Self-Assembled Protein–Aromatic Foldamer Complexes with 2:3 and 2:2:1 Stoichiometries

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ABSTRACT: The promotion of protein dimerization using the aggregation properties of a protein ligand was explored and shown to produce complexes with unusual stoichiometries. Helical foldamer 2 was synthesized and bound to human carbonic anhydrase (HCA) using a nanomolar active site ligand. Crystal structures show that the hydrophobicity of 2 and interactions of its side chains lead to the formation of an HCA-2-2 complex in which three helices of 2 are stacked, two of them being linked to an HCA molecule. The middle foldamer in the stack can be replaced by alternate sequences 3 or 5. Solution studies by CD and NMR confirm left-handedness of the helical foldamers as well as HCA dimerization.

T he recognition of protein surface features by smaller molecules is a means to mask and inhibit interactions in which proteins may engage, in particular with other proteins. A possible effect of a surface-binding ligand is thus a disruption of protein assemblies, an outcome that can serve therapeutic or pharmacological purposes. Conversely, inducing homo- or heteromeric protein assembly can be useful when activity results from the association of several proteins. This is generally achieved by presenting multitopic protein surface ligands that can simultaneously bind two identical or two different proteins, and eventually enhance activity. Multitopic ligands can also serve in the fabrication of ordered protein arrays. An alternate, essentially unexplored approach, to induce protein assembly into discrete objects would be to use monotropic ligands that themselves possess self-assembly properties. Along this line, we have recently reported the characterization of a 2+2 complex between two human carbonic anhydrase (HCA) molecules and two molecules of helical aromatic foldamer 1 (Figure 1). This complex is stabilized by protein–foldamer, foldamer–foldamer, and protein–protein interactions. We now have extended this exploratory work and report the intriguing formation of HCA–aromatic foldamer complexes comprised of two proteins and either three identical foldamers (2:3 stoichiometry), or two identical and a third, different, foldamer (2:2:1 stoichiometry).

Several groups have been investigating aromatic foldamers as rigid helical or linear scaffolds that can be equipped with proteinogenic side chains to recognize protein, nucleic acid, or saccharide surfaces. For example, 1 is a stable helical oligoamide of 8-amino-2-quinolinecarboxylic acid. It was identified and characterized while studying interactions between HCA and related foldamer sequences bearing varied side chains protruding from the helix in position 4 of each quinoline unit. These foldamers possess an appended arylsulfonamide HCA nanomolar ligand of the enzyme active site that confines them at the protein surface. Specific interactions with the protein may then be detected by the induction of a handedness bias in the achiral helix backbone that otherwise exists as a racemic mixture of interconverting right-handed (P) and left-handed (M) enantiomers. In the case of HCA-1, circular dichroism (CD) at pH 7.4 in a phosphate buffer shows a positive band in the quinoline absorption region characteristic of P helicity (Figure 1c). This observation triggered structural investigations both in the solid state (Figure 1b) and in solution that eventually showed that HCA-1 is not a 1:1 but an unprecedented 2:2 complex in which the hydrophobic nature of the aromatic helix cross section promotes foldamer–foldamer interactions that ultimately result in dimerization. Among over 400 HCA structures found in the protein databank (PDB), the packing and intercomplex protein–protein contacts of the (HCA-1)_2 structure are unique. Compound 2 was prepared (see SI) using solid phase synthesis within a series of variants of 1 to test the scope of the formation of (HCA-1)_2. In the sequence of 2, two monomers of 1 have been swapped. HCA-2 also shows a positive CD band indicative of P helicity at pH 7.4 in phosphate buffer (Figure 1c). However, the HCA-2 complex failed to crystallize from the Zn-rich medium in which crystals of (HCA-1)_2 grow. Other crystallization conditions were eventually found using commercial sparse-matrix screens and a structure of HCA-2 was resolved at 1.8 Å resolution and revealed several distinct features (Figure 2, Figure S7).

The structure shows a higher aggregate containing two HCA molecules and not two but three molecules of 2 arranged in a
stack, again mediated by contacts between the aromatic helices’ cross sections. The two peripheral helices of the stack have their ligand bound to an HCA active site while the ligand of the central helix is not bound and actually not visible in the electron density map due to disordered positions. The complex has an overall noncrystallographic pseudo-C$_2$ symmetry.

Consequently, the central foldamer is in fact present in two degenerate orientations within the stack and was modeled as such. Other remarkable differences exist between the (HCA-1)$_2$ and HCA$_2$-2$_3$ structures: (i) the axis of the stack of helices is parallel to the proteins’ surfaces in the latter, whereas it is perpendicular in the former. The foldamer side chains thus point directly toward the protein surfaces in HCA$_2$-2$_3$ and establish remarkable contacts (Figure S7, see also the similar HCA$_2$-2$_2$-3 complex in Figure 3); (ii) these contacts include interactions with N-terminal His3 which (along with Ser2) is visible in the electron density map whereas it is usually not seen in HCA crystal structures; (iii) the relative orientation of the two HCA molecules is completely different; (iv) foldamers are P-helical in (HCA-1)$_2$ and M-helical in HCA$_2$-2$_3$. This latter feature first appeared to contradict the sign of the CD band observed in phosphate buffer. However, it was then realized that the crystallization medium was in fact quite different (see SI). A few controls established that M-helicity is favored not by salt or the nature of the buffer but by a lower pH and that it can indeed be observed in solution (Figure 1c). $M$ and $P$ helicities can be induced alternatively by changing pH (Figure S16). In contrast, the CD band of the (HCA-1)$_2$ complex is still positive at pH 4.6. The low pH must not only disfavor interactions with the P helix as this would result in no CD signal at all but also favor interactions with the M helix. In the HCA$_2$-2$_3$-3 complex, a salt bridge is indeed observed between the terminal foldamer

