Foldaxanes: Rotaxane-like Architectures from Foldamers

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CONSPECTUS



Mechanically interlocked molecules such as rotaxanes and catenanes contain free-moving components that cannot dissociate and have enabled the investigation and control of various translational and rotational molecular motions. The architecture of pseudo-rotaxanes and of some kinetically labile rotaxanes is comparable to that of rotaxanes but their components are reversibly associated and not irreversibly interlocked. In other words, pseudo-rotaxanes may fall apart. This account focuses on a peculiar family of rotaxane-like architectures termed foldaxanes. Foldaxanes consist of a helically folded oligomer wound around a rod-like dumbbell-shaped guest. Winding of the helix around the rod thus entails an unwinding-rewinding process that creates a kinetic barrier. It follows that foldaxanes, albeit reversibly assembled, have significant life-times and may not fall apart while defined molecular motions are triggered. Foldaxanes based on helically folded aromatic oligoamide hosts and oligo alkyl carbamate guests can be designed rationally through the inclusion of complementary binding motifs on the rod and at the inner rim of the helix so that helix length and rod length match. Single helical foldaxanes (bimolecular species) and double helical foldaxanes (trimolecular species) have thus been produced as well as poly-[n]-foldaxanes, in which several helices bind to long rods with multiple binding stations. When the binding stations differ and are organized in a certain sequence, a complementary sequence of different stacked helices, each matching with their binding station, can be assembled, thus reproducing in an artificial system a sort of translation process. Foldaxane helix handedness may be controlled by stereogenic centers on the rod-like guest. Handedness can also be transmitted from helix to helix in poly-foldaxanes. Foldaxane formation has drastic consequences on the rod properties, including its stiffening and the restriction of the mobility of a macrocycle already interlocked on the rod. The fast translation - without dissociation - of helices along rod-like guests has been demonstrated. Because of the helical nature of the hosts, translation may be accompanied by rotation in various sorts of screwlike motions. The possibility, on longer timescales, for the helix to dissociate from and re-associate to the rod has allowed for the design of complex, kinetically controlled, supramolecular pathways of a helix on a rod. Furthermore, the design of helices with a directionality, *i.e.* with two distinct termini, that bind to non-symmetrical rod-like guests in a defined orientation makes it possible to also control the orientation of molecular motion. Altogether, foldaxanes constitute a distinct and

full-of-potential family of rotaxane-like architectures that possess designer structures and allow to orchestrate the time scales of various supramolecular events.

KEY REFERENCES

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1. INTRODUCTION

Mechanically interlocked molecules have opened up considerable capabilities in molecular machinery.^{4,5,6,7} For example, rotaxanes (Figure 1A) have emerged as an archetypical architecture enabling the controlled motion of rings mechanically bound to dumbbell-shaped molecular rods.^{8,9,10} The moving parts of a rotaxane, the ring and the rod, cannot dissociate and various strategies have been devised to produce such objects.¹¹ In contrast, pseudo-rotaxanes allow for the dissociation of their subcomponents because the rod is not dumbbell-shaped, thus allowing ring unslippage (Figure 1B).¹² Kinetically labile rotaxanes have also been described (Figure 1C).^{13,14,15} Pseudo-rotaxanes possess the advantage that they can be produced through self-assembly by simply mixing their components. Their existence rests on the ring affinity for the rod. However, they can dissociate which makes their use in molecular machinery complicated: a machine may preferably not be allowed to fall apart to achieve work.

This account focuses on foldaxanes, a peculiar family of rotaxane-like architectures¹⁶ in which the ring is replaced by a helically folded oligomer, *i.e.* a foldamer,^{17,18,19,20} (Figure 1D). A one-turn helix may be assimilated to a ring closed by non-covalent interactions. It is a dynamic object but its time constants, *e.g.* the rates at which it winds and unwinds, may substantially vary depending on its inherent rigidity and upon adding helix turns. Using stable helices, long-lived foldaxanes may be produced, that is, helix-rod complexes that do not dissociate while some molecular motion takes place. Designing foldaxane structural characteristics thus enables the orchestration of the time scales of the various processes involved and create new opportunities for controlling molecular motions.



Figure 1. (A) Irreversible formation of a [2]rotaxane and (B-D) reversible formation of various types of architectures: pseudo-rotaxanes (B), kinetically labile rotaxanes (C), foldaxanes (D) and pseudo-foldaxanes (E).

