Temperature Control of Sequential Nucleation-Growth Mechanisms in Hierarchical Supramolecular Polymers

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Abstract: Upon cooling in solution, chiral triarylamine tris-amide unimers produce organogels by stacking into helical supramolecular polymers, which subsequently bundle into larger fibers. Interestingly, circular dichroism, vibrational circular dichroism, and AFM imaging of the chiral self-assemblies revealed that monomolecular $P$-helical fibrils formed upon fast cooling, whereas bundled $M$-superhelical fibers formed upon slow cooling. The mechanistic study of this structural bifurcation reveals the presence of a strong memory effect, reminiscent of a complex stepwise combination of primary and secondary nucleation-growth processes. These results highlight the instrumental role of sequential self-assembly processes to control supramolecular architectures of multiple hierarchical order.

Introduction

Supramolecular polymers[1–8] which involve stacking of π-conjugated molecules are of particular interest to design new materials for organic electronics[9–10] and photonics.[11] Beyond their molecular structure, the functional properties and efficiencies of such soft materials in devices are strongly dependent on their morphologies at all scales.[12–14] Therefore, the fundamental understanding of supramolecular polymerization mechanisms is of critical importance to get control over the structuring processes occurring from molecular level to mesoscale. Efforts from various research groups worldwide are currently showing that supramolecular polymerization can go much further than the simplest isodesmic growth regime, and that it can make use of a number of more complex kinetic pathways for giving rise to highly cooperative processes and thermal hystereses.[15–18] For instance, living supramolecular polymerization[19–22] and access to supramolecular block copolymers[23–27] have been recently demonstrated by precisely engineering molecular structures of monomers and by tuning experimental conditions. In addition, the presence of competing kinetic and thermodynamic self-assembly pathways has been shown to potentially bias the expression of supramolecular chirality of unidimensional helical polymers.[28–30] For instance, Meijer and co-workers have shown the possibility to invert the absolute configurations of single helical fibrils made of oligo(p-phenylenevinylene) by acting on the primary nucleation step and the subsequent primary growth of single polymer chains.[31] Up to now, these observations have been limited to unidimensional supramolecular polymers, although one may expect that such pathway complexity could also affect higher levels of hierarchical organization comprising multiple nucleation steps.

We became interested in approaching such systems by using supramolecular polymers made of $C_7$-symmetric triarylamine tris-amides (TATAs).[32] They were recently shown to present helical stacked structures with a possible hierarchical bundling of their primary fibrils up to the formation of organogels.[33,34] From a functional point of view, this new family of supramolecular polymers[35–38] presents unusual electronic,[39–41] photonic,[42] and plasmonic properties.[43,44] In addition, TATA-based supramolecular polymers display high cooperativity in the expression of their chirality at the level of the single fibrils. For instance, unusually low amounts of enantiopture TATAs can drive the homochiral twisting of monocolumnar helices by a sergeant and soldier mechanism.[45] Even circularly polarized light was reported to switch the helicity of fibrils made of non-chiral TATAs.[46] By using chiral side chains to produce 1) enantiopture TATAs,[47] 2) enantiopture TATAs having reversed amide orientations,[48,49] or 3) enantiopture carbonyl-bridged TATAs,[50] it was also demonstrated that temperature or solvent can induce an inversion of helicity within the corresponding monocolumnar supramolecular polymers thanks to an intermolecular rearrangement of the hydrogen-bond network.

In the present work, we investigate for the first time the hierarchical supramolecular polymerization mechanism of $C_7$-symmetric TATAs organogelators decorated either with chiral...
alkyl chains [(S)-TATA] or with achiral ones [(C₉)-TATA and (C₁₄)-TATA] (Figure 1). By a combination of circular dichroism (CD), vibrational circular dichroism (VCD), transmission and scanning electron microscopies (TEM, SEM), and atomic force microscopy (AFM), we show that, intriguingly, the hierarchical supramolecular structure and chirality of the (S)-TATA fibers can be modified by simply controlling the cooling rate of the initial solutions, because of a stepwise combination of primary and secondary nucleation-growth mechanism.

