for microorganism growth, that is advectively transported upward into the ice with velocity  $U$  from a constant concentration source in the sea compartment. Osmolyte production function  $P$  was chosen as in (1), with  $r_O = 0.2 \text{ h}^{-1}$ . Again, see Appendix C for other parameter values. The nutrient usage term  $R$  in equations (A4) and (A5) is chosen to be of Monod form, commonly chosen for microbial systems,

$$R(S_{brine}, O_{brine}, C_{brine}) = r_B \frac{C_{brine}}{K + C_{brine}} e^{-(S_{brine} + Y_{sal} O_{brine})/\lambda}, \quad (2)$$

where  $K$  is a half-saturation (set to a large value in order to maintain first-order nutrient kinetics). The exponential term is the same effective salinity inhibition factor as in (1), for consistency. That is, inhibition of microbial activity is assumed to affect osmolyte production and nutrient usage equally. See Appendix A6 for additional details. Note that we did not see a significant nutrient depletion in the results, though, even with  $K$  large enough so that first order kinetics were maintained. This is likely in part due to ice formation inducing brine concentration. Also note that the presence of nutrients in sea ice, in the layer near the interface with the sea, has been reported to be correlated with high biomass concentrations, referred to in Roukaerts et al. (2021) as the sea-ice nutrient paradox. Here the association arises as a consequence of microbially-induced convection.

Initial conditions are set as in the biofilm model computation described in Section 4.2, except with  $B_{bulk}(z, 0) = |z|/L$ . The form of the initial biomass, with microbes concentrated more towards the ice-sea interface, is so as to allow the microbes a chance to form and sustain a convective layer. It also, ~~quite-though~~ roughly, approximates the distribution of microorganisms that might be expected ~~to a developed ice sheet~~ in a developed sea ice column.

The planktonic model is, as is the biofilm model, capable of formation of a convective layer at the bottom of the ice ~~sheet-column~~, though only for sufficiently large rate of osmolyte production – otherwise microorganisms are excluded to the sea compartment only. In fact, setting  $r_O = 0.1 \text{ h}^{-1}$ , as in the biofilm computation, results in washout and no convective layer persisting. See Fig. 3, right column, for a representative simulation with  $r_O = 0.2 \text{ h}^{-1}$  and resulting convective layer forma-



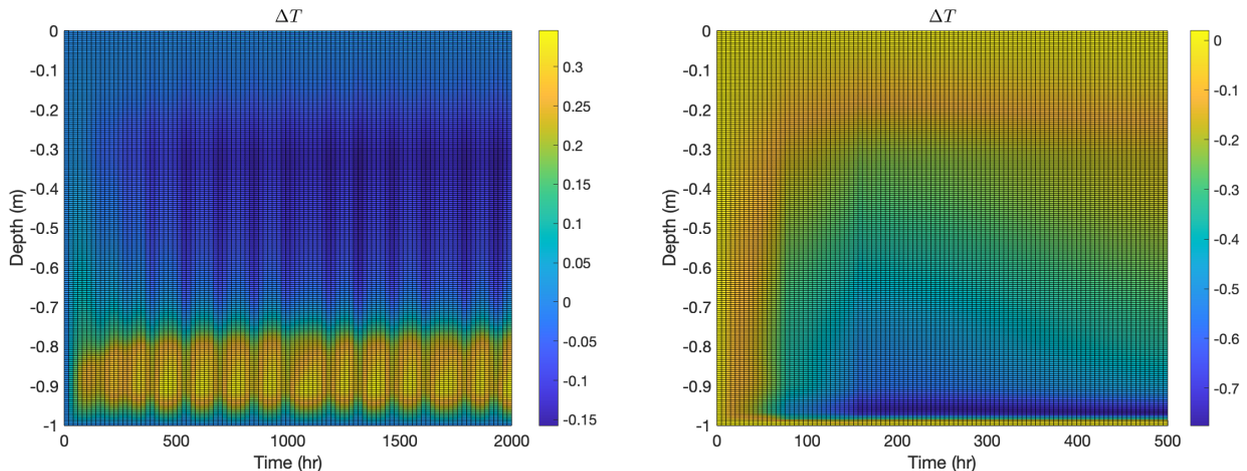
**Figure 3.** Computational results for a 1 m thick ice sheet-column with biofilm population for 2000 h (left) and planktonic population for 500 h (right). Vertical axes indicate depth from the ice-air interface at  $z = 0$ . Initial conditions for both computations consist of an abiotic system at approximate steady state, with biological perturbation added. Transients seen in the early hours of the simulation are consequences of this transition. (Top) vertical velocity (m/h). (Middle) bulk osmolyte concentration (ppt). (Bottom) bulk salt-concentration-salinity (ppt).

tion. Results are different in some important ways from those in the biofilm example. Note that an approximately 3-5 cm thick, steady convective layer forms at the very bottom of the ice sheet-column, with thickness here determined largely by the Rayleigh condition through the height  $h$  (see (A14)) of the top of the layer. That is, the top of the convective layer occurs approximately where the gravitational potential energy, from increased density, and decreased fluid viscosity, from increased permeability, is sufficient so as to trigger a convective instability. Because microorganisms advect with flow, unlike in the biofilm case, they then wash out back into the sea compartment. That is, in the planktonic model, microorganisms can only penetrate ice where upward flow advects them, whereas in the biofilm case, organisms can sit in non-convective ice regions, for a time, producing osmolyte until concentrations become large enough to trigger fluid instability, thus allowing, periodically, a significantly thicker advective layer.

