**An atoms-to-mesoscale approach to ice-vapor surface dynamics with a quasi-liquid interface**

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**Abstract**

*We explore the hypothesis that a key factor in determining the dynamics and morphology of faceted ice-vapor surfaces is the quasi-liquid layer that forms at this interface at temperatures above . We do so by modeling the ice surface as a system of reaction-diffusion equations in which the time scales of quasi-liquid freezing and melting, horizontal diffusion, and exchanges with the vapor phase are made explicit. Model parameterizations are informed by atomistic (molecular dynamics) simulations, mesoscale simulations of the vapor field around growing and ablating ice crystals, and quantitative ice surface morphologies derived from scanning electron microscope experiments. The outcome is a more unified, predictive, and experimentally grounded picture of the dynamics and morphology of faceted ice-vapor surfaces than has previously been presented.*

**Plain Language Summary**

This is a mathematical exploration of the texture of ice surfaces on a microscopic scale. The main hypothesis is that this texture is governed by the behavior of a thin layer of water, intermediate between ice and liquid, that is known to form at the ice-air interface. We carry out this exploration by constructing a set of mathematical relationships that, when solved on a computer, reveal how parts of this quasi-liquid layer migrate to other regions of the ice surface, freeze, or evaporate into the air. The form and parameters of these equations are informed by insights from images of ice surfaces from scanning electron microscope experiments, and molecular-level modeling. We believe the insights obtained from this effort will be useful not only from a fundamental perspective, but also practically, especially in connection to the role of ice clouds in Earth's climate system.

1. **Introduction**

Why we care about ice crystal morphology … the crystals that make up cirrus clouds modulate Earth’s climate by reflecting or scattering sunlight before it reaches Earth’s surface, but the extent to which they do that depends on the morphology of those crystals. Hexagonal prisms dominate, but there are lots of variations on that theme: long and thin hexagonal needles, short and wide plates, sometimes indented at the ends, or hollowed, or even dendritic forms, like snowflakes. Their surfaces can be smooth or rough on a scale that matters to light of comparable wavelength (Järvinen et al. 2023).

Numerous models of ice surface morphology and dynamics during vapor depositional growth have been presented. Harrington and Pokrifka (2021) provide an excellent review. Of these, models rooted in atomistic structure and processes are appealing because of the evident connection between molecular structure on the one hand, and high-resolution microscopic observations (such as scanning electron microscopy) on the other; a familiar example is that crystals of water ice are understood to owe their hexagonal shape to the hexagonal structure of the unit cell of an ice lattice. Molecular Dynamics studies have contributed greatly to this effort (see, e.g., (Llombart, Noya, and MacDowell 2020))

Of particular interest here is the Burton-Cabrera-Frank theory of crystal growth, also called classical nucleation theory, or CNT. CNT’s atomistic view of the process of crystal growth goes along the following lines: when a gas-phase molecule (e.g., a water molecule) encounters a crystalline surface, it initially becomes attached to that surface as an “admolecule.” Not yet part of the crystal’s lattice, this admolecule diffuses across the surface until it meets one of two fates: either it fills an unoccupied position in the crystal lattice (often visualized as a step between ledges), or else it detaches from the surface and re-enters the gas phase. New layer formation is governed by a 2D nucleation process in which the step free energy plays a crucial role (Kuroda and Lacmann 1982).

CNT has long framed how we think about ice crystal growth from the vapor phase, but it is not reliable when it comes to describing the behavior of crystals placed in an inhomogeneous vapor field on a mesoscopic scale. To be specific, when a growing faceted ice crystal is situated in a supersaturated vapor field, it is easy to show that the surrounding water vapor concentration will be drawn down in such a way that facet intersections (i.e., crystal corners) will experience persistently higher vapor pressures than facet centers. CNT predicts that this would result in faster growth at those corners, leading ultimately to dendritic forms (e.g., snowflakes). However, under typical cirrus cloud conditions, it is known that hexagonal prisms resist that tendency, and instead maintain their faceted, hexagonal form.