Results and Discussion

Synthesis and characterization of TATA-based supramolecular polymers

Compounds (S)-TATA, (C₉)-TATA and (C₁₄)-TATA were obtained in a few synthetic steps with high yields by using straightforward procedures from the literature, and they were characterized by ¹H, ¹³C NMR spectroscopies and ESI-MS (see Supporting Information, section S1.2). These three compounds were found to spontaneously self-assemble in a variety of organic solvents, such as chloroform, toluene, and acetonitrile. A first indication of their self-assembly was revealed by the lack of all ¹H NMR resonance signals from the triarylamine core at room temperature, and their further appearance at higher temperatures. For instance, in [D₈]toluene, upon gradual heating (from 25 to 105 °C) of a 5 mM solution, the recovery of enantiopure (S)-TATA resonance signals was observed, suggesting a transition from the assembled to the molecularly dissolved states (see Supporting Information, Figure S1). Subsequent cooling of this solution to room temperature resulted in the formation of an opaque gel with again the disappearance of the ¹H NMR resonance signals. The onset gelation concentration (OGC) for (S)-TATA in toluene was measured around 0.56 g L⁻¹ (0.75 mM, Figure S2). The formation of thermoresponsive physical organogels was also observed in other aromatic solvents as well as in chlorinated solvents, but at much higher concentrations (OGC of 2.8 mM and 9.0 mM in dichloroethane and dichloromethane, respectively), whereas no gelation occurred in polar solvents such as DMSO, DFM, THF or methanol/toluene mixture even at high concentrations (Supporting Information, section S2.2, Table S1). Compounds (C₉)-TATA and (C₁₄)-TATA were also found to form thermoresponsive gels in toluene at OGC similar to (S)-TATA (0.70 and 0.72 mM respectively).

We then probed the structures of these gels by polarized optical microscopy (POM) at concentrations of 0.1 and 1 mM in toluene (Figures 2A and Figure S3 in the Supporting Information). For both of these concentration regimes, that is, below and above the OGCs of the three TATA molecules, we observed large fibers up to several hundreds of micrometers in length. At 1 mM, compared to (S)-TATA, fibers made of (C₉)-TATA...
appear shorter and more rigid, whereas those made of \((\text{C}_{14} \text{~TATA})\) appear thinner and shorter (Figure S3, Supporting Information). The morphology of the fibers was then studied by TEM (Figures 2B and Figure S4 in the Supporting Information) and SEM (Figures 2C and Figure S5 in the Supporting Information). Micrographs show that TATA-based polymers self-assemble into bundles of fibers up to several micrometers in length and up to hundreds of nanometers in diameter. The internal organization of TATA molecules within these self-assemblies was first studied by Fourier transform infrared (FTIR) spectroscopy (Supporting Information, section S2.4). This technique has been widely used to identify and differentiate molecular packing through different types of hydrogen bonds in supramolecular polymers.\(^{33, 48, 51–53}\) FTIR spectra of \((\text{S})\)-TATA, \((\text{C}_9 \text{~TATA})\), and \((\text{C}_{14} \text{~TATA})\) in both the solid state after the synthesis and after self-assembly from a freeze-dried solution (at an initial concentration of 1 mM in toluene) show well-defined bands at around 3290 and 1653 cm\(^{-1}\) corresponding to the stretching vibrations of the \(\text{N}–\text{H}\) and \(\text{C}–\text{O}\) bonds, respectively (Figures 2D and Figure S7 in the Supporting Information). These observations clearly indicate the presence of intermolecular hydrogen bonds between molecules as compared with the frequencies of the free \(\text{N}–\text{H}\) and \(\text{C}–\text{O}\) stretching vibrations (3500–3400 and \(\approx 1685\) cm\(^{-1}\) respectively).\(^{34}\) The high symmetry of the amide and carbonyl stretching vibrations also suggests the existence of a single type of hydrogen bonds involving a regular columnar packing of the TATAs.\(^{51}\) In addition, taking into account the MM2 geometry optimization model of \((\text{S})\)-TATA pentamer (Figure S8, Supporting Information) and our previously reported DFT calculations,\(^{33}\) we assigned these frequencies to the formation of threefold helical intermolecular hydrogen bonding between neighboring TATA molecules within a single columnar stack. AFM imaging of \((\text{S})\)-TATA at low concentrations (0.1 mM in toluene) confirmed the formation of thin fibrils with a minimal diameter of 1.5 nm that corresponds to the diameter of an individual columnar stack (Figure 2F).\(^{13, 34}\)