#### 4.4 Model: Transport Effects

Change in temperature  $T(z, t) - T(z, 0)$  is shown in Figure 4. This quantity is the difference between the ice sheet-temperature, with biofilm of planktonic populations (at time  $t$ ) and the temperature of the initial  $t = 0$  ice sheet-column approximate steady state, computed without biofilm (left) or planktonic (right) populations. In the biofilm case (left), as might be expected, temperature increases in the convective layer due to the influx of relatively warm seawater. Perhaps less expectedly, temperature above the convective layer actually decreases with introduction of biofilm activity. This seems to be a consequence of an increase in brine volume fraction there due to osmolyte production, which results in a drop in thermal conductivity, see Appendix A1. Note, as a consequence, heat flux density  $-k(\partial/\partial z)T$  through the non-convective part of the ice ~~sheet is reduced~~. (Thermal column is reduced (thermal diffusivity  $k$  is nearly unchanged in the upper parts of the ~~ice-sheet-column~~ since there is very little osmolyte present there.)). That is, the model predicts microbial activity can reduce heat transport through the ice sheet-a bit, despite advective transport in the convective layer. This prediction, though, may be dependent on model assumptions such as the form of the salinity inhibition factor in (1), and also may be overestimated since the liquidus relation used in computations tends to over estimate brine volume fraction at lower temperatures (see Appendix A3). In the planktonic model case (right), the temperature increment  $T(z, t) - T(z, 0)$  is negative throughout the ice sheet-column, even in the convective layer, again apparently as a consequence of an increase in brine volume fraction due to osmolyte production, which results in a drop in thermal conductivity. As in the biofilm model case, heat flux density through the non-convective part of the ice sheet-column is reduced.

In Figure 5 ~~we plot shows a plot of~~ the ratio  $\Lambda$  of outflux to osmolyte production,  $\Lambda = \alpha \max(0, Ra - Ra_{\text{crit}})/P$ , see equations (A3) and (A16), for both the biofilm and planktonic model computations. Note that in both cases, the flows seen in Fig. 3 are largely driven by thin layers where the Rayleigh condition  $Ra > Ra_{\text{crit}}$  is satisfied. It is also interesting to note that, in these layers, the ratio  $\Lambda \approx 1$  suggesting that the flow rate is effectively determined by the osmolyte production rate. That is, microorganisms can regulate the flow rate through the ice (and hence nutrient supply and byproduct expulsion) by their osmolyte production rate. This is not surprising: if flow rate is smaller than osmolyte production rate, then osmolyte will accumulate which will increase permeability, leading to higher flow rate, and vice-versa in the case that flow rate is higher than osmolyte production rate. In this way the ice community can function something like a self-regulated chemostat: by setting

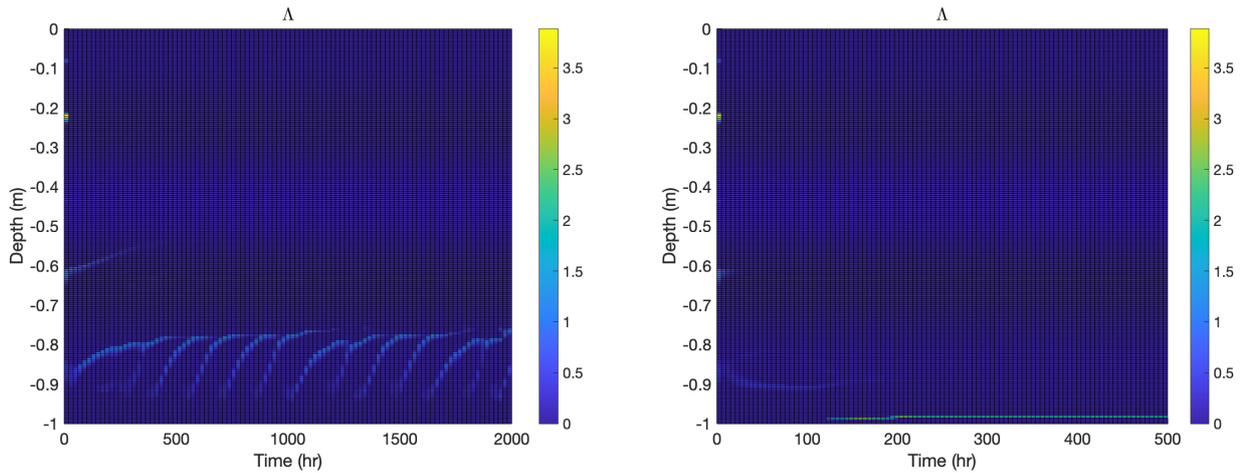


**Figure 4.** Temperature change  $T(z, t) - T(z, 0)$  after addition of microbes for the same computations as in Fig. 3, again with biofilm population for 2000 h (left) and planktonic population for 500 h (right). Vertical axes indicate depth from the ice-air interface at  $z = 0$ . In the biofilm case, (1) temperature increases in the convective region because of inflow of relatively warm seawater and (2) temperature decreases in the mid-ice region likely because of smaller diffusivity (due to increased brine volume fraction). In the planktonic case, temperature decreases throughout the ice [sheetcolumn](#), although the decrement is relatively small in the thin convective region at the bottom of the [sheetcolumn](#).

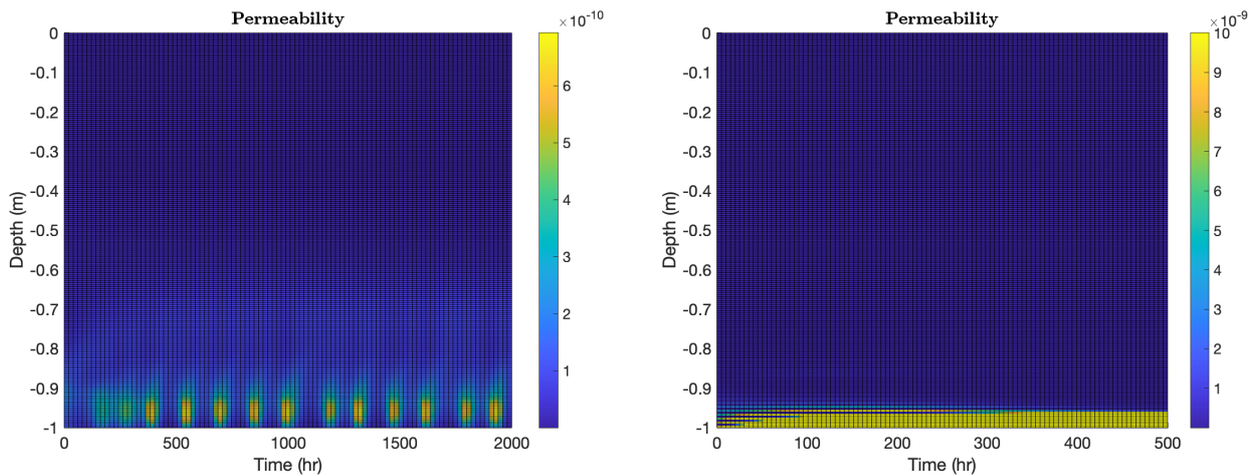
the osmolyte production rate, they also set the convective flux rate and, as a consequence, the rates of inflow of nutrients and outflow of byproducts.

