Introduction

The structures of amyloid fibrils and oligomers represent a vast frontier, of yet unknown scope. The fibrils and aggregates that amyloidogenic peptides and proteins form are rich in β-sheets, and their structures are tremendously important in amyloid diseases. Many structures of amyloid fibrils have been discovered by solid-state NMR spectroscopy of amyloidogenic peptides and proteins and by X-ray crystallography of smaller fragments. Studying amyloid oligomer structures at high resolution is challenging, because amyloid oligomers are heterogeneous and dynamic, forming various species of different sizes and morphologies. Although a few structures of amyloid oligomers have been discovered in the last decade, there are not enough to provide a full understanding of amyloid assemblies.

Our laboratory has pioneered the use of macrocyclic β-sheets as a tool for exploring the structures of amyloid oligomers. In collaboration with the Eisenberg group, we began using X-ray crystallography to elucidate the oligomeric assemblies of macrocyclic β-sheets. Subsequently, our laboratory has independently elucidated oligomer structures formed by macrocyclic β-sheet peptides derived from Aβ, α-synuclein, and β2-microglobulin, which are associated with Alzheimer’s disease, Parkinson’s disease, and hemodialysis-related amyloidosis.

Transthyretin (TTR) is a tetrameric protein that is associated with amyloid diseases, including senile systemic amyloidosis and familial amyloid polyneuropathy. Two monomers of TTR come together in an edge-to-edge fashion to form dimers. The dimers further pack in a face-to-face fashion to form tetramers (Fig. 1). The TTR tetramers transport thyroid hormone and retinol-binding protein. Dissociation of the tetramers into monomers leads to formation of insoluble fibrils and toxic oligomers, which are associated with TTR-related amyloid diseases.

Development of small molecule drugs has thus become an important approach for treating familial amyloid polyneuropathy by stabilizing the tetramers with a small molecule drug called tafamidis. Tafamidis acts by binding to the dimerization interface of two dimers and preventing dissociation into monomers.

Inspired by the importance of the dimerization interface in stabilizing the TTR assembly and the propensity of TTR to form fibrils, we set out to elucidate the assembly of the β-hairpin that comprises the dimerization interface of the TTR tetramer. We incorporated the two β-strands from the TTR β-hairpin into a macrocyclic β-sheet and studied its assembly by X-ray crystallography. To our surprise, we discovered that the peptide exhibits a new mode of supramolecular assembly, forming square channels consisting of an extended network of β-sheets.

Results

Design, synthesis, and structure determination of peptide 1

We incorporated residues 106–112 from strand G of TTR and residues 115–121 from strand H into a macrocyclic β-sheet (Fig. 2). We used two δ-linked ornithine (δOrn) turn units to connect these two β-strands and form a macrocycle. We replaced the proline-tyrosine turn (residues 113 and 114) with a δOrn turn to promote β-hairpin formation. We connected residues 106 and 121 with another δOrn turn to reinforce the β-hairpin structure. We also incorporated an N-methyl group on alanine 109 (N-Me-A109) to prevent uncontrolled
We replaced tyrosine 116 with \( p \)-iodo-phenylalanine (FI\(_{\text{116}}\)) to determine the X-ray crystallographic phases by single-wavelength anomalous diffraction (SAD) phasing. Peptide\(_1\) was synthesized as the trifluoroacetic acid (TFA) salt using standard Fmoc-based solid-phase peptide synthesis, solution-phase cyclization, and RP-HPLC purification. The TFA salt is readily soluble in water, but crystallizes or aggregates in buffer, depending on the conditions. Peptide\(_1\) was screened in 288 crystallization conditions using Hampton Research crystallization kits (Crystal Screen, Index, and PEG/Ion). Peptide\(_1\) grew rod-shaped crystals suitable for X-ray diffraction after two weeks in 0.1 M NaOAc buffer at pH 5.3, 0.2 M CaCl\(_2\), and 31% isopropanol. X-ray diffraction data were collected to 2.08 Å using an X-ray diffractometer with a Cu rotating anode. The X-ray crystallographic structure of peptide\(_1\) was solved and refined in the \( P4_212_1 \) space group, with one macrocycle in the asymmetric unit.

**X-ray crystallography of peptide\(_1\)**

The X-ray crystallographic structure of peptide\(_1\) reveals a hierarchical supramolecular assembly. The peptide folds to form a \( \beta \)-hairpin monomer. The \( \beta \)-hairpin assembles to form extended \( \beta \)-sheets. Four extended \( \beta \)-sheets form a square channel. The square channels pack into a pattern that resembles tilted windows.