An analogous circumstance arises when faceted crystals are situated in a subsaturated vapor field, in which case water vapor ablated from the surface distributes itself in such a way that crystal corners experience persistently *lower* vapor pressure compared to facet centers. CNT predicts that this would result in faster ablation at facet corners, eventually leading to a rounded crystal geometry. But our laboratory studies show that under such circumstances crystals typically maintain their faceted shape (albeit roughened on a micrometer scale), despite lower vapor conditions at corners.

Both cases just described can be summed up as a resilience of hexagonal prismatic faceting despite persistent inhomogeneities in the overlying vapor field. Classical nucleation theory has no satisfactory explanation for this resilience. (However, see Jayaprakash et al (1983)).

To add to that criticism is the lack of fealty of CNT to what is known about the molecular structure of the ice-vapor interface. When the temperature of ice rises above , both experiment and theoretical studies have shown that the interface is entirely covered by a quasi-liquid layer (QLL); molecular dynamics studies have shown that water molecule striking the surface are thermalized by the QLL efficiently and quickly (on a picosecond time scale) (Neshyba et al. 2009). On the other hand, recent experimental work has shown that there really are steps and ledges *underneath* the QLL, i.e., at the interface between the QLL and the underlying ice (Murata, Nagashima, and Sazaki 2019).

Bridging the gap between atomistic and mesoscale levels, therefore, would seem to require an atomistic model that begins with the existence of a QLL (above ), that preserves relevant parts of CNT as much as possible. One such attempt was presented by some of the authors in 2016 (Neshyba et al. 2016) (henceforth N2016), in the form of a dynamic quasi-liquid continuum model, referred to here as QLC-1. Because that model forms the foundation of the revision presented here (“QLC-2”), we summarize it next.

QLC-1 frames the problem of ice surface dynamics in terms of two mesoscale variables, and , which represent the thickness of the ice surface and its quasi-liquid part, respectively, at each point on the surface (see Fig. 1). Time evolution of these variables is governed by a pair of reaction-diffusion equations that represent the three processes indicated in Fig. 1, namely, (i) exchanges (deposition and ablation) of QLL molecules with the vapor phase, (ii) horizontal diffusion of QLL molecules across the ice surface, and (iii) interconversion of QLL molecules to/from the underlying ice.

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| **Figure 1**. Visual representation of mesoscale variables , , and , and processes affecting them, in QLC-1 (as well as the present revision, QLC-2) model. Dashed arrows represent processes affecting how these variables evolve over time. |

The main insight afforded by QLC-1 is that it provides a mechanism by which faceted ice crystal growth can occur. At the heart of that mechanism is a process N2016 termed “diffusive slowdown,” which can be summarized as follows:

1. At the micrometer level, the QLL can be thought of as consisting of a continuum of microstates, ranging from a thin, less-volatile microstate labeled surface I, to a thick, more-volatile one labeled surface II. The difference in these volatilities is quantified in QLC-1 as a difference in equilibrium supersaturation, . Evidence for the existence of these microstates, and their difference in volatility, has emerged mainly from molecular dynamics simulations (N2016).
2. In a growing ice crystal, each time a new layer forms, a new pair of these microstates appears on the surface. Since new layers typically form at facet corners (where the water vapor concentration is highest), it is also the case that the horizontal distance between newly-formed ice layers is smaller at facet corners compared to facet centers. In this paper, we designate this distance as “”; in Fig. 2(a) we see that .

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| **Figure 2**. An ice surface covered by QLL, as simulated by QLM-2. |

1. Horizontal diffusion moves quasi-liquid away from thicker, surface II-like regions of the surface, and toward thinner, surface I-like regions. However, because surface I accounts for a smaller fraction of the total surface area (as seen in Fig. 2(b)), diffusion has a greater proportional effect on it. As a consequence, diffusion leads to an increase in the average volatility of the surface, causing the surface as a whole to experience a net “diffusive slowdown” in its growth rate. (The reader is referred to N2016 for a quantitative version of this argument.)

A second process identified in N2016 is that diffusive slowdown is not homogeneous across a facet. Instead, in a growing ice crystal, more diffusive slowdown occurs at facet corners. The reason is as follows:

1. In regions where is small, QLL thickness gradients are large. In Fig. 2(b), for example, it is clear that the gradient in QLL thickness at I’ is greater than at I. It follows that, in a growing ice crystal, more diffusive slowdown occurs at facet corners.