265 In the context of transport, it is also interesting to look at microbially-induced changes in ice permeability, see Figure 6. Permeability relates flow rate through a material to flow driving force, see Darcy’s equation (A13), and thus is the central material property for understanding fluid flow through sea ice. We see significant impacts on ice structure in both the biofilm and planktonic cases. Microbes, at least as predicted by the model presented here, can produce qualitative changes in ice structure with regards to fluid flow properties, and do this simply through production of osmolytes. Changing the local effective salinity  
 270 regulates brine volume fraction, which is an important regulator for permeability, which in turn is an important regulator for transport rate of nutrients and biproducts.

The two model cases are different, though. “Biofilm” ice permeability exhibits the same temporal periodicity as observed in the induced conductive flow (the activity bursts in Fig. 6, left panel, match well with those in Fig. 3, top left panel), but note that smaller permeability changes also occur well into the ice [sheetcolumn](#), following biofilm activity. Permeability changes  
 275 in “planktonic” ice are limited to a small bottom layer where microbes are found. On the other hand, permeability levels in that layer are relatively large, due in large part to active convection of warm ocean water. Note that permeability impacts actually spread beyond the convective regions, in both cases, largely because the harmonic averaging used in formula (A15) is somewhat blurring in nature.



**Figure 5.** The ratio  $\Lambda = \alpha \max(0, Ra - Ra_{crit})/P$  for the two computations shown in Figures 3 and 4. In both cases, flow is caused by thin regions, where  $\Lambda \approx 1$ , that drive Rayleigh-Taylor convection, suggesting that microbes can control convective transport rate through osmolyte production  $P$ .



**Figure 6.** Ice permeability ( $m^2$ ), computed as in formula (A15), for the same computations as in Fig. 3, again with biofilm population for 2000 h (left) and planktonic population for 500 h (right). Vertical axes indicate depth from the ice-air interface at  $z = 0$ . In the biofilm case, (1) temperature increases in the convective region because of inflow of relatively warm seawater and (2) temperature decreases in the mid-ice region likely because of smaller diffusivity (due to increased brine volume fraction). In the planktonic case, temperature decreases throughout the ice sheetcolumn, although the decrement is relatively small in the thin convective region at the bottom of the sheetcolumn.

## 5 Conclusions

280 We have presented a simplified model of microbially-induced convective flow within sea ice, based on microbial osmolyte production. The new model is an augmented version of a previous, abiotic one for density-driven salt draining that relies on entrainment of new seawater via advancing ice sheet-column growth. Here we suggest that microbes, through production of osmolytes or substances with osmolytic properties, can induce continuing convection in a bottom layer of sea ice by themselves, essentially, even after inorganic salt would have largely drained out of biologically inert ice. This effect is observed for simple  
285 models of both biofilm and planktonic microbial communities, and potentially provides a mechanism to obtain nutrients and remove byproducts at a self-regulated rate. At the same time, a side effect of this process is a decrease in overall ice sheet column thermal conductivity with the consequence of reduced heat transport, though this effect may be small.

The proposed mechanism is relatively simple: sea ice organisms produce osmolytes that increase local effective salinity, which in turn increase local brine volume fraction, opening the ice to fluid flow. It is also passively and stably self-regulating:  
290 the fluid flow rate through the ice must match, on average, the osmolyte production rate. If not, overly slow fluid flow would allow osmolytes to accumulate which further increases ice permeability, while overly fast fluid flow would flush osmolytes too fast and thus would lower permeability.

In support, using a *C. neogracilis* culture, a diatom observed in the field in sea ice, we measured a decrease in the freezing point of approximately 1° C. Although the *in situ* community is rarely monospecific, this species is frequently reported as dominant, e.g. ~~(Katsuki et al., 2009; Crawford et al., 2018). While this result is suggestive, further studies looking at other species or species assemblages, the composition and role of osmolytes, and the direct role of cells on the freezing point are needed. Also, notably,~~ Katsuki et al. (2009); Crawford et al. (2018). Notably, however, we performed short-term measurements (30-40 min) using lab cultured organisms, reducing opportunities for adaptation and acclimation, while conditions closer to those found *in situ* ~~could have produced~~ may produce a different response ~~from the microalgae. So, while the result is suggestive,~~  
300 further studies looking at other species or species assemblages, the composition and role of osmolytes, and the direct role of cells on the freezing point are needed.

The simulations presented set initial conditions using a simulation for an established, non-advecting (abiotic) ice sheetcolumn, thus imitating to an extent a winter-spring transition from biologically inactive to biologically active ice. This differs from what one might expect in the late fall during formation of first year ice, where a microbially-induced convective layer would form  
305 along with the growing ice sheetcolumn. We did not investigate this latter possibility here because of its coupling with ice sheet column growth, which introduces additional complications that can obscure the main point. Nevertheless, we would expect that a similar microbially-influenced convective layer could be present, at least in model simulations.

While the model significantly simplifies what are likely complicated fluid dynamics, and we cannot rule out dependence on model simplifications, the underlying mechanism nevertheless seems robust; microbial osmolyte production can eventually  
310 increase ice permeability until advective flow is triggered, and the two should balance over time. Another prediction of note: the model indicates that biofilm, i.e., ice-attached organisms, increase the advective layer thickness in comparison to planktonic

ones, as planktonic organisms tend to be washed out by the advection. Again, while this observation may depend on details of the fluid mechanics, it seems plausible.

315 Conversely, more quantitative predictions, e.g., thicknesses of convective layers, are likely generally dependent on parameter choices at least some of which are uncertain, e.g., osmolyte production rate  $r_O$  and osmolyte salinity yield  $Y_{\text{sal}}$ . Though to the extent possible we attempt to choose reasonable values, and we note that quantitative predictions in turn appear reasonable, nevertheless we would be cautious about quantitative results. This is already true within the constraints of the 1D model, even before considering the significant simplifications made in constructing it. It is also worth noting that we assume a steady environment for the purposes of demonstrating that the model can predict constant or, in the biofilm case, periodic behavior  
320 induced by microbial activity. Even leaving aside model predictions, the actual sea ice environment is far from steady, exhibiting temperature variations on various time scales, for example. Mechanical stressing by the underlying sea dynamics, not included in the model, are also likely to have significant impact.

