Antifreeze proteins (AFPs) have many potential applications, ranging from cryobiology to aerospace, if they can be incorporated into materials. Here, a range of engineered AFP mutants were prepared and site-specifically conjugated onto RAFT polymer-stabilized gold nanoparticles to generate new hybrid multivalent ice growth inhibitors. Only the SNAP-tagged AFPs lead to potent ‘antifreeze’ active nanomaterials with His-Tag capture resulting in no activity, showing the mode of conjugation is essential. This versatile strategy will enable the development of multivalent AFPs for translational and fundamental studies.

AFPs have been observed to lead to oligomerization to a tetramer and the onset of hyperactivity, linked to its supramolecular assembly. Davies and coworkers assembled 6–11-mers of AFP type III on PAMAM dendrimers to mimic this oligomerization. On a per-protein basis there was only a small increase in IRI, but the ability to span multiple ice faces increased the TH activity. Synthetic mimics of AFPs also show strong molecular weight dependence on activity, with longer polymers (such as poly(vinyl alcohol) having significantly higher IRI activity) as do supramolecular safranine-O based mimics.

Despite the evidence for increasing the valency of AFPs to modulate activity there remain few reports of multivalent display, in part due to the challenges of site-specific protein conjugation. Traditional approaches to combine polymers with proteins involve targeting unpaired cystine residues. Unnatural amino acids for bioconjugation can be incorporated in a site-specific fashion by the AMBER stop codon. Johnson and co-workers have developed recombinantly expressible ‘SNAP-tags’ based on O’alkylguanine-DNA alkyltransferase. By attaching the tag as a fusion protein a covalent bond can be formed to any surface bearing benzylguanine without any unnatural amino acids. The commonly used hexa-histidine purification tag can also facilitate ionic-conjugation.

Here AFPs are conjugated, by site-specific methods, onto nanoparticles to generate hybrid ice growth inhibiting materials to mimic the multivalent presentation of hyperactive antifreeze proteins and aid the application of these exciting proteins (Fig. 1).

RAFT (reversible addition fragmentation chain transfer) polymerization was employed to synthesize telechelic poly(hydroxyethyl acrylamide), pHEA, bearing a pentafluorophenyl (PFP) ester at the ω-terminus and a trithiocarbonate at the α-terminus. These polymers were characterized by SEC, 1H, 19F NMR and IR (Table 1). The PFP group was substituted by addition of amino-benzylguanine (BG) (for SNAP conjugation) or tris-NTA amine (for His-Tag capture). Successful conjugation was confirmed by 19F NMR (Fig. 2B) as well as by
By using an excess of the amine, the RAFT agent end group was displaced to reveal a thiol for gold particle conjugation. It was also attempted to introduce a maleimide onto the particles, but this did not lead to stable particles (see ESI† for details) so was not taken further.

4 nm citrate-stabilized gold nanoparticles (AuNP) were synthesized by NaBH4 reduction of HAuCl4 and subsequently functionalized with thiol-terminated polymers by mixing, followed by centrifugal dialysis.35 The nanoparticles were characterized by transmission electron microscopy (Fig. 3A–C), dynamic light scattering (Fig. 3E) and UV-Vis confirming successful addition of the polymer. X-Ray photoelectron spectroscopy (XPS) showed a clear increase in the nitrogen N 1s signal following coating (Fig. 3D). Screening experiments showed that NTA-pHEA gave more stable particles when \( M_n = 3000 \text{ g mol}^{-1} \) and for BG-pHEA = 10 000 g mol\(^{-1} \) and hence these chain lengths were used from this point onwards.

Recombinant AFP type III (from ocean pout) were produced by expression in *Escherichia coli*, and purified by nickel affinity chromatography and HPLC. Three mutants were produced bearing N-terminal modifications of hexa-histidine (His-AFP), SNAP (SNAP-AFP) and cysteine (Cys-AFP). Direct conjugation of the SNAP-AFP was achieved by incubation with the BG-pHEA45@AuNP4 in buffer. NTA-pHEA17@AuNP4 was first activated by addition of NiCl2 followed by His-AFP. Direct Cys-AFP immobilization onto AuNPs was unsuccessful (see ESI†). Particles were further purified by centrifugal dialyses and removal of non-conjugated residual protein confirmed by nanodrop absorption analysis on residual washings. Following

---

**Table 1** Precursor polymers synthesized

| Polymer     | [M]/[CTA] (−) | Conv.\(^a\) (%) | DP\(_n\) (−) | \( M_{n(THF)}^a \) (g mol\(^{-1} \)) | \( M_{n(SEC)}^b \) (g mol\(^{-1} \)) | \( D^b \) (−) |
|-------------|--------------|-----------------|-------------|---------------------------------|---------------------------------|-------------|
| PFP-pHEA95  | 103          | 93              | 95          | 11 000                          | 15 000                          | 1.3         |
| PFP-pHEA45  | 47           | 94              | 45          | 5700                           | 8300                            | 1.1         |
| PFP-pHEA17  | 26           | 64              | 17          | 2500                           | 3000                            | 1.1         |

\(^a\) Determined by \(^1\)H NMR. \(^b\) Determined by SEC against PMMA standards.