In summary, a flat facet exposed to supersaturated vapor will initially experience a reduction in growth rate at its center because of decreased vapor concentration there. But that same reduction leads to a higher step density at facet corners, as new layers of ice form more readily there, hence a reduction in the growth rate at corners. When these effects become balanced – which can (and does) occur as an emergent property of the equations of motion defining QLC-1 (and QLC-2, as we will show here), the result is equal growth rates across the entire facet. That steady state, in turn, is interpreted at the mesoscale (e.g., in a high-resolution optical or scanning electron microscopy experiments) as faceted growth.

QLC-1 suffered from some structural deficiencies, however, of which the most important for our present purpose is that the time scale of process (iii) illustrated in Fig. 1, the interconversion of quasi-liquid and ice, was fixed relative to processes (i) and (ii). In real crystal facets, these time scales are expected to vary from facet to facet, or as a function of temperature and vapor pressure. These time scales should therefore be adjustable quantities within the theory.

Our goal in this communication is to evaluate strengths and weaknesses of a revised quasiliquid continuum model for ice crystal growth and ablation designed to address this deficiency. Section 2 presents such a model, referred to here as QLC-2. Section 3 (with details given in Appendix 1) summarizes scanning electron microscopy (SEM) image processing algorithms that can be used to construct quantitative ice surface morphologies. Section 4 (with details given in Appendix 2) summarizes results from gas-phase simulations that can be used to inform parameterizations of QLC-2 having to do with the vapor field in contact with growing ice crystals. Section 5 describes two strategies for parametrizing QLC-2 simulations. Section 6 presents simulations and SEM experiments in pursuit of five lines of investigation, using both strategies. Section 7 discusses implications of these results in other contexts, including cirrus ice crystal morphologies and ideas from nonlinear dynamics.

1. **QLC-2: A revised quasi-liquid reaction-diffusion model**

Like QLC-1, QLC-2 represents an ice surface as two mesoscale variables and three processes, as shown in Fig. 1. The governing equations are

(1a)

(1b)

Some notes about this model are as follows, with differences between it and QLC-1 noted:

1. represents surface diffusion of the QLL; the underlying ice is considered immobile on time scales considered here.
2. is the rate at which vapor-phase water molecules strike the quasi-liquid; it is assumed that these stick with 100% efficiency, and thermalize on a picosecond time scale (i.e., instantaneously within the time scales of a QLC-2 simulation).
3. prescribes the thickness of quasi-liquid when it is in equilibrium with the underlying ice, according to

(2)

This formulation ensures that the QLL thickness varies continuously from the thin microstate (“surface I”) with thickness , to the thick microstate (“surface II”) with thickness . (Note the connection to work of (Benet et al. 2019, Eq. 8), that also shows a sinusoidal dependence.)

1. Because surfaces I and II have different volatilities, the surface supersaturation at a given point on the surface (designated in Eq. 1a) is a function of both the microstate and the water vapor concentration above it. To compute , we define a variable that quantifies the degree to which a given surface is similar to surface I or II,

(3)

With this definition, surface I will have , while surface II will have . We then express the surface supersaturation as

(4)

where is a measure of the difference in the equilibrium vapor pressure of surfaces I and II, and is the supersaturation relative to surface I, mentioned previously. We will assume here that both (a scalar quantity) and are fixed parameters of a given solution. Eq. 4 is at slight variance with, and simpler than, the corresponding expression in QLC-1.

1. Typically, a parabolic form is used to approximate appearing in Eq. 4,

(5)

where is a “center reduction”, the fractional reduction of supersaturation at facet centers relative to facet corners,

(6)

Negative values of are also used in the simulations presented here, to represent the enhancement of water vapor concentration at facet centers when a crystal is ablating because of exposure to subsaturated conditions.

1. is a first-order relaxation constant describing the time scale at which quasi-liquid/ice equilibrium is achieved. That is, if we imagine a surface with quasi-liquid amount , then equilibration after a time occurs according to

(7)

If one takes the time derivative of Eq. 6, and assumes that is small, the second term on the right-hand side of Eq. 1b results.

