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# Effect of H-Bonding on Order Amplification in the Growth of a Supramolecular Polymer in Water

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*Supramolecular polymer, self-assembly, molecular modeling, molecular simulation, H-bonding, order amplification, cooperativity, ordered aggregation, 1D assembly*

**ABSTRACT:** While a great deal of knowledge on the roles of hydrogen bonding and hydrophobicity in proteins have resulted in the creation of rationally designed and functional peptidic structures, the roles of these forces on purely synthetic supramolecular architectures in water have proven difficult to ascertain. Focusing on a 1,3,5-benzenetricarboxamide (BTA) based supramolecular polymer, we have developed a molecular modeling approach to dissect the energetic contributions involved in the self-assembly (enthalpic, entropic, electrostatic, hydrophobic, etc.) upon growth of both ordered BTA stacks and random BTA aggregates. Utilizing this set of simulations, we have unraveled the cooperative mechanism for polymer growth, where a critical size must be reached in the aggregates before emergence and amplification of order into the experimentally observed fibers. Furthermore, we have found that the formation of ordered fibers is favored over disordered aggregates solely on the base of electrostatic interactions. Detailed analysis of the simulation data suggests that H-bonding is a major source of this stabilization energy. Experimental and computational comparison with a newly synthesized 1,3,5-benzenetricarboxyester (BTE) derivative, lacking the ability to form the H-bonding network, demonstrated that this BTE variant is also capable of fiber formation, albeit at a reduced persistence length. This work provides unambiguous evidence for the key 1D driving force of hydrogen bonding in enhancing the persistency of monomer stacking and amplifying the level of order into the growing supramolecular polymer in water. Our computational approach provides an important relationship directly linking the structure of the monomer to the structure and properties of the supramolecular polymer.

Supramolecular polymers can mimic many of the properties of conventional covalent polymers, while allowing for dynamic, bioinspired and adaptive properties.<sup>1</sup> Formed via transient non-covalent interactions between monomers, dynamic behavior and reversibility are encoded into their molecular structure. These inherent properties resulted in the development of many functional materials with enhanced processing characteristics, self-healing behavior, stimuli responsiveness, and novel electronic properties.<sup>2</sup>

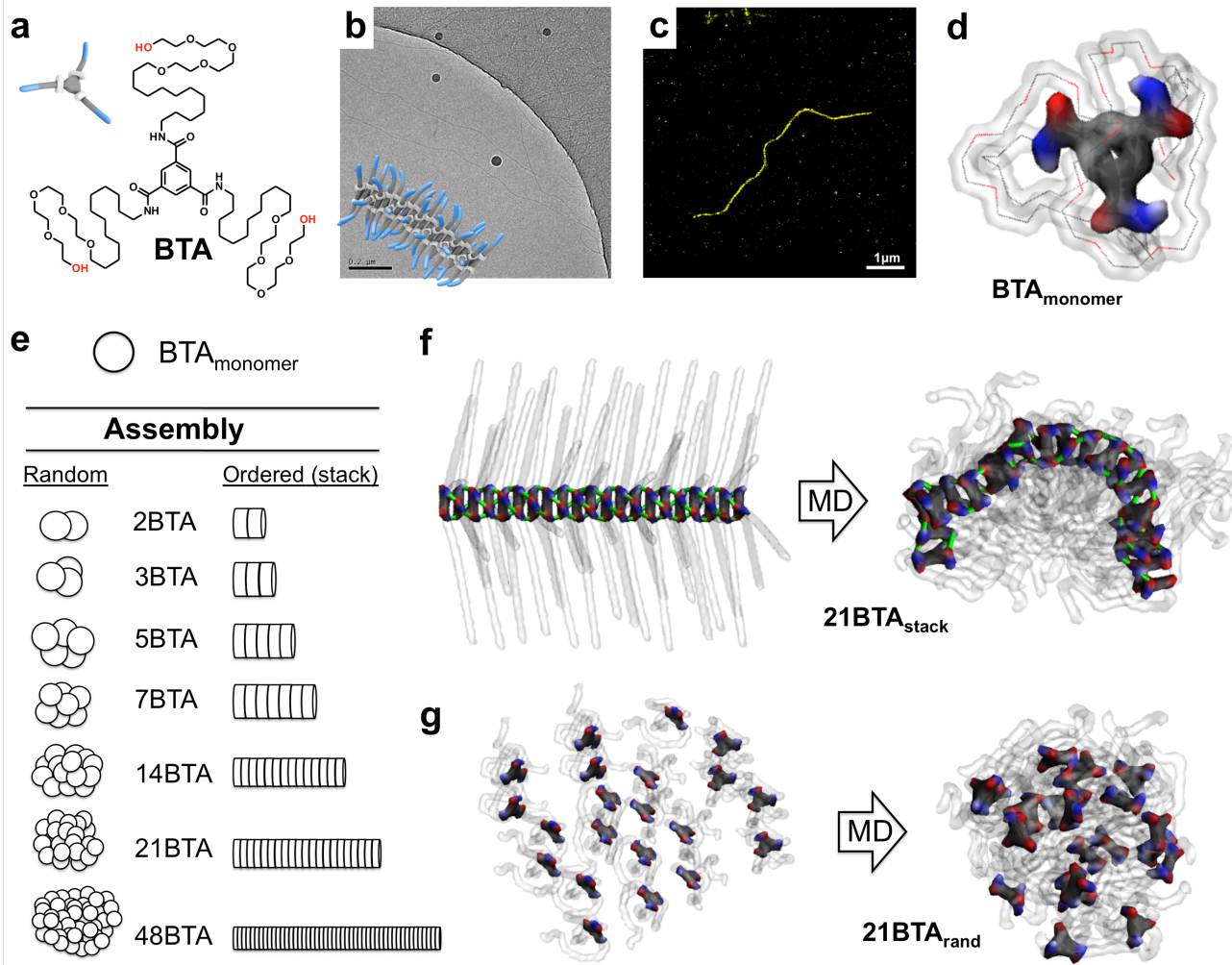
Supramolecular polymers are ideally suited to build new biomaterials<sup>3</sup> that can mimic or interact with dynamic, biological environments. Attaining fundamental understanding on the interplay of different types of non-covalent interactions in relation to the dynamics of the formed materials could allow the design of functional water-soluble supramolecular polymers<sup>4</sup> as dynamic as cells and the extra-cellular matrix (ECM).<sup>5</sup> Recently, this led to the development of bioinspired supramolecular polymers based on peptide amphiphiles,<sup>6-8</sup> ureidopyrimi-

dinones,<sup>9</sup> cyclodextrins,<sup>10-11</sup> cucurbiturils,<sup>12-13</sup> dendrimers,<sup>14</sup> and functionalized gold nanoparticles<sup>15</sup> to name a few.

In this framework, 1,3,5-benzenetricarboxamide (BTA) based supramolecular polymers, in which the BTA monomers self-assemble directionally due to well defined three-fold hydrogen bonding and stacking of cores, are an ideal scaffold for fundamental studies of supramolecular polymers.<sup>16</sup> BTA self-assembly is well characterized in a variety of molecular architectures and environments.<sup>16</sup> By engineering a hydrophobic dodecyl spacer ( $C_{12}$ ) connected to a water-compatible tetraethylene glycol tail ( $EG_4$ ) around the BTA core (Figure 1a), we have recently shown that these monomers form high-aspect ratio supramolecular fibers in aqueous solution (Figures 1b,c).<sup>17</sup> This BTA analog has been used to study the mechanism of monomer exchange,<sup>18</sup> the ability to reorganize and cluster monomers in response to binding with multivalent recruiters,<sup>19</sup> and how small changes in the monomer structure affect the exchange dynamics.<sup>20-21</sup> Despite these advances, the polymerization behavior of water-soluble BTAs (and many

aqueous H-bonding based supramolecular polymers) remains extremely difficult to interpret at a fundamental level.<sup>21</sup> Due to the hydrophobicity of the monomers, the efficient depolymerization of such amphiphilic polymers in aqueous solution using temperature or organic solvent is prohibitively difficult. Consequently, information as to the behavior, importance and interplay of interactions such as H-bonding and hydrophobicity

than isodesmic polymerization if the free-energy of monomer addition to the growing chain is the same.<sup>22</sup> Traditionally, BTA based supramolecular polymers are cooperative in the formation of the H-bonding motif in organic solvents.<sup>16</sup> While some preliminary results suggested cooperativity in a sugar functionalized BTA in water,<sup>23</sup> little is known about the ability of assembled BTAs to maintain their cooperative H-bonding



**Figure 1.** Modeling BTA assembly. (a) Molecular structure of the water-soluble BTA monomer studied herein. (b,c) TEM (b) and STORM (c) images of BTA fibers in water. (d) Equilibrated BTA monomer in water solution obtained from the MD simulation. (e) Modeling strategy adopted in this study. Comparison between ordered and disordered BTA assemblies of the same size (same number of BTA monomers) allows studying the modulation of the interactions leading to the growth of ordered supramolecular polymers in water. (f) Starting (0 ns) and final (400 ns) configurations of the **21BTA<sub>stack</sub>** simulated system. (g) Starting (0 ns) and final (400 ns) configurations of the **21BTA<sub>rand</sub>** simulated system. In the snapshots, the BTA side chains are transparent, the BTA cores are colored per atom (C: grey, O: red and N: blue) and H-bonds are colored in green. Water molecules are not shown for clarity.

become difficult to ascertain with conventional experimental techniques.