The optical properties of all TATA molecules were then studied by UV/Vis and fluorescence spectroscopies (Supporting Information, section S2.5). The general trends observed are: 1) solvatochromism, that is, bathochromic shift of absorption and emission maxima with solvent polarity (Figure S10A, Supporting Information); 2) temperature-independent emission and absorption spectra at concentrations of TATA below which self-assembly occurs (i.e., 0.01 mM in toluene; Figures S10 and S17, Supporting Information); 3) at room temperature and at higher concentrations (i.e., at 1 mM when self-assembly occurs in toluene) shifting of the absorption and emission maxima towards lower wavelengths together with a decrease of fluorescence intensity and an increase in absorption intensity (Figure S9, Table S2, Supporting Information). Such hypsochromic shifts result from \(\pi\)–\(\pi\) stacking interactions between the aromatic rings of TATA molecules. Compound \((\text{S})\)-TATA was further characterized by circular dichroism in order to provide information on the chirality of its supramolecular structure.\(^{31, 55–57}\) All CD spectra were recorded after a fast cooling of hot solutions of \((\text{S})\)-TATA to room temperature and in different solvents at a concentration of 0.5 mM (Figure S18, Supporting Information). The formation of chiral aggregates was demonstrated in solvents such as toluene, xylene, and chlorobenzene.

Having characterized the supramolecular structures of these self-assemblies, we further investigated their mechanism of supramolecular polymerization by using different temperature-dependent spectroscopic techniques and mainly focusing on \((\text{S})\)-TATA.

**Study of the supramolecular polymerization mechanism**

Temperature-dependent UV/Vis spectroscopy measurements were performed to investigate the self-assembly mechanism of TATA molecules (see details in the Supporting Information, section S2.5.1).\(^{33}\) Figure 3A shows UV/Vis spectra of a 0.5 mM solution of \((\text{S})\)-TATA in toluene when decreasing temperature from 338 to 283 K (and with increments of \(-5\) K). A decrease in absorption intensity is observed together with a hypsochromic shift of the absorption maximum from 322 to 310 nm. These changes indicate a transition from the molecularly dissolved state to the aggregated state \((\text{S})\)-TATA\(_{agg}\). The corresponding cooling trace (Figure 3B) shows a decrease of the absorption intensity measured at 315 nm with temperature. The non-sigmoidal character of the curve, together with the presence of two temperature regimes, clearly suggest a cooperative aggregation mechanism for the self-assembly process.\(^{58}\) Melting curves \(\alpha_{agg} = r(T)\) were also obtained by recording the absorption intensity at 315 nm during the cooling and heating processes at a rate of 1 K min\(^{-1}\) for concentrations going from 0.1 to 1 mM (i.e., below and above the OGC (0.75 mM), Figure S13 in the Supporting Information). These experiments reveal for all concentrations the existence of a thermal hysteresis characterized by a difference between the critical elongation temperature during the cooling process \((T_c)\) and the heating process \((T_h)\), thus suggesting the existence of a kinetic barrier in the self-assembly of \((\text{S})\)-TATA during the cooling process (Figure 3C).\(^{15}\) The thermal hysteresis was observed at all cooling and heating rates (Figures S11, Supporting Information) and with a decrease of the difference between \(T_c\) and \(T_h\) \((\Delta T)\) when increasing concentrations (Figure 3D). Interestingly, elongation temperatures in both heating and cooling processes increase with concentrations, thus confirming that \((\text{S})\)-TATA molecules follow a highly cooperative polymerization process in toluene (Figure S13, Supporting Information). Fitting the melting curves in the heating process, with models of supramolecular polymerization optimized by Meijer and co-workers,\(^{19}\) confirmed the cooperative mechanism (Figures S12, S13 and Table S3, Supporting Information).\(^{58}\)

