Poly(vinyl alcohol) Molecular Bottlebrushes Nucleate Ice

Panagiotis G. Georgiou, Nina L. H. Kinney, Ioanna Kontopoulou, Alexander N. Baker, Steven A. Hindmarsh, Akalabya Bissoyi, Thomas R. Congdon, Thomas F. Whale,* and Matthew I. Gibson*

**ABSTRACT:** Ice binding proteins (IBPs) have evolved to limit the growth of ice but also to promote ice formation by ice-nucleating proteins (INPs). IBPs, which modulate these seemingly distinct processes, often have high sequence similarities, and molecular size/assembly is hypothesized to be a crucial determinant. There are only a few synthetic materials that reproduce INP function, and rational design of ice nucleators has not been achieved due to outstanding questions about the mechanisms of ice binding. Poly(vinyl alcohol) (PVA) is a water-soluble synthetic polymer well known to effectively block ice recrystallization, by binding to ice. Here, we report the synthesis of a polymeric ice nucleator, which mimics the dense assembly of IBPs, using confined ice-binding polymers in a high-molar-mass molecular bottlebrush. Poly(vinyl alcohol)-based molecular bottlebrushes with different side-chain densities were synthesized via a combination of ring-opening metathesis polymerization (ROMP) and reversible addition-fragmentation chain-transfer (RAFT) polymerization, using “grafting-to” and “grafting-through” approaches. The facile preparation of the PVA bottlebrushes was performed via selective hydrolysis of the acetate of the poly(vinyl acetate) (PVAc) side chains of the PVAc bottlebrush precursors. Ice-binding polymer side-chain density was shown to be crucial for nucleation activity, with less dense brushes resulting in colder nucleation than denser brushes. This bio-inspired approach provides a synthetic framework for probing heterogeneous ice nucleation and a route toward defined synthetic nucleators for biotechnological applications.

**INTRODUCTION**

Ice-binding proteins recognize and bind ice faces, leading to control over their formation (ice-nucleating proteins, INPs) or growth (antifreeze proteins, AFPs). AFPs are well studied and have defined ice-binding sites to engage ice either directly or via an anchored mechanism. INPs, in contrast, are less well understood despite being commercialized as extracts from the bacterium *Pseudomonas syringae*. IBPs have many potential applications across a range of biotechnological and engineering areas where it would be advantageous to control ice formation. The study of INPs is challenging due to the low solubility of the INP and the need for the transmembrane/lipid anchored domains for activity. Davies and co-workers reported recombinant mutated INPs, but it was essential to study activity while attached to the host organism (*E.coli*) to obtain ice nucleation functional data.

In addition to IBPs, it has emerged that synthetic materials can also bind ice, reproducing the macroscopic properties of IBPs—presenting opportunities not only for biomedical applications but also to understand IBP function. Synthetic polymers can be produced with excellent control over architecture and composition, allowing molecular shape and composition to be tuned to probe the material/ice interface. The most active and widely studied ice-binding polymer is poly(vinyl alcohol) (PVA). PVA’s ice recrystallization inhibition (IRI) activity can be tuned by varying molecular weight, shape, and assembly into particles. PVA binds to ice via hydrogen bonding of the hydroxyl groups, but is unique in that most other hydroxylated polymers do not show this function. Antifreeze (glyco)proteins, in contrast, have hydrophobic segments that bind to ice (rather than the glycan hydroxyls), underpinning the concept that a molecular mechanism can lead to similar measurable macroscopic effects.

Considering the success of mimicking the macroscopic property of ice recrystallization inhibition, there remain remarkably few nonprotein ice nucleators, none of which have been designed from the bottom-up. Silver iodide, which has a very similar structure to hexagonal ice, is probably the best known. Feldspar (an aluminosilicate) is a potent ice nucleator, and its function appears to be caused by defects rather than the bulk structure. Graphene nanomaterials can...
also nucleate ice,\textsuperscript{25} and the size of the flakes proves crucial.\textsuperscript{26} The critical ice nucleus size is predicted to be \(\sim 20 \text{ nm}\) for nucleation at \(-5 \text{ °C}\),\textsuperscript{27} supporting the need for aggregates or molecularly large species. Arrays of cholesterol crystals\textsuperscript{28} and fractions of lignin (a heterogeneous material) have also been reported to have some activity,\textsuperscript{29} alongside marine biogenic particles.\textsuperscript{30} Ice-nucleating macromolecules of protein and polysaccharide origin have also been reported,\textsuperscript{31,32} but their chemical structures remain unknown.

In general, effective ice nucleators are thought to have a good lattice match to ice, meaning that water molecules bind to the nucleator in spatial locations similar to those they would occupy in ice. According to classical nucleation theory (CNT), the barrier to ice nucleation is due to the energetic cost of forming an interface between supercooled water and ice. By templating ice, a nucleator allows the formation of a smaller ice critical cluster reducing the energy barrier to nucleation and causing ice to form at warmer temperatures. For ice nucleation to occur at warmer temperatures, more significant nucleation sites possessing a lattice match to ice, or at least an ability to bind water in an icelike structure, are required.\textsuperscript{3,27}

There are currently no design rules for ice nucleators, but some key observations have been made. There is evidence that when the size of an AFP increases, it can become an INP supported by experimental\textsuperscript{33} and modeling evidence.\textsuperscript{3} Small IBPs (\(\sim 2 \text{ nm}\)) nucleate ice at cold temperatures, near to where homogeneous ice nucleation occurs\textsuperscript{12} but ice-nucleating proteins (INPs) maintain a coherent lattice match to ice across nucleation sites spanning as many as 30 individual INPs, nucleating ice at \(-2 \text{ °C}\).\textsuperscript{3} This corresponds to 50 nm \times 50 nm nucleation sites. Ogawa et al. reported that linear poly(vinyl alcohol), which is known to bind ice, is capable of nucleating ice at temperatures a few degrees above the homogeneous nucleation temperature but did not show a molecular-weight effect in the range tested.\textsuperscript{33}

Taken together, the above shows an exciting possibility that a chemically defined synthetic ice binder could be engineered to become an ice nucleator if sufficient size, density, and placement of the ice binders can be accessed. A synthetic ice nucleator would enable the first true structure–property relationships to be determined, without using heterogeneous mixtures.

Here, we introduce an ice nucleator platform based on well-defined poly(vinyl alcohol) molecular brushes obtained\textsuperscript{via} a combination of ring-opening metathesis polymerization (ROMP) and reversible addition–fragmentation chain-transfer (RAFT) polymerization, using “grafting-to” and “grafting-through” approaches. We demonstrate that densely grafted PVA bottlebrush polymers exhibit ice nucleation activity and are the first example of a synthetic polymer ice nucleator. Lower-density brush copolymers had reduced activity, demonstrating that confinement of the PVA is essential. These results show that synthetic ice nucleators can be accessed by macromolecular engineering.