Of particular importance to understanding supramolecular polymers is their mechanism of growth. The two major mechanisms to date are isodesmic and cooperative polymerizations.<sup>22</sup> While in an isodesmic polymerization the addition of each monomer to a growing polymer occurs with the same release of free-energy, in a cooperative polymerization monomer addition becomes increasingly favorable with the growth of the polymer, usually after a nucleation event. In general, cooperative supramolecular polymerization forms longer polymers

motif in the presence of water as a competitive solvent.

Driven by the clear challenges in obtaining insight at a molecular level into these supramolecular polymers computer simulations have been recently introduced to complement experimental work. Previous studies based on atomistic and DFT simulations have been conducted to model stacks of smaller BTA derivatives with shorter side chains in the gas-phase or in organic solvents, focusing on the study of self-assembly cooperativity in the BTA stacks.<sup>24-27</sup> These efforts provided important insight on H-bonding and dipole-dipole interactions in an intrinsically ordered condition, in which the short side

chains and the absence of a polar solvent emphasize the level of order in the system and the role of H-bonding. However, in aqueous solution the increased structural complexity of the water-soluble monomers (Figure 1a) and the presence of important solvophobic effects make the computational study of these supramolecular fibers drastically more complicated.<sup>20</sup>

All-atom molecular dynamics (MD) simulations recently allowed the study of self-assembled fibers of peptide amphiphiles<sup>6</sup> and water-soluble BTA monomers<sup>20</sup> in explicit water. In particular, our MD simulations permitted detailed inspection of a BTA-based supramolecular polymer in water at atomic scale and provided insight into the tertiary structure, hydrophobic vs. H-bonding interactions and penetration of water into the structure. Our models of infinite water-soluble BTA fibers showed that these supramolecular polymers are far from being perfectly extended and ordered in water solution. Moreover, while H-bonding remains active in the BTA polymer, the strong folding of the BTA side chains (primary folding) and of the fibers themselves (secondary folding) during the simulations demonstrate the predominant character of the hydrophobic effects.<sup>20</sup> Comparing models of chiral and achiral monomer assemblies also allowed us to untangle the effect of a subtle mutation in the monomer on the polymer structure, which provided plausible explanations for differences in experimentally observed monomer exchange dynamics between the achiral and chiral assemblies.<sup>20</sup>

A thorough understanding of the behavior of BTA-based supramolecular polymers in water is fundamental toward the rational design of self-assembled BTA materials in biologically relevant environments. In such a complex framework, we are interested in answering fundamental questions such as: What are the factors controlling the directionality of monomers' self-assembly? What interactions control/favor the emergence of order in the polymer? What is the exact role of H-bonding in this process? While in organic solvent monomer self-assembly has been suggested to proceed in rather uniform way during polymer growth, is this the same in water? Does a precise limit exist (critical size) over which order emerges in the supramolecular structure while below this threshold disordered aggregates are formed?

Herein we report a comprehensive atomistic modeling approach to understand the factors controlling the growth of ordered supramolecular BTA polymers in water to answer these questions. Systematic all-atom MD simulations of ordered (stacked) one-dimensional (1D) BTA assemblies of incremental size, and comparison with randomly ordered BTA aggregates of the same size, allow us to obtain unique insight on the amplification of order and the interactions governing self-assembly in the growing polymer (Figure 1). Finally, experimental and computational comparison to a structurally analogous 1,3,5-benzenetriester (BTE) derivative, having the amides simply replaced by esters and lacking the ability to form a strong H-bonding network, permits us to unambiguously define the role of H-bonding on the growth of an ordered BTA supramolecular polymer in water.

## RESULTS AND DISCUSSION

**Cooperativity in BTA self-assembly in water.** The cooperative self-assembly behavior of BTAs in the gas-phase or in organic media has been well studied, both experimentally<sup>28</sup> and computationally.<sup>25-27,29</sup> However, in water experimental

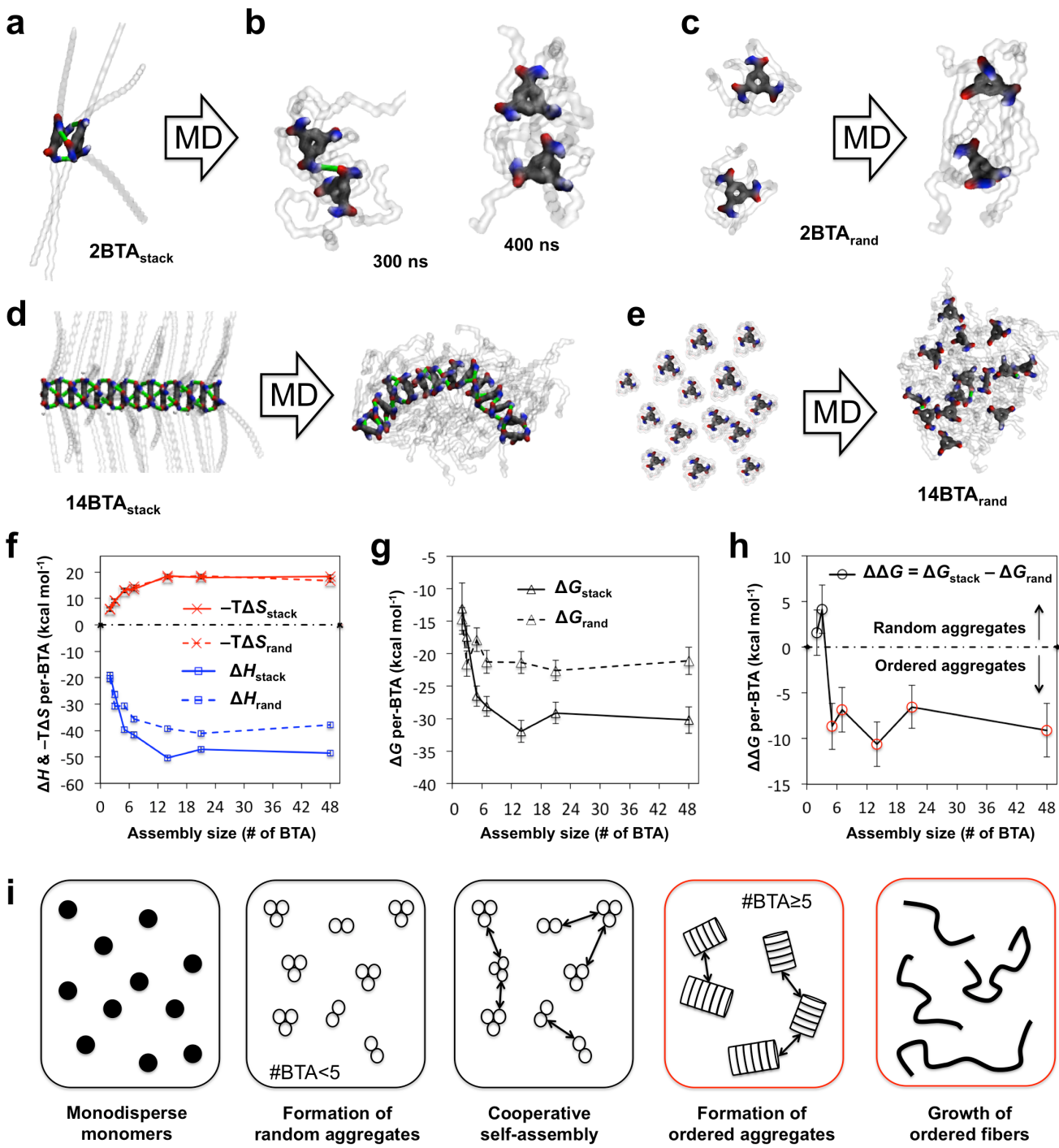
limitations prevented the unambiguous assignment of a cooperative self-assembly mechanism for the BTA described here.