Results suggest plausibility of significant microbial impact on sea ice structural and material properties, beyond albedo effects, even though the presentation here is conservative in some ways. Notably, we impose a relatively large 18°C temperature  
325 difference between the air and ocean interfaces to argue robustness. This localizes effects to the bottom of the ice [sheetcolumn](#), but for less extreme temperature differences it may be possible, even, for microbes to open channels and maintain activity through the entire ice [sheetcolumn](#), which could have important implications on sea ice both as a microbial habitat and also as a nutrient cache. Sea ice may indeed be a rich and productive microbial environment. Indeed, recent observations support this possibility (Audh et al., 2023). On another note, model results suggest the possibility of significant amounts of microbially pro-  
330 duced osmolyte. Indeed, as an example, high concentrations of DMSP and DMS have been observed in sea ice environments, and their biological origins has attracted attention of astrobiologists (Madhusudhan et al., 2025).

In the broader picture, moving from static sea ice models to dynamic, mushy-layer sea-ice in climate models has shown significant influence on results, including increasing snow-ice and coastal ice production as well as enhancement of surface ocean salinity due to brine rejection (Turner and Hunke, 2015; Bailey et al., 2020; Singh et al., 2021; DuVivier et al., 2021), and also  
335 more accurate predictions of both low-level cloud cover around Antarctic coasts and atmospheric energy input (DuVivier et al., 2021). Results reported here suggest that including biological processes, specifically microbial modification of ice properties, may also merit attention.

## Appendix A: Equations

### A1 Conserved Scalar Fields

340 The model equations are largely standard (although we track bulk enthalpy  $H_{\text{bulk}}$  rather than, directly, temperature  $T$ , following Notz (2006); Griewank and Notz (2013)) except with the addition of transport equations for osmolyte, nutrient, and

biomass concentrations ( $O_{\text{bulk}}$ ,  $C_{\text{bulk}}$ , and  $B_{\text{bulk}}$  respectively). These quantities are governed by transport equations

$$\frac{\partial}{\partial t}(\rho H_{\text{bulk}}) = -\nabla \cdot (\rho H_{\text{brine}} \mathbf{U}) + \nabla \cdot (k \nabla T), \quad (\text{A1})$$

$$\frac{\partial}{\partial t} S_{\text{bulk}} = -\nabla \cdot (S_{\text{brine}} \mathbf{U}) + \nabla \cdot (\kappa_S \nabla S_{\text{brine}}), \quad (\text{A2})$$

$$345 \quad \frac{\partial}{\partial t} O_{\text{bulk}} = -\nabla \cdot (O_{\text{brine}} \mathbf{U}) + \nabla \cdot (\kappa_O \nabla O_{\text{brine}}) + P(S_{\text{brine}}, O_{\text{brine}}, C_{\text{brine}}, B_{\text{bulk}}), \quad (\text{A3})$$

$$\frac{\partial}{\partial t} C_{\text{bulk}} = -\nabla \cdot (C_{\text{brine}} \mathbf{U}) + \nabla \cdot (\kappa_C \nabla C_{\text{brine}}) - R(S_{\text{brine}}, O_{\text{brine}}, C_{\text{brine}}) B_{\text{bulk}}, \quad (\text{A4})$$

$$\frac{\partial}{\partial t} B_{\text{bulk}} = -\nabla \cdot (B_{\text{brine}} \mathbf{U}) + \nabla \cdot (\kappa_B \nabla B_{\text{brine}}) + (Y_B R(S_{\text{brine}}, O_{\text{brine}}, C_{\text{brine}}) - \gamma_B) B_{\text{bulk}}, \quad (\text{A5})$$

where  $k$  is a thermal diffusivity and the various  $\kappa$ 's are material diffusivities. Yield coefficient  $Y_B$  translate substrate usage into production of new biomass, and  $\gamma_B$  is a microbial decay rate. Functions  $P$  and  $R$  are osmolyte production and limiting nutrient  
 350 reaction terms, respectively, discussed further below, with examples of particular choices given in (1) and (2) respectively. Note that both can be inhibited at sufficiently high concentrations of salt and osmolyte, as a result of microbial activity inhibition. The last term in equation (A5) is the net biomass growth term, with growth term  $Y_B R$  supposed proportional to nutrient uptake  $R$  and including first order decay with rate  $\gamma_B$ . Definitions for bulk and brine quantities are discussed below.

We neglect the diffusion terms in (A2)-(A5), as is common practice for an ideal mushy layer (Worster, 1992, 1997). Veloc-  
 355 ity  $\mathbf{U}$  is discussed below. The thermal conductivity depends on brine volume fraction  $\phi_b$ , which we approximate in volume averaged form as  $k = k_b \phi_b + k_i (1 - \phi_b)$  where  $k_b$ ,  $k_i$  are the thermal conductivities in brine and ice, respectively. Note that  $k_i$  is roughly a factor of 10 larger than  $k_b$  (Kannuluik and Carman, 1951; Yen, 1981; Thomas and Dieckmann, 2008), so that an increase in brine volume fraction  $\phi_b$  results in decreased thermal conductivity. Density  $\rho$  in equation (A1) is also in principle a weighted average over brine and ice densities, but we neglect the difference between the two densities (approximately 10%),  
 360 again as is frequently done (Worster, 1997). A consequence of this supposition is that we can impose  $\nabla \cdot \mathbf{U} = 0$ .

Equations (A2)-(A5) take no-flux boundary conditions at the ice-air interface (i.e., normal derivatives across the interface are zero), and, at the ice-ocean interface, flux is set to the inflowing advective flux, e.g.  $S_{\text{sea}} \mathbf{U} \cdot \mathbf{n}$  for equation (A2) where  $\mathbf{n}$  is an in-pointing interface normal vector. Equation (A1) takes Dirichlet boundary conditions using equation (A9) together with given values for air and sea temperatures ( $T = -20^\circ \text{C}$  and  $T = -2^\circ \text{C}$ , respectively). Note that these conditions depend on  
 365 brine volume fraction  $\phi_b$ , but are approximated here using  $\phi_b = 1$  at the ice-ocean interface and  $\phi_b = 0$  at the ice-air interface.