Figure 1. Foldamer–HCA interactions. (a) Formulas of HCA ligand 4 and foldamers 1, 2, 3 and 5. (b) Crystal structure (left) of the (HCA-1)$_2$ dimeric complex previously reported$^6$ and schematic view (right) of the foldamer–protein interface in that complex. Proteins are shown as ribbons (left) or as solvent-accessible surfaces (right). Foldamers are shown in the same color (yellow or green) as the color of the protein to which their ligand moiety is bound. Zinc ions at the bottom of the ligand binding pocket and at the foldamer–protein interface are shown as purple spheres. (c) CD spectra of HCA-1 and HCA-2 complexes in different buffers.

Figure 2. Crystal structure of the HCA$_2$-2$_3$ complex (PDB# 5L3O). (a) Asymmetric unit showing two HCA molecules in green and yellow and three foldamer helices in CPK representation. The green and yellow foldamers have their ligand bound to an HCA active site. Native Zn$^{2+}$ ions are shown as red spheres. A third foldamer (purple and magenta for the two orientations) is sandwiched between the first two. (b) View down the foldamers’ helix axis of the complex showed in panel a.
carboxylate function and His3. Whether this interaction is critical to M helicity remains to be demonstrated. We sought to confirm the assembly features of the HCA-2 complex in solution by using NMR spectroscopy as established for HCA-1 (Figure 4, Figures S13−15). At pH = 4.6, 15N HSQC spectra of [15N]HCA and [15N]HCA-4 demonstrated that the protein is stable, well-folded and fully bound by 4 (Figure 4a, note the shift of the Trp208 amide cross-peak located near to the sulfomanide moiety). Intermolecular contacts were then identified by comparing 1H−15N HSQC spectra of 2 or 4 bound to [15N]HCA (Figure 4a,c). Compound 4 lacks a foldamer helix and chemical shift perturbations (CSPs) in the spectrum of HCA-2 as compared to HCA-4 can thus be attributed mainly to foldamer−protein contacts. Quite remarkably, CSPs were principally located at residues involved in protein−foldamer and protein−protein contacts of the HCA-2 crystal structure (Figure 4e). Size estimate of the HCA-2 complex by measuring 1HN T2 of a 500 μM sample revealed a correlation time (τc) of 31.3 ns (Table S2), consistent with a dimeric state (two proteins) but the possible presence of a third foldamer could not be established. At 100 μM, dissociation results in a significantly reduced correlation time of 23.2 ns. In the alternate phosphate condition at pH 7.4, 1H−15N HSQC spectra again confirm protein folding and complex formation (Figure 4b). In contrast to pH 4.6, the comparison of HCA-4 and HCA-2 spectra at pH 7.4 show limited CSPs suggesting a weaker protein−foldamer interaction. Size estimates by 1H T2 indicate a mainly monomeric state (Table S2) in which helix handedness induction nevertheless takes place (Figure 1c) as for HCA-1.

Because the middle foldamer in the stack of helices does not need to possess an HCA ligand and does not engage in the same interactions with HCA as the peripheral helices, we explored the possibility to replace it by other foldamer sequences. Compound 3 is an analogue of 2 that lacks the HCA ligand. When HCA was set to crystallize in the presence of 2 (1 equiv) and 3 (0.5 equiv) a structure similar to that of HCA-2 was obtained (Figure 3, Figure S9), yet with a better resolution (1.6 Å). LC-MS analysis after dissolving the crystals ascertained the presence of 3 in the structure, presumably in the middle of the helical stacks (Figure S12). However, given the similarity between 2 and 3, and the absence of their flexible chains in the electron density maps, they cannot be distinguished in the structure. Next, compound 5 was designed which possesses a C2 symmetrical helical structure and contains twice the two N-terminal residues of 2 and 3. HCA crystals were grown in the presence of 2 (1 equiv) and 5 (0.5 equiv) and a structure at 1.4 Å was obtained (Figure S10, PDB #5L6K).
SLVS). The structure unambiguously shows the presence of S in the middle of the helical stack. Because of its symmetry, its two degenerate positions are identical. The electron density map indicates that a small amount of I occupies the middle position but this could not be modeled accurately (Figure S11). Overall, although the contribution of the middle foldamer to the overall stability of the multimeric assembly is not established, the fact that it can be substituted ascertain the existence of a driving force for that site not to remain unoccupied.

In summary, we have established that linking helical aromatic foldamers to a protein surface leads to the formation of protein complexes with intriguing stoichiometries. Simply swapping two foldamer residues causes subtle changes in foldamer–protein surface interactions, resulting in a major change in assembly behavior at different pH values. The stacking of hydrophobic helical cross sections constitutes a recurrent pattern and presumably a driving force that adds to foldamer–protein and protein–protein interactions. An alternate way to control protein association may be to exploit deliberately the self-assembly properties of aromatic amide helices that show a propensity to form not stacks, but multistranded triple or double helices, including in water.14

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/jacs.7b00184.

Experimental details for synthetic procedure, spectroscopic and crystallographic data (PDF)

Model for SL3O (PDB)

Model for SL6k (PDB)

Model for SLVS (PDB)

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Notes

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