Consistent with previous DFT and atomistic MD studies on BTA stacking in the gas-phase<sup>24,26-27</sup> and organic solvent (nonane),<sup>25,29</sup> we built atomistic models of pre-stacked ordered assemblies of a water-soluble BTA (Figure 1a) in which the size of the stacks was systematically increased. In particular, we built model stacks composed of 2, 3, 5, 7, 14 and 21 initially extended BTA monomers. This series was completed by including a previously reported "infinite" fiber model composed of 48 stacked BTAs replicated along the main axis with periodic boundary conditions.<sup>20</sup> These seven pre-stacked assemblies (**2BTA<sub>stack</sub>** to **48BTA<sub>stack</sub>** – Figure 1e) were equilibrated as immersed in a simulation box filled with TIP3P<sup>30</sup> explicit water molecules along 400 ns of MD simulation in NTP conditions using the AMBER 12 software.<sup>31</sup> During this time, all BTA assemblies rearranged and successfully reached the equilibrium in the MD regime (computational details are provided in the SI).

The MD simulations show that while larger BTA stacks bend, but remain stable (e.g., Figure 1f: **21BTA<sub>stack</sub>**; Figure 2d: **14BTA<sub>stack</sub>**), core stacking was found unstable in the smaller aggregates. Seen in Figures 2a,b for example, the initial stacking in the **2BTA<sub>stack</sub>** system disassembled early during the MD run. H-bonding between the two BTA monomers fails and a single H-bond only intermittently appears in the system (Figure 2b). In this case, the structural rearrangement and side chain folding due to hydrophobic collapse are evidently stronger than the H-bonding. Similar behavior is seen in the **3BTA<sub>stack</sub>** system (see SI). Increasing the size of the stacks was found to result in increased stability of the core-core stacking. In particular, BTA stacking is found stable in aggregates with sizes  $\geq$ **5BTA<sub>stack</sub>**, but only on a local basis (short-range), while globally BTA stacking was still found non-uniform and discontinuous. On the other hand, long, uniform and persistent BTA stacks are obtained for assemblies with size  $\geq$ **14BTA<sub>stack</sub>**.

Energetic and structural analyses of the MD simulations provide useful data to quantify these observations. From the equilibrated phase of each MD simulation (the last 100 ns) we calculated the self-assembly free-energies ( $\Delta G$ ) of each system as composed of an enthalpic and an entropic term:  $\Delta G = \Delta H - T\Delta S$ . All self-assembly  $\Delta G$  were calculated as the free-energy gain for  $n$  monomers to stay in an assembled state rather than disassembled in solution (see Methods section) – the more negative/favorable the  $\Delta G$ , the stronger the assembly (values are calculated per-BTA monomer in order to compare between different size assemblies).

The enthalpic ( $\Delta H$ ) term captures both non-bond solute-solute ( $\Delta E_{\text{gas}}$ ) and solute-solvent ( $\Delta G_{\text{solv}}$ ) interactions. Particularly interesting for this work,  $\Delta E_{\text{gas}}$  includes the van der Waals ( $\Delta E_{\text{vdW}}$ ) and electrostatic ( $\Delta E_{\text{ele}}$ ) non-bonding interactions between the BTA monomers. In general the assemblies modeled herein are characterized by favorable enthalpic variation ( $\Delta H < 0$ : enthalpy of the assembly is more favorable than that of the disassembled monomers, as the interactions between the BTAs are augmented in the assembly compared to a disperse solution), and an unfavorable entropic term ( $-T\Delta S > 0$ ), as the monomers lose degrees of freedom in the assembled compared to the disassembled state.



**Figure 2.** Self-assembly energies. (a,b) Starting (extended) and equilibrated structures taken from the MD simulation of **2BTAs<sub>stack</sub>** system. During the run the initial core stacking (a) disappears and a single H-bond (b) appears only intermittently between the BTAs. (c) Starting and final MD structure of **2BTAs<sub>rand</sub>** system. (d,e) Initial and equilibrated (400 ns) configurations of **14BTAs<sub>stack</sub>** (d) and **14BTAs<sub>rand</sub>** (e) systems. (f) Per-BTA  $\Delta H$  (blue) and  $-T\Delta S$  (red) data for ordered (**BTAs<sub>stack</sub>**: solid lines) and disordered BTA assemblies (**BTAs<sub>rand</sub>**: dotted lines) as a function of assembly size. (g) Per-BTA self-assembly free-energies ( $\Delta G$ ) for ordered (**BTAs<sub>stack</sub>**: solid line) and disordered BTA assemblies (**BTAs<sub>rand</sub>**: dotted line) as a function of assembly size. (h) Per-BTA  $\Delta\Delta G$  values. Negative  $\Delta\Delta G$  values indicate that formation of ordered BTA aggregates (stacks) is energetically favored over disordered (random) ones. (i) Scheme illustrating the mechanism of formation of ordered BTA supramolecular polymers in water deduced from the MD data.

Figure 2f,g (solid black, blue and red lines) shows the  $\Delta G_{\text{stack}}$ ,  $\Delta H_{\text{stack}}$  and  $-T\Delta S_{\text{stack}}$  collected values for the different size pre-stacked BTA assemblies. The energy data demon-

strate self-assembly cooperativity. In particular, the per-BTA self-assembly free-energy  $\Delta G_{\text{stack}}$ ,  $-13 \text{ kcal mol}^{-1}$  for the smaller **2BTAs<sub>stack</sub>** system, is seen to become rapidly more favorable

for increasing stack sizes. Starting from the **14BTA<sub>stack</sub>** system,  $\Delta G_{\text{stack}}$  converges to a value per-BTA of  $\approx 30 \text{ kcal mol}^{-1}$  (Figure 2g). Cooperativity is also reflected by the enthalpic and entropic terms reported in Figure 2f, where favorable  $\Delta H_{\text{stack}}$  and unfavorable  $-T\Delta S_{\text{stack}}$  terms respectively converge to  $\approx -50 \text{ kcal mol}^{-1}$  and  $\approx +20 \text{ kcal mol}^{-1}$  for sizes  $\geq \textbf{14BTA}_{\text{stack}}$ . Interestingly,  $\Delta G_{\text{stack}}$ ,  $\Delta H_{\text{stack}}$  and  $-T\Delta S_{\text{stack}}$  values for **14BTA<sub>stack</sub>** and **21BTA<sub>stack</sub>** are identical to those of **48BTA<sub>stack</sub>**, modeling an infinite BTA fiber. This means that above  $\approx 14$  BTA monomers the system is in “polymer bulk conditions”.

The enthalpic term ( $\Delta H$ ) contains the BTA-BTA interaction energies (van der Waals and electrostatic interactions, also including H-bonding) but also the interaction of the BTAs with the solvent (hydrophobic effects). While we learn from the  $\Delta H_{\text{stack}}$  plot in Figure 2f that globally these interactions become more favorable growing larger stacks up to **14BTA<sub>stack</sub>**, little can be said at this stage on the modulation of the fundamental interactions in the assemblies and on the role that these play on the growth of an ordered supramolecular BTA polymer in solution.

**Ordered vs. disordered BTA assemblies.** MD simulation of an individual BTA monomer in water solution shows that the long side chains fold around the BTA core to minimize the hydrophobic surface exposed to the solvent (Figure 1d).<sup>20</sup> To compare to the ordered BTA polymers, we systematically built additional BTA model systems in which multiple copies of individual and disassembled BTA monomers were randomly placed in a simulation box filled of explicit water molecules. In this way, we created molecular systems containing 2, 3, 5, 7, 14, 21 and 48 pre-equilibrated BTA monomers initially dispersed in solution (**2BTA<sub>rand</sub>** to **48BTA<sub>rand</sub>**). These systems also underwent 400 ns of MD simulation in NPT conditions. During this simulation time, the BTA monomers aggregated in solution forming chaotic/disordered BTA aggregates (Figures 1g, 2c and 2e) that remained stable during the MD runs (see SI).