## EXPERIMENTAL SECTION

### Materials

All chemicals were used as supplied unless otherwise stated. ROMP catalyst G3 ([H1Mes]2[ppt]2(CH)Ru = CHPH), cis-5-norbornene-exo-2,3-dicarboxylic anhydride (95%), exo-5-norbornene carboxylic acid (97%), N-boc-ethylendiamine (≥98%), 1-hydroxypyrrolidone-2,5-dione (98%), ethyl vinyl ether (EVE) (≥99%), trifluoroacetic acid (TFA, 99%), mPEG\textsubscript{108}, succinimidyl carboxymeth-yl ester (NHS-mPEG, \(M_n \sim 10,000 \text{ Da}\), poly(ethylene glycol) methyl ether methacrylate (PEGMA, \(M_n \sim 1000 \text{ Da}\)), 4-cyan-4-(phenylcarboxothioiyloxy)pentanoic acid (97%), 4,4’-azobis(4-cya-

\begin{align*}
\text{Photo-Polymerization of Vinyl Acetate Using 2-(Ethoxycarbo-} \nonumber \\
\text{nothioylthio)-2-methylpropanoic Acid NHS-Ester.} \nonumber \\
\text{2-(Ethoxycarbo-} \nonumber \\
\text{nothioylthio)-2-methylpropanoic Acid NHS-Ester.} \nonumber \\
\text{Synthesis of Poly(amino exo-norbornene imide), P(NB-NH}_\text{2})_\text{)} \nonumber \\
\text{Homopolymer Precursors via Ring-Opening Metathesis} \nonumber \\
\text{Polymerization (ROMP). A typical procedure for the synthesis} \nonumber \\
\text{of P(NB-NBoc)_2 homopolymer via ROMP is described. A} \nonumber \\
\text{stock solution of 100 mg/mL of G3 in CHCl}_3 \text{ and a solution of 173} \nonumber \\
\text{mg of NB-Boc (50 eq. 0.57 mmol) in 3.3 mL of CHCl}_3 \text{ were prepared.} \nonumber \\
\text{Then, 100 \muL of G3 stock solution (10 mg, 1 eq. 0.011 mmol) was} \nonumber \\
\text{added to the vial of NB-NBoc monomer solution with} \nonumber \\
\text{rapid stirring, and polymerization was allowed to proceed at room} \nonumber \\
\text{temperature for 1 h (final [NB-NBoc] = 50 mg/mL, final [G3] = 2.9} \nonumber \\
\text{mg/mL]). The polymerization was then quenched by the addition of} \nonumber \\
\text{a few drops of ethyl vinyl ether, and P(NB-NBoc)_2 homopolymer} \nonumber \\
\text{was precipitated from diethyl ether, isolated by vacuum filtration, and} \nonumber \\
\text{dried under vacuum prior to} \text{H NMR and SEC analyses. The} \nonumber \\
\text{procedure was repeated for [NB-NBoc]/[G3] ratios of 100, 200, and} \nonumber \\
\text{400.} \text{H NMR (400 MHz, CDCl}_3 \text{ M}_\text{NMR} = 15,300 \text{ g mol}^{-1}.} \nonumber \\
\text{SEC (DMF} \nonumber \\
\text{+ 5 mM NH}_3 \text{BF}_4 \text{ M}_{\text{SEC}} = 17,600 \text{ g mol}^{-1},} \nonumber \\
\text{and D}_{\text{SEC}} = 1.6. \text{Next, P(NB-NBoc)_2 homopolymer (100 mg, 0.003 mmol) was} \nonumber \\
\text{dissolved in a 1.1 TFA/DCM solution (5 mL) and stirred at room} \nonumber \\
\end{align*}
temperature for 4 h. The polymer reaction mixture was concentrated to dryness under reduced pressure to afford a brown residue that was precipitated by the addition of diethyl ether. Removal of the solvent resulted in the isolation of the amino-deprotected polymer product as a brown solid \( (P(NB-NH_2))_{50} \). The resulting polymer was then dissolved in DI water and dialyzed against DI water (dialysis membrane MWCO = 3.5 kDa) for approximately 14 h and lyophilized to afford the polymer products as brown solid. The

**Figure 1.** (A) Design and synthesis of graft PVA-based brush copolymers generated via “grafting-to” procedure. Schematic representation of the synthetic route followed for the preparation of \( P((P(NB-NH)_{g}-PVAc)_{210}) - \text{stat} - P(NB-NH_2)_n \) \((n = 50, 100, 200, 400)\) graft copolymers. (B) Normalized SEC-RI chromatograms for \( P((P(NB-NH)_{g}-PVAc)_{210}) - \text{stat} - P(NB-NH_2)_n \) copolymer precursors in DMF + 0.1% LiBr and solution characterization of resulting \( P((P(NB-NH)_{g}-PVAc)_{210}) - \text{stat} - P(NB-NH_2)_n \) \((n = 50, 100, 200, 400)\) in water via (C) DLS analysis (insets: representative photograph of 1 mg mL\(^{-1}\) aq solutions of \( P((P(NB-NH)_{g}-PVAc)_{210}) - \text{stat} - P(NB-NH_2)_n \) \((n = 50, 100, 200, 400)\)). (D) Dry-state TEM image obtained from a 1 mg mL\(^{-1}\) solution of \( P((P(NB-NH)_{g}-PVAc)_{210}) - \text{stat} - P(NB-NH_2)_n \) \((n = 50, 100, 200, 400)\) in water. (E) Representative AFM image obtained from a 5 mg mL\(^{-1}\) solution of \( P((P(NB-NH)_{g}-PVAc)_{210}) - \text{stat} - P(NB-NH_2)_n \) graft copolymer in water.
procedure was repeated for PNB(PNBS), 100, 200, and 400 homopolymers. SEC characterization of the resulting depolymerized polymer side chain) was not possible due to high sample–column interactions. 'H NMR (400 MHz, DMF-d6): δ (ppm) = 8.04–7.92 (br s, CH2NH of polymer side chain), 5.73–5.56 (br m, CH=CH2 of polymer backbone), 5.56–5.33 (br m, CH=CH2 of polymer backbone), 3.74–3.48 (br s, CH2CH2NH of polymer side chain), 3.48–3.24 (br m, CH2CH2NH of polymer side chain), 3.24–2.83 (br s, CH3C(O)CH3 of polymer backbone), 2.83–2.56 (br s, CH2CH2CH of polymer backbone), 2.13–1.80 (br s, CH2CH2CH of polymer backbone), 1.62–1.32 (br m, CH2CH2CH of polymer backbone).

