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Assessing Stabilization Through \( \pi-\pi \) Interactions in Aromatic Oligoamide \( \beta \)-Sheet Foldamers

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Supporting Information Placeholder

ABSTRACT: We have recently introduced aromatic oligoamide \( \beta \)-sheet foldamers based on rigid turn units and short linear strands that undergo intramolecular \( \pi-\pi \) stacking. We now report that conformational stability in these structures can be reached using less rigid turn units and more extensive \( \pi-\pi \) interactions between longer linear strands. For this study, two-stranded sheets of variable length were prepared. Their conformation was assessed in solution by \( ^1\)H NMR and in the solid state by x-ray crystallography.

In comparison with helical synthetic foldamers, sheet-like structures have much less frequently been observed. A possible reason for this may be that, when removed from the context of a tertiary structure as found in proteins, sheets tend to extensively aggregate and precipitate which makes them difficult to isolate and characterize. In contrast, helices generally are well-behaved discrete and soluble objects that fulfill their potential for non-covalent interactions intramolecularly. Indeed, synthetic foldamer sheets based on hydrogen-bonding are often designed to be dimeric in order to keep aggregation and solubility under control. Thus, the design of a soluble multistranded \( \beta \)-sheet foldamer would entail the orchestration of non covalent interactions so that they operate preferentially in an intramolecular fashion, thereby preventing extensive aggregation. Following this principle, we recently introduced aromatic oligoamide \( \beta \)-sheet foldamers stabilized by \( \pi-\pi \) interactions between short linear segments and rigid turn units such as 5 (Chart 1) that hold them in a face-to-face orientation. A key feature of 5 lies in its four methyl groups ortho to the aromatic amine functions holding the two xylol rings parallel to each other and perpendicular to the dinitrobenzene unit, as reflected by strong ring current effects which cause a large upfield shift of the NMR signal of the proton meta to both nitro groups pointing towards the interior of the structure (H\(_{\text{int}}\)). As an extension to this initial design, we hereby demonstrate that rigidity at the turn units can be loosened and compensated by enhanced \( \pi-\pi \) interactions between extended linear aromatic oligoamide segments, without causing aggregation.

Chart 1. Two stranded aromatic oligoamide \( \beta \)-sheets possessing linear segments of increasing length.
Trimeric turn 1 was first elongated using terephthalic acid monomers to form pentamer 2. In order to substantially enhance face-to-face intramolecular $\pi-\pi$ stacking, it was anticipated that larger units than simple benzene derivatives would be necessary. For this purpose, we introduced a new monomer, 6-amino-4-isobutoxy-7-methoxy-2-quinolinecarboxylic acid. As described in the supporting, its synthesis follows routes developed for other amino-2-quinolinecarboxylic acid monomers. The main feature of the new monomer is that the relative orientation of its 6-amine and 2-acid functions will confer a linear structure to oligoamide segments in which it is inserted. Elongation of pentamer 2 proceeded via the saponification of its terminal esters, and the subsequent coupling of two quinoline monomers or two quinoline dimers to yield 3 and 4, respectively.

The chemical shift values of the H$_{\text{int}}$ protons was monitored in CDCl$_3$ as a probe of the conformation of the new oligomers in this solvent (Fig. 1) as they were found to undergo significant variations while $\delta$$_{\text{int}}$ remained essentially unchanged. In rigid turn 5, this resonance is strongly upfield shifted ($\delta$$_{\text{int}}$ = 4.91 ppm), while the absence of methyl group in 1 makes its structure be more flexible and much reduce intramolecular ring current effects ($\delta$$_{\text{int}}$ = 6.68 ppm). Upon elongation of trimer 1 into pentamer 2 and heptamer 3, no major change is observed; H$_{\text{int}}$ resonances actually undergo minor downfield shifts. In contrast, a strong upfield shift is observed between heptamer 3 ($\delta$$_{\text{int}}$ = 6.08 ppm) and nonamer 4 ($\delta$$_{\text{int}}$ = 6.08 ppm, $\Delta$$\delta$ = 0.72 ppm). The effect on $\delta_{\text{int}}$ of the additional quinoline monomers at the end of the linear strands from 3 to 4 is very remote and suggests a better defined conformation of 4 leading to stronger ring current effects upon H$_{\text{ext}}$. The non linear trend of $\delta_{\text{int}}$ values from 1 to 4 is indicative of cooperative effects.