Notably, the disordered BTA aggregates produced by the MD runs are local minimum energy conformations (in the real system these should reconfigure into ordered stacks) in which the systems are trapped due to the intrinsic sampling limitations of the MD technique. Nevertheless, these disordered aggregates offer an interesting “far-from-equilibrium” comparison with the ordered **BTA<sub>stack</sub>** systems. In fact, while in **BTA<sub>rand</sub>** systems the BTA monomers self-assemble mainly due to hydrophobic effects and non-directional non-bond interactions (e.g., van der Waals), no persistent H-bonding between the monomers is formed in these chaotic aggregates. On the other hand, in **BTA<sub>stack</sub>** systems H-bonding intrinsically plays a key role.

Analysis of the self-assembly energies extracted from the MD simulations of these systems provided us with the  $\Delta G_{\text{rand}}$ ,  $\Delta H_{\text{rand}}$  and  $-T\Delta S_{\text{rand}}$  data reported in Figure 2f,g (dotted black, blue and red lines). Traces of cooperative self-assembly are also seen in the case of disordered aggregates. This is reasonable, as some cooperativity can be expected due to the reduction in solvent-accessible surface areas.<sup>21b,32</sup> The self-assembly energy for the random aggregates ( $\Delta G_{\text{rand}}$ ), converging to  $\approx -22 \text{ kcal mol}^{-1}$  per-BTA (Figure 2g: dotted black line), is found less favorable than for the ordered assemblies ( $\approx -30 \text{ kcal mol}^{-1}$ ). This is largely due to a reduced  $\Delta H_{\text{rand}}$  term ( $\approx -40 \text{ kcal mol}^{-1}$  for aggregates larger than 14 BTAs) compared to that of

**BTA<sub>stack</sub>** systems ( $\approx -50 \text{ kcal mol}^{-1}$ ). Interestingly, the self-assembly entropic term ( $-T\Delta S$ ) is found invariant between ordered and disordered assemblies (Figure 2f: superimposed solid and dotted red lines), meaning that the BTA monomers lose the same number of degrees of freedom while self-assembling in ordered rather than disordered way. Here we find an interesting analogy with the component of self-assembly directly imputable to hydrophobic effects, generally thought of as to be due to entropy variations.<sup>33</sup> In fact, hydrophobic effects are reasonably the same upon formation of ordered or disordered BTA assemblies (see also below for additional evidences), consistent with a nearly identical self-assembly entropic term.

Comparison of the  $\Delta G_{\text{stack}}$  and  $\Delta G_{\text{rand}}$  data provides another important result. The difference between the self-assembly free-energies,  $\Delta\Delta G = \Delta G_{\text{stack}} - \Delta G_{\text{rand}}$ , indicates whether and to what extent formation of ordered BTA assemblies is energetically favored over that of disordered ones. Seen in Figure 2h,  $\Delta\Delta G$  is found  $> 0$  for smaller aggregates (number of BTA  $\leq 3$ : black circles), while it drops to  $\Delta\Delta G \approx -6.7\text{--}10 \text{ kcal mol}^{-1}$  for assembly sizes  $\geq 5$  BTAs (red circles). This interesting evidence suggests that below a certain critical size,  $\approx 5$  BTA based on our setup, formation of disordered BTA assemblies is favored over that of small ordered BTA stacks. On the other hand, above this threshold ( $\geq 5$  BTA) there is a rather constant free-energy gain, on average  $\Delta\Delta G = -8.4 \text{ kcal mol}^{-1}$  per-BTA monomer, favoring formation of stacked assemblies over disordered ones. Importantly, this effect is found to be entirely ascribed to enthalpic effects (Figure 2f). Figure 2i reports a scheme illustrating the growth mechanism of ordered BTA supramolecular polymers in water that can be deduced purely based on these thermodynamic observations, and regardless of the kinetic effects that, although important in the self-assembly mechanism, cannot be ascertained from such atomistic-resolution simulations: self-assembly of initially dispersed BTA monomers (i) induces formation of small disordered BTA aggregates (ii) that, when (iii) reaching the critical size ( $\approx 5$  BTA), evolve toward ordered oligomers (iv). Further cooperative self-assembly of these ordered oligomers results into growth of a supramolecular polymer (v).

This mechanism finds consistency with experimental observations. In fact, upon injection of the BTA monomers from methanol (molecularly dissolved) into water an initial UV spectrum is observed, which changes in time into the final spectrum obtained for the BTA polymers.<sup>17</sup> Also, recent temperature dependent studies on sugar-decorated BTA variants showed that at high temperature small aggregates are formed, which upon cooling are then converted into supramolecular polymers.<sup>23</sup> These examples support the general mechanism for polymer growth shown in Figure 2i.

**Key interactions in BTA self-assembly.** Seeing that the emergence and amplification of order in the BTA assemblies are controlled by enthalpic effects, and considering that  $\Delta H$  encompasses both solute-solute and solute-solvent interactions, immediate questions arising at this point are: Is order amplification in the polymer mainly controlled by hydrophobic effects or by BTA-BTA interactions? Is this mostly due to van der Waals or electrostatic interactions? What is the exact role of H-bonding in the growth of an ordered polymer?

The first step was to identify indicators that could be unambiguously ascribed to hydrophobic effects or to the different types of BTA-BTA interactions. A useful indicator of the strength of the hydrophobic effect is the solvent accessible

surface area (SASA) of the BTAs, and in particular the variation of the BTA SASA in the assembled or disassembled states. In fact, while aggregating in water the BTA monomers reduce the amount of surface exposed to the solvent (Figure 3a: SASA shrinkage). The SASA data reported in Figures 3b,c are extracted from the MD simulations of the smaller (Figure 3b: **2BTA**<sub>stack</sub> in red and **2BTA**<sub>rand</sub> in blue) and the larger BTA systems (Figure 3c: **48BTA**<sub>stack</sub> in red and **48BTA**<sub>rand</sub> in blue).

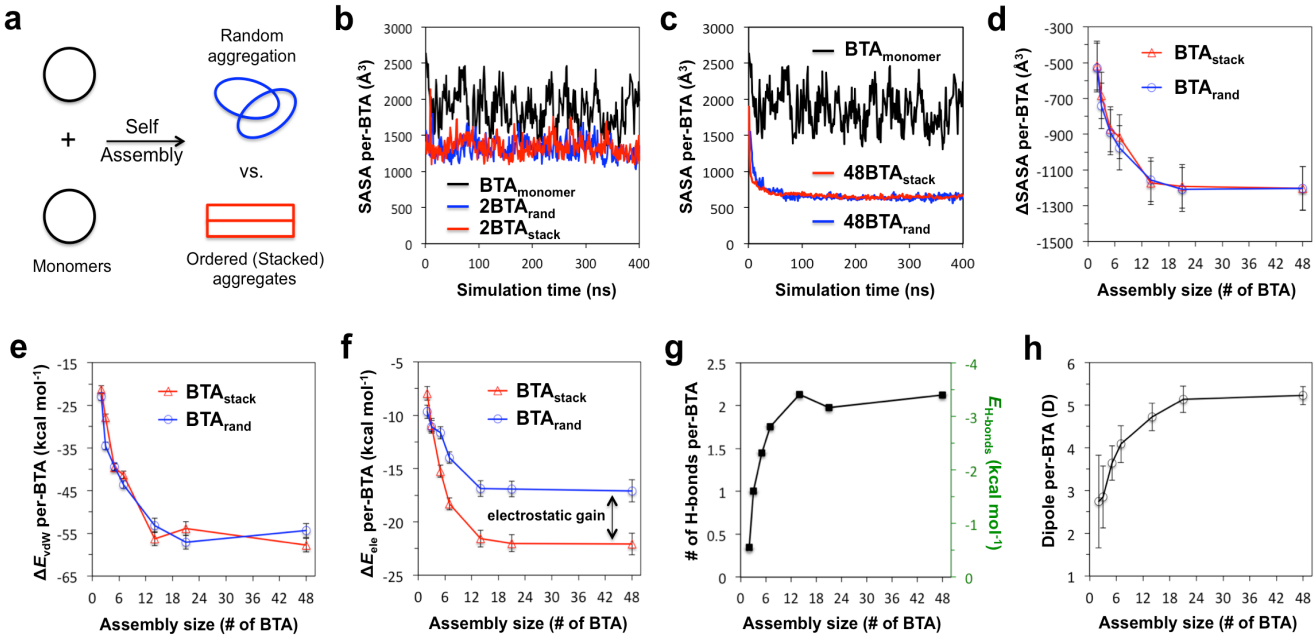
It is interesting to compare these data with the SASA of the disassembled BTA monomer (in black). It is evident that the BTAs reduce their SASA upon aggregation, and this effect is much stronger in the larger assemblies than in the smaller ones (SASA data for all simulated systems are provided in the SI). Shown in Figure 3d, the per-BTA SASA shrinkage accompanying self-assembly (calculated as:  $\Delta\text{SASA} = \text{SASA}_{\text{assembly}} - \text{SASA}_{\text{monomer}}$ ) is plotted as a function of the aggregate size for all simulated systems.  $\Delta\text{SASA}$  becomes more negative while increasing the size of the aggregate and converges for aggregate sizes  $\geq 14\text{BTA}$ , demonstrating the cooperativity expected for hydrophobic aggregation mentioned above. Interestingly, this analysis demonstrates that the  $\Delta\text{SASA}$  data (SASA shrinkage) is nearly identical in the ordered and disordered assemblies, which is also consistent with our conclusions on hydrophobic aggregation and entropy variations. These results suggest that hydrophobic and entropic effects taken alone are not responsible for preferential formation of ordered directional BTA aggregates *vs.* disordered random ones in solution, thus indicating that the origin for the growth of ordered su-