Synthesis of P((PNB-NH)-g-PPEG210)-stat-P(NPB-NH)3 graft Copolymers via "Grafting-To" Approach. A representative synthesis of P((PNB-NH)-g-PPEG210)-stat-P(NPB-NH)3 was as follows: NHS-PVAc210 (2.0 mg, 0.0002 mmol, 1 eq), P(NB-NH)3 (0.34 mg, 0.018 mmol, 100 eq), and EDC.HCl (3.5 mg, 0.018 mmol, 100 eq) were dissolved in 6.5 mL of anhydrous DMF, followed by the addition of Et3N (2 μL, 0.018 mmol, 100 eq), and the reaction mixture was stirred at 70 °C for 16 h. An aliquot of the crude reaction mixture was then taken for SEC analysis in DMF (+0.1% LiBr). The resulting polymer was precipitated twice in diethyl ether and was dried under vacuum.

Our initial hypothesis was that densely packed PVA side chains, to mimic the regular arrays of IBPs which lead to nucleation, were essential. A "grafting-to" approach was first employed, which results in low grafting densities due to steric repulsions between bulky side chains (Figure 1A). 3,4 Poly(norbornene) bearing primary amines were first synthesized via an N-Boc-protected intermediate monomer (NB-NBoc, Figures S1 and S2), initiated by Grubbs third-generation bispyridyl complex (G3), [M]/[G3] of 50, 100, 200. Full characterization by 'H NMR and size exclusion chromatography (SEC) analysis is shown in Table 1 and Figures S17 and S18. Deprotection of P(NB-NBoc), with trifluoroacetic acid (TFA) gave the desired poly(amino norbornene imide) P(NB-NH)2 homopolymers (Figure S19).

\[ \text{Polymerization of Poly(ethylene glycol) Methyl Ether Methacrylate Using 4-Cyano-4-(phenylcarboxyloxy)pentanoic Acid.} \]

A vial was charged with 4-cyano-4-(phenylcarboxyloxy)pentanoic acid (5.0 mg, 0.018 mmol, 1 eq), PEGMA (3.58 g, 3.6 mmol, 200 eq), ACVA (1.0 mg, 0.004 mmol, 0.2 eq), and 17 mL of DMF. The vial was then sealed and deoxygenated using three successive cycles of freeze–pump–thaw to remove O2(g). The vial was placed into an aluminum heating block, which had been preheated to 70 °C to initiate polymerization. After 24 h, the polymerization was then quenched by exposing the vial to air and submerging it into liquid N2. An aliquot was withdrawn for the determination of monomer conversion by 'H NMR spectroscopy. The polymer was precipitated into diethyl ether from dioxane twice to yield a white polymer product that was further dried under vacuum. 'H NMR (400 MHz, methanol-d4): δ (ppm) = 3.88–3.51 (br s, 16,000H, CH2CH2 of polymer side chain), 2.48–1.58 (br m, 400H, C(CH3)2 of polymer backbone), 1.41–0.76 (br m, 600H, C(CH3)2 of polymer backbone), Msh = 200,000 g mol⁻¹. SEC (5 mM NH4BF4 in DMF) M₅,SEC,RI = 134,300 g mol⁻¹, D₅,SEC,RI = 2.0.

### RESULTS AND DISCUSSION

- **Grafting-to Approach**: A representative synthesis of a PNB-g-PPEG210 brush polymer by ROMP "grafting-through" as follows: NB-PPEG210 (100 mg, 0.006 mmol, 20 equiv) was dissolved in 1.9 mL of anhydrous THF in a 3 mL vial equipped with a magnetic stirring bar. To the vial was added 100 μL of a 2.0 mg mL⁻¹ stock solution of G3 in THF (final [NB-PPEG210]/[G3] = 20, [NB-PPEG210] = 50 mg mL⁻¹). The reaction mixture was stirred at room temperature for 2 h. The polymerization was then quenched via addition of a few drops of ethyl vinyl ether. An aliquot of the crude reaction mixture was then taken for SEC analysis in DMF (+0.1% LiBr). The resulting polymer, PNB-g-PPEG210, was isolated by precipitation from diethyl ether and was dried under vacuum.

- **Synthesis of PNB-g-PPEG210 Bottlebrush Polymers via ROMP “Grafting-Through” Approach**: A representative synthesis of a PNB-g-PPEG210 bottlebrush polymer by ROMP "grafting-through" as follows: NB-PPEG210 (100 mg, 0.006 mmol, 20 equiv) was dissolved in 1.9 mL of anhydrous THF in a 3 mL vial equipped with a magnetic stirring bar. To the vial was added 100 μL of a 2.0 mg mL⁻¹ stock solution of G3 in THF (final [NB-PPEG210]/[G3] = 20, [NB-PPEG210] = 50 mg mL⁻¹). The reaction mixture was stirred at room temperature for 2 h. The polymerization was then quenched via addition of a few drops of ethyl vinyl ether. An aliquot of the crude reaction mixture was then taken for SEC analysis in DMF (+0.1% LiBr). The resulting polymer, PNB-g-PPEG210, was isolated by precipitation from diethyl ether and was dried under vacuum.
characterized by SEC analysis revealing bimodal peaks at high-molecular-weight direction. Figure 1B shows corresponding SEC curves of the NHS-PVAc$_{210}$ side chain (dashed black curve) and the resultant graft copolymers, in which the peak positions of the graft copolymers completely shifted to higher molecular weights. A significant amount of unfunctionalized NHS-PVAc$_{210}$ was also apparent for all graft copolymer formulations. Grafted side chains (%) of the resulting graft copolymers from SEC analysis were estimated to be ca. 20–40% (Table 2), due to steric repulsions between bulky side chains, limiting grafting density. The acetate protecting groups were quantitively removed by an excess of hydrazine hydrate, as confirmed by $^1$H NMR and FTIR spectroscopic analysis (Figures S20 and S21). The solution behavior of the resulting PVA graft copolymers was investigated by dynamic light scattering (DLS), transmission electron microscopy (TEM), and atomic force microscopy (AFM). DLS analysis showed that graft copolymer samples exhibited an increase in length upon increasing the backbone of P(NB-NH$_2$)$_n$ (Figure 1C). The fact that these traces are multimodal can be rationalized by considering additional relaxations occurring, which contributed to the scattering profile due to the anisotropy of the bottlebrush polymers in solution. Dry-state TEM images of P((P(NB-NH)$_2$)-g-PVA$_{210}$)-stat-P(NB-NH$_2$)$_n$ confirmed the presence of small cylindrical nanostructures (Figures 1D and S22). Importantly, no large aggregates were observed in the TEM images, indicating that neither graft copolymers were spontaneously self-assembled into higher-order structures when dissolved in aqueous media at room temperature. AFM analysis also showed the formation of some spherical particles (Figures 1E and S23). While the PVA graft copolymers appear to be spherical in the AFM images, they likely exist as cylindrical nanostructures in aqueous environments (as confirmed by TEM analysis) due to their anisotropic dimensions, as has been reported for brush polymers in solution measured using X-ray or neutron scattering analyses. Moreover, brush polymers have been shown to de-wet various substrates due to unfavorable surface interactions, causing them to present as globules in AFM images.