The development of foldaxanes stemmed from the use of helical foldamers with sizeable cavities as containers for molecular recognition.^{21,22,23,24,25,26,27,28} While this background mostly concerns the recognition of guests small enough to entirely fit in a helix cavity, an increasing number of reports target rod-like guests that protrude from the helix, thus defining a foldaxane or pseudo-foldaxane architecture (Figure 1,D,E). With few exceptions,^{29,30} foldaxanes involve relatively rigid aromatic helices. Following early reports based on oligo-*meta*-phenyleneethynylenes^{31,32,33} and on naphthyridine oligomers,³⁴ a range of foldaxane has been

reported that exploit the modularity and synthetic access of aromatic oligoamide foldamers.^{21,35,36,37} Depending on the properties of the foldamer host and on the rod-like guest, single or double helical foldaxanes may be produced. Their design principles and their intriguing properties are presented in detail in the following.

2. SINGLE HELICAL FOLDAXANE DESIGN

Aromatic oligoamide foldamers adopt stable conformations such as helices or sheets that can be simply predicted because they constitute the direct outcome of local preferences at each rotatable bond that define the relative orientation of consecutive units.³⁵⁻³⁷ Over the years, a toolbox of diamine, diacid and amino acid heterocyclic aromatic monomers that differ by their size and the position of their amine and acid substituents have been developed. A sampling of these monomers is shown in Figure 2A, and typical interactions that govern backbone conformation are shown in Figure 2B. Depending on the monomer used, different strand curvatures may be imparted that will give rise to helical structures of varying diameter. For example, Q_n sequences (Figure 2B) form highly curved helices with no cavity. In contrast, Q^F_n sequences (Figure 2C) form less curved helices and possess a small cavity in which an alkyl chain may fit. Upon combining monomers that code for a wide helix diameter in the center of a sequence and short Q_n segments at both ends, a helical capsule can be produced that can completely seclude a guest molecule from the surrounding medium (Figure 2D).²¹ Helical containers based on these principles have been shown to bind a broad range of guests, from anions,³⁸ cations³⁹ to organic acids^{24,40} and carbohydrates.^{41,42}



Figure 2. (A) Toolbox of heterocyclic amino acid, diacid or diamine aromatic building blocks of helical aromatic oligoamide foldamers, and their letter and color code. Folding principles of (B) Q_2 and (C) Q^{F_2} . Aryl-amide pseudo-conjugation, hydrogen bonds (dashed lines) and electrostatic repulsions (arrows) contribute to the stabilization of a preferred conformation at each aryl-amide bond, giving rise to a helix in sufficiently long sequences. Intramolecular aromatic stacking comes as an additional, solvent dependent, stabilizing force. (D) Encapsulation of a guest molecule (yellow sphere) in a molecular helical capsule (blue tube) possessing a reduced diameter at both ends, leading to a host-guest complex *via* a transient local unfolding of the helix.

It was devised that by removing the end-caps from a capsule sequence, *i.e.* the highly curved extremities, guest molecules may be allowed to protrude from the helix to produce foldaxanes and pseudo-foldaxanes. Capsule sequence 1^{43} (Figure 3A) was selected for this purpose. It comprises two P^NP^CP^N polar clefts that can hydrogen bond to a guest's polar function (Figure 3B), a central Q^F₂A^FQ^F₂ cylindrical cavity to accommodate a short alkyl chain, and terminal Q₃ caps that close the cavity. Sequence 1 was rationally designed to bind to guest molecules such as 1,4-butanediol or 4-amino-1-butanol and successfully does so (Figure 3E) with a *K_a* of *ca*. 10³ L.mol⁻¹ in chloroform.⁴³ Using 1 as a starting point, the two Q₃ caps were deleted to let longer guests protrude from the cavity and allow for the binding of dumbbell-shaped molecules. Accordingly, sequences