---

IR. By using an excess of the amine, the RAFT agent end group was displaced to reveal a thiol for gold particle conjugation. It was also attempted to introduce a maleimide onto the particles, but this did not lead to stable particles (see ESI† for details) so was not taken further.

4 nm citrate-stabilized gold nanoparticles (AuNP) were synthesized by NaBH4 reduction of HAuCl4 and subsequently functionalized with thiol-terminated polymers by mixing, followed by centrifugal dialysis.35 The nanoparticles were characterized by transmission electron microscopy (Fig. 3A–C), dynamic light scattering (Fig. 3E) and UV-Vis confirming successful addition of the polymer. X-Ray photo electron spectroscopy (XPS) showed a clear increase in the nitrogen N 1s signal following coating (Fig. 3D). Screening experiments showed that NTA-pHEA gave more stable particles when \( M_n = 3000 \text{ g mol}^{-1} \) and for BG-pHEA = 10 000 g mol\(^{-1} \) and hence these chain lengths were used from this point onwards.

Recombinant AFP type III (from ocean pout) were produced by expression in *Escherichia coli*, and purified by nickel affinity chromatography and HPLC. Three mutants were produced bearing N-terminal modifications of hexa-histidine (His-AFP), SNAP (SNAP-AFP) and cysteine (Cys-AFP). Direct conjugation of the SNP-AFP was achieved by incubation with the BG-pHEA45@AuNP4 in buffer. NTA-pHEA17@AuNP4 was first activated by addition of NiCl2 followed by His-AFP. Direct Cys-AFP immobilization onto AuNPs was unsuccessful (see ESI†). Particles were further purified by centrifugal dialyses and removal of non-conjugated residual protein confirmed by nanodrop absorption analysis on residual washings. Following
With these site-specifically attached AFP/nanoparticle conjugates to hand, their IRI activity could be assessed by the 'splat' assay. In brief, a polynucleated wafer of ice crystals is seeded from a solution containing the compounds of interest and the ice crystals annealed at $-8 \, ^\circ \text{C}$ for 30 minutes. After this time the average ice crystal size is evaluated and compared to a negative control of buffer alone: smaller ice crystal sizes signify more IRI activity. The two AFP mutants were very potent inhibitors, preventing all growth below 0.01 mg mL$^{-1}$. There was no significant difference between the two mutants, showing the fusions were tolerated in terms of retention of function. His-AFP-Ni-NTA-pHEA$_{17}$@Au$_4$ was found to be surprisingly inactive as an ice growth inhibitor with no activity up to 1 mg mL$^{-1}$ (AFP concentration). This could be due to the dynamic nature of the His-NTA interaction or incorrect orientation of the AFP on the surface. This also shows that designing a multivalent AFP is non-trivial, with the correct chemistry being essential. Previous studies suggest excess charge does not affect AFP activity and was ruled out as the cause.$^{36}$ In contrast, the covalent hybrid nanoparticle SNAP-AFP-BG-pHEA$_{45}$@Au$_4$ was found to be a very potent IRI, Fig. 4B. In terms of total AFP concentration (to enable multivalent effects to be seen) the nanoparticles essentially identical activity compared to SNAP-AFP alone. If considered on a molar basis (as has been done by Davies et al. for dendrimer AFPs$^{22}$) then the particles are more active. AFPs are known to become hyperactive when they oligomerize, but such enhancement was not seen here due to the different 3-D placement of the AFPs.$^{21}$ Using a nanoliter osmometer the thermal hysteresis (TH) gap (non-colligative freezing point depression) was determined. TH is closely linked to the ability to bind specific ice faces (in this case the prism plane) hence providing additional information about the functionality of the particles; the TH is unique to antifreeze proteins, and is defined as the difference between the freezing and melting points. It was observed that the AFP and SNAP-AFP nanoparticles have very similar TH gaps, of 1.4 °C at 7.6 mg mL$^{-1}$. As with the IRI measurements, this confirms that all activity is retained despite the major structural modifications and that the placement of the AFPs via-polymeric linkers is tolerated. As the effective AFP concentration in solution when immobilized on NPs is lower, this suggests that the particles can bind multiple planes of ice simultaneously. To probe this dynamic ice shaping experiments were conducted in sucrose solution (as described previously$^{28}$). Compared to SNAP-AFP alone (Fig. 4H), the SNAP–AuNP conju-