A series of multidimensional NMR experiments (HSQC, HMBC, TOCSY and ROESY) allowed us to fully assign the $^1$H NMR spectrum of 4. As shown in Fig. 2, ROESY experiments revealed contacts that unambiguously establish a folded two-stranded hairpin structure of 4 in solution. In particular interstrand (as opposed to intrastand) correlations are demonstrated when they occur between protons that are too distant to establish a contact if they would belong to the same aromatic ring such as, for example, 7-methoxy protons and the proton in position 5 of the quinoline rings (OMe/H$_3$ and OMe/H$_{10}$ correlations in Fig. 2). In contrast, no such correlations were observed in the spectra of heptamer 3. Thus, the greater flexibility of turn 1 as compared to turn 5 can be compensated by extensive intramolecular $\pi-\pi$ interactions between long linear strands. All observed correlations, and in particular OMe/H$_2$ and OMe/H$_{10}$ (Fig. 2), are consistent with an antiparallel orientation of the two strands of 4, with each ring head-to-tail with respect to the ring on which its stacks, as observed previously with multistranded sheets having shorter linear segments. Yet the existence of conformers with a parallel orientation of the two strands of 4 cannot be completely ruled out from solution studies. Indeed, the interstrand NOE correlations expected in a parallel arrangement may not be distinguished from intrastand correlations, unless the two strands are strongly offset in which case interstrand correlations may match with those expected in an antiparallel arrangement of the two strands. Spectra recorded over a wide temperature range (-50°C to 40°C) did not reveal any major change (Figure S2), suggesting that the same structure prevails over this range. Even at low temperature, the diastereotopicity of side chain isobutoxy protons does not appear in the spectrum, indicating fast rotational dynamics of each linear strand with respect to the other. As a last point worth noting, it is unclear whether the H$_{\text{ext}}$ resonance in the folded conformation of 4 remains at lower field than in 5 because the latter is still better organized, because the former adopts a conformation at the turn in which ring current effects are weaker, or because ring currents are simply less intense in the former due to the different substitution patterns of its $\text{para}$-phenylenediamine rings.

![Figure 1. Part of the 300 MHz $^1$H NMR spectra in CDCl$_3$ at 298K of: a) hairpin turn trimer 1; b) hairpin turn pentamer 2; c) hairpin turn heptadecamer 3; d) hairpin turn nonamer 4; and e) hairpin turn trimer 5. Signals of H$_{\text{ext}}$ are marked with white circles. Signals of H$_{\text{ext}}$ are marked with full black circles. Stars indicate signals belonging to an impurity.](image)

![Figure 2. Schematic representation of $^1$H-$^1$H correlations observed in 400 MHz ROESY-2D spectra of nonamer 4 at 25 °C in CDCl$_3$. Blue arrows represent correlations within the hairpin turn; pink arrows represent interstrand correlations; other correlations are represented by grey arrows.](image)
Å) is too large for tight aromatic stacking to occur in short sequences such as 1 and 2. Only the long strands of 4 compensate for this impediment, allowing its quinoline rings to stack at a distance of 3.4 Å.

The packing of 4 in the crystal shows extended head-to-tail stacks suggestive of a multi-stranded sheet-like aggregation mode (Fig. 3e). Nevertheless, this organization is restricted to the solid state, and compound 4 shows good solubility (> 13 mM in CDCl₃) and concentration independant chemical shift values up to 13 mM indicating limited aggregation in solution. Thus, the face-to-face π-π interactions that stabilize the two-stranded hairpin-turn structure of 4 are strong enough to be effective intramolecularly and weak enough not to prevail intermolecularly.

In summary, we have shown that π-π stacking may direct the folding of extended two-stranded β-hairpin structures in an organic solvent even when a relatively flexible turn unit is used. The balance between intra- and intermolecular interactions allows folding to occur and aggregation to be prevented. These objects are reminiscent or related foldamer structures including some crescent-like macrocycles, zippers based on oligoanthranilamides, as well as dimeric, trimeric and tetrameric aromatic oligoamide β-helices. It is anticipated that folding and aggregation of such β-hairpins will operate differently in protic media due to the very strong solvophobic component that makes π-π stacking be much stronger in these solvents. Research along this line is in progress and will be reported in due course.

ASSOCIATED CONTENT
Supporting Information
Synthetic schemes, experimental procedures, full characterization of new compounds, crystallographic data, detailed NMR investigations including complete ¹H NMR assignment and solution structure elucidation of 4. This material is available free of charge via the Internet at http://pubs.acs.org.”

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Author Contributions
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Notes
The authors declare no competing financial interest.

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