## A2 Salinities and Volume Fractions

Seawater is a solution of water and many salt species, as well as other dissolved and undissolved contaminants, but we simplify by dividing into water, microbially produced antifreeze chemicals (e.g., DMSP) lumped together under the designation “osmolyte”, and other dissolved chemical species lumped together under the designation “salt”. Undissolved contaminants,  
 370 e.g., microbial cells, extracellular polymers, etc., are neglected here though could certainly effect freezing properties. In a unit

control volume  $V$ , we define (partial) bulk salt and osmolyte salinities (ppt) as

$$S_{\text{bulk}} = \frac{m_{\text{salt}}}{m_{\text{water},\ell} + m_{\text{water},s}} \cdot 10^{-3} \text{ ppt},$$

$$O_{\text{bulk}} = \frac{m_{\text{osmo}}}{m_{\text{water},\ell} + m_{\text{water},s}} \cdot 10^{-3} \text{ ppt},$$

where  $m_{\text{salt}}$  and  $m_{\text{osmo}}$  are the control volume masses of salt and osmolyte respectively, and  $m_{\text{water},\ell}$  and  $m_{\text{water},s}$  are the  
 375 control volume masses of liquid water and ice, respectively. Then (partial) brine volume fractions are defined as

$$S_{\text{brine}} = \frac{m_{\text{salt}}}{m_{\text{water},\ell}} \cdot 10^{-3} \text{ ppt} = \phi_{b,m}^{-1} S_{\text{bulk}},$$

$$O_{\text{brine}} = \frac{m_{\text{osmo}}}{m_{\text{water},\ell}} \cdot 10^{-3} \text{ ppt} = \phi_{b,m}^{-1} O_{\text{bulk}},$$

where

$$\phi_{b,m} = \frac{m_{\text{water},\ell}}{m_{\text{water},\ell} + m_{\text{water},s}}$$

380 is the brine mass fraction. Note that we are neglecting  $m_{\text{salt}}$  and  $m_{\text{osmo}}$  contributions to total mass as small. Finally, we define  $\phi_b$  to be the brine volume fraction, i.e., the ratio of brine volume to total volume in the unit control volume, and make the approximation  $\phi_b = \phi_{b,m}$ .

These definitions, written in terms of mass, can also be written in terms of densities by multiplying the right-hand side by  $V^{-1}/V^{-1}$ , where  $V$  is a unit volume. Also note that, with the approximate equivalence of liquid and ice water volume,  
 385 the quantity  $V^{-1}(m_{\text{water},\ell} + m_{\text{water},s})$  can be considered constant (in both space and time), and hence  $S_{\text{bulk}}$  and  $O_{\text{bulk}}$  are, approximately, functions of salt and osmolyte concentrations only. A similar observation holds for volume fraction  $\phi_b$ .

### A3 Liquidus Relations

Solvent (here water) freezing temperature is generally lowered in the presence of a solute – briny water has a lower freezing point than pure water. Freezing temperature and brine salinity in a simple solution (a solution with a single solvent species) have  
 390 been connected through a so-called *liquidus relation*  $S_{\text{brine}} = \mathcal{L}(T)$  (Worster, 1992, 1997; Feltham et al., 2006; Wells et al., 2011, 2019) where  $T$  is freezing temperature at concentration  $S_{\text{brine}}$ . Abiotic seawater is often approximated in this context as a simple solution of water and generic “sea salt” though seawater is more complicated, even without considering its insoluble component. A liquidus relation can still be applied, though it depends on the nature of the solution, and approximations of it have been constructed for seawater as a function of salinity (Notz, 2006).

395 The liquidus function  $\mathcal{L}$  is generally determined empirically for a given solution. For computations, we use here a linear approximation

$$\mathcal{L}(T) = -T/0.05411 \tag{A6}$$

which generally works well for seawater at temperatures near or above  $-6^\circ\text{C}$ , approximately; for lower temperatures; (A6) over estimates brine concentration (Notz, 2006), but the effect on results here is small because ice permeability is too low, even  
 400 with more accurate liquidus approximations, to allow significant flow channels to occur in regions with low temperatures.

Microbes are able in certain cases to produce osmolytic compounds to protect themselves from freezing. We suppose that these osmolytes can be found extracellularly, though we don't distinguish between active and passive (e.g., through cell lysing) excretion. For reference, note the reported wide-spread and significant concentrations of the compound DMSP associated with antarctic ocean algae and cyanobacteria communities - we aren't aware of evidence that it is, or isn't, actively excreted, but  
 405 it has been measured in large concentrations regardless. To introduce effects of microbes, we divide solutes into two types, abiotic (e.g., inorganic salts) and biotic. In fact, nonsoluble biotic contaminants might also impact freezing temperature, but are not separately considered. That is, we consider a brine system with two solute types, one of which is subject to microbial control.

Extension of the liquidus relation to two (or more) solute types requires a short computation, as follows. Denoting abiotic  
 410 and biotic concentrations by  $S_{\text{brine}}$  and  $O_{\text{brine}}$  respectively, we use the liquidus approximation as in the abiotic case in the form

$$S_{\text{brine}} + Y_{\text{sal}}O_{\text{brine}} = \mathcal{L}(T) \quad (\text{A7})$$

where  $Y_{\text{sal}}$  is a salinity yield coefficient. That is, we don't assume that the biotic osmolyte has the same effect on freezing point, per mass, as the abiotic one. Note then the pointwise requirement  $S_{\text{brine}}/O_{\text{brine}} = S_{\text{bulk}}/O_{\text{bulk}}$ , together with (A7), results in  
 415 the pair of liquidus-like relations

$$O_{\text{brine}} = \frac{O_{\text{bulk}}}{S_{\text{bulk}} + Y_{\text{sal}}O_{\text{bulk}}} \mathcal{L}(T), \quad S_{\text{brine}} = \frac{S_{\text{bulk}}}{S_{\text{bulk}} + Y_{\text{sal}}O_{\text{bulk}}} \mathcal{L}(T). \quad (\text{A8})$$

**Note 1.** The value of the salinity yield  $Y_{\text{sal}}$  is unknown (and, indeed, effective salinity is an introduced idea here). Based on the results of Section 4.1, we suppose, lacking other measurements, that biologically-induced osmolyte is relatively effective in comparison to abiotic salinity and so have set  $Y_{\text{sal}} = 5$  in computations. Note that we have not observed significant qualitative  
 420 differences for other  $O(1)$  choices of  $Y_{\text{sal}}$  in computations.