For instance, we determined an elongation enthalpy \(\Delta H_e\) of \(-66\pm 1\) kJ mol\(^{-1}\) and a critical elongation temperature \(T_e\) of 313.8 K at a concentration of \(1.0 \times 10^{-4}\) M. Over the range of concentrations studied, \(T_e\) decreased linearly according to the Van ‘t Hoff plot (Figure 3E). For \((\text{S})\)-TATA in toluene, the standard enthalpy change \(\Delta H_f\) and the standard entropy change \(\Delta S_f\) for the elongation process were estimated to be \(-72\pm 1\) kJ mol\(^{-1}\) and \(-153\pm 4\) J mol\(^{-1}\) K\(^{-1}\) respectively, leading to an elongation equilibrium constant \(K_e\) of \(4.0 \times 10^6\) M\(^{-1}\) at 298 K (Figure S14 and Table S4, Supporting Information). The enthal-
The existence of thermal as a function of $T$ (Table S3, Supporting Information). Typical values for in toluene measured at different temperatures. At 317 K, the gradual increase of the absorption intensity is an artefact due to the slow evaporation of toluene from the cuvette over the time of the experiment.

Additionally, for (S)-TATA, the elongation temperatures in the heating ($T_e$) and cooling ($T'_e$) processes along with the elongation enthalpy $\Delta H_e$ were found to increase in a mixture of toluene/MCH compared to pure toluene at a concentration of 0.1 mM (Table S3, Supporting Information). Typical values for elongation constants were found in a range going from $4 \times 10^6$ m$^{-1}$ (in toluene) to $10^8$ m$^{-1}$ (in toluene/MCH 1:1 (v/v)).

Finally, we evaluated how temperature can influence the kinetics of the supramolecular polymerization process. As reported by Würtzner and co-workers, the existence of a thermal hysteresis implies that the monomers can be kinetically inactivated between $T_r$ and $T_r'$, that is, that the supramolecular polymerization can be lagged depending on the temperature at which the self-assembling process is monitored. To verify this hypothesis, we followed the evolution of the absorbance of (S)-TATA with time at different temperatures (Figure 3F). For a 0.21 mM solution, we found that 1) for temperatures below $T_r$, polymerization occurs immediately and thermodynamic equilibrium is reached within around 1 hour; that 2) for temperatures between $T_r$ and $T_r'$, the aggregation of TATA molecules is delayed with increasing time as the temperature of polymerization approaches $T_r'$; and that 3) above $T_r$ (for instance at 317 K), no polymerization occurs over the 6 hours of the experiment. These experiments also show that temperature can be used as a trigger to kinetically bias the supramolecular polymerization process towards sigmoidal growths. \[36\]
Influence of the cooling rate on the hierarchical ordering

We then studied the chirality of the aggregation process of (S)-TATA by first using temperature-dependent CD spectroscopy. We focused on the self-assembly behavior in toluene at different concentrations and for different cooling rates between 50 °C and 20 °C. In all experiments, the linear dichroism (LD) was also measured, and in all cases no significant contribution was observed (i.e., LD was never higher than one thousandth of the CD signal).