2a-2d are composed of a central cylindrical cavity $Q^{F}_{m}A^{F}Q^{F}_{m}$ of variable length, and all possess two $P^{N}P^{C}P^{N}$ hydrogen bond donor clamps. Gratifyingly, these foldamers were shown to bind α, ω alkanediamine-derived dicarbamate guests, thus producing an ensemble of helix-rod complexes.^{1,2} Thin guests such as **5** can thread themselves into the cavity (Figure 3B,F). Complexes also form with dumbbell-shaped guests such as **3** and **4** (Figure 3F) which require another mechanism of formation. As depicted in Figure 1D, it is inferred that the helix must wind around the rod. However, aromatic oligoamide helices are particularly stable and may not completely unfold for this purpose.⁴⁴ A sort of screwing around the rod while the helix keeps its shape may be envisaged. Thus, bulky stoppers do not alter the binding energy (see for example $2c \supset 4b$ vs. $2c \supset 5$ in Figure 3D). However, bulky stoppers alter the binding and release mechanisms and thus the associated kinetics. Threading typically takes place fast and cannot be monitored by NMR, whereas winding may take days to reach equilibrium.^{1,3}



Figure 3. (A) Sequences **1** and **2a-2d**. (B) Formulae of rod-like guests possessing two carbamate functional groups (orange) with bulky end groups (**3a-3j**, **4a-4b**) or without (**5**). (C) Equilibria between a dumbbell-shaped rod (R), a single (SH) or double (DH) helical foldamer and a foldaxane (FD). (D) K_a values in CDCl₃ at 318 K of various foldaxanes and one pseudo-foldaxane, as determined by ¹H NMR spectroscopy. Crystal structures of complexes (E) **1**⊃4-amino-1-butanol and (F) **2b**⊃**4a**, **2c**⊃**4b** and **2c**⊃**5**, from left to right. ^aFor a homogeneous presentation, association constants (K_a) for **2d** have been recalculated at equilibrium using the equation proposed in C.

The formation constants of foldaxanes made of different helix and rod lengths revealed two main trends (Figure 3D). First, a rather strict match between helix length and guest length is required for optimal binding. Removing or adding a single methylene unit within the rod may have drastic consequences on foldaxane stability. Second, foldaxane stability increases for longer helices and rods, which culminated with a $K_a = 345\ 000\ \text{L.mol}^{-1}$ for $2d\supset 3f$. This suggests that foldaxane stability is not solely determined by the hydrogen bonds between the carbamate and the $P^NP^CP^N$ polar clefts but that contacts between the alkyl chains and the fluorine atoms of Q^F and A^F units are also favorable.

Another aspect is that helices **2a-2d** bind selectively to dicarbamates of α , ω -alkanediamines. The P^{NPC}P^N polar clefts hydrogen bond to the carbamate carbonyl group and binding affinities collapse if the carbamate orientation is inverted by swapping amino and oxy functions, or if it is replaced by an amide or an ester. This selectivity offered the possibility to fine tune the system. Thus, the P^{NPC}N polar cleft where the terminal pyridine was replaced by a naphthyridine was designed to selectively recognize an amide instead of a carbamate (Figure 4).⁴⁵ The outcome is that dissymmetrical sequence **6**, which possesses both P^{NPCPN} and P^{NPCN} segments selectively binds to dissymmetrical guests like **7c** with a K_a of 28 000 L.mol⁻¹. Complexes with the amide and carbamate mismatching the two polar clefts are transiently observed before disappearing below detection levels of ¹H NMR. Such complexes make it possible to control the relative orientation of the helix and the rod, a property that has been achieved with few rotaxanes.^{46,47,48,49}



Figure 4. (A) Sequence of dissymmetrical helical foldamer 6 and formulae of dissymmetrical guests **7a-7g**. (B) Crystal structure of $6 \supset 7c$ with zooms on the binding modes between either the P^NP^CP^N cleft to the carbamate of the guest (left) or P^NP^CN to the amide (right).