**Table 2**  Nanoparticle characterization

| Particle                        | $D_h$  | $D_{TEM}$   | $\zeta$ potential |
|--------------------------------|--------|-------------|-------------------|
| Citrate@Au$_4$                 | 4.3    | 3.9 ± 0.7   | $-19.4 ± 2.7$     |
| NTA-pHEA$_{17}$@Au$_4$         | 8.5    | 3.7 ± 0.8   | $-15.9 ± 4.8$     |
| Ni-NTA-pHEA$_{17}$@Au$_4$      | 7.7    | 3.9 ± 0.8   | $-6.16 ± 1.1$     |
| His-AFP-Ni-NTA-pHEA$_{17}$@Au$_4$ | 8.9    | 4.3 ± 0.9   | $-7.03 ± 1.3$     |
| BG-pHEA$_{45}$@Au$_4$          | 16.5   | 4.0 ± 1.1   | $-10.2 ± 1.9$     |
| SNAP-AFP-BG-pHEA$_{45}$@Au$_4$ | 8.92   | 4.0 ± 1.1   | $-4.35 ± 0.9$     |

$^a$ Hydrodynamic diameter from DLS. $^b$ Gold core diameter, average of >100 particles by TEM.
gates resulted in more faceting of the ice crystals, consistent with them being able to bind more ice faces simultaneously.22

Conclusions

In conclusion, we report the synthesis of hybrid nanoparticle–AFP conjugates, making use of polymer-stabilized gold nanoparticles and protein engineering to enable precision-conjugation. Hetero-telechelic polymers were obtained by RAFT, enabling a nanoparticle immobilization moiety (thiol) and distal protein capture (nickel NTA or O6-benzyl-guanine) moiety to be installed. The polymer coating is essential to enable conjugation in buffer, avoiding nanoparticle aggregation. It was found that the SNAP-tag (covalent) approach lead to the most active particles compared to oligohistidine capture. The observed ice recrystallisation inhibition and thermal hysteresis activity was comparable to free antifreeze proteins on a valency-corrected basis confirming retention of activity in this format. Other conjugation methods based on ionic interactions did not give stable particles or resulted in loss of activity. These results demonstrate that antifreeze protein can be incorporated into more complex assemblies to facilitate their application in colloid science or for fundamental studies exploiting nanoparticle cores for bioimaging and tracking.

Data access statement

The research data supporting this publication can be found at http://wrap.warwick.ac.uk.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

We acknowledge funding from the ERC (638661 and 789182), Leverhulme Trust RPG (2015 0194), BBSRC (BB/R506588/1) and the University of Warwick. Peter Davies and Robert Eves (both Queens University, Canada) are thanked for the TH measurements and providing the AFP type III plasmids.