**\( \beta \)-Hairpin monomer.** Peptide\(_1\) forms a hydrogen-bonded antiparallel \( \beta \)-sheet (Fig. 3). The \( \beta \)-sheet is relatively flat, with two distinct surfaces. One surface displays side chains from residues T\(_{106}\), A\(_{108}\), L\(_{110}\), S\(_{112}\), S\(_{115}\), S\(_{117}\), T\(_{119}\), and V\(_{121}\); the other surface displays side chains from I\(_{107}\), N-Me-A\(_{109}\), L\(_{111}\), FI\(_{116}\), T\(_{118}\), and A\(_{120}\). We term these two surfaces the major and minor faces. The major face contains mostly polar side chains and the minor face contains mostly hydrophobic side chains.

**Extended \( \beta \)-sheets.** Peptide\(_1\) forms an extended network of \( \beta \)-sheets (Fig. 4). The \( \beta \)-sheets are not fully aligned, but rather are shifted by four residues toward the C-terminus. The upper strand of one \( \beta \)-hairpin monomer forms four hydrogen bonds with the lower strand of the adjacent \( \beta \)-hairpin monomer. The \( \delta \)-NH of the ornithine that connects S\(_{112}\) and S\(_{115}\) hydrogen bonds with the amide oxygen of T\(_{118}\); the amide oxygen of A\(_{112}\) hydrogen bonds with the \( \alpha \)-NH of T\(_{115}\); the \( \alpha \)-NH of L\(_{111}\...
hydrogen bonds with the amide oxygen of A120; the amide oxygen of N-Me-A110 hydrogen bonds with the δ-NH of the ornithine that connects T106 and V121.

**Square channels.** Four extended β-sheets make the sides of a square channel (Fig. 5). Each side of the square channel is approximately 22 Å long, and the inside of the channel is hollow. The side chains of I107, N-Me-A109, L111, F116, and A120 point into the square channel, creating a hydrophobic interior. The side chains of T106, S112, S115, S117, and T119 point outward, making the outer surface of the square channel mostly polar. Fig. 5B shows a top view of the square channel and illustrates the hydrophobic interior and the polar outer surface.

Each square channel is stabilized by extended networks of hydrogen bonds between the four sides. The upper strand of each β-hairpin monomer in one side forms four hydrogen bonds with the lower strand of each β-hairpin monomer in the adjacent side (Fig. 5C and D). The amide oxygen of A108 hydrogen bonds with the χ-NH of the ornithine that connects S112 and S115; the χ-NH of I107 hydrogen bonds with the amide oxygen of S115; the amide oxygen of T106 hydrogen bonds with the χ-NH of F116; the χ-NH of the ornithine that connects T106 and V121 hydrogen bonds with the amide oxygen of S117.

These two networks of hydrogen bonds—those within the extended β-sheets (Fig. 4) and those between the extended β-sheets (Fig. 5C and D)—stabilize the square channels. All of the amide NH and carbonyl groups participate in hydrogen bonding, except the χ-NH of T118. To visualize how these two hydrogen-bonding networks act together to stabilize the square channels, one can imagine breaking all of the hydrogen bonds between two extended β-sheets at the corner of the square channel and unrolling and flattening the square channel (Fig. 6).
Tilted windows. The square channels further assemble in a “tilted windows” pattern (Fig. 7). The square channels do not align perfectly in the crystal lattice, but rather are tilted. Each channel makes limited hydrophobic contact with the neighboring channel. Although the outer surface is composed mostly of solvated polar side chains that are not in direct contact, hydrophobic contacts between V121 on adjacent square channels stabilize the tilted windows packing.

Discussion

In designing peptide 1 to mimic the TTR tetramerization interface, we anticipated that a tetramer would form (Fig. 8A). Instead of recapitulating the natural mode of assembly of TTR, peptide 1 forms a new mode of supramolecular assembly that has not been observed previously by X-ray crystallography—square channels. In the dimer subunit of the native TTR tetramer, strand H of one monomer pairs with strand H of another monomer. In the extended β-sheet formed by peptide 1, strand H pairs with strand G (Fig. 8B). The β-hairpins are shifted by two residues toward the N-terminus in the native TTR, while the β-hairpins in peptide 1 are shifted by four residues toward the C-terminus. The differences of the two assemblies may arise because other interactions, in addition to the β-sheet interactions, also stabilize the native TTR tetramer.