3. FOLDAXANES AND DOUBLE HELICES

The key modification to transform a capsule sequence such as 1 (Figure 3A,E) into a helix prone to foldaxane formation is the removal of the Q₃ segments that cap the helix cavity. Yet, this change has another consequence than forming an open-ended helix. Helices with a wide diameter have a strong propensity to self-assemble into double helical dimers because of the lower energy cost of spring-like extension when helix diameter is larger – as for macroscopic metal springs.⁵⁰ Q₃ segments are more curved and disfavor this process (whereas Q₂ may accommodate it).⁵¹ It follows that foldaxane assembly competes with double helix formation (Figure 3C). Nevertheless, despite the generally high stability of these double helices, this competition does not overwhelm foldaxane formation. The reason is that one double helix can dissociate to form two foldaxanes. The overall equilibrium is thus defined by the ratio between the constant of dimerization and the square of the constant of formation of the foldaxane from a single helix. Nevertheless, double helix formation complicates measurements. Because the single helix is involved in two equilibria, it is undetectable in NMR spectra at equilibrium. This makes it more difficult to directly measure the constant of formation of the foldaxane from the single helix. Instead, we calculate the constant of formation of two foldaxanes from the double helix (Figure 3C). Furthermore, the kinetics of double helix formation are slow, particularly so for long oligomers, and it may take days to bring foldaxane formation to equilibrium when mixing a rod guest and a double helix. Fortunately, a majority of the aromatic oligoamide strands can be precipitated in their out-of-equilibrium single helical form from methanol. Producing foldaxanes from the single helix is much faster and equilibrium is reached rapidly.

Double helix formation can thus be a practical impediment, but it can also be turned to an advantage. Just like double helical capsules have been produced,⁵² we endeavored to design double helical foldaxanes. Sequences **8a-8c**, composed of a Q^F_n segment and a single P^NP^CP^N segment were designed for this purpose (Figure 5A).^{2,53,542} They assemble into double helical dimers in solution which may be parallel or anti-parallel. The latter configuration, in which one P^NP^CP^N is found at each extremity of the duplex, was shown to produce double helical foldaxanes similar to the single helical foldaxanes presented above. The crystal structures of (**8a**)₂ \supset **3b**, (**8a**)₂ \supset **3d** and (**8a**)₂ \supset **3e** (Figure 5D) confirmed the stoichiometry, symmetry and structure of these double-helical foldaxanes, in agreement with NMR data.⁵³ Again, a complex structural pathway must be invoked for the winding of a double helix around a dumbbell rod.



Figure 5. (A) Sequences of double-helical foldamers $(8a) -(8c)_2$. (B) Formulae of dicarbamate guests. Guest 9b is terminated by pyrene and Ru(bpy)₃²⁺ chromophores (hexafluorophosphate counter anions are omitted for clarity). Guest 9a is its precursor. Guests 10a-10b possess only one bulky end group and thread themselves in double helix (8c)₂. (C) Formation of a [3]foldaxane from a double helix (green) wrapped around anchor points (orange) of a dumbbell-shaped rod. Crystal structures and association constants in CDCl₃ of foldaxanes (D) (8a)₂ \supset 3b, (8a)₂ \supset 3d and (8a)₂ \supset 3e (left to right), (E) (8b)₂ \supset 9a and (F) (8c)₂ \supset 10a and (8c)₂ \supset 10b (left to right). Doubles helices (8a)₂, (8b)₂ and (8c)₂ are shown in red/black, green/black and blue/black, respectively.

An interesting feature of these double helical foldaxanes is the lower dependence of their stability on the length of the rod than for single helical foldaxanes. The crystal structures of Figure 5D involve the same double helix $(8a)_2$ and three different rods and revealed that the two strands of the double helix may screw into one another to a variable extent to adjust their conformation to

the guest. The structures of $(8c)_2 \supset 10a$ and $(8c)_2 \supset 10b$ (Figure 5F) also show an adjustment of the double-helix to the length of the rod *via* screwing/unscrewing motions.³ It follows that the transfer of a double helix from a guest to another guest of different length involves a screwing or unscrewing of the duplex (Figure 5C). This notion has relevance to controlled molecular motion because it combines a translation and a rotation, two motions that are generally dissociated in molecular systems.^{55,56,57,58}

4. SHUTTLING MOTIONS BASED ON FOLDAXANE ARCHITECTURES

The fact that helix winding around dumbbell rods to form foldaxanes takes place slowly, *i.e.* on time scales ranging from minutes to days depending on their length, enabled their use in a manner similar to that of macrocycles in rotaxane-based molecular shuttles. Dumbbell-shaped rod **11a** was conceived for a proof-of-principle study (Figure 6A).¹ It contains two carbamate functions and one urea, that is, three hydrogen bond acceptors. Thus, **11a** possesses two binding stations but only one may be occupied at a given time. Solution and solid state data confirmed the exclusive formation of 1:1 single helical foldaxane **2c** \supset **11a**.¹ Using exchange NMR spectroscopy, it could be demonstrated that single helix **2c** transfers from one station to the other and that this occurs at rates (~3 min⁻¹) incompatible with the time necessary for unwinding (2 day⁻¹) and rewinding the helix.¹ Thus, actual sliding takes place without dissociation. Molecular dynamics simulation provided insights about energetic barriers and structural changes during the sliding process: on top of translation and rotations, the foldamer helix was also found to undergo a series of swelling and shrinking.⁵⁹