**Note 2.** The liquidus relation (A6) is an empirical one, intended for seawater relatively near to  $-1.8^\circ\text{C}$  in temperature. We don't know how accurate it is when osmolyte is included as in (A7), even in this same temperature range, as we don't have available a liquidus relation for the combined seawater plus microbially-produced osmolyte system. This is another reason to be cautious particularly about quantitative model predictions.

#### 425 **A4 Formulas for Brine Volume Fraction and Concentrations**

We divide the various fields by slow and fast time scales. Bulk conserved quantities  $H_{\text{bulk}}$ ,  $S_{\text{bulk}}$ ,  $O_{\text{bulk}}$ ,  $C_{\text{bulk}}$ ,  $B_{\text{bulk}}$  are governed by equations (A1)-(A5), and change on the relatively slow diffusive and advective time scales. Locally, it is commonly assumed that an ice-brine balance, consistent with liquidus relations (A8), is approached on a much faster time scale. That is, relations (A8) are imposed as constraints. Additionally, enthalpy and temperature are related by the approximation (Notz, 2006)

$$430 \quad H_{\text{bulk}} = c_s T - (1 - \phi_b)L_0, \quad (\text{A9})$$

where  $c_s$  is ice heat capacity, approximated to be a constant, independent of temperature and salinity, and  $L_0$  is the latent heat of fusion. Note that  $H_{\text{bulk}}|_{T=0} = 0$ . Lastly, assuming that the ice phase is pure water, then

$$S_{\text{bulk}} = S_{\text{brine}}\phi_b, \quad O_{\text{bulk}} = O_{\text{brine}}\phi_b, \quad B_{\text{bulk}} = B_{\text{brine}}\phi_b. \quad (\text{A10})$$

435 Combining these relations with (A7), we obtain

$$\phi_b = \frac{S_{\text{bulk}} + Y_{\text{sal}}O_{\text{bulk}}}{\mathcal{L}(T)} \quad (\text{A11})$$

$$H_{\text{bulk}} = c_s T - L_0 + L_0 \frac{S_{\text{bulk}} + Y_{\text{sal}}O_{\text{bulk}}}{\mathcal{L}(T)} \quad (\text{A12})$$

Given  $H_{\text{bulk}}$ ,  $S_{\text{bulk}}$  and  $O_{\text{bulk}}$ , the nonlinear equation (A12) is solved for  $T$ , and then (A11) for  $\phi_b$ , and then relations (A10) can be used to compute brine concentrations.

## 440 A5 Velocity

Dynamics within the sea ice are commonly modeled via a Darcy flow (Worster, 1992)

$$\mu \mathbf{U} = \Pi(-\nabla p + (\rho - \rho_0)\mathbf{g}) \quad (\text{A13})$$

where  $\mu$  is the dynamics viscosity,  $\mathbf{g} = -g\hat{\mathbf{z}}$  ( $g$  is the gravitational constant),  $\rho_0$  is a reference density (e.g., density of seawater just below the ice layer),  $\Pi$  is the sea ice permeability, and  $p$  and  $\mathbf{U}$  are pressure and velocity, respectively. Pointwise ice  
445 permeability  $\Pi$  is modeled as  $\Pi = \Pi(\phi_b) = \Pi_0\phi_b^3$ , although an average is used, see (A15) below. The velocity field  $\mathbf{U}$ , with units of length per time, can be understood in the context of Darcy flow as volume flux, with units of volume per time per area, obtained by averaging over a local flux surface through a porous medium.

Following Griewank and Notz (2013); Turner et al. (2013); Griewank and Notz (2015), we suppose that convective flow, induced by a buoyancy driven Rayleigh-Taylor instability, can occur in the ice [sheetcolumn](#). Presence/absence of such a flow  
450 is determined by the Raleigh number  $Ra$  (see Section A6), which can be defined to be the ratio of two time scales:

1. Diffusive time scale  $t_D$  is the time scale for diffusive transport of heat from sea to local brine, approximately  $t_D = h^2/\kappa$ , where  $\kappa = k/c\rho$  is the thermal diffusivity and  $h$  is the height above the ice-sea interface. Note: diffusive transport of heat, rather than salt, is appropriate here, as heat diffuses much more quickly and can reduce salinity by melting ice.
2. Convective time scale  $t_C$  is the time scale for local brine, at height  $h$  above sea, to reach the ocean by buoyancy-driven  
455 flow. Darcy flow speed is estimated as  $g\Delta\rho\Pi/\mu$  where  $\Delta\rho$  is density difference between local brine and seawater, so that  $t_C = h\mu/(\Pi g\Delta\rho)$ .

We then define a local Raleigh number  $Ra$  as the nondimensional ratio

$$Ra = \frac{t_D}{t_C} = \frac{g\Delta\rho\Pi h}{\kappa\mu} = \frac{gc\rho\Delta\rho\Pi h}{k\mu} \quad (\text{A14})$$

where  $\Delta\rho$  is the density difference between local brine and seawater. Permeability  $\Pi = \Pi(\mathbf{x})$  is spatially variable. We follow  
 460 here the recommendation in Jones and Worster (2014) and replace  $\Pi$  in our 1D computations by its harmonic average

$$\Pi_{\text{harm}}(z) = \left[ \frac{1}{z_0 - z} \int_z^{z_0} \frac{1}{\Pi(\phi_b(\hat{z}))} d\hat{z} \right]^{-1} \quad (\text{A15})$$

where  $z_0$  is the z-coordinate of the sea-ice interface.

We employ a Boussinesq approximation, which suppose  $\rho$  constant (again, also neglecting change in density between solid  
 and brine phases) except for buoyancy. That is,  $\rho$  is constant except for computation of buoyancy, in which case  $\rho = \rho_0(1 +$   
 465  $\beta\Delta S)$ , i.e.,  $\Delta\rho = \beta\rho_0\Delta S$  (where  $\rho_0$  is density at  $S = 0$  and  $\beta$  is a buoyancy conversion factor from ~~salt concentration~~ salinity  
 to density), so

$$\text{Ra} = \frac{gc\rho\beta\rho_0\Delta S\Pi h}{k\mu},$$

again using  $\Pi = \Pi_{\text{harm}}$  in our 1D computations. Here  $S$  is total ‘‘salt’’, i.e.,  $S = S_{\text{brine}} + O_{\text{brine}}$ ; recall that the osmolyte  
 concentration is normalized so that it has the same density properties as salt (the two are distinguished, rather, by effects  
 470 on freezing temperatures through the yield  $Y_{\text{sal}}$ ). Note that there is a conservation of mass issue – osmolyte is, ultimately,  
 generated from material present in the brine, and so new osmolyte might not change brine density. However, osmolyte solute  
 properties likely differ from those of its chemical constituents and so can have different effects on brine density. The inclusion  
 of osmolyte in the total salt  $S$  does not appear to have a large effect on results in any case.