The square channels formed by peptide 1 resemble β-barrels, because both are tubular structures that are composed of networks of hydrogen-bonded β-strands. Eisenberg and coworkers found that an amyloidogenic peptide from αB crystallin forms a six-stranded β-barrel, which they termed a “cylindrin” (Fig. 9). The square channels formed by peptide 1 resemble the cylindrin in that the interior is hydrophobic. The
structures differ in that the square channels are hollow and run the length of the crystal, while the cylindrin forms a discrete hexamer with a tightly packed interior.

The interfaces between the square channels formed by peptide 1 resemble the layered β-sheet structures that compose amyloid fibrils. In amyloid fibrils, adjacent β-sheets form laminated layers in which the β-strands are nearly orthogonal to the fibril axis. The interfaces formed by the square channels in the “tilted windows” pattern are not as tightly laminated as many amyloid interfaces, such as the “steric zippers” discovered by Eisenberg and coworkers (Fig. 10). The β-strands of the square channels are not orthogonal to the channel axis, but rather are at about a 40° angle.

Developing new peptide supramolecular assemblies enhances the understanding of peptide and protein interactions and also facilitates the creation of new molecular devices and functional materials. The supramolecular assemblies formed by peptide 1 are particularly reminiscent of a few important peptide nanostructures formed by β-sheets. Ghadiri and coworkers designed cyclic peptides that form hollow nanotubes in the solid state and in lipid bilayer membranes. The peptides are composed of alternating P- and L-amino acids and assemble to form extended β-sheet-like structures. Lynn and coworkers found that peptide fragments from the central region of Aβ, Aβ16–22, assemble to form large nanotubes composed of β-sheets. Schneider and coworkers developed peptide hydrogels as drug delivery vehicles. The hydrogels are composed of β-hairpin peptides with alternating hydrophobic and polar residues.

Although we do not yet understand all of the rules governing the supramolecular assembly of β-sheets, it is clear that the hydrogen-bonding edges impart Lego-like assembly. What makes the square channels formed by macrocyclic β-sheet peptide 1 especially interesting is that we did not design the peptide to assemble in this fashion, but rather this mode of assembly emerged from the structure and sequence of the peptide. The incorporation of a single N-methyl group helps reduce the near-infinite ways in which a pre-organized β-sheet can assemble and allows a single mode of self-assembly to emerge. As we continue to study the supramolecular assembly of β-sheet peptides containing different amino acid sequences, we anticipate better understanding how the structure and sequence of a peptide dictates its mode of supramolecular assembly.

Conclusion
In our efforts to explore the supramolecular assembly of β-sheets, we have designed a macrocyclic peptide derived from TTR. Rather than recapitulating the assembly of the native TTR tetramer, the peptide formed square channels that comprise extended networks of hydrogen-bonded β-sheets. This unanticipated result is intriguing, because the assembly represents an emergent property of the peptide. We anticipate that this new assembly of β-sheets will serve as a piece of puzzle toward understanding the rules of β-sheet supramolecular assembly.

Acknowledgements
This work was supported by the National Science Foundation (NSF CHE-1507840).

Notes and references

Fig. 10 (A) Interface between square channels in the tilted windows assembly. (B) Steric zipper formed by a peptide from the yeast prion protein Sup35 (PDB 1YJP).
19 A related cyclic peptide derived from the dimerization interface of the TTR tetramer has been found to inhibit Aβ aggregation: P. Y. Cho, G. Joshi, M. D. Boersma, J. A. Johnson and R. M. Murphy, ACS Chem. Neurosci., 2015, 6, 778–789.
22 If this Orn turn is omitted, the resulting acyclic peptide (H-Thr-Ile-Ala-N-Me-Ala-Leu-Leu-Ser-5Orn-Ser-PheI-Ser-Thr-Ala-Val-OH) does not crystallize under the conditions that peptide 1 crystallizes. It also does not aggregate in thioflavin T fluorescence assays.
24 If the N-methyl group of peptide 1 is omitted, a gel forms when the cyclic peptide is deprotected. The placement of the N-methyl group on alanine 109 is important for crystallization; homologues with alternative placements of the N-methyl group do not crystallize under the conditions that peptide 1 crystallizes.
26 Thioflavin T fluorescence assays show that peptide 1 aggregates over the course of hours. Circular dichroism shows that the aggregates exhibit β-sheet structure. For details of these experiments, see the ESI†.