Figure 6. (A) Formulae of rods **11a-11b** and **12a-12b**. (B) Equilibria that govern the pH-controlled shuttling of single helix **2c** (blue) along rod **11b**. (C) Unidirectional fast shuttling (with a non-bulky spacer) or slow unwinding/rewinding (with a bulky spacer) of double helix (**8c**)₂ (green) from the α -station to the ω -station of rods such as **12a** (tetraethylene glycol spacer) or **12b** (*p*-xylyl spacer).

In the case of rod **11a**, the helix may move from one binding station to the other with equal probability. A step beyond consists in controlling this motion by triggering repulsive interactions for one station. We used dissymmetrical rod **11b** (Figure 6A) which possesses one carbamate/urea station with a heptyl segment (noted α) and the other with a diethylamine segment (noted ω). Single foldaxane **2c** \supset **11b** was produced and its NMR showed two sets of signals with slightly different proportions (58:42) meaning that single helix **2c** spends a bit more time on one station than on the other. Upon titrating **2c** \supset **11b** with an acid to protonate the amine-containing station, one set of signals disappeared, and only one remained. This reflects the fact that helix **2c** is trapped around the α station and repelled from the ω station because the helix cavity does not accommodate

the ammonium function (Figure 6B). Adding a base reverts the process, eventually producing a foldaxane-based pH-controlled molecular shuttle.

In a subsequent step, helix motions around rods was organized into kinetically controlled multi-step pathways. We designed rod 12a (Figure 6A) possessing successively an α -binding station known to have a weak affinity for double helix $(8c)_2$, a non-bulky spacer, an ω -binding station known to have a strong affinity for $(8c)_2$ and a bulky stopper.³ Rod 12b only differs from 12a in that it has a bulky p-xylyl spacer. Upon adding an excess of 12a or 12b to a solution of (8c)₂, NMR showed that a single new species was immediately and quantitatively formed that corresponds to the fast and quantitative threading of the double helix (8c)₂ around the α -station of the rod, leading to kinetic supramolecular products $(8c)_2 \supset 12a \cdot \alpha$ and $(8c)_2 \supset 12b \cdot \alpha$, respectively (Figure 6C). Over time, the emergence of a second set of resonances was observed by NMR and attributed to the thermodynamic products $(8c)_2 \supset 12a - \omega$ and $(8c)_2 \supset 12b - \omega$. Foldaxane $(8c)_2 \supset 12a - \omega$ ω was produced within ~3 hours which indicates a shuttling motion. The tetraethylene glycol spacers does not hamper the passage of the helix. In contrast, $(8c)_2 \supset 12b - \omega$ took about ~15 days to form, indicating that the bulky spacer of 12b imposes a dissociation of $(8c)_2 \supset 12a - \alpha$ and the rewinding of the helix around the ω -station (Figure 6C).³ Similar pathways can also be implemented with a control over the orientation of the rod and helix, using the building blocks shown in Figure 4.45