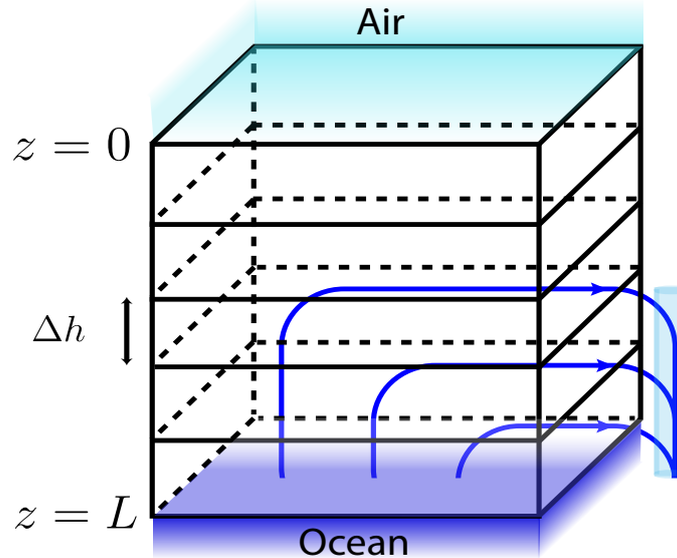
## A6 Parameterized Convection

475 The details of flow in sea ice generated by Rayleigh-Taylor instability are complex so instead we follow Griewank and Notz  
 (2013); Turner et al. (2013); Griewank and Notz (2015) and replace the full flow description  $\mathbf{U}$  in (A13) used in transport  
 equations (A1)-(A5) with a parameterized model of the form  $\mathbf{u}(\mathbf{x}, t) = (0, 0, U(z, t))$  where

$$U(z) = \int_{z_0}^z \nabla \cdot \mathbf{u}(\hat{z}) d\hat{z}, \quad \nabla \cdot \mathbf{u}(\hat{z}) = \alpha \max(0, \text{Ra} - \text{Ra}_{\text{crit}}). \quad (\text{A16})$$

Here  $z_0$  is the z-coordinate of the sea-ice interface, and  $\text{Ra}_{\text{crit}}$  is the critical Rayleigh instability value, see Table A1. Note  
 480 that  $\mathbf{u}$  neglects much of the detail of the full flow by only tracking upward velocity. The motivation and intuition is that cold,  
 salty downward flow occurs mostly in relatively large channels (formed because the temperature in colder downflow quickly  
 equilibrates with that in surrounding warmer ice, and then excess salinity causes that surrounding ice to melt) that empty rather  
 directly into the sea below with warmer less salty replacement fluid seeping upwards through surrounding mushy ice, see  
 Figure A1. The vertical velocity component  $U(z)$  accounts, in an averaged sense, for this upward replacement flow.

485 When convective flow is present, the planktonic microbial model (recall Section 4.3) includes a sea compartment microbe  
 biomass  $B_{\text{sea}}(t)$ , tracking microbes close to the bottom of the ice sheetcolumn, which receives influx from convectively drained



**Figure A1.** Parameterized flow diagram: flow moves upward (only) to account for drainage. (Adaptation from Turner et al. (2013).)

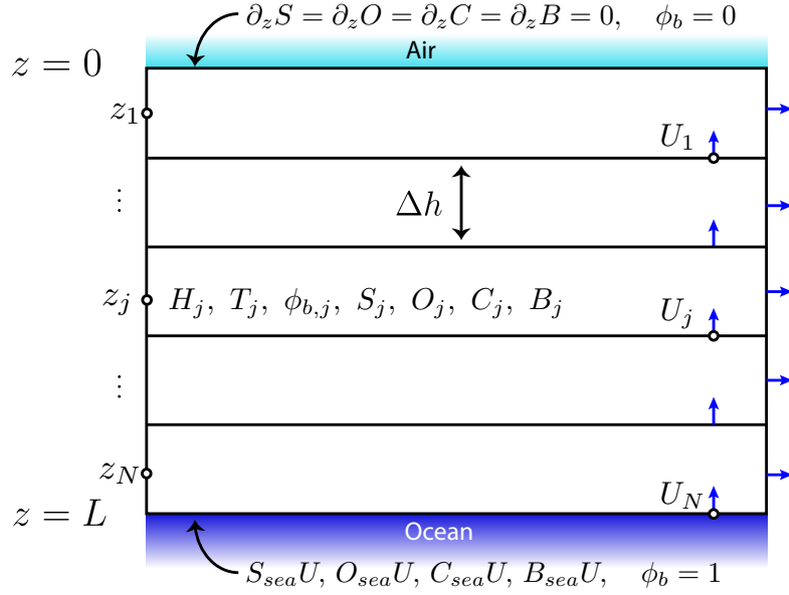
flow according to

$$\frac{d}{dt} B_{\text{sea}}(t) = \int_0^L B_{\text{brine}}(z, t) \nabla \cdot \mathbf{u}(z) dz + \delta(B_{\text{sea},0} - B_{\text{sea}}),$$

where  $\delta$  is a dilution coefficient. The second term represents mixing between the near ice sheet-column organisms and the background, given by constant  $B_{\text{sea},0}$ .