pramolecular polymers in water must be searched elsewhere.

Then we focused on the non-bond terms of the global solute-solute BTA interaction ( $\Delta E_{\text{gas}}$ ). Shown in Figure 3e, the van der Waals interactions ( $\Delta E_{\text{vdW}}$ ), while cooperative, do not discriminate between ordered and disordered assemblies, similar to entropic and hydrophobic terms. On the other hand, BTA-BTA electrostatic interactions ( $\Delta E_{\text{ele}}$ ) are augmented in the ordered (stacked) assemblies compared to disordered ones – by  $\approx 30\%$  for sizes  $\geq 14\text{BTA}$  (Figure 3f). This electrostatic gain ( $\Delta\Delta E_{\text{ele}}$ ) of  $\approx 5 \text{ kcal mol}^{-1}$  per-BTA captures most of the global  $\Delta\Delta G$  of Figure 2h, revealing the main energetic factor responsible for the growth of ordered BTA stacks in solution.

While  $\Delta\Delta E_{\text{ele}}$  identifies the electrostatic gain to ordered BTA assemblies, clearly accounting for the presence of the threefold H-bonding between the BTA cores, this also includes other factors such as a more favorable electrostatic environment generated by the initial ordered displacement of the BTA cores, amplification of dipole-dipole interactions and formation of macrodipoles.<sup>25,27,34</sup>

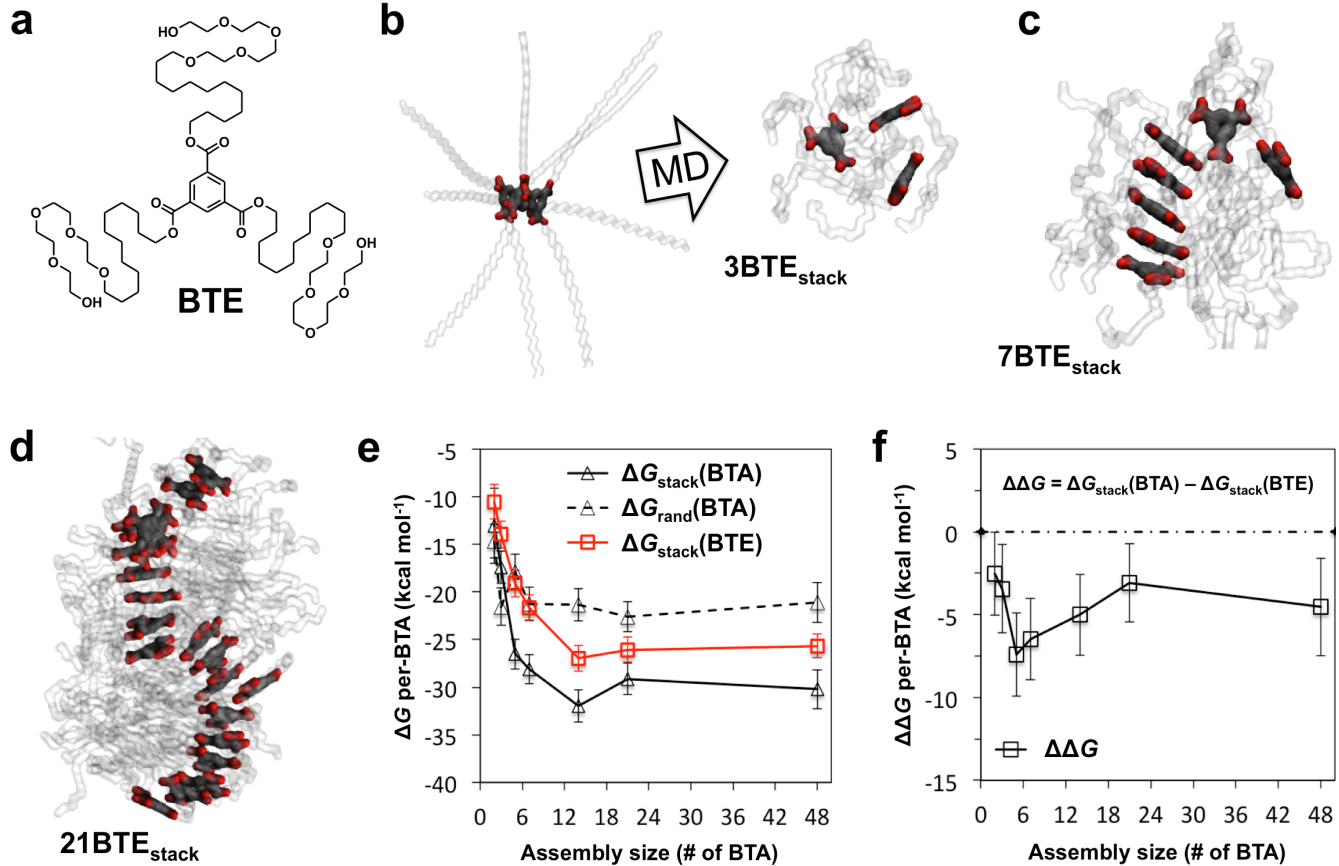
We calculated from the MD simulation of the stacked assemblies the average number of H-bonds per-BTA in the stacks and the associated H-bonding energy contribution ( $E_{\text{H-bonds}}$ ). As the atomistic force field used for this study does not contain an explicit term for H-bonding,  $E_{\text{H-bonds}}$  was estimated by multiplying the average number of H-bonds in each system for the energy per-single H-bond in aqueous solution for peptidic structures ( $\approx 1.58 \text{ kcal mol}^{-1}$ ).<sup>20,35</sup> Figure 3g shows that the average number of H-bonding per-BTA rapidly increases



**Figure 3.** Dissecting fundamental interactions in BTA self-assembly. (a) Conceptual scheme used to interpret hydrophobic aggregation. Dispersed BTA monomers aggregate hydrophobically to decrease the surface (black circles) exposed to the surface. (b,c) Two examples: average solvent accessible surface area (SASA) of the BTAs in (a) ordered **2BTA**<sub>stack</sub> (red) and disordered **2BTA**<sub>rand</sub> (blue), or (c) ordered **48BTA**<sub>stack</sub> (red) and disordered **48BTA**<sub>rand</sub> (blue) assemblies compared to the SASA of the BTA monomer (black). Analogous plots for all other size systems are in the SI. (d) SASA variation per BTA ( $\Delta\text{SASA}$ ) for all cases as a function of the size of the assembly. (e) Van der Waals interactions ( $\Delta E_{\text{vdW}}$ ) between the BTAs in the assemblies. As with  $\Delta\text{SASA}$  (d), also  $\Delta E_{\text{vdW}}$  data shows characteristic cooperativity while growing larger aggregates, and negligible difference between ordered (**BTA**<sub>stack</sub>: red) and disordered (**BTA**<sub>rand</sub>: blue) aggregates. (f) Electrostatic interactions ( $\Delta E_{\text{ele}}$ ) between the BTAs in the assemblies, showing clear differences between the two systems. (g) Average number of H-bonding per-BTA in the **BTA**<sub>stack</sub> assemblies (black axis) and related H-bonding energy estimated by the average energy per-single H-bond in aqueous solution for peptidic structures ( $\approx 1.58 \text{ kcal mol}^{-1}$ ).<sup>20,35</sup> (g) Amplification of the dipole moment of the BTAs as a function of the size of the assembly calculated for **BTA**<sub>stack</sub> systems.