Notes and references

1 P. L. Davies, Trends Biochem. Sci., 2014, 39, 548–555.
2 M. M. Harding, P. I. Anderberg and A. D. J. Haymet, Eur. J. Biochem., 2003, 270, 1381–1392.
3 A. Hakim, J. B. Nguyen, K. Basu, D. F. Zhu, D. Thakral, P. L. Davies, F. J. Isaacs, Y. Modis and W. Meng, J. Biol. Chem., 2013, 288, 12295–12304.
4 C. Sidebottom, S. Buckley, P. Pudney, S. Twigg, C. Jarman, C. Holt, J. Telford, A. McArthur, D. Worrall, R. Hubbard and P. Lillford, Nature, 2000, 406, 256.
5 K. R. Walters, A. S. Serianni, T. Sfrorno, B. M. Barnes, J. G. Duman and J. G. Duman, Proc. Natl. Acad. Sci. U. S. A., 2009, 106, 20210–20215.
6 O. Parent and A. Ilinca, Cold Reg. Sci. Technol., 2011, 65, 88–96.
7 W. O. Valarezo, F. T. Lynch and R. J. McGhee, J. Aircr., 1993, 30, 807–812.
8 C. I. Biggs, T. L. Bailey, B. Graham, C. Stubbs, A. Fayer and M. I. Gibson, Nat. Commun., 2017, 8, 1546.
9 I. K. Voets, Soft Matter, 2017, 13, 4808–4823.
10 A. P. Esser-Kahn, V. Trang and M. B. Francis, J. Am. Chem. Soc., 2010, 132, 13264–13269.
11 K. Liu, C. Wang, J. Ma, G. Shi, X. Yao, H. Fang, Y. Song and J. Wang, Proc. Natl. Acad. Sci. U. S. A., 2016, 113, 14739–14744.
12 Y. Tachibana, G. L. Fletcher, N. Fujitani, S. Tsuda, K. Monde and S. I. Nishimura, Angew. Chem., Int. Ed., 2004, 43, 856–862.
13 C. P. Garnham, R. L. Campbell and P. L. Davies, Proc. Natl. Acad. Sci. U. S. A., 2011, 108, 7363–7367.
14 K. Meister, S. Straadaike, A. L. DeVries, S. Lotze, L. L. C. Olijve, I. K. Voets and H. J. Bakker, Proc. Natl. Acad. Sci. U. S. A., 2014, 111, 17732–17736.
15 J. A. Raymond, P. Wilson and A. L. DeVries, Proc. Natl. Acad. Sci. U. S. A., 1989, 86, 881–885.
16 L. L. C. Olijve, K. Meister, A. L. DeVries, J. G. Duman, S. Guo, H. J. Bakker and I. K. Voets, Proc. Natl. Acad. Sci. U. S. A., 2016, 113, 3740–3745.
17 C. B. Marshall, G. L. Fletcher and P. L. Davies, Nature, 2004, 429, 153.
18 S. P. Gaethert, M. J. Kulper, S. M. Gagné, V. K. Walker, Z. Jia, B. D. Sykes and P. L. Davies, Nature, 2000, 406, 325–328.
19 A. J. Scotter, C. B. Marshall, L. A. Graham, J. A. Gilbert, C. P. Garnham and P. L. Davies, Cryobiology, 2006, 53, 229–239.
20 K. Mochizuki and V. Molinero, J. Am. Chem. Soc., 2018, 140, 4803–4811.
21 S. Mahatabuddin, Y. Hanada, Y. Nishimiya, A. Miura, H. Kondo, P. L. Davies and S. Tsuda, Sci. Rep., 2017, 7, 42501.
22 C. A. Stevens, R. Drori, S. Zalis, I. Braslavsky and P. L. Davies, Bioconjugate Chem., 2015, 26, 1908–1915.
23 C. Budke, A. Dreyer, J. Jaeger, K. Gimpel, T. Berkemeier, A. S. Bonin, L. Nagel, C. Plattner, A. L. Devries, N. Sewald and T. Koop, Cryst. Growth Des., 2014, 14, 4285–4294.
24 T. Congdon, R. Notman and M. I. Gibson, Biomacromolecules, 2013, 14, 1578–1586.
25 C. Stubbs, J. Lipecki and M. I. Gibson, Biomacromolecules, 2017, 18, 295–302.
26 R. Drori, C. Li, C. Hu, P. Raiteri, A. Rohl, M. D. Ward and B. Kahr, J. Am. Chem. Soc., 2016, 138, 13396–13401.
27 N. Stephanopoulos and M. B. Francis, Nat. Chem. Biol., 2011, 7, 876–884.
28 C. Stubbs, L. E. Wilkins, A. E. R. Fayer, M. Walker and M. I. Gibson, Langmuir, 2018, DOI: 10.1021/acs.langmuir.8b01952.
29  D. Bontempo, K. L. Heredia, B. A. Fish and H. D. Maynard, 
J. Am. Chem. Soc., 2004, **126**, 15372–15373.

30  M. W. Jones, R. A. Strickland, F. F. Schumacher, 
S. Caddick, J. R. Baker, M. I. Gibson and D. M. Haddleton, 
Chem. Commun., 2012, **48**, 4064–4066.

31  C. H. Kim, J. Y. Axup and P. G. Schultz, Curr. Opin. Chem. 
Biol., 2013, **17**, 412–419.

32  A. Keppler, S. Gendreizig, T. Gronemeyer, H. Pick, 
H. Vogel and K. Johnsson, Nat. Biotechnol., 2003, **21**, 86–89.

33  H. Y. Cho, M. A. Kadir, B. S. Kim, H. S. Han, 
S. Nagasundarapandian, Y. R. Kim, S. B. Ko, S. G. Lee and 
H. J. Paik, Macromolecules, 2011, **44**, 4672–4680.

34  S.-J. Richards and M. I. Gibson, ACS Macro Lett., 2014, 3, 
1004–1008.

35  N. S. Ieong, K. Brebis, L. E. Daniel, R. K. O’Reilly and 
M. I. Gibson, Chem. Commun., 2011, **47**, 11627–11629.

36  R. C. Deller, B. M. Carter, I. Zampetakis, F. Scarpa and 
A. W. Perriman, Biochem. Biophys. Res. Commun., 2018, **495**, 
1055–1060.