## Appendix B: Numerical Methods

Computations are conducted on a 1D domain  $z \in [0, L]$ ,  $x, y \in \mathcal{R}$ , see Figure B1. All quantities are assumed to depend spatially only on  $z$ , i.e., the system is effectively 1D, with  $z = 0$  corresponding to the upper, ice-air interface, and  $z = L$  corresponding to the lower, ice-sea interface. The domain is discretized into  $N$  uniform subintervals of length  $\Delta h$  (with  $N\Delta h = L$ ). In the computations illustrated in Figures 3-5, we set  $N = 200$ . Quantities of interest are discretized according to subintervals. Offset discretization is employed for the velocity  $U$  and for thermal conductivity, with values taken on subinterval boundaries, while other quantities (enthalpy, temperature, local Rayleigh number, brine volume fraction, and concentrations) are assigned values at each subinterval's midpoint  $z_i$ ,  $1 \leq i \leq N$ . Thermal conductivity on the interface is computed as a harmonic average of the thermal conductivity in the neighboring subintervals, following (Notz, 2006). Equations (A1)-(A5) are integrated using central differencing for second order terms, and using offset central differencing for advective terms, with first order explicit time integration. Due to use of an explicit method, a small time step  $\Delta t = 5 \cdot 10^{-6}$  hr was used.



**Figure B1.** Representation of the computational grid. Velocity and thermal conductivity take values on the interfaces between discretized intervals. Other quantities take values inside the intervals. No-flux and in-flux boundary conditions are indicated, along with Dirichlet conditions for  $\phi_b$ .

Each iteration consists of the following sequence of computations: given “current” values of each bulk field (the conserved, slowly varying quantities) either from initial conditions or from the previous iteration,

1. Temperature profile is updated using the current enthalpy profile via (A12).
- 505 2. Using the current profile  $S_{\text{bulk}} + Y_{\text{sal}}O_{\text{bulk}}$ , brine volume fraction is computed using (A11).
3. Current brine concentration fields  $S_{\text{brine}}$ ,  $O_{\text{brine}}$ ,  $C_{\text{brine}}$ , and  $B_{\text{brine}}$  are computed using (A10).
4. The local Rayleigh number for each subinterval is updated using the new  $S_{\text{brine}}$  profile and (A14) with, for subinterval  $i$ ,  $h = L - z_i$ .
5. The parameterized velocity profile is updated using (A16).
- 510 6. The slow fields  $H$ ,  $S_{\text{bulk}}$ ,  $O_{\text{bulk}}$ ,  $C_{\text{bulk}}$ , and  $B_{\text{bulk}}$  are updated using discretizations of (A1)-(A5).

Initial conditions for abiotic fields are set to be profiles obtained by running the code without microbes and osmolyte to an approximate steady state. Initial conditions for biomass bulk concentration  $B_{\text{bulk}}$  are  $B_{\text{bulk}}(z, 0) = 1$  for the biofilm case and  $B_{\text{bulk}}(z, 0) = |z|/L$  for the planktonic case. Initial osmolyte bulk concentrations are  $O_{\text{bulk}}(z, 0) = 0$  for the biofilm case and  $O_{\text{bulk}}(z, 0) = 5|z|/L$  for the planktonic case.

Symbol	Name	Value	Units	References
$B_{sea,0}$	Background microorganism density	0.001	kg/m <sup>3</sup>	
$c_b$	Brine heat capacity	3500	J/(kg K)	Notz (2006)
$c_s$	Ice heat capacity	2112	J/(kg K)	Notz (2006)
$g$	Gravitational acceleration constant	9.8	m/s <sup>2</sup>	
$k_b$	Thermal conductivity of brine	0.523	W/(m K)	Yen et al. (1991)
$k_s$	Thermal conductivity of ice	220	W/(m K)	Yen et al. (1991)
$K$	Half saturation constant	0.5	kg/m <sup>3</sup>	Yen et al. (1991)
$L_0$	Latent heat of fusion	333500	J/kg	Weast (1981)
$r_O$	Osmolyte production rate	various	1/h	
$Ra_{crit}$	Critical Rayleigh number	4.89	-	Griewank and Notz (2015)
$S_{sea}$	Sea salinity	34	ppt, (or g/kg)	
$Y_B$	Biomass yield coefficient	1	-	
$Y_{osmo}$	Osmolyte yield coefficient	1	-	
$Y_{sal}$	Salinity yield coefficient	5	-	
$\alpha$	Brine removal rate coefficient	0.03	1/h	
$\beta$	Solutal expansion coefficient	0.78237	kg/(m <sup>3</sup> ppt)	Turner et al. (2013)
$\gamma_B$	Microbe decay coefficient	0.02	1/h	
$\gamma_{osmo}$	Osmolyte decay coefficient	0.005	1/h	
$\delta$	dilution coefficient	0.1	1/h	
$\lambda$	Salinity inhibition coefficient	68	ppt, (or g/kg)	
$\mu$	Dynamic viscosity (water)	$2 \cdot 10^{-3}$	Pa s	Kestin et al. (1978)
$\Pi_0$	Permeability coefficient	$1 \cdot 10^{-8}$	m <sup>2</sup>	Freitag (1999); Jones and Worster (2014)
$\rho_b$	Density of seawater	1028	kg/m <sup>3</sup>	Millero and Poisson (1981)

**Table A1.** Model parameters. Some parameters are weakly temperature dependent – we approximate with (constant) values appropriate for seawater temperature.

## 515 Appendix C: Parameters and Fields

See Table A1 for a list of parameters, and Table A2 for a list of fields.

Symbol	Name	Units
$h$	Height above sea interface	m
$B_{\text{brine}}$	Brine biomass concentration	kg/m <sup>3</sup>
$B_{\text{bulk}}$	Bulk biomass concentration	kg/m <sup>3</sup>
$C_{\text{brine}}$	Brine limiting nutrient concentration	kg/m <sup>3</sup>
$C_{\text{bulk}}$	Bulk limiting nutrient concentration	kg/m <sup>3</sup>
$H$	Specific enthalpy	J/kg
$O_{\text{brine}}$	Brine osmolyte concentration	ppt, <u>(or g/kg)</u>
$O_{\text{bulk}}$	Bulk osmolyte concentration	ppt, <u>(or g/kg)</u>
$P$	Permeability	–
Ra	Local Rayleigh number	–
$S_{\text{brine}}$	Brine <del>salt concentration</del> <u>salinity</u>	ppt <u>(or g/kg)</u>
$S_{\text{bulk}}$	Bulk <del>salt concentration</del> <u>salinity</u>	ppt <u>(or g/kg)</u>
$U$	Vertical velocity	m/h
$\phi_b$	Brine volume fraction	–

**Table A2.** Model fields (all are functions of  $z$  and  $t$ ).

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520 *Competing interests.* The authors declare that they have no conflicts of interest.

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