Equations 7 and 1b represent the primary departure of QLC-2 from QLC-1. With this revision, we are able to specify the rate of quasi-liquid/ice equilibration relative to processes (i) and (ii). Specifying a small value for , for example, would represent the idea that quasi-liquid/ice equilibration is fast compared to those processes, while large would mean the opposite. We do not have reliable observational values of , but we do have a guidepost: because the “diffusive slowdown” mechanism for stabilization of faceted ice growth described above requires that quasi-liquid/ice equilibration be slow compared to surface diffusion, we should not be surprised if we find that large leads to stable growth dynamics. We return to this topic below.

1. **ESEM/GNBF retrievals**

Environmental SEM of imaging of ice crystals has seen considerable development in recent years, including the ability to image actively growing and ablating crystals by manipulating the temperature and pressure inside an SEM chamber (Pfalzgraff, Hulscher, and Neshyba 2010; Zimmermann et al. 2007). In tandem with those developments are computer codes for generating quantitative surface morphologies using a Gauss-Newton in a Bayesian Framework (GNBF) algorithm (Butterfield et al. 2017). The combination – a process we will refer to here as “ESEM/GNBF retrieval” – provides opportunities for vetting model predictions of surface morphology against experiment at resolutions that are not quite commensurate with one another, but approaching that level. Details are given in Appendix 1.

1. **Vapor field simulations**

QLC-2 requires numerous parameterizations. Some of these were provided, in N2016, by molecular dynamics simulations, and some by experimental observations. Unexploited in N2016, however, was the use of gas-phase simulations to define properties of , the vapor concentration overlying a given ice surface. These simulations show that the parabolic form for , given in Eq. 5, is an accurate representation of the surface water vapor concentrations under conditions of constant water vapor concentration far from the crystal; they also provide a means of parameterizing of as a function of the size of the crystal (), as well as the ambient temperature and air pressure. Details about these gas-phase simulations are given in Appendix 2.

1. **Parameterizations**

For the purpose of exploring solutions to the QLC-2 equations of motion, it is necessary to define values of the parameters appearing in Eqs. 1-7. To do so, we employ here two separate strategies.

**Strategy A – Standalone QLC-2**

In Strategy A, model parameters are explicitly specified as inputs,

* , the distance from facet center to corner.
* , the surface diffusion coefficient
* , the Hertz-Knudsen deposition velocity
* , the ice-QLL equilibration time constant.
* and , which describe a parabolic representation of .

The chief advantage of Strategy A is its simplicity. A disadvantage is that in real life, some of these values are actually related to one another. A prominent example is that the surface diffusivity and the kinetic deposition rate ( and ) both increase with increasing temperature, whereas in Strategy A they are treated as independently specifiable quantities.

**Strategy B – Integrated QLC-2 + Vapor Field**

In Strategy B, some model parameters are explicitly specified (as in Strategy A), but others are made functions of ambient conditions, or decided upon on the basis of consistency with the vapor surrounding the crystal. Explicitly specified parameters are

* , distance from facet center to corner.
* and , temperature and pressure surrounding the crystal.
* supersaturation of water vapor far from the crystal.
* , the ice-QLL equilibration time constant.

while others are obtained as follows:

* The surface diffusion coefficient is obtained from the ambient temperature,

(8)

with and (based on a fit to diffusion data of (Price, Ide, and Arata 1999)).

* The Hertz-Knudsen deposition speed is also temperature-dependent,

(9)

where is water’s equilibrium vapor pressure, and is its molar mass.

* The diffusivity of water vapor though air (required by the Vapor Field code) is a function of ambient temperature *and* ambient pressure (see Appendix 2 for details)

(10)

* , the growth rate of the ice surface, is supplied iteratively, as follows:
  1. A Vapor Field simulation takes growth rate of the ice surface () as an input, along with , , , and , and predicts and .
  2. A QLC-2 simulation takes and as inputs, along with , , , and , and predicts a new growth rate of the ice surface ().
  3. Steps 1 and 2 are repeated until and agree within some tolerance level.\

The chief disadvantage of Strategy B is its relative complexity, and the fact that the iteration just described makes the computation slower. Its main advantage is that connections to physical conditions are more transparent and self-consistent.

