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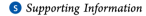
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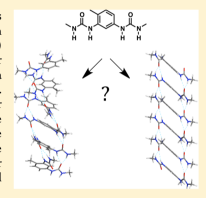
Conformational Plasticity of Hydrogen Bonded Bis-urea Supramolecular Polymers

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ABSTRACT: We report a detailed structural investigation of supramolecular polymers formed by hydrogen bonded self-assembly of bis-urea monomers. The careful exploration of the energy landscape by molecular mechanics/molecular dynamics (MM/MD) simulations has allowed us to identify three distinct self-assembled structures of similar stabilities. These structures have been compared to X-ray crystal data. We observe that a slight change in the molecular structure can favor a particular structure over the others. Detailed analysis shows that hydrogen bonds stabilize all three structures to a similar extent. Therefore, it is the interactions among the lateral substituents, and with the filament environment, that are the decisive factors in the competition between the possible self-assembled structures. This study constitutes a clear reminder that the conformation of a supramolecular polymer is a sensitive function of the molecular structure and may significantly differ from the solid-state conformation of a model compound.



■ INTRODUCTION

Supramolecular polymers are chains of small molecules held together through reversible noncovalent interactions. 1-4 Compared to usual covalent polymers, the dynamic character of the interaction is responsible for the appearance of new properties, such as self-healing behavior or improved processability. Moreover, their high molar mass structure is maintained only by weak interactions, which makes them highly sensitive to small perturbations. Therefore, supramolecular polymers are particularly interesting systems enabling the quantification of subtle self-assembly features such as chiral amplification⁵ or solvation effects.^{6,7}

Advances in this field are directly linked to our ability to rationally improve the design of a given supramolecular polymer, which in turn requires one to know its precise structure. However, this is a difficult task because of the lack of long-range order in these systems and because of their dynamic character. A popular approach to circumvent this problem is to use modern molecular simulation tools validated by experimental data. 8-10 Often, a model compound similar to the supramolecular polymer is studied experimentally, for instance by X-ray diffraction on crystals, to obtain structural information. 11-15 These data can then be used to select and validate a simulation method to assess the conformation of the supramolecular polymer. However, this is still a significant challenge, because the conformation of the model compound in the crystal is not necessarily the most stable conformation of the solvated assembly, as solid-state packing effects distort the energy landscape. This approach therefore requires a comprehensive exploration of the energy landscape of the supramolecular assemblies to identify the most relevant conformations and arrangements.

Besides determining the most stable structures, the molecular modeling approach also allows one to highlight the richness of the structural variability of a given class of polymers and can provide a description of those structures with a unique level of details. This is particularly interesting for the class of bis-urea based supramolecular polymers studied in this work. In principle, the assembly of these molecules can be favored/ controlled by hydrogen-bonding between the urea groups, π stacking between the toluene units and interactions between the hydrocarbon side groups.

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In nonpolar solvents (such as toluene or dodecane), 2,4-bis(2-ethylhexylureido)toluene (bis-urea 1) (Figure 1) has

Figure 1. Structure of the bis-urea monomers examined in this study; the torsion angles φ_1 and φ_2 are noted on 1 (the arrows indicate in which direction the torsion angle is considered positive).

been shown to self-assemble by hydrogen bonding into two distinct supramolecular structures, which are stable over a wide range of concentrations and temperatures, and are in dynamic exchange with the monomer. 16-18 A small-angle neutron scattering (SANS) study established that both structures are long and fibrillar, the high temperature structure being thinner than the low temperature structure. Throughout this work, we focus exclusively on the high temperature structure, which has been shown by SANS to consist of single stranded filaments. Quantitative measurements of the strength of hydrogen bonds by IR spectroscopy and isothermal titration calorimetry (ITC), and comparison to data for monoureas revealed that both urea groups of a given monomer are hydrogen-bonded to the next monomer in a cooperative fashion, i.e., random association of urea groups can be excluded. 19,20 On the basis of those data and the structure of a monolayer probed by scanning tunneling microscopy (STM), ^{21–23} a schematic ladder-like supramolecular arrangement has been proposed. 16 However, many degrees of freedom are present in this molecule, for instance, those related to the flexibility of the side groups and the relative orientation of the methyl groups on adjacent toluene units in the assemblies; as a result, considerable uncertainty remains concerning its organization in the solvated assembly.