5. POLY-FOLDAXANES AND HELIX HANDEDNESS CONTROL

The high stability of single and double helical foldaxanes and the match between helix and rod length invited to challenge the formation of poly-foldaxanes on rods that would possess multiple binding stations. Such assemblies might for example enable to organize in space various functional groups attached to each helical component. Double helix $(8b)_2$ binds to rod 13a which possesses a single dicarbamate binding station (Figure 7A, $K_a = 1$ 700 L.mol⁻¹ in CDCl₃).² It also binds to longer dumbbell-shaped rods 13b to 13d (Figure 7A), possessing respectively two, three and five binding stations identical to that of 13a. This eventually produces poly-foldaxanes $(8b)_4 \supset 13b$, $(8b)_6 \supset 13c$ and $(8b)_{10} \supset 13d$ (Figure 7C), the latter having a length of 9.1 nm and a mass of 22 kDa.² Neither the host nor the guest possess stereogenic centers. Thus, (8b)₂ exists as a racemate of right- (P) and left-handed (M) enantiomeric conformers. Upon assembling multiple helices on a rod, diastereomeric complexes may result. For example, $(8b)_4 \supset 13b$ may have PP, *MM* or *PM* (*meso*) stereochemistry. However, ¹H NMR and crystallography demonstrate that the poly-foldaxane are all homochiral. Only racemates of $(P-\mathbf{8b})_{2n} \supset \mathbf{13}$ and $(M-\mathbf{8b})_{2n} \supset \mathbf{13}$ are produced, indicating quantitative helix-helix handedness communication between the helices bound to the same rod. The different dicarbamate stations of 13b-13d are separated by an ethylene glycol spacer which plays an important role in poly-foldaxane design. Indeed, if the spacer between two binding stations is too short, steric hindrance between adjacent double helices may prevent all stations from being occupied. Conversely, if the spacer is too long, an absence of contacts between adjacent double helices would hamper helix-helix handedness communication.

In addition to relative helix-helix handedness control, absolute handedness control was achieved through the incorporation of a chiral group at both ends of a binding station. For example, rods 14a-(*S*,*S*) and 14b-(*S*,*S*) possess two (*S*) phenethyl groups (Figure 7A). Circular dichroism and ¹H NMR showed that the single helix 2c binds to 14a-(*S*,*S*) to give [2]foldaxane *P*-2c \supset 14a-(*S*,*S*) with a *d.e.* of 93 %. Similarly, double helix (8b)₂ binds to 14b-(*S*,*S*) to give [3]foldaxane *P*-(8b)₂ \supset 14b-(*S*,*S*) with a *d.e.* of 56 %. X-ray crystal structures allowed for the unambiguous assignment of the stereochemistry of these complexes (Figure 7B).



Figure 7. (A) Achiral rods 13a-13d and chiral rods 14a-14b and 15. (B) Crystal structures of *P*- $2c \supset 14a$ -(*S*,*S*) (left) and *P*-(8b)₂ $\supset 14b$ -(*S*,*S*) (right) showing rod-to-helix chirality induction. (C) Crystal structures of poly-foldaxanes (8b)₄ $\supset 13b$ (top, left), (8b)₆ $\supset 13c$ (top, right) and energy-minimized molecular model, using MMFFs, of the [11]foldaxane (8b)₁₀ $\supset 13d$ (bottom). (D) Crystal structure of the homochiral [4]foldaxane *M*-(8c)₂•*M*-2d $\supset 15$ with a zoomed view of the

helix-helix interface (E). Helices **2c**, **2d**, (**8b**)₂ and (**8c**)₂ are colored in brown, purple, blue and green, respectively. Carbamate groups and ethylene glycol spacers are represented in orange and pink, respectively.

Geared toward template rods containing more information, compound **15** was made up successively of an (*R*)-naphthylethyl bulky group, a binding station known to bind (**8c**)₂, an ethylene glycol spacer, a binding station known to bind **2d** and finally an (*R*)-chiral phenethyl bulky group (Figure 7A). Solution and solid state studies showed the formation of a homochiral [4]foldaxane M-(**8c**)₂•M-**2d** \supset **15** (Figure 7D).² Additional rods made of multiple binding stations of variable lengths allowed for the faithful spatial arrangement of up to five single or double helices of three different kinds.² This type of assembly can be assimilated to a translation process where a sequence of alkyl carbamates is translated into a sequence of stacked aromatic helices. Translation of chemical information from one molecular type to another has reached exquisite levels in natural systems, *e.g.* the translation of mRNA into proteins by the ribosome. In contrast, synthetic systems that achieve translation are few and far from being optimized.^{60,61,62} The possibility to translate polycarbamates into aromatic oligoamides thus stands as an intriguing and original perspective.