Next, a series of high-density PVA bottlebrush polymers were prepared using ROMP “grafting-through” approach using a norbornene-end poly(vinyl acetate) macromonomer (NB-PVAc), Figure 2A. Exo-S-norbornene-2-methyamine was synthesized in this case (Figures S8–S15; see the Synthetic Procedures section in the Supporting Information), as it was demonstrated by Matson and co-workers that the choice of anchor group dramatically affects overall ROMP propagation rates with norbornene anhydride-based macromonomers leading to poorer chain extensions and conversions when targeting high $M_n$ bottlebrushes. A norbornyl xanthate chain-transfer agent (NB-CTA) of 2-(ethoxycarbonothioylthio)-2-methylpropanoic acid exo-S-norbornene-2-methylamide was synthesized and used to prepare an NB-PVAc$_{208}$ macromonomer via photo-initiated RAFT/MADIX polymerization (NB-PVAc$_{208}$ Conv. = 67%, $M_n$SEC RI = 17.9 kDa, $D_M$ = 1.4). Bottlebrush polymers were synthesized through ROMP of the norbornene amide end group on the PVAc macromonomer by varying the molar ratio of G3 catalyst ([NB-PVAc$_{208}$]/[G3]) to achieve PNB$_{n}$/g-PVAc$_{208}$ bottlebrush polymers ($n = 20, 30$ and 40).

SEC analysis revealed low-molecular-weight shoulders, which may correspond to unreacted NB-PVAc$_{208}$ macromonomer. However, the long low-molecular-weight fraction could also be explained by the premature termination of ROMP polymers. Large $D_M$ values can be primarily attributed to the employment of a high $M_n$ of PVAc macromonomer with a relatively large dispersity (Figure 2B and Table 3) as well as the above-mentioned premature termination events. In addition, number-average molecular weight values ($M_n$) determined by SEC analysis were evidently lower than theoretically expected values determined from polymerization conversions. This discrepancy in theoretical and observed $M_n$ likely originates from the branched nature of the bottlebrush polymers, leading to reduced hydrodynamic volumes relative to linear polymers of the same molecular weight and thus increased retention on the SEC columns. Acetate protecting groups were removed by hydrazine hydrate, as confirmed by $^1$H NMR and FTIR analyses (Figures S24 and S25) to yield PNB$_{n}$/g-PVAc$_{208}$ ($n = 20, 30, 40$) bottlebrush polymers that were dialyzed (MWCO = 300 kDa) to ensure removal of unreacted NB-PVAc$_{208}$ macromonomer.

### Table 1. Molecular Characteristics of P(NB-NBoc)$_n$ Prepared via Solution ROMP Using Different Initial [NB-NBoc]/[G3] Ratios, as Determined by $^1$H NMR Spectroscopy and SEC Analysis

| [NB-NBoc]/[G3] | % conv. | $M_n$SEC (kDa) | $M_n$SEC (kDa) | $D_M$SEC |
|----------------|---------|----------------|----------------|----------|
| 50             | >99     | 15.3           | 17.6           | 1.11     |
| 100            | >99     | 30.6           | 34.1           | 1.11     |
| 200            | >99     | 61.3           | 64.4           | 1.15     |
| 400            | >99     | 122.4          | 103.8          | 1.23     |

$^a$Monomer conversion calculated from $^1$H NMR spectroscopy in CDCl$_3$. $^b$Calculated from conversion. $^c$M$_n$ and $D_M$ values calculated from PMMA standards using DMP + 5 mM NH$_4$BF$_4$ as the eluent.

### Table 2. Molecular Characteristics of P((P(NB-NH)$_2$)-g-PVAc$_{210}$)-stat-P(NB-NH$_2$)$_n$ (n = 50, 100, 200, 400) Graft Copolymers as Determined by SEC Analysis

| sample | $M_n$SEC (kDa) | $D_M$SEC | grafted side chains (%) |
|--------|---------------|----------|-------------------------|
| NHS-PVAc$_{210}$ | 22.8 | 1.56 | ------ |
| P((P(NB-NH)$_2$)-g-PVAc$_{210}$)-stat-P(NB-NH$_2$)$_{50}$ | 378.6 | 1.10 | 33.0 |
| P((P(NB-NH)$_2$)-g-PVAc$_{210}$)-stat-P(NB-NH$_2$)$_{100}$ | 557.4 | 1.23 | 24.0 |
| P((P(NB-NH)$_2$)-g-PVAc$_{210}$)-stat-P(NB-NH$_2$)$_{200}$ | 1013.4 | 1.21 | 22.0 |
| P((P(NB-NH)$_2$)-g-PVAc$_{210}$)-stat-P(NB-NH$_2$)$_{400}$ | 1539.4 | 1.44 | 17.0 |

$^a$M$_n$ and $D_M$ values calculated from PMMA standards using DMP + 0.1% LiBr as the eluent. $^b$Calculated from SEC curves of reaction mixture before and after modification. $^c$Grafted side chains (%) = ($M_n$SEC RI graft copolymer − ($M_n$(NB-NH$_2$)$_{stat}$))/$M_n$SEC RI, NHS-PVAc$_{210}$. 

https://doi.org/10.1021/acs.biomac.2c01097

Biomacromolecules XXXX, XXX, XXX–XXX
DLS analysis also confirmed the formation of anisotropic particles and exhibited an increase in size upon increasing the backbone, as expected (Figure 2C). Dry-state TEM images of PNB_{n-g-PVAc_{208}} (n = 20, 30, 40) also confirmed the presence of small cylindrical nanostructures (Figures 2D and S26). AFM analysis again showed the formation of spherical particles, which is likely due to unfavorable surface interactions of PVA side chains with silicon surface, causing them to present as globules (Figures 5 and S27). Since PVA’s side-chain length is by far larger than bottlebrush backbones, it may appear spherical/ellipsoid, but further investigation is beyond the scope of this study as the extended local bottlebrush structure is shown (below) to be sufficient for the intended application.