Common to both strategies are the following fixed values:

* and , representing the difference in equilibrium number density between surfaces I and II (fixed at and ).
* , the amount by which the equilibrium vapor supersaturation of surface II exceeds that of surface II (fixed at ).
* , the thickness of a single layer of ice (fixed at ).

1. **Results**

Here we describe the results of five lines of investigation we have pursued, each focusing on a particular topic or question. ESEM/GNBF and gas-phase modeling results are presented alongside, as relevant.

*I. Effect of variation in the time scale of modeled ice-quasiliquid equilibration*

[still working on this, preliminary work says that bigger stabilizes the formation of steady states, but otherwise has little effect on the shape of the steady-state profiles] … See Fig. 3 …

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| **Figure 3**. Effect of varying the ice-QLL equilibration timescale. |

*II. Curvature of faceted surfaces undergoing growth and ablation*

Figure 4 shows a modeled ice crystal surface under growing and ablating conditions. The growth scenario on the left of the figure resulted from supersaturated water vapor concentrations, distributed as shown in Fig. 4(a). Figures 4(b) and 4(c) show that these conditions lead to steady state, “V”-shaped profiles, in which the surface is dominated by primarily surface I – like microstates. This scenario exhibits more tightly bunched (smaller ) at facet boundaries, which in turn (as described above in the summary of diffusive slowdown) leads to a net increase in volatility of the surface as a whole, hence faceted growth.

The ablating scenario on the right of Fig. 4 resulted from subsaturated water vapor amounts, distributed as shown in Fig. 4(d). Figures 4(e) and 4(f) show that these conditions also lead to steady state, although in this case the profile is “” shaped (i.e., rounded), the surface is dominated by surface I – like microstates, and the layer bunching leads to *reduced* volatility of the surface near the corners, hence faceted ablation.

The model results shown in Fig. 4 also suggest the following general pattern: growing ice facets possesses convex curvature, whereas ablating ice facets possess concave curvature. A useful metric for describing the curvature of steady state profiles such as those appearing in Fig. 4 is the local slope of the surface. Here we quantify that slope as a mean horizontal distance between successive molecular layers, defined as

(11)

For example, both facet profile in Fig. 4 are characterized by .

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| **Figure 4**. Steady states of growing (left panels) and ablating facets (right panels), for baseline parameters constructed according to Strategy B (see Table 1). Top: vapor supersaturation adjacent to a facet spanning ; Middle: time evolution of the number of steps, beginning with a flat surface (#steps = 0) to steady state; Bottom: steady-state profiles of QLL-covered faceted ice surfaces. | |

It is well-known that real ice crystals exhibit faceted growth, but are they also capable of faceted ablation? Figure 5 displays ESEM images of an ice crystal observed under growing and ablating conditions. Since the ablating crystal retains its flat surface, we can conclude that faceted ablation has indeed occurred. The figure shows, moreover, that faceted ablation occurs even when the surface is rough (e.g., the prismatic facets in the figure). In fact, we observe faceted ablation quite frequently in ESEM images of ablating ice crystals.

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| **Figure 5**. An ice crystal under growing (left) and ablating (right) conditions. | |

Do real faceted ice crystals exhibit curvature like that indicated by the QLC-2 predictions of Fig. 4? Fig. 6(a) displays an ESEM image of a crystal and a GNBF construction of a portion of its basal facet. The crystal is known to be growing, since subsequent images taken of this crystal revealed expanding boundaries against the metal substrate to which the crystal is attached. The GNBF construction, displayed in Fig. 6(b), reveals a distinct convexity, on the order of 1000s of layers over the horizontal span analyzed () … which corresponds to .

Turning to ablation, Figs. 6(c-d) show an ESEM image of an ablating, faceted crystal, and its GNBF-constructed surface. *The GNBF reconstruction hopefully reveals concavity ….* .*]*

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| **Figure 6**. Top panels: SEM images of a growing crystal (left) and an ablating crystal (right). Squares show outlines of areas of targeted for GNBF surface retrievals. Lower panels: show GNBF height reconstructions of those areas. | |

Thus, the pattern predicted by QLC-2 is borne out: growing ice facets do indeed exhibit facet convexity, while ablating ice facets exhibit facet concavity – although we hasten to point out that the values are far smaller in the observations than in the model.