It is thus the aim of the present study to reach a detailed structural description of the supramolecular filaments and an evaluation of their energetic stability, via a joint approach combining experimental structural studies and molecular mechanics/molecular dynamics (MM/MD) simulations of model compounds for the filaments.

Needle-like crystals suitable for X-ray diffraction have been obtained for the closely related molecules 2 and 3 (Figure 1). On one hand, these two compounds can be used to set up and

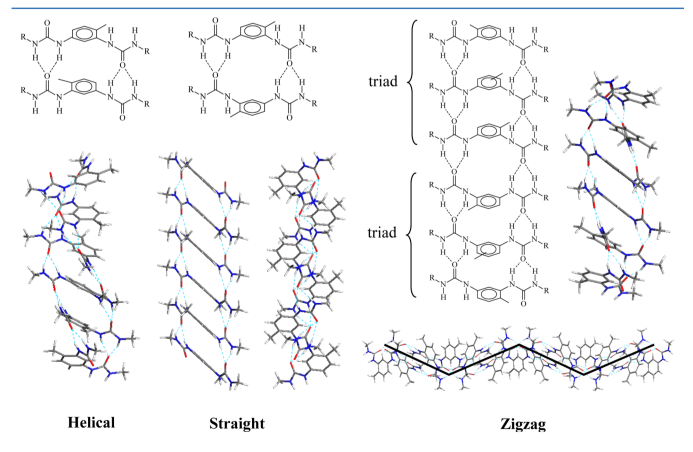


Figure 2. Orientation of the bis-urea molecules in the filament morphologies discussed in this study. The 3D displays correspond to bis-urea 4, i.e., where the substituent R of the schematic motif is a methyl, and were obtained by an MM simulation. For the straight filament, two views are displayed, along two perpendicular directions.

validate a simulation methodology to study assemblies of bisurea 1 molecules. On the other hand, they allow to extend the investigation to bis-urea supramolecular polymers differing by the nature of the substituents. Finally, we considered bis-urea 4, which has a very small side group, i.e., a methyl group, and should thus reveal the most stable conformation of the bis-urea core in the absence of any steric hindrance from the substituents.

METHODOLOGY

The bis-ureas 1–4 were modeled with the Dreiding²⁴ force field using the Materials Studio modeling package. The validity of Dreiding for this study was evaluated on single molecules versus a quantum chemical method. Special attention was dedicated to the behavior of the torsions, which are the internal coordinates most influencing the supramolecular organization. The comparative analysis (available as Supporting Information) indicates the suitability of Dreiding, which is further reinforced by the fact that this force field correctly reproduces the experimental solid-state morphology of bis-ureas 2 and 3 (see below).

After validating the force field, a methodology to generate filament structures as exhaustively as possible has been devised. To that aim, we performed conformational searches for a single molecule, then for dimers, and then for octamers, i.e., for assemblies of increasing size, to funnel the search toward the most stable structures having a complete hydrogen bond network. That approach was applied to the simplest compound, bis-urea 4, to assess the dominant role of the hydrogen bonds (among the other intermolecular interactions) in stabilizing molecular assemblies.

First, the potential energy surface of bis-urea 4 was explored by systematically varying the orientation of the urea groups with respect to the aromatic unit (φ_1 and φ_2 in Figure 1) in a single isolated molecule. That work also provided the stable conformations of the bis-urea core in the absence of intermolecular interactions. Then, the stable conformations obtained were used to build hydrogen-bonded dimers. Among them, however, only a few have an arrangement that can be extrapolated to a fully hydrogen-bonded filament; thus only a few octamers with a periodic motif were built subsequently from these dimers.

When building the octamers, the stacked molecules were placed so that the methyl groups on adjacent toluene rings have alternating orientation, ²⁵ in order to reduce steric hindrance (see the motif of helical and straight structures in Figure 2). That strategy produces only filaments with a repetitive motif of at most two molecules. We also included the repetitive motif of six molecules that is experimentally observed in the crystal structure of bis-urea 2 (Figure 2, zigzag structure). In total, five conformations were obtained for each bis-urea compound, but only three of them keep their structure during MD simulations. We will mainly discuss these three persistent structures, which are referred to according to the shape of their morphology: helical, straight, or zigzag. The two unstable structures are discussed in the Supporting Information.

Once these stable morphologies have been obtained, the influence of the steric effects from the side groups was investigated. The three filament configurations were systematically studied for bis-ureas 1 to 4, generating 12 different structures in total. Filaments of 32 molecules were built, geometry-optimized by MM and then submitted to 1500 ps MD simulations. Data for analysis were collected every 3 ps

after the initial 150 ps of trajectory, when most strong fluctuations have disappeared. To analyze the systems, the distribution of torsions is displayed as an average of 5 bins of 1° each, and the evolution of the potential energy is displayed as block averages calculated on 10 structures.