During the handling of these materials, LCST (lower critical solution temperature) behavior was also observed for dense bottlebrush polymers prepared by ROMP “grafting-through”. This has not previously been reported, due to the absence of high-density PVA copolymers, and was hence explored. It is, however, anticipated that the backbone hydrophobicity for bottlebrushes prepared by “grafting-through” will lead to some self-assembly which may contribute to this behavior. Variable-temperature ultraviolet–visible (UV–vis) spectroscopy
Table 3. Molecular Characteristics of PNB-g-PVAc208 (n = 20, 30, 40) Bottlebrush Polymers Prepared via ROMP Using Different Initial [NB-PVAc208]/[G3] Ratios, as Determined by 1H NMR Spectroscopy and SEC Analysis

| [NB-PVAc208]/[G3] | % conv. | Mn,th (kDa) | Mn,SEC (kDa) | D3,SEC |
|-------------------|---------|-------------|-------------|--------|
| 20                | >99     | 358.0       | 162.6       | 2.11   |
| 30                | >99     | 537.0       | 241.1       | 2.41   |
| 40                | >99     | 716.0       | 340.3       | 2.39   |

“Monomer conversion calculated from 1H NMR spectroscopy in CDCl3. 6Calculated from conversion. 7Mn and D3 values calculated from PMMA standards using DMF + 0.1% LiBr as the eluent.

showed a cloud point of approximately 51 °C for PNB20-g-PVAc208 and 54 °C for PNB40-g-PVAc208 (Figure 3A). No apparent LCST behavior was observed for PVA graft copolymers prepared by “grafting-to” except for P((P(NB-NH)20-g-PVAc208)-stat-P(NB-NH2))400 that exhibited a decrease in transmittance at 75 °C, confirming that the side chain density is having a significant impact on the solution behavior of these polymers. DLS temperature ramp experiment for dense bottlebrush polymers showed a stepwise increase in the size of the dissolved nanostructures that occurred as the solution was heated above 40 °C (Figure 3B). Linear PVA homopolymers are known to possess LCST behavior in aqueous media close to water’s boiling point (∼100 °C), which is reduced by alkylation and increases with molecular weight, which agrees with these observations here.34 The close packing of polymer chains in this bottlebrush state facilitates the exclusion of water upon heating, lowering the temperature at which this transition occurs, and confirms that the solution properties compared to linear polymers are distinct.45

With this panel of bottlebrush polymers prepared, and their solution properties evaluated, the ice-binding interactions were then investigated, by assessment of their IRI (ice recrystallization inhibition) activity (Figure 4A). This was achieved by the “splat assay”, whereby small ice crystals are formed at −78 °C, then allowed to grow at −8 °C, and their growth was measured.11,46 Smaller MGS (mean grain size) means more ice recrystallization inhibition activity. Figure 4A(I) shows dose-dependent IRI of the polymer library vs controls of linear PVA. In all cases, there was a small but significant increase in IRI activity, with the bottlebrushes (high and low density) inhibiting below 0.1 mg mL−1. It has been previously reported that increasing the molecular weight of PVA increases IRI activity11 explaining these observations, although architecture-related enhancements cannot be fully ruled out. Voets and co-workers have previously prepared an acrylate-based PVA bottlebrush polymer using ATRP, which also showed no significant change in IRI compared to linear.13 In this previous work, there was no investigation of ice nucleation activity (see below). Note it was not possible to compare against linear PVA of the same MW, as linear PVA becomes increasingly insoluble in aqueous solutions at high MW, justifying our nonlinear synthetic approach to access PVA, which could not be obtained by conventional methods. The ROMP-derived backbones P(NB-NH2)20 (n = 50, 100, 200, 400) were tested as a control (Figure 4A(II)) showing weak IRI. It should be noted that we have previously shown that the facial amphiphilicity of ROMP-derived polymers can lead to observable IRI activity. The magnitude of IRI of the control amino-functional polymers P(BN-NH2) was significantly less than this previous report but is not negligible.42

A PEG-based graft copolymer (as a non-PVA, nonlinear control) was also synthesized using an N-hydroxy succinimide-poly(ethylene glycol) (NHS-PEG227) and grafted into the presynthesized poly(aminooxobornene imide) P(NB-NH2)200 homopolymer to yield P((P(NB-NH)2-g-PEG227)-stat-P(NB-NH2))200 (see the Synthetic Procedures section in the Supporting Information, Figures S28 and S29). A PEG methacrylate monomer (PEGMA20) was also employed to prepare P(PEGMA20)200 homopolymer (Figures S30 and S31). In both cases, nonlinear PEG controls were shown to have no appreciable IRI activity. Ice shaping (in sucrose solutions) showed that the PVA graft copolymers and bottlebrushes lead to more faceting of ice crystals, compared to linear PVA homopolymer (Figures 4B and S32–S35), as the large polymers can span multiple crystals as seen previously for
Figure 4. Assessment of ice recrystallization inhibition activity and ice-binding. (A) IRI activity summary of (I) P((P(NB-NH))_g-PVA_{310})-stat-P(NB-NH)_{320} (n = 50, 100, 200, 400) graft copolymers, PNB_{2100} (n = 20, 30, 40) bottlebrush polymers and (II) P(NB-NH)_{320} (n = 50, 100, 200, 400) and PEG-based control samples. Error bars are ±SD from a minimum of three repeats. The percent mean grain size (MGS) was reported relative to PBS control. (B) Modified “sucrose sandwich” ice shaping assay images for no additive (I) and 1 mg mL\(^{-1}\) linear and bottlebrush PVA samples were prepared so that each emulsion droplet will contain more than a million of even the largest bottlebrush. As shown in Figure 5 linear PVA\(_{3000}\) increased \(T_f\) by 3.5 °C compared to pure water (i.e \(\Delta T_f = 3.5 \) °C). For linear PVA\(_{310}\) a \(\Delta T_f\) of 3.2 °C was determined. This is qualitatively consistent with the findings of Ogawa et al., who found that the mass proportion of PVA present in an experiment determined the shift in nucleation temperature, rather than molar concentration (Figure S39).\(^{31}\)

The \(\Delta T_f\) observed for linear PVA is larger than that measured by Ogawa et al. who found a maximum value of approximately 2 °C for 2 mg mL\(^{-1}\) PVA solutions. The reason for this disparity is not clear. We observed that low-density bottlebrushes nucleated at lower temperatures than linear PVA, while high-molecular-weight high-density bottlebrushes nucleated at warmer temperatures, with a maximum \(\Delta T_f\) of 4.5 °C for PNB\(_{4000}\)-PVA\(_{3000}\) equivalent to a nucleation onset of −32.4 °C. For both high- and low-density bottlebrushes, a trend to higher nucleation temperatures was observed as the total molecular weight of the polymers increased, noting the large error in SEC of bottlebrushes.

To investigate the nucleation ability of larger quantities of the PVA, microcrometer droplet freezing assays\(^{36}\) were conducted. In this assay, pure water droplets usually freeze at around −33 °C, above the homogeneous nucleation temperature. This is due to the inevitable presence of ice-nucleating impurities in the water and supporting substrates employed.\(^{51}\) As with the DSC measurements, 2 mg mL\(^{-1}\) solutions were prepared, and each droplet contains >10\(^{12}\) individual PVA molecules. The results are shown in Figure SA.