We first monitored the evolution of the CD signal at a fixed concentration of 0.5 mM and for a cooling rate of 1 K min⁻¹ (that will be referred hereafter as “slow cooling”). The CD signal showed increased intensities for the positive maximum at 318 nm and for the negative one at 344 nm (Figure 4A). Surprisingly, when a hot solution of (S)-TATA was cooled with a rate of 10 K min⁻¹ (that will be referred hereafter as “fast cooling”), the CD signal displayed an increasing negative intensity maximum at 292 nm, together with a second increasing negative intensity at 344 nm (Figure 4B). Importantly, no signal corresponding to molecular CD was observed at high temperature, probably because of the high dynamics of TATA propeller-like chirality in solution. However, upon aggregation, the freezing of the propeller conformation results in the appearance of a molecular-propeller contribution to the CD spectrum. It is also expected that the aggregation of (S)-TATA in such hierarchical supramolecular systems should produce a complex CD signal resulting from the combination of 1) molecular CD, 2) first-order aggregation-induced CD (i.e., CD caused by the spatial organization of chromophores in the primary aggregates, such as single-chain polymers described here, for instance), and 3) aggregation-induced CD of higher orders (i.e., CD caused by further aggregation of the single chain polymers into superstructures, such as chiral bundle of fibers and superhelices for instance, and as observed in the literature for a few other systems). At the same concentration of 0.5 mM, we were also able to perform VCD experiments (Figure 4C; see details in the Supporting Information, section S2.7). Interestingly, a clear inversion of the bisignate peak for the amide I band at 1650 cm⁻¹ was observed (-/+ for fast cooling, and +/− for slow cooling), in line with the CD measurements. Thus, the results reported in Figure 4A–C suggest the presence of two
self-assembly pathways that diverge between the slow and fast cooling regimes and that lead to distinct thermodynamic and kinetic supramolecular structures, respectively. In addition, the stability of the CD signal (sign and intensity) remains for at least 2 days for both 0.1 and 0.5 mM concentrations and fast cooling, indicating the high stability of the kinetic self-assembly at room temperature, a situation not yet reported in the literature and of particular fundamental and practical interests.

To test our hypothesis, we studied how concentration influences the kinetics of aggregation for the two cooling regimes (see Figure 4D (for 0.1, 0.5, and 1 mM); and Figure S19 (for 0.25, 0.32, and 0.75 mM)). On the one hand, at all concentrations and cooling rates, UV monitoring of the polymerization process at 315 nm confirmed the presence of a thermal hysteresis with \( T_c \) and \( T_r \) increasing with increasing concentrations (Figures 3C,D, and Figures S11 and S13–S15 in the Supporting Information). On the other hand, although monitoring the CD signal of (S)-TATA solutions in toluene confirmed the nucleation growth mechanism of the polymerization (Figure S20, Supporting Information), the data demonstrate in addition the presence of different self-assembling pathways depending on the cooling rate. Negative CD at 315 nm occurs mainly in relatively diluted solutions ([(S)-TATA] \( \leq 0.32 \) mM) in the slow cooling, and up to moderate concentrations ([(IS)-TATA] \( \leq 0.75 \) mM) in the fast cooling conditions. Positive CD at the same wavelength is observed mainly at high concentrations ([(IS)-TATA] = 1.0 mM) in the fast cooling, and at moderate and high concentrations (0.5 mM \( \leq [(S)-TATA] \leq 1.0 \) mM) in the slow cooling conditions. These observations suggest that, at low concentrations and for fast cooling rate, primary chiral assemblies form corresponding to monomolecular TATA fibrils. Interestingly, at slow cooling rates, a mixture of single fibrils with higher-order aggregates, such as superhelical structures, are possibly obtained (as the intensity of the negative signal is only of 7 mdeg, compared to the 70 mdeg for the fast cooling). At a higher concentration of 1 mM, primary fibrils self-organize into higher-order aggregates, such as superhelical structures, irrespective of the cooling rate. Another very interesting and important result highlighted at 1 mM, especially for the fast cooling regime, is the presence of a non-monotonic variation of the sign of the CD as a function of temperature, revealing a negative signal that turns into a positive one upon cooling, as well as the reappearance of a negative signal before going to zero upon heating. This memory effect strongly suggests that the superhelical structures are made from single helices, and that their primary helical structure is not lost when forming the superstructures. In the medium concentration range (0.