from  $\approx 0.3$ – $1$  in the case of the smaller and unstable stacks ( $2\text{BTA}_{\text{stack}}$  and  $3\text{BTA}_{\text{stack}}$ ) to  $\approx 2.2$  for stacks sizes  $\geq 14\text{BTA}_{\text{stack}}$ . Consistently, in the latter cases the approximated H-bonding energy contribution was found to reach a maximum of  $\approx 3.4$  kcal mol $^{-1}$  per-BTA, equal to  $\approx 70\%$  of the global electrostatic gain  $\Delta E_{\text{ele}}$ .

for full synthetic information). Compared to the BTA monomers studied herein, BTE monomers lack the ability to form a H-bonding network between the cores in the supramolecular polymer while preserving the same hydrophobicity and structure of the cores. We compared the assemblies formed by these monomer variants in solution by means of MD simula-



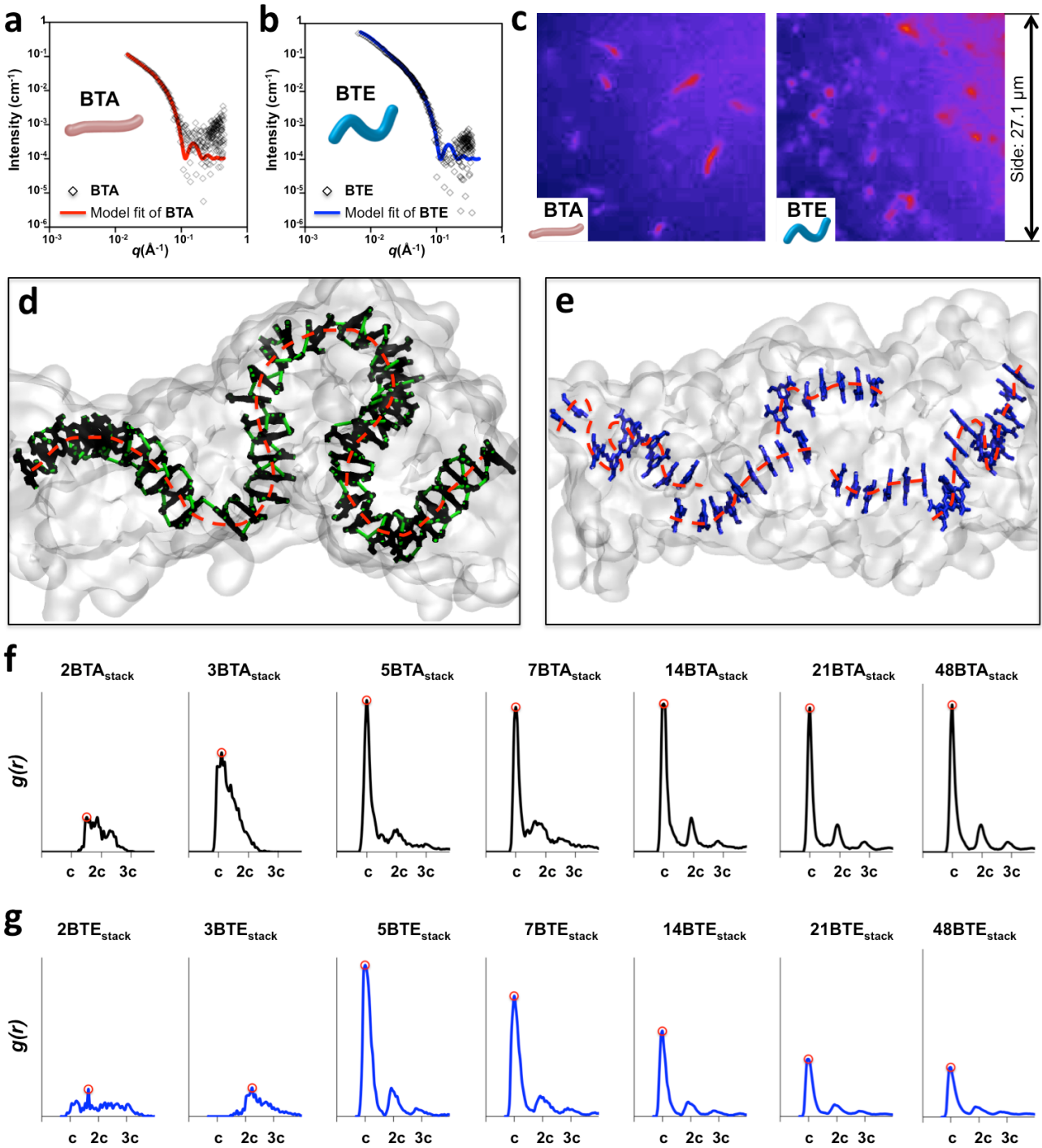
**Figure 4.** Modeling BTE stacks. (a) Molecular structure of the BTE monomer. (b) Starting (extended) and equilibrated structures taken from the MD simulation of  $3\text{BTE}_{\text{stack}}$  system. (c,d) Equilibrated structures obtained from the MD simulations of  $7\text{BTE}_{\text{stack}}$  (c) and  $21\text{BTE}_{\text{stack}}$  (d) models. (e) Per-monomer self-assembly free-energies ( $\Delta G$ ) for the different size simulated  $\text{BTE}_{\text{stack}}$  assemblies (red).  $\text{BTA}_{\text{stack}}$  (solid black line)  $\text{BTA}_{\text{rand}}$  (dotted black lines)  $\Delta G$  data are provided for comparison. (f) Per-BTA difference ( $\Delta \Delta G$ ) between the self-assembly free-energies of  $\text{BTA}_{\text{stack}}$  ( $\Delta G_{\text{stack}}(\text{BTA})$ ) and  $\text{BTE}_{\text{stack}}$  ( $\Delta G_{\text{stack}}(\text{BTE})$ ) assemblies of various sizes. Negative  $\Delta \Delta G$  values indicate the favorable contribution to the stacked assemblies brought by H-bonding.

From the MD simulations we also calculated the average dipole moment of the BTAs in the stacks. Consistent with the H-bonding data and with previous simulations of smaller BTA derivatives in an apolar solvent,<sup>25,27</sup> the BTA dipole moment is seen to be amplified while growing with the stack size until a plateau is reached from  $14$ – $21\text{BTA}_{\text{stack}}$  (Figure 3h). While these analyses clearly indicate the amplification of electrostatic interactions along the stacks as the main factor leading to the growth of ordered BTA supramolecular polymers in water, we performed a last step to unambiguously isolate the effect of H-bonding in the process.

**The effect of H-bonding.** We synthesized and investigated a 1,3,5-benzenetriester (BTE) derivative identical to BTA but having the amides replaced by ester groups (Figure 4a, see SI

tions and experiments).

We built molecular models for BTE stacks ( $\text{BTE}_{\text{stack}}$ ) of analogous sizes to those created and simulated for the BTA case ( $\text{BTA}_{\text{stack}}$ ). In this way, we obtained seven initially extended models for BTE stacks from  $2\text{BTE}_{\text{stack}}$  to  $48\text{BTE}_{\text{stack}}$ . Analogous to  $48\text{BTA}_{\text{stack}}$ , the larger  $48\text{BTE}_{\text{stack}}$  system was also built to model an infinite BTE polymer through periodic replication of the 48 initially extended BTE monomers along the main axis of the fiber. The force field parameters for the BTE monomers were obtained according to the same procedure previously used for the BTA variants<sup>20</sup> and other similar three-branched stacking discotic units (see SI for details).<sup>36</sup> Each BTE stack model was simulated for 400 ns of NPT MD in explicit water molecules as done for the  $\text{BTA}_{\text{stack}}$  systems.