Due to the variability of background measurements in this type of experiment, great care needs to be applied in interpreting results. The two linear PVA\(_{4000}\) and PVA\(_{310}\) and P(PEGMA\(_{100}\))\(_{300}\) nucleated at temperatures slightly warmer temperatures than the instrument background. As neither of these polymers can bind ice, they provide a negative control for the PVA bottlebrushes. In contrast, the various PVA bottlebrushes were observed to all nucleate ice at temperatures higher than the freezing background and higher than the linear PVA, showing that this macromolecular engineering approach to high-density PVA does lead to polymeric ice nucleating agents.

Supporting Information) were frozen in a differential scanning calorimeter (DSC) by cooling at 1 °C/min, using a technique similar to that of Marcolli et al.\(^{19}\) Figure 5B shows the DSC curves obtained, with the onset point of the exothermic peak caused by water crystallization defined as the ice nucleation temperature, \(T_f\). Marcolli et al. found a \(T_f\) of −36.5 °C for droplets for emulsions containing pure water droplets of roughly the same size, while Ogawa et al. found a \(T_f\) of −37.5 °C for a pure water emulsion of smaller droplets. We determined \(T_f\) for pure Milli-Q water to be −36.9 °C. This temperature is taken to represent the onset of homogeneous ice nucleation in our system. Similar emulsions containing 2 mg mL\(^{-1}\) linear and bottlebrush PVA samples were prepared so that each emulsion droplet will contain more than a million of even the largest bottlebrush. As shown in Figure 5 linear PVA\(_{3000}\) increased \(T_f\) by 3.5 °C compared to pure water (i.e \(\Delta T_f = 3.5 \) °C). For linear PVA\(_{310}\) a \(\Delta T_f\) of 3.2 °C was determined. This is qualitatively consistent with the findings of Ogawa et al., who found that the mass proportion of PVA present in an experiment determined the shift in nucleation temperature, rather than molar concentration (Figure S39).\(^{31}\)

The \(\Delta T_f\) observed for linear PVA is larger than that measured by Ogawa et al. who found a maximum value of approximately 2 °C for 2 mg mL\(^{-1}\) PVA solutions. The reason for this disparity is not clear. We observed that low-density bottlebrushes nucleated at lower temperatures than linear PVA, while high-molecular-weight high-density bottlebrushes nucleated at warmer temperatures, with a maximum \(\Delta T_f\) of 4.5 °C for PNB\(_{4000}\)-PVA\(_{3000}\) equivalent to a nucleation onset of −32.4 °C. For both high- and low-density bottlebrushes, a trend to higher nucleation temperatures was observed as the total molecular weight of the polymers increased, noting the large error in SEC of bottlebrushes.

To investigate the nucleation ability of larger quantities of the PVA, microcrometer droplet freezing assays\(^{36}\) were conducted. In this assay, pure water droplets usually freeze at around −33 °C, above the homogeneous nucleation temperature. This is due to the inevitable presence of ice-nucleating impurities in the water and supporting substrates employed.\(^{51}\) As with the DSC measurements, 2 mg mL\(^{-1}\) solutions were prepared, and each droplet contains >10\(^{12}\) individual PVA molecules. The results are shown in Figure SA.

Due to the variability of background measurements in this type of experiment, great care needs to be applied in interpreting results. The two linear PVA\(_{4000}\) and PVA\(_{310}\) and P(PEGMA\(_{100}\))\(_{300}\) nucleated at temperatures slightly warmer temperatures than the instrument background. As neither of these polymers can bind ice, they provide a negative control for the PVA bottlebrushes. In contrast, the various PVA bottlebrushes were observed to all nucleate ice at temperatures higher than the freezing background and higher than the linear PVA, showing that this macromolecular engineering approach to high-density PVA does lead to polymeric ice nucleating agents.

The less densely grafted P((P(NB-NH))_g-PVA\(_{310}\))-stat-P(NB-NH)\(_{320}\) (n = 50, 100, 200, 400) showed intermediate activity, while the densely grafted PNB\(_{4000}\)-PVA\(_{3000}\) (n = 20, 30, 40) were more active, suggesting that dense presentation of PVA side chains is essential for nucleation activity—these trends are shown in Figure S39. With the much smaller droplets frozen in the DSC measurements, where each droplet contains molecules of the order of 10\(^9\), the low-density bottlebrushes nucleated ice less well than linear PVA, while the...
high-density bottlebrushes nucleated ice better than linear PVA. High-density PNB-\(g\)-PVA\(_{208}\) was observed to nucleate ice at average temperatures as warm as \(-19.2\) °C, which is unprecedented for a fully synthetic soluble material. We have previously reported that large (>200 nm) polymer nanoparticles with a poly(vinyl pyrrolidone), PVP, corona also nucleated ice at similar temperatures.\(^{32}\) The large size and dense display in those particles may be related, but the particles did not have variable density coronas so the structure–property relationship was not clear. These particles did show weak ice binding (and IRI) and hence agree with the results here. Soluble extracts from pollen, which are thought to be polysaccharides, have been reported to nucleate ice, but these particles did not have variable density coronas so the contribution of a small number of larger aggregates is impossible to exclude, in this initial report.

There is evidence that ice nucleating protein activity is related to other ice-binding proteins based on their size and packing.\(^{3,32,33}\) The mechanism(s) of heterogeneous ice nucleation is not fully understood, even for the long-studied ice nucleation protein from \textit{Pseudomonas syringae}. This is in part due to the challenge of studying membrane-anchored proteins, and that isolation of these as pure protein domains does not recapitulate the 3D assembly. Protein expression techniques only allow for linear extension (in a single protein), whereas this polymeric strategy provides defined macro-molecules with multiple ice binding domains in a constrained manner, therefore providing experimental evidence that crowding and size are essential for nucleation, in agreement with modeling of IBPs.\(^{3}\) CNT suggests that a nucleation site must have a diameter of between 2 and 5 nm, to nucleate ice at \(-19\) °C, with the size depending on the strength of the interaction with the ice critical cluster. The bottlebrushes are each at least 100 nm across and therefore larger than the ice cluster. It seems likely that the occasional patch of bottlebrush surface assumes a conformation that is sufficiently icelike to act as a nucleation site. In our ice nucleation experiments, we kept the mass of polymer per droplet the same while varying the size of the polymers.

This means that our experiments using larger bottlebrushes, which induce ice nucleation at higher temperatures, had less polymer surface area in contact with water. Simulations suggest that curved surfaces nucleate ice less well than flat surfaces.\(^{34}\) We tentatively suggest that the lesser curvature of the larger bottlebrushes increases the likelihood that they harbor an active site, and that for more rigid conformation, high-density bottlebrushes similarly favor the emergence of larger flat areas of icelike structure in the polymer surfaces. While the magnitude of ice nucleation activity observed here is much less than that for ice nucleating proteins, the ability to sequentially modify a bottom-up synthesized chemical nucleator will enable new avenues of discovery in the study of ice nucleators, not possible with biological materials, or the poorly defined inorganic nucleator.