*III. Characteristic length scales of growing and ablating surfaces*

Here we investigate the possibility that growing and ablating ice crystals possess intrinsic length scales. Focusing first on observations, we note that a distinct growth/ablation asymmetry appears in the roughening evident in the ESEM images displayed in Fig. 5. …

Figure 7 shows the dependence of the mean horizontal layer separation, (introduced above), on the parameter , defined by

(12)

where (as described above) is the surface diffusion coefficient, is the edge length of the crystal, and is the kinetic deposition velocity. In constructing Fig. 7, we employed Strategy B to arrive at a self-consistent set of parameterizations for the baseline scenario, but for all other points freely assuming a range of values of , , and , as indicated in the caption.

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| **Figure 7**. Mean horizontal layer separation () as a function of parameter , with , , and varied independently, and all other parameters held fixed (see Table 1). The large square symbol marks the baseline scenario. |  |

It is clear from Fig. 7 that a good parameterization is a straight line,

(13a)

with best-fit parameters , . The approximation is due to the fact that is small. Best Analysis of ablating scenarios also yields a straight line,

(13b)

with slightly different parameters: and .

Figure 8 shows values of for a range of corner supersaturations (), holding parameter constant. We see that when conditions begin to become supersaturated, starts off at about and declines monotonically, ultimately leading (we surmise) to hollowed or dendritic growth at high supersaturations. On the left-hand side are shown results when conditions are subsaturated. We see that under these conditions, starts much higher than on the supersaturated side, at about , and declines monotonically with increasing subsaturation, ultimately leading (we surmise) to facet rounding.

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| **Figure 8**. Mean horizontal layer separation () as a function of corner supersaturation according to Eqs. 14a and 14b. |

We have carried out a best-fit analysis of the supersaturated-condition points of Fig. 8 in the factorized form , which, with the approximation indicated in Eq. 13a, becomes

(14a)

Similarly, for ablating facets, and using the approximation indicated in Eq. 13b,

(14b)

where

(15)

with , , and .

Figure 8 shows the results of applying this functional form to the numerical data, using best-fit parameters , , , The fit is pretty good … And what it means is, we can predict steady-state properties over a very big range of physical variables (crystal length, surface diffusivity, , and imposed supersaturations).

*IV. Dependence on crystal size*

Figure 9 shows simulation results when the crystal edge length is varied, employing Strategy B. Fig. 9(a) shows that the steady-state concentration of water vapor at crystal corners declines with increasing crystal size. This is expected, since a larger growing crystal means there is more crystal surface area drawing water vapor out of the air.

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| (a) | (b) |
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| **Figure 9**. Steady-state results using Strategy B, with increasing crystal size: (a) Supersaturation, , at crystal corners; and (b) Percent reduction in at facet center relative to crystal corners (Eq. 6). | |

Fig. 9(b) shows the percent reduction in surface vapor concentration at facet center relative to facet corner (see the definition of in Eq. 6). is seen to be a smoothly increasing function of crystal size, well-described by

(16)

Other simulations (not shown) show that higher far-field vapor concentrations () do not greatly alter this parameterization. However, yet other simulations show that this parameterization is greatly affected by lowered temperature (which increases ), and by increasing ambient pressure (which also increases ). Lower ambient temperatures and higher ambient pressures are both associated with smaller vapor diffusion coefficients, . Smaller *,* in turn, manifests as a more steeply curved profile across the crystal surface, compared to the example shown in Fig. A1(b): it is harder for water vapor to diffuse across the facet surface when is small.

The expected consequence of the foregoing is that lower ambient temperatures and higher ambient pressures will tend to promote excess growth at facet corners, ultimately leading to greater facet convexity – and, we conjecture, at sufficiently small , dendritic forms such as snowflakes.