The MM energy minimizations were performed with a conjugate gradient algorithm until a convergence criterion of 0.001 kcal/mol·Å was reached. The long-range interaction cutoff distance was set to 14 Å with a spline width of 3 Å. The charges were assigned from the pcff force field.

The MD simulations were performed in the NVT ensemble (i.e., particle number, volume, and temperature are kept constant). The Nosé thermal bath coupling scheme²⁶ was used to maintain the temperature at 300 K, with a coupling constant of 0.1. A Verlet velocity algorithm²⁷ was used to integrate the equations of motion with a 1 fs time step. All structures in our study were relaxed prior to MD simulations.

RESULTS

Experimental Structure of Filaments in the Solid State. The first clues about the organization of bis-urea 1 in filaments and the driving forces leading to them were obtained by analyzing the X-ray diffraction data of the closely related systems, bis-ureas 2 and 3. Bis-urea 3 crystallizes in a monoclinic cell (a = 17.566 Å, b = 23.795 Å, c = 8.651 Å, β = 117.40°), forming straight filaments with a structural motif of two molecules repeating along c. A given molecule has both urea groups tilted with respect to the phenyl core by the same absolute value (129.3°) and with the same sign, but the sign alternates from one molecule to the next (see Figure 1 to localize the torsions φ_1 and φ_2 that describe the orientation of the urea groups). This type of filament morphology is referred to as straight, and is displayed in Figure 2. (Note that all morphologies represented in Figure 2 were those obtained by simulation and correspond to bis-urea 4 filaments, which are used to describe characteristics common to all four bis-urea compounds and common to both crystalline and simulation conditions. For the representation of the actual crystal structures of bis-urea 2 and 3, see the Supporting Information.)

Bis-urea 2 crystallizes in a monoclinic cell (a = 19.628 Å, b =18.277 Å, c = 24.896 Å, $\beta = 112.503^{\circ}$). The structural motif has six molecules repeating along c, with kinks appearing every three molecules, thus leading to zigzagging filaments. This type of filament morphology is thus referred to as zigzag, and is displayed in Figure 2. (Note that in the example displayed, the zigzag shape is more visible when looking at the hydrogen bond network; see also Figure S3). Each motif is composed of two triads of molecules. In the first triad, the central molecule has both urea groups tilted by 131.8°. For the other two molecules, one urea is tilted by 125.7° (φ_2), the other one by 137.6° (φ_1). In the second triad, the torsions are similar, except that the sign is negative. As the sign changes when going from one triad to the next, the two frontier molecules are organized as in the straight motif (compare the straight motif and the two central molecules of the zigzag motif in Figure 2). In each triad, the position of the methyl group on the toluene ring of the central molecule is undefined in the crystal structure data. This orientational disorder is consistent with the fact that the orientation of the methyl group on the central molecule alternates with the methyl group of any of the two neighboring molecules, but not with the methyl group of both molecules.

In summary, these two bis-urea-based monomers crystallize differently due to the nature of their side groups, but,

remarkably, both structures in the solid state are filaments where all urea moieties are involved in hydrogen bond. As experimental evidence suggests that bis-urea 1 also forms filaments in solution (although crystal data are not available), these results point out that filament structures have a strong tendency to form for this class of supramolecular polymers, mainly driven by the formation of intermolecular hydrogen bonds.

Simulations of Single Filaments. The potential energy map of a single bis-urea 4 molecule (Figure 3) reflects the

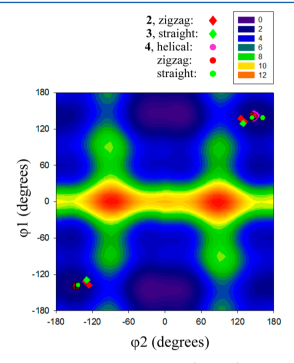


Figure 3. Map of relative potential energy (kcal/mol) of an isolated bis-urea 4 molecule built by varying φ_1 and φ_2 , showing the preferential orientations of the urea moieties with respect to the core. The values corresponding to the helical, zigzag, and straight conformations for a simulated single filament of bis-urea 4, and the zigzag and straight conformations for the experimental crystalline filaments of bis-ureas 2 and 3, respectively, are located on the map.

preferential orientations of the urea moieties with respect to the core in the absence of intermolecular interactions. The potential energy is minimal for $|\varphi_1|=150^\circ$ and $|\varphi_2|=20^\circ$, but around that minimum, conformations requiring less than the thermal energy to be accessible at room temperature are found for φ_1 between -130° and -160° , and between 130° and 160° , and φ_2 between -40° and 40° ; in other words, the global minimum is in a potential energy valley with a large flat bottom, where the urea groups can be almost coplanar with the ring or tilted by up to a few tens of degrees.