**CONCLUSIONS**

Here, we report a synthetic polymer capable of nucleating ice, inspired by how ice-binding proteins assemble into ice-nucleating structures. Our design was based on molecular bottlebrushes with (ice binding) PVA side chains, with the side-chain density tuned using ROMP to enable “grafting-through” (high density) or “grafting-to” (low density). The bottlebrush polymers were fully characterized, confirming the confinement of the PVA chains in the higher-density brushes, which also gave rise to LCST-type behavior, not seen for linear equivalents. The bottlebrush polymers were found to bind ice and retain their potent ice-recrystallization inhibition activity, with a slight increase relative to linear PVA. However, due to the limited solubility of high-MW linear PVA, this effect could be due to the size increase, rather than a pure architectural effect. Using ice nucleation assays, we observed that high-density PVA bottlebrushes significantly increased the ice nucleation temperature compared to other materials. Lower-
density PVA brushes also favored nucleation in microliter droplet freezing assays, although they nucleated ice at colder temperatures than linear PVA in an emulsion freezing experiment using picoliter scale droplets. We hypothesize that larger bottlebrushes are more likely to possess an ice nucleation site due to their low degree of curvature and that the conformational rigidity of high-density bottlebrushes is again expected to favor the formation of flat, icelike water–polymer interfaces that can act as ice nucleation sites. The role of aggregates of these brushes could not be excluded. Our strategy of controlling side-chain density would be challenging with protein engineering tools alone. These materials may guide the development of new ice nucleators for cryobiology as synthetic probes to elucidate fundamental mechanisms of action and show that programmable ice nucleation from bottom-up synthesis may be achievable.

**ASSOCIATED CONTENT**

Supporting Information
The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acs.biomac.2c01097.

Materials and additional synthetic methods; experimental procedures; synthetic procedures; supplementary NMR, SEC, FTIR, DLS, TEM, and AFM data; and additional ice shaping/nucleation data (PDF)

**AUTHOR INFORMATION**

**Corresponding Authors**
Thomas F. Whale – Department of Chemistry, University of Warwick, CV4 7AL Coventry, U.K.; Email: tom.whale@warwick.ac.uk
Matthew I. Gibson – Department of Chemistry, University of Warwick, CV4 7AL Coventry, U.K.; Division of Biomedical Sciences, Warwick Medical School, University of Warwick, CV4 7AL Coventry, U.K.; orcid.org/0000-0002-8297-1278; Email: m.i.gibson@warwick.ac.uk

**Authors**
Panagiotis G. Georgiou – Department of Chemistry, University of Warwick, CV4 7AL Coventry, U.K.; orcid.org/0000-0001-8968-1057
Nina L. H. Kinney – Department of Chemistry, University of Warwick, CV4 7AL Coventry, U.K.; orcid.org/0000-0003-3123-7361
Ioanna Kontopoulou – Department of Chemistry, University of Warwick, CV4 7AL Coventry, U.K.
Alexander N. Baker – Department of Chemistry, University of Warwick, CV4 7AL Coventry, U.K.
Steven A. Hindmarsh – Department of Physics, University of Warwick, CV4 7AL Coventry, U.K.
Akalabya Bissoyi – Department of Chemistry, University of Warwick, CV4 7AL Coventry, U.K.
Thomas R. Congdon – Department of Chemistry, University of Warwick, CV4 7AL Coventry, U.K.

Complete contact information is available at: https://pubs.acs.org/10.1021/acs.biomac.2c01097

Notes
The authors declare no competing financial interest.

**ACKNOWLEDGMENTS**

This project (MIG) received funding from the European Research Council (ERC) under the European Union’s Horizon 2020 research and innovation program (grant agreement nos. 866056 and 899872) and the Royal Society for an Industry Fellowship (191037) joint with Cytica. This project also received funding from the European Union’s Horizon 2020 research and innovation program under the Marie Skłodowska-Curie Grant Agreement No. 814236. T.F.W. thanks the Leverhulme Trust and the University of Warwick for supporting an Early Career Fellowship (ECF-2018-127). N.K. thanks the Natural Environment Research Council (NE/S007350/1) for a PhD studentship. The Warwick Polymer Research Technology Platform is acknowledged for SEC analysis, and the Warwick Electron Microscopy Research Technology Platform is acknowledged for TEM and AFM. The authors also acknowledge the University of Warwick Advanced Bioimaging Research Technology Platform supported by BBSRC ALERT14 Award BB/M01228X/1 and Dr. S. Bakker for TEM. Dr. D.J Fox is thanked for assistance with the (safe handling) of the LiAlH₄ reduction step. For the purpose of open access, the author has applied a Creative Commons Attribution (CC BY) license to any author-accepted manuscript version arising from this submission.

**REFERENCES**

(1) Bar Dolev, M.; Braslavsky, I.; Davies, P. L. Ice-Binding Proteins and Their Function. *Annu. Rev. Biochem.* 2016, 85, 515–542.
(2) Meister, K.; Straazdaite, S.; DeVries, A. L.; Lotze, S.; Olijve, L. L. C.; Voets, I. K.; Bakker, H. J. Observation of Ice-like Water Layers at an Aqueous Protein Surface. *Proc. Natl. Acad. Sci. U.S.A.* 2014, 111, 17732–17736.
(3) Qiu, Y.; Hudait, A.; Moliner, V. How Size and Aggregation of Ice-Binding Proteins Control Their Ice Nucleation Efficiency. *J. Am. Chem. Soc.* 2019, 141, 7439–7452.
(4) Venketesh, S.; Dayananda, C. Properties, Potentials, and Prospects of Antifreeze Proteins. *Crit. Rev. Biotechnol.* 2008, 28, 57–82.
(5) Forbes, J.; Bissoyi, A.; Eickhoff, L.; Reicher, N.; Hansen, T.; Bon, C. G.; Walker, V. K.; Koop, T.; Rudich, Y.; Braslavsky, I.; Davies, P. L. Water-Organizing Motif Continuity Is Critical for Potent Ice Nucleation Protein Activity. *Nat. Commun.* 2022, 13, No. 5019.
(6) Voets, I. K. From Ice-Binding Proteins to Bio-Inspired Antifreeze Materials. *Soft Matter* 2017, 13, 4808–4823.
(7) Biggs, C. I.; Bailey, T. L.; Graham, B.; Stubbs, C.; Fayer, A.; Gibson, M. I. Polymer Mimics of Biomacromolecular Antifreezes. *Nat. Commun.* 2017, 8, No. 1546.
(8) He, Z.; Liu, K.; Wang, J. Bioinspired Materials for Controlling Ice Nucleation, Growth, and Recrystallization. *Acc. Chem. Res.* 2018, 51, 1082–1091.
(9) Inada, T.; Lu, S. S. Inhibition of Recrystallization of Ice Grains by Adsorption of Poly(Vinyl Alcohol) onto Ice Surfaces. *Cryst. Growth Des.* 2003, 3, 747–752.
(10) Budke, C.; Koop, T. Ice Recrystallization Inhibition and Molecular Recognition of Ice Faces by Poly(Vinyl Alcohol). *ChemPhysChem* 2006, 7, 2601–2606.
(11) Congdon, T.; Notman, R.; Gibson, M. I. Antifreeze (Glyco)Protein Mimetic Behavior of Poly(Vinyl Alcohol): Detailed Structure Ice Recrystallization Inhibition Activity Study. *Biomacromolecules* 2013, 14, 1578–1586.
(12) Congdon, T. R.; Notman, R.; Gibson, M. I. Synthesis of Star-Branched Poly(Vinyl Alcohol) and Ice Recrystallization Inhibition Activity. *Eur. Polym. J.* 2017, 88, 320–327.
(13) Olijve, L. L. C.; Hendrix, M. R. M.; Voets, I. K. Influence of Polymer Chain Architecture of Poly(Vinyl Alcohol) on the Inhibition of Ice Recrystallization. *Macromol. Chem. Phys.* 2016, 217, 951–958.
Biomacromolecules