*IV. Resilience of steady states*

In ESEM experiments, facets exhibit a certain resilience, in that an initially faceted surface, after it is roughened by some perturbation, can usually be restored to its initial smooth faceted state after the initial conditions are restored. An example is shown in Fig. 9, in which the fraction of smooth, faceted area is seen to increase as a result of lowered temperature.

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| **Figure 9**. SEM images showing the fraction of smooth, faceted area increasing as a result of lowered temperature (top to bottom). |  |

What resilience, if any, is exhibited by modeled facets? Figure 10 is an examination of this question.

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| **Figure 10**. Examination of model facet resilience. Curves in the upper-left graph show timelines of that characterize the surface when subjected to the supersaturation curves shown in the upper-right insets, over the indicated time intervals. |

The sequence of images in Fig. 10 begins with an initially-flat profile, which is then subjected to the following sequence:

1. From to , the surface is been exposed to the supersaturated water vapor curve shown in the inset located in the upper right part of the figure. By , the profile has evolved to the faceted, steady-state profile labeled “A”.
2. From to , a perturbation is introduced in the form of the subsaturated water vapor curve shown in the inset located in the middle-right part of the figure. During this time, the surface evolves into the highly perturbed state labeled “B”.
3. From to , the initial supersaturated water vapor regime is restored. During this time, the surface recovers its pre-perturbation faceted profile labeled “C”.

Conclusion … qualitatively, QLC-2 exhibits resilience analogous to that of real crystal facets, such as the one shown in Fig. 9. We should hasten to add, however, the perturbed states exhibit vastly different vertical scales. That is, in Fig. 10, the perturbed profile seen at , caused by the perturbation begun at , is non-faceted (i.e., “rough”) only on the order of a few dozen monolayers of ice, whereas the actual crystal shown in Fig. 9 exhibits ridge-to-valley depths on the order of thousands of monolayers.

1. **Summary and discussion**

The QLC-2 model presented here offers significant technical improvements over the QLC-1 model reported in N2016, in that its numerical solutions are more stable, and its parameterizations are better constrained by independent numerical simulations of the overlying vapor field, i.e., the dependence of the center reduction parameter () on crystal size ().

More fundamentally, QLC-2’s equations of motion embody a more faithful atomistic representation of ice/QLL freeze/melt equilibration. That connection, in turn, enables the theory to represent a more unified picture of ice surface dynamics than has previously been possible. The model predicts, for example, facet convexity and concavity under growth and ablation conditions (respectively), a pattern that is echoed in reconstructions of SEM-grown ice crystals. The model also provides a mechanism by which real ice crystals resist dendritic geometries when subjected to supersaturation conditions (i.e., faceted growth), and how they also resist rounding when subjected to subsaturation conditions (faceted ablation). It is, essentially, a theory of faceting that rests fundamentally on atomistic variations in the thickness and volatility of the quasi-liquid layer that are exposed as a crystal grows or ablates.

The model also predicts a certain resilience to perturbations – also echoed by real ice crystals – in that faceted surfaces that have been disrupted in some way recover when conditions favorable to faceting are restored.

QLC-2 makes an unexpected qualitative prediction about characteristic distances, quantified here as , the mean distance between adjacent layers of ice. We find that separates mathematically as a product of two functions: one is a linear function of variable, , and the other is a highly nonlinear function of the surface supersaturation. We find that the latter is an asymmetrical function of the surface supersaturation, such that growth conditions lead to smaller values of , while ablation conditions lead to larger values of , for comparable departures from equilibrium. We have no ready explanation for this behavior, but we do note that SEM observations are consistent with these observations in the sense that ridges in rough facets of growing crystals are spaced more closely than in rough facets of ablating crystals. As noted previously, this coincidence is subject to a very big caveat, namely, that the depths of these characteristic distances are vastly different in QLC-2 compared to experiment: in the former it is a few monolayers, while in the latter it is thousands of monolayers.