On this map the filament conformations determined from X-ray diffraction data or from simulation are visualized as diamonds or circles, respectively. None of them are located in the area of minimal potential energy of the monomer. This means that to form the stabilizing hydrogen-bond network, the urea moieties have to deviate from their preferential orientation in isolated molecules. It is actually in the second lowest potential energy valley, where $|\varphi_1|$ is about $130-150^\circ$ and $|\varphi_2|$ is about $140-170^\circ$, that the zigzag, straight, and helical morphologies are located.

It is remarkable to have similar torsions in these three morphologies, considering the differences between them: the size and shape of the side groups, the environment (crystal or vacuum), the morphology of the filament itself (helical, zigzagging, or straight). It is thus likely that both the intramolecular interactions associated to the orientation of the urea moieties and the intermolecular interactions associated to hydrogen bonding synergistically stabilize the assemblies when $|\varphi_1|$, $|\varphi_2| \sim 125-160^\circ$. Within that constraint, the morphology still conserves some freedom to vary and respond to packing or steric effects induced by the substituents, by tuning the frequency of the φ_1 , φ_2 sign change along the filament. This sign alternation is seen on the energy map (Figure 3), where the zigzag and straight morphologies have two well-defined, symmetrically positioned populations of φ_1 , φ_2 : the sign alternates from half a motif to the next (i.e., each molecule or each triad for the straight and zigzag morphologies, respectively). Helical filaments, on the contrary, have only one population of torsion angles (all φ_1 and $\varphi_2 > 0$ in a M-helix or all φ_1 and φ_2 < 0 in a P-helix).

The relative adaptability of the filament morphologies to different conditions is further illustrated during the MD simulations performed on the unstable bis-urea 4 filaments: as mentioned in the Methodology section, two of the initial five filament morphologies are unstable during the simulations, and one of them evolves toward the more stable helical morphology in a few picoseconds (see Supporting Information). However, in the course of this morphology conversion, a few bis-urea molecules do not follow the main path and adopt conformations with both φ_1 and φ_2 close to 60° (Figure 4). As a result, the filament folds in two. What is remarkable is that only a few torsional "defects" introduced in the helical morphology are sufficient to fold the filament, while conserving the full integrity of the hydrogen bond network. This observation suggests that bis-urea filaments have the possibility to fold and adapt their morphology to changes of their environment without much energy penalty.

Figures 5 and 6 show the H-bond energy and the total potential energy, respectively, for the filaments of the four bisurea compounds in the course of the MD simulations, which allows one to estimate the relative stability of the helical, zigzag, and straight morphologies.

As far as H-bond energy is concerned, bis-urea 4 is the system that has the strongest hydrogen-bond network, with similar strength for the three morphologies (Figure 5). This is probably due to the small size of the substituents, which limits steric hindrance. However, larger substituents do not destabilize much the H-bond network: at most 1 kcal/mol per molecule, or ~10% of the energy of an optimal H-bond. For a given bisurea molecule, the H-bond energy difference between the three morphologies is even smaller. The largest difference, 0.7 kcal/mol, is found for bis-urea 2, between the helical structure on one hand, and the zigzag and straight structures on the other hand. In all cases, the H-bond network persists throughout the MD simulations.

Concerning $\pi-\pi$ stacking interactions, the situation is very dependent on the morphology. In the helical morphology, adjacent aromatic rings are placed in close contact, with a distance down to 3.5 Å, and are therefore stabilized by $\pi-\pi$ stacking interactions. This is also the case for the zigzag morphology, but only within triads (see Supporting Information). By contrast, in the straight morphology, adjacent bisureas position their aromatic groups on opposite sides of the

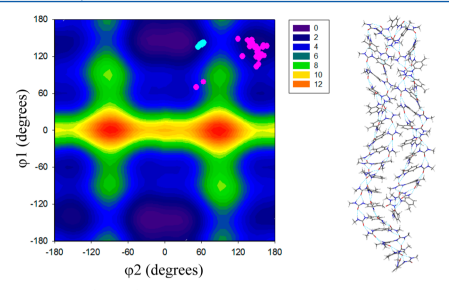


Figure 4. Map of relative potential energy (kcal/mol) of an isolated bis-urea 4 molecule built by varying φ_1 and φ_2 . The values for a simulated unstable filament of bis-urea 4 are located on the map before (cyan dots) and after (magenta dots) 1500 ps dynamics. The folded structure obtained after MD simulation is displayed on the right.