**Figure 5.** BTE vs. BTA supramolecular polymers. (a,b) Experimental SAXS profiles of (a) BTA (0.45 wt% in H<sub>2</sub>O) and (b) BTE (0.45 wt% in H<sub>2</sub>O) assemblies, fit with the Schurtenberger–Pedersen form factor. (c) Fluorescence microscopy images of BTA and BTE assemblies from 10  $\mu\text{M}$  aqueous solutions diluted to 0.5  $\mu\text{M}$  for imaging. (d,e) Detail of BTA core stacking taken from the MD simulation of the  $48\text{BTA}_{\text{stack}}$  system. BTA cores are colored in black, H-bonding in green and the fiber is represented as transparent surface. (d) Detail of BTA core stacking taken from the MD simulation of the  $48\text{BTE}_{\text{stack}}$  system. BTE cores are colored in blue. Red dotted lines are provided to guide the eye. (f,g) Radial distribution functions  $g(r)$  of the BTA (f: black) and BTE (g: blue) cores along the fibres calculated from the equilibrated phase MD trajectories (the last 100 ns of each MD run). Intercore spacing  $c$  equals to 3.4  $\text{\AA}$ . The  $g(r)$  peaks indicative of stacking between neighbor BTA cores –  $g(c)$  – are identified by red circles.

Initially, we guessed that the lack of H-bonding would have made BTE monomers produce only random non-directional aggregates in water, and that this would have been reflected by

strong instability in the BTE stacks during the MD simulations. However, while complete stacking destabilization was seen for the smaller BTE assemblies ( $\leq 3\text{BTE}_{\text{stack}}$ : Figure 4b),

larger BTE stacks ( $\geq 5\text{BTE}_{\text{stack}}$ ) were found to possess a certain level of intrinsic stability during the MD simulations, albeit to a lower extent compared to  $\text{BTA}_{\text{stack}}$  systems. For example, it is interesting to note in Figures 4c,d that while core-core stacking is present in  $7\text{BTE}_{\text{stack}}$  and  $21\text{BTE}_{\text{stack}}$  systems, this seems to persist well only at short distance (stable stacks of  $\approx 5$  BTEs), while defects appear in various spots along larger stacks. Due to the lack of the H-bonding network, stack bending during the MD simulations produces stacking instabilities and local slipping of the BTE cores respect to each other.

As previously done for the BTA case, we calculated from the MD simulations the self-assembly free-energies for the  $\text{BTE}_{\text{stack}}$  systems. The plot of the  $\Delta G_{\text{stack}}(\text{BTE})$ , in red in Figure 4e, is shown to be nearly identical to that of the random BTA assemblies ( $\text{BTA}_{\text{rand}}$ ) for sizes  $\leq 7$  monomers. Above this limit ( $\geq 14\text{BTE}_{\text{stack}}$ ),  $\Delta G_{\text{stack}}(\text{BTE})$  is found lying in between the plots of  $\Delta G_{\text{rand}}(\text{BTA})$  and  $\Delta G_{\text{stack}}(\text{BTA})$ . This is reasonable, as  $\text{BTE}_{\text{stack}}$  stacks certainly lack the contribution of H-bonding compared to  $\text{BTA}_{\text{stack}}$  assemblies, but compared to  $\text{BTA}_{\text{rand}}$  systems these benefit from a better arrangement of the cores (favorable electrostatic environment). This result indicates that formation of ordered assemblies is probable also in the case of BTE.

The difference ( $\Delta\Delta G$ ) between the self-assembly free-energies of  $\text{BTA}_{\text{stack}}$  and  $\text{BTE}_{\text{stack}}$  systems is found to converge to  $\approx -4$  kcal mol $^{-1}$  on average for stacks greater than 14 monomers (Figure 4f), well compatible with the contribution of H-bonding ( $E_{\text{H-bonds}} \approx -3.4$  kcal mol $^{-1}$ ) found for the BTA stacks (*vide supra*, Figure 3g). Thus, the difference in free-energy between  $\text{BTA}_{\text{stack}}$  and  $\text{BTE}_{\text{stack}}$  systems can be imputed in good approximation to the effect of H-bonding. On the other hand, the difference in free-energy between  $\text{BTE}_{\text{stack}}$  and  $\text{BTA}_{\text{random}}$  systems ( $\Delta\Delta G$  also  $\approx -4$  kcal mol $^{-1}$ ), both lacking H-bonding, can be ascribed to the beneficial effect of order into the core assembly (stacking), which makes the formation of BTE ordered assemblies favored over disordered ones also in the absence of H-bonding.

Experimentally, BTA and BTE self-assembly were compared in the milli- to micromolar regime using small-angle X-ray scattering (SAXS) and fluorescence microscopy experiments (Figure 5a–c). Both BTA and BTE variants form directional fibers in aqueous solution, yet the BTE fibers are found to be less rigid than the BTA.

The SAXS profiles obtained for BTA and BTE fibers (Figure 5a,b) are both found to fit well with a form factor describing semi-flexible rods (Pedersen and Schurtenberger worm-like chain model).<sup>37</sup> This model yields a cross-sectional radius of  $\approx 3.3$  nm for both fibers, while their persistence lengths differ considerably. The BTE fibers exhibit a relatively small Kuhn length ( $\approx 8$  nm), while we obtain  $\approx 27$  nm for BTA fibers, indicating that the BTE fibers are significantly more flexible than BTA fibers. This is further confirmed by the observation that BTA fibers can also be reliably fit by a rigid rod model, indicating a persistence length on the order of the experimental resolution ( $\approx \pi/q_{\min} = 45$  nm, see SI for details).

The differences in persistence length shown via SAXS are further supported by fluorescence microscopy. While previous fluorescence imaging of BTA fibers relied on (dynamic) covalent attachment of dyes,<sup>17–18</sup> here we used Nile Red to label the hydrophobic interior of the assembled supramolecular systems in a non-covalent fashion.<sup>23</sup> Time-lapse imaging (see SI for the

movies) shows that BTE self-assembly produces less stiff, less ordered and more polydisperse fibrous aggregates. Attempting to illustrate this via a snapshot (Figure 5c) one can see long, stiff and defined fibers of the BTA system on the left in comparison to the poorly defined BTE system on the right. Surprisingly, during the BTE imaging several long ( $\mu\text{m}$ ) and highly flexible fibers passed through the field of view; however, the majority of the sample was dominated by short and ill-defined aggregates.

The BTE self-assembly was also investigated utilizing UV-vis spectroscopy, dynamic light scattering (DLS), and cryo-TEM. The UV-vis showed little change upon self-assembly, the DLS experiments showed a mono-modal distribution, while structures could not be visualized in cryo-TEM. This data and further discussion are reserved for the SI, but is attributed to the non-uniform self-assembly of the BTE derivative.

Taken altogether, these data reveal that while formation of directional assemblies and supramolecular polymers in water is possible also in the absence of H-bonding, the latter has noticeable effect on the persistency and rigidity of the supramolecular polymer.

Additional analysis of the  $\text{BTA}_{\text{stack}}$  vs.  $\text{BTE}_{\text{stack}}$  molecular models also support the above conclusion. The radial distribution functions ( $g(r)$ ) of the BTA and BTE cores extracted from the equilibrated phase MD simulations (the last 100 ns of each run) are useful indicators of the levels of order in monomer stacking.<sup>20,36</sup> As the  $g(r)$  measures the relative probability of finding neighbor BTA cores at stacking distance ( $c$ : closest neighbor,  $2c$ : second neighbor, etc.), the relative height of the  $g(r)$  plots at distance  $c$ ,  $2c$ ,  $3c$ , etc. provides interesting insight on the amplification of order in the assemblies with increasing aggregate size. In general, the higher the  $g(r)$  peaks at stacking distances  $c$ ,  $2c$ ,  $3c$ , the more ordered, stable and persistent the core stacking in the assembly.