D. Structure and Dynamics of an Alpha-Helical Antifreeze Protein. Clathrate Waters Bind Antifreeze Proteins to Ice. Nature Nanosheets. Chem. Lett. of Active Sites. High-Speed Imaging of Ice Nucleation in Water Proves the Existence L. The Importance of Feldspar for Ice Nucleation by Mineral Dust in Baustian, K. J.; Carslaw, K. S.; Dobbie, S.; Sullivan, D. O.; Malkin, T. 16, 8915 (2019), 19, 2742–2753.

(37) McKenzie, T. G.; Fu, Q.; Uchiyama, M.; Satoh, K.; Xu, J.; Boyer, C.; Kamigaito, M.; Qiao, G. G. Beyond Traditional RAFT: Alternative Activation of Thioacrylonitrile Compounds for Controlled Polymerization. Adv. Sci. 2016, 3, No. 1500394.

(38) Aragón, S. B.; Pecora, R. Theory of Dynamic Light Scattering from Large Anisotropic Particles. J. Chem. Phys. 1977, 66, 2506–2516.

(39) Zhang, Z.; Carrillo, J.-M. Y.; Ahn, S.; Wu, B.; Hong, K.; Smith, G. S.; Do, C. Atomicistic Structure of Bottlebrush Polymers: Simulations and Neutron Scattering Studies. Macromolecules 2014, 47, 5808–5814.

(40) Pesek, S. L.; Xiang, Q.; Hammouda, B.; Verduzzo, R. Small-Angle Neutron Scattering Analysis of Bottlebrush Backbone and Side Chain Flexibility. J. Polym. Sci., Part B: Polym. Phys. 2017, 55, 104–111.

(41) Gallayamov, M. O.; Tartsch, B.; Mela, P.; Potemkin, I. L.; Sheiko, S. S.; Börner, H.; Matyjaszewski, K.; Khokhlov, A. R.; Möller, M. Vapor-Induced Scattering Dynamics of Adsorbed Linear and Brush-like Macromolecules as Observed by Environmental SEM: Polymer Chain Statistics and Scaling Exponents. J. Polym. Sci., Part B: Polym. Phys. 2007, 45, 2368–2379.

(42) Radzinski, S. C.; Foster, J. C.; Chapleski, R. C.; Troya, D.; Matson, J. B. Bottlebrush Polymer Synthesis by Ring-Opening Metathesis Polymerization: The Significance of the Anchor Group. J. Am. Chem. Soc. 2016, 138, 6998–7004.

(43) Blosch, S. E.; Scannelli, S. J.; Alabaliar, M.; Matson, J. B. Complex Polymer Architectures Using Ring-Opening Metathesis Polymerization: Synthesis, Applications, and Practical Considerations. Macromolecules 2022, 55, 4200–4227.

(44) Congdon, T.; Shaw, P.; Gibson, M. I. Thermoresponsive, Well-Defined, Poly(Vinyl Alcohol) Co-Polymers. Polym. Chem. 2015, 6, 4749–4757.

(45) Varlas, S.; Hua, Z.; Jones, J. R.; Thomas, M.; Foster, J. C.; O’Reilly, R. K. Complementary Nucleobase Interactions Drive the Hierarchical Self-Assembly of Core–Shell Bottlebrush Block Copolymers toward Cylindrical Supramolecules. Macromolecules 2020, 53, 9747–9757.

(46) Knight, C. A.; Hallett, J.; DeVries, A. L. Solute Effects on Ice Recrystallization: An Assessment Technique. Cryobiology 1988, 25, 55–60.

(47) Graham, B.; Fayer, A. E. R.; Houston, J. E.; Evans, R. C.; Gibson, M. I. Facially Amphiphatic Glycopolymer Reactor Inhibits Ice Recrystallization. J. Am. Chem. Soc. 2018, 140, 5682–5685.

(48) Stevens, C. A.; Drori, R.; Zalis, S.; Braslavsky, I.; Davies, P. L. Dendrimer-Linked Antifreeze Proteins Have Superior Activity and Thermal Recovery. Bioconjugate Chem. 2015, 26, 1908–1915.

(49) Adar, C.; Reicher, N.; Braslavsky, I.; Rudich, Y.; Koop, T. Contrasting Behavior of Antifreeze Proteins: Ice Growth Inhibitors and Ice Nucleation Promoters. J. Phys. Chem. Lett. 2019, 10, 966–972.

(50) Ogawa, S.; Koga, M.; Osanai, S. Anomalous Ice Nucleation Behavior in Aqueous Polyvinyl Alcohol Solutions. Chem. Phys. Lett. 2009, 480, 86–89.
Enhancement of Ice Recrystallization Inhibition. *J. Am. Chem. Soc.* 2021, 143, 7449−7461.

(53) Bissoyi, A.; Reicher, N.; Chasnitsky, M.; Arad, S.; Koop, T.; Rudich, Y.; Braslavsky, I. Ice Nucleation Properties of Ice-Binding Proteins from Snow Fleas. *Biomolecules* 2019, 9, No. 532.

(54) Lupi, L.; Hudait, A.; Molinero, V. Heterogeneous Nucleation of Ice on Carbon Surfaces. *J. Am. Chem. Soc.* 2014, 136, 3156−3164.