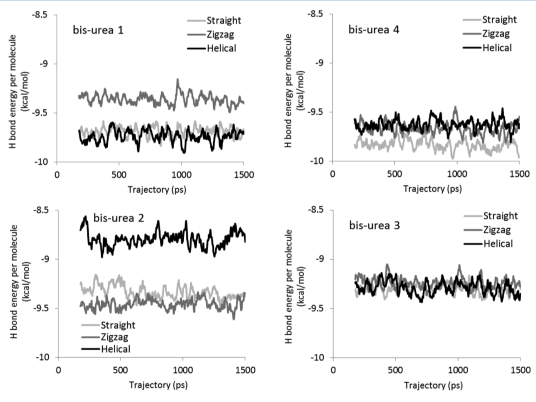


Figure 5. Plots of the H-bond energy per molecule during the MD simulations for the helical, zigzag, and straight morphologies of the four ureabased systems. The energy values are block averages over 10 successive MD structures. See Table S5 for averages and standard deviations.

hydrogen bonded urea network (Figure 2), so that $\pi - \pi$ stacking is impossible.

The influence of $\pi-\pi$ interactions on the stability of the assemblies is expected to be most visible for the system where there is minimal influence of the organization of side groups, which could overcome any change of $\pi-\pi$ interactions from one morphology to the next. It is thus in bis-urea 4, the system with the smallest side group, that we will search any energetic effect due to change of $\pi-\pi$ interactions between assemblies. Figure 6 shows the potential energy of the assemblies, including all intra- and intermolecular interactions. The stability decreases

from continuous π stacking (helical), to partial π stacking (zigzag), to absent π stacking (straight), thus suggesting that $\pi-\pi$ interactions do stabilize the assemblies that have them. However, these energy differences are indeed dwarfed by the interactions between side groups, when their size is larger. (See in Figure 6, bis-urea 1, where the energy difference between the morphologies is too small to hierarchize them; or bis-ureas 2 and 3, where the influence of the side group is clearly dominant.)

In the rest of the discussion, we will thus focus on the influence of the side groups. From that point of view, a

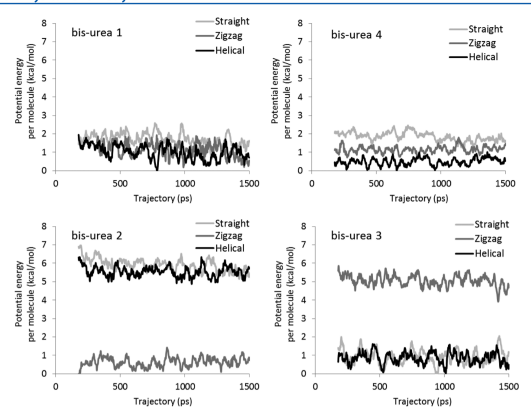


Figure 6. Plots of the potential energy per molecule during the MD simulations for the helical, zigzag, and straight morphologies of the four systems. The energy values are block averages over 10 successive MD structures, and the zero of the energy scale corresponds to the lowest-energy block average found during the MD runs on the three morphologies. See Table S6 for averages and standard deviations.

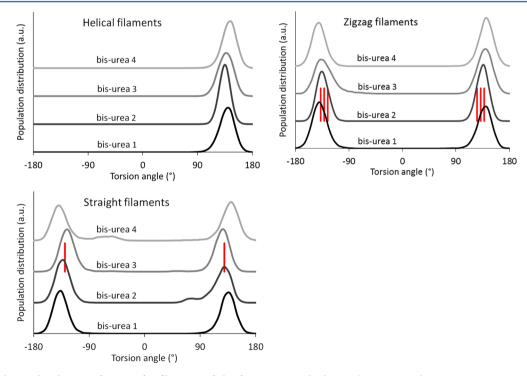


Figure 7. Population distribution of φ_1 , φ_2 for filaments of the four compounds during the MD simulations. By comparison, the population distribution corresponding to the two X-ray experimental structures are represented as red bars.