Qualitative implications of this work for cirrus cloud particles can be summarized as follows: As such particles fall through Earth’s atmosphere, such crystals will encounter increased pressure, hence smaller , but also increased temperatures (unless there is an atmospheric inversion), hence larger . When the effect of increased pressure dominates, QLC-2 predicts enhanced growth at facet corners, hence greater facet convexity, and therefore a greater propensity toward hollowed crystal structures. When the effect of increased temperature dominates, however, we can expect more regular hexagonal shapes. Although there is no single observational datum that would help us resolve these predictions, we can comment that in exceptionally cold regions (such as the Antarctic Plateau), or even in mid-latitudes where high-altitude cryo-capture of ice crystals on ground-launched balloons is possible, observations have shown that cirrus clouds are frequently hollowed, suggesting dominance of increased pressure distinctive (Magee et al. 2014, 2021; Walden, Warren, and Tuttle 2003).

We should note that the results presented here do not consider variations in parameters , and , because these values are highly uncertain; more detailed molecular dynamics calculations could provide plausible values, but those studies have not been reported. We can speculate that because different facet types (basal, prismatic, or pyramidal) have distinct underlying crystal cell structures, their quasi-liquid properties will also be Exploratory numerical studies varying the thickness of a single “layer” of ice has shown that a proportional increase in results. Preliminary numerical experiments varying and have shown that …

A separate speculation concerns the observation that the dependence of values exhibited by QLC-2 is the same as in Turing patterns. In one sense this should come as no surprise, since Turing’s theory, like QLC-2, is based on a reaction-diffusion equation. But there are also very big differences, including the fact that Turing’s analysis proceeds from an analysis of sensitivity to perturbations to an initially homogeneous distribution of chemical species, whereas the patterns in QLC-2 emerge as steady states of the equations of motion.

Finally, we note that the atoms-to-mesoscale approach represented in QLC-2 is not as fully integrated as we would like it to be … maybe some insight from Jake’s heterogeneous multiscale approach is the way forward on this (Shohet et al. 2020).

**Appendix 1 – ESEM/GNBF retrieval**

Here is a summary of key equations and constraints in the ESEM/GNBF retrieval …

**Appendix 2 – Vapor field code**

Simulation of the water vapor partial pressure, , in the space surrounding a square-shaped crystal shown in Fig. A1(a), was achieved by integrating the two-dimensional diffusion equation

(A1)

where is the diffusion coefficient of water vapor through air, which is computed by (see Air-Diffusion Coefficients of Gases in Excess of Air),

(A2)

Equation A1 is solved subject to Neumann boundary conditions that represent depletion of water vapor in the layer adjacent to the crystal surface due to crystal growth. This rate of depletion is given by

(A3)

where

* + , the specified growth rate of the ice surface;
  + , the mass density of ice; and
  + , the molar mass of water

Dirichlet conditions representing the far-field vapor concentration (), are imposed at the outside boundary of the simulation space. Integration over time was performed using Euler’s method, parameters for which are given in Table A1.

|  |  |
| --- | --- |
| **Table A1. Parameters for simulation of the vapor field around a growing, square-shaped ice crystal** | |
| Simulation space dimensions |  |
| Time step for integration |  |
| Time interval for integration |  |
| Spatial discretization |  |
| Diffusion coefficient at , |  |
| Ambient temperature |  |
| Ambient pressure |  |
| Diffusion Temperature-correction exponent |  |
| Diffusion coefficient under ambient conditions |  |
| Far-field water vapor partial pressure |  |
| Far-field water vapor supersaturation |  |
| Far-field distance from the origin |  |
| Mass density of ice |  |
| Growth rate of ice surface |  |

The resulting vapor concentration contours shown in Fig. A1(a) are seen to decrease with proximity to the crystal, as expected since the growing crystal is drawing water vapor out of the surrounding air. This reduction is greater at facet center compared to facet corners, in a roughly parabolic fashion, as shown in the profile in Fig. A1(b).

|  |  |
| --- | --- |
| (a) | (b) |
|  |  |
| **Figure A1**. (a) Simulation of vapor partial pressures, , in units , around a growing ice crystal. The black-outlined box at the center indicates the surface of the crystal, in this case spanning . Contours outside the crystal show values of at steady state. Dirichlet conditions fix a partial pressure of at the far-field distance of , while Neumann conditions representing ice growth are imposed at (see text). (b) along the surface of the crystal. | |

**Appendix 3 – Numerical considerations**

Python, accelerated with Numby. Code and data are available on Github.

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Other notes …