Shown in Figure 5f, the  $g(r)$  plots of the BTA cores present the characteristic three peaks  $g(c)$ ,  $g(2c)$  and  $g(3c)$  (red circles) typical of ordered stacking for aggregate size  $\geq 14\text{BTA}_{\text{stack}}$ . Looking at the smaller BTA aggregates, absence of the  $g(c)$  and  $g(2c)$  peaks indicates that no ordered stacking is present in  $2\text{BTA}_{\text{stack}}$  and  $3\text{BTA}_{\text{stack}}$ , consistent with the fact that these small stacks are seen to disassemble during the MD simulations. Starting from  $5\text{BTA}_{\text{stack}}$  the first  $g(c)$  peak increases to a maximum level that is conserved also in the larger aggregates. It is interesting to note that the  $g(r)$  plots of  $5\text{BTA}_{\text{stack}}$  and  $7\text{BTA}_{\text{stack}}$  possess clear first  $g(c)$  peak, but no clear  $g(2c)$  and  $g(3c)$  peaks, indicating that these two systems possess short-range stacking order, but still not a long-range one. Based on our setup, these results demonstrate that to see real order amplification the BTA stacks need to grow above 14 BTA monomers. Above this limit, consistency of the  $g(r)$  plots and of  $g(c)$ ,  $g(2c)$  and  $g(3c)$  peaks (red circles) in  $14\text{BTA}_{\text{stack}}$ ,  $21\text{BTA}_{\text{stack}}$  and  $48\text{BTA}_{\text{stack}}$  systems demonstrates that order is uniformly amplified along the fiber for stack sizes  $\geq 14\text{BTE}_{\text{stack}}$ .

Interestingly, the same analysis for the  $\text{BTE}_{\text{stack}}$  systems (Figure 5g) shows that while for assembly sizes  $\leq 5\text{BTE}_{\text{stack}}$  the  $g(r)$  plots are in good approximation consistent with those of  $\text{BTA}_{\text{stack}}$  ones, starting from sizes  $\geq 7\text{BTE}_{\text{stack}}$  the heights of the  $g(r)$  plots drop dramatically. This observation is fully consistent with the experimental data, indicating that stacking persistency into BTE fibers is reduced compared to BTA ones. As

seen in Figure 4c,d, BTE stacks seem to preserve a short-range order ( $\approx$ 5 BTE) imputable to the above mentioned order effect, but failing in real long-range order amplification. Figures 5d,e provide visual inspection inside the **48BTA<sub>stack</sub>** and **48BTE<sub>stack</sub>** simulated systems, showing that while core stacking is uniform inside the BTA polymer, the BTE core stacking breaks into shorter segments. This effect is consistent with the enhanced flexibility seen in the experiments for BTE assemblies compared to BTA ones, and is directly imputable to the lack of H-bonding in the BTE assembly.

## CONCLUSIONS

Focusing on 1,3,5-benzenetricarboxamide (BTA), a self-assembling motif forming supramolecular polymers in water, we have devised a computational approach to study in detail the factors triggering and controlling the formation of an ordered assembly in aqueous environment, where an intricate interplay of hydrophobic and monomer-monomer non-bond interactions takes place. Systematic comparison of ordered (stacked) and disordered BTA assemblies of the same size (number of monomers) indicate that the aggregates need to reach a critical size to favor the formation of ordered stacks over disordered aggregates. Detailed decomposition of the self-assembly energies demonstrates that the amplification of order in the assemblies and consequent growth of an ordered supramolecular polymers in water are exclusively due to electrostatic effects. In order to isolate the effect of H-bonding, we compared the BTA assemblies with those formed by a 1,3,5-benzenetricarboxyester (BTE) variant by means of experiments and MD simulations. BTE monomers differ only by having the amide groups replaced by esters, thus lacking the ability to form the H-bonding network between the monomers. Nevertheless, these were also found to form fibrous assemblies in water, but BTE fibers were found more flexible and less persistent than BTA supramolecular polymers. This work provides a unique picture of the direct role of H-bonding in the self-assembly of supramolecular polymers in water. Our results unambiguously demonstrate that order amplification in the fibers and the growth of a persistent directional supramolecular polymer in water solution is greatly controlled by H-bonding. This approach allows directly relating the structure of the monomer to the structure and properties of the supramolecular polymer that these form in water.

## METHODS

**Computational energetic analysis.** The molecular model for the water-soluble BTA studied herein was taken from our previous work.<sup>18</sup> The water-soluble BTE model was built and parametrized accordingly (see SI). The pre-stacked (**BTA<sub>stack</sub>** and **BTE<sub>stack</sub>**) and the random (**BTA<sub>rand</sub>**) assemblies were built as explained above. All assembled systems, as well as the disassembled monomers, were equilibrated for 400 ns of MD in periodic boundary NPT condition in explicit TIP3P<sup>28</sup> water molecules at the experimental temperature of 20°C and 1 atm of pressure. The last 100 ns of each MD were chosen as representative of the equilibrium condition for each system in the MD regime (see SI for simulation details) and used for energetic analysis.

All energy variations ( $\Delta E$ ) reported herein (e.g.,  $\Delta G$ ,  $\Delta H$ ,  $\Delta S$ ,  $\Delta E_{ele}$ ,  $\Delta E_{vdw}$ , etc.) are calculated as:

$$\Delta E = E_n - E_{monomer} \quad (1)$$

where  $E_n$  is the average energy of the BTA monomers in the different size aggregates (i.e., the energy of the  $n$ th assembly divided by the

number  $n$  of BTA monomers in the aggregate) and  $E_{monomer}$  is the energy of the disassembled monomer in solution. For example, the global self-assembly free-energy,  $\Delta G = G_n - G_{monomer}$ , of **21BTA<sub>stack</sub>** measures the per-monomer  $\Delta G$  gain for the BTAs to stay stacked into a 21-mer rather than disassembled in solution.

According to this scheme, the  $\Delta G$  for the various simulated assemblies were calculated with the MM-PBSA approach<sup>38</sup> as:

$$\Delta G = \Delta H - T\Delta S \quad (2)$$

where  $\Delta H$  and  $-T\Delta S$  are respectively the enthalpic and the entropic terms of the free-energy  $\Delta G$  of the various assemblies, respectively providing information on the per-monomer enthalpic and entropic gain (or penalty) for  $n$  monomers to stay assembled (in stacked/ordered or random/disordered configuration) rather than disassembled in solution.

The  $\Delta H$  can be calculated as the sum of the molecular mechanics solute-solute interactions and the solvation energy of the systems as in Eq. 3:

$$\Delta H = \Delta E_{gas} + \Delta G_{solv} \quad (3)$$

$$\Delta E_{gas} = \Delta E_{internal} + \Delta E_{ele} + \Delta E_{vdW} \quad (4)$$

$$\Delta G_{solv} = \Delta G_{PB} + \Delta G_{SA} \quad (5)$$

$\Delta E_{gas}$  is the gas-phase (in vacuum) molecular mechanical energy provided by the force field, which includes, together with the internal energy  $\Delta E_{internal}$  (sum of bond, angle and torsion terms), the electrostatic ( $\Delta E_{ele}$ ) and van der Waals ( $\Delta E_{vdW}$ ) non-bond interactions (Eq. 4).  $\Delta G_{solv}$  is the total solvation energy (Eq. 5), which is the sum of a polar term ( $\Delta G_{PB}$ : calculated according to the Poisson-Boltzmann<sup>39</sup> approach) and a non-polar solvation term ( $\Delta G_{SA}$ : related to the reduction of the SASA due to solvophobic effects). All energies, including the entropic terms of the free-energies ( $-T\Delta S$ ) were calculated using the *MMPSA.py* module of AMBER 12.<sup>40</sup> Additional details on the computational procedures are provided in the SI.

## ASSOCIATED CONTENT

### Supporting Information

The Supporting Information is available free of charge on the ACS Publications website.

Detail of the simulated molecular systems, simulation and data analyses procedures, and additional material from the MD simulations; synthesis procedure for BTE monomers, SAXS and fluorescent microscopy methods are provided as Supporting Material (PDF).

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### Author Contributions

G.M.P., M.B.B., A.R.A.P. and E.W.M. conceived the project and contributed to the concept of the manuscript. G.M.P. designed the computational strategy. M.G. and G.M.P. performed the MD simulations and analyzed the computational results. M.B.B., C.M.A.L., I.K.V. and L.A. performed the synthesis of the BTE monomers and the experiments. G.M.P. and M.B.B. wrote the

manuscript and managed the project. All authors have given approval to the final version of the manuscript.

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