distinction can be made between bis-ureas 1 and 4 on one hand, and 2 and 3 on the other hand. For 1 and 4, the range of total potential energy within which the morphologies lie is merely 0.7 and 1.4 kcal/mol, respectively (Figure 6 and Table

S6). Since the energy differences are quite small between the three morphologies, it is likely that the substituents have room to organize and do not exert strong steric constraints. Despite small energy differences between them, each morphology is, however, quite stable, i.e., they do not convert into each other, nor evolve into a mixing of morphologies during the MD simulations. It is a feature also shared by the other bis-urea systems studied here. Similarly, the population distributions of φ_1 and φ_2 are narrow for all systems (Figure 7), meaning that the conformation of the bis-urea cores is maintained during the MD simulations, and they are similar whatever the side groups and whatever the environment (for the simulated single filaments or the two experimental crystalline filaments). These observations reflect the steepness of the potential energy valley in which these morphologies lie once the filament is formed. The stability of the assemblies is attributed to cooperative effects, in particular to cooperative H-bonds, between neighboring molecules.

Based on these simulations, the three morphologies could exist for bis-ureas 1 and 4, and the actual morphology adopted by these systems in a given environment most probably results from a subtle balance of intrafilament and extra-filament interactions. For instance, experimentally, it was found that filaments of enantiomerically pure bis-urea 1 in solution show no CD signal, ²⁸ suggesting that the morphology is nonhelical, i.e., straight or zigzag.

For bis-ureas 2 and 3, the difference of potential energy between the morphologies is about 5 and 4 kcal/mol, respectively (Figure 6 and Table S6). This is a large value, which is also much larger than the differences in the H-bond energy (below 1 kcal/mol) between the morphologies. This result indicates that the structure of the substituents has here a strong influence on the morphology adopted by the filament, and that some morphologies are clearly destabilized by steric interactions

As bis-ureas 1 and 2 are structural isomers, the differences between the two systems in terms of stability of the three morphologies are especially noteworthy. For bis-urea 2, the zigzag morphology is much more stable than the straight and helical morphologies, by about 5 kcal/mol. The zigzag morphology is also the morphology that is found experimentally in crystals. Thus, the intrafilament interactions are here highly selective, as they give rise to the same organization both in the presence of interfilament interactions (in the experiment) and in the absence of such interactions (in the MD simulations). The presence of a quaternary carbon directly attached to the urea moieties for bis-urea 2, instead of a secondary carbon for bis-urea 1, probably limits the range of orientations of the molecules for which steric hindrance is small and simultaneously the hydrogen bond network is strong, which leads to the "selection" of a single morphology (i.e., zigzag). The same reasons can be invoked for explaining the widening of the energy differences between the morphologies of bis-urea 3 when compared to those of bis-urea 1. In bis-urea 3, the second atom of the side group is a quaternary carbon, instead of a tertiary carbon in bis-urea 1. For bis-urea 3, the simulated straight and helical morphologies have similar characteristics in terms of potential energy and hydrogen bond energy. The fact that it is the straight morphology that is found experimentally in crystals indicates that other interactions not present in single filaments are significant enough to favor one of the two morphologies in the solid state. In that selection, interfilament interactions likely play a significant role. In solution, however, interactions with the solvent are likely the dominant interactions tilting the energy balance toward one of the two morphologies. Globally, these results show the

importance of the branching and size of the side group in determining the actual morphology of the filament.

CONCLUSION

The careful exploration of the energy landscape of bis-urea supramolecular polymers has allowed to identify three distinct self-assembled structures of similar stabilities, two of which are also found in X-ray crystal data. In fact, a slight change in the molecular structure can favor a particular structure over the others. Detailed analysis shows that hydrogen bonds stabilize all three structures to a similar extent, whatever the side groups bulkiness. However, the total potential energy is strongly affected by the nature of the side groups. Therefore, interactions among the lateral substituents, and with the filament environment, appear to be the decisive factor in the competition between the possible self-assembled structures. This study constitutes a clear reminder that the conformation of a supramolecular polymer is a sensitive function of the molecular structure and may significantly differ from the solidstate conformation of a model compound.

ASSOCIATED CONTENT

S Supporting Information

Synthesis, crystal structure data, and additional simulation results. This information is available free of charge via the Internet at http://pubs.acs.org

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Notes

The authors declare no competing financial interest.

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