6. EFFECT OF FOLDAXANE FORMATION ON ROD PROPERTIES

In all the systems presented above, foldaxane formation amounts to trapping an otherwise highly flexible guest containing an alkyl chain into the rigid cylindrical cavity of a multi-turn aromatic helix. The extent of this rigidification, *i.e.* how well does the helix keep the two ends of the guest away from each other, was investigated using the distance dependence of reversible electronic energy transfer (REET).⁵⁴ To achieve this, a bulky Ru(bpy)₃²⁺ (bpy = 2,2'-bipyridine) and a pyrene were installed at the ends of rod **9b** (Figure 5B) containing a binding station for (**8b**)₂.

The emissive ³MLCT state of the Ru(bpy)₃²⁺ chromophore is quasi-isoenergetic with the triplet state of the pyrene unit. This similarity allows the energy to shuttle back and forth between the two chromophores of the flexible thread all the more so that they can come in close proximity and a delayed luminescence was observed. After foldaxane formation, the chromophores are kept at a distance and effectively decoupled. Indeed, the structure of $(8b)_2 \supset 9a$ shows an interchromophore distance ≥ 1.8 nm (Figure 5E). This translates into characteristic photophysical signatures.

Another intriguing interplay between helix and rod was discovered while extending foldaxane design to rods other than alkyl carbamate guests. Applying the same principles that led to the design of sequence 2 from the structure and molecular recognition properties of capsule 1 (Figure 3A), we endeavored to produce foldaxanes derived from the ability of capsule sequence 16 (Figure 8A) to bind to tartaric acid 18 with high affinity, selectivity and diastereoselectivity (K_a $> 10^{6}$ L.mol⁻¹ in CDCl₃/DMSO - 99:1).⁴⁰ These molecular recognition properties stem from the larger cavity size of 16, imparted by the long pyridine-pyridazine-pyridine central segment (Figure 2A) and from the presence of 7-amino-1,8-naphthyridine units which hydrogen bond to the carboxylic acid guest. Thus, we synthesized sequence 17, an analogue of 16 lacking terminal Q₃ caps while preserving molecular recognition features, and rod 19, a dumbbell-shaped tartaric acid derivative.⁶³ Mixing 17 and 19 led to the rapid formation of the expected 1:1 foldaxane $17 \supset 19$ (Figure 8B). There again, helix handedness was quantitatively biased by stereogenic centers of the guest. This complex was found to be long-lived but it is in fact a kinetic supramolecular byproduct. Upon prolonged incubation, it slowly transformed in the 2:2 host-guest complex $(17)_2 \supset (19)_2$ in which two guest molecules bound at the extremities of a double helical (19)₂ (Figure 8B). We commented above that foldaxane most often outcompetes double helix formation (Figure 3C) because one duplex can produce two foldaxanes. This no longer holds true in the case of 17 and

19 because the rod-like guest also binds to the ends of the double helix through hydrogen-bonding (Figure 8B). Coincidentally, **19** also quantitatively biases the helicity of $(17)_2$ but with the opposite handedness to that it induces in **17**. Thus, mixing **17** and **19**-(*D*) leads first to the formation of *P*-**17** \supset **19**-(*D*) then to the formation of *M*-(**17**)₂ \supset (**19**-(*D*))₂.



Figure 8. (A) Sequences of 16 and 17 and formulae of 18 and 19. (B) Energy minimized molecular model, using MMFFs, of *M*-17 \supset 19-(*L*) (left) and crystal structure of (*P*-17)₂ \supset (19-(*L*))₂ (right). Tartaric acid moieties and bulky end groups are represented in orange and grey, respectively. (C) Formula of [2]rotaxane 20. (D) Crystal structure of the foldarotaxane 2d \supset 20 comprised of a DB24C8 macrocycle around an ammonium moiety and helix 2d wrapped around a dicarbamate

binding station; (left) zoom on the thread highlighting a kink due to a hydrogen bond between the amide (H-bond donor) and the central carbamate function (H-bond acceptor). The ammonium, amide and carbamate functional groups are represented in green, dark purple and orange, respectively. The macrocycle is shown in red whereas the helix is color coded as in Figure 2.

Recent works exploited the ability of fact that alkyl-carbamate rods that form foldaxanes may themselves be rotaxanes.⁶⁴ For instance, [2]rotaxane **20** (Figure 8C) possesses a dicarbamate binding station on which a helical foldamer may wind. It also contains an ammonium group as a binding site for a dibenzo-24-crown-8 (DB24C8) macrocycle. As evidenced by NMR, single helix **2d** wraps around the dicarbamate station of the rotaxane **20** to form $2d \supset 20$ ($K_a = 207\ 000\ L.mol^{-1}$ in CDCl₃), a new architecture which was called a foldarotaxane. The crystal structure of $2d \supset 20$ confirms that each building block binds to its complementary interacting site (Figure 8D). Upon deprotonation of the ammonium, the macrocycle may occupy other locations of the rod. Helix **2d** still winds around the rod to form a foldarotaxane, but a significant decrease of affinity was observed ($K_a = 9100\ L.mol^{-1}$ in CDCl₃). The drop is even more pronounced upon subsequent *t*-butyl-carbamoylation of the amine ($K_a = 3800\ L.mol^{-1}$). Overall, the presence and the localization of the macrocycle modulates the association between the helix and the thread and, reciprocally, the foldamer can act as a supramolecular auxiliary to compartmentalize the macrocycle around a region of the molecular rod for which it may have a lesser affinity.

7. SUMMARY AND OUTLOOK

In summary, foldaxane development has made considerable progress owing to the use of designer aromatic oligoamide sequences. Structural parameters such as host-guest affinity and selectivity, host-guest relative orientation, foldaxane single or double helicity, as well as relative

and absolute handedness can be controlled according to simple construction rules. Furthermore, the timescales of assembly and disassembly through different mechanisms, and of various translation and screw-like molecular motions, can also be orchestrated by means of structural features. Foldaxanes thus extend earlier work on rotaxanes and pseudo-rotaxanes into new directions. Given the rich literature on molecular recognition within helical containers,²¹⁻²⁶ one may anticipate other foldaxanes than the alkyl-carbamate-guest/aromatic-oligoamide-host to emerge. In this respect, foldaxanes based on tartaric acid guests,⁶³ or on helices not derived from aromatic amides²⁹⁻³⁴ constitute early examples of a growing family. One may also conceive foldaxanes using other foldamer shapes than helices. For example, some aromatic bent sheets also possess a cavity.⁶⁵ The combination of rotaxanes and foldaxanes also has potential for development. An interesting prospect would be to use of a foldamer helix as a controlled molecular auxiliary to confine another interlocked subunit (e.g. a macrocycle) along a molecular rod. Helixrod host-guest complexes may also open new possibilities in the field of topologically complex molecules, e.g. knots. In a multi-turn foldaxane, the helix and the rod are arranged with multiple crossing points. Upon appropriately covalently connecting the ends of the rod with the ends of the helix, a topologically non-trivial structure may result, as has been proposed for the synthesis of knots from double helices.^{66,67} Progress along these lines is being made in our laboratories and will be reported in due course.

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Arundhati Roy received her PhD degree in Supramolecular Chemistry from IISER Pune (India) under the mentorship of Prof. Pinaki Talukdar in 2017. Following a 3-year postdoctoral experience with Prof. Huaqiang Zeng at A*STAR (Singapore), she moved to Ludwig Maximilian University Munich, Germany as an Alexander von Humboldt research fellow with Prof. Ivan Huc in 2020. Her current research interests focus on the development of aromatic foldamers for molecular recognition and transport.

Ivan Huc has been since 2017 a full Professor at the Ludwig-Maximilians-Universität (Munich, Germany). He obtained his doctorate from the University of Paris VI in 1994 for work performed jointly at Ecole Normale Supérieure (France) and MIT (USA). He subsequently worked as post-doc, next as CNRS researcher at the University of Strasbourg until 1998, when he was appointed group leader at the European Institute of Chemistry and Biology at the University of Bordeaux. His group focuses on the design, synthesis and characterization of aromatic foldamers and their applications including pharmacological aspects.

Yann Ferrand is a CNRS research director and group leader at the Institute of Chemistry and Biology of Membranes and Nano-objects of the University of Bordeaux (France). He studied chemistry at the University of Rennes (France) where he received his PhD degree in 2005. Then, he worked as a post-doc in the School of Chemistry of the University of Bristol (United Kingdom) until 2007. His research focuses on the design and synthesis of synthetic macromolecules (*e.g.* foldamers) for the recognition of complex molecular targets and their use as sensors.

Author Contributions

The manuscript was written through contributions of all authors. All authors have given approval to the final version of the manuscript. ‡ V. K. and A. R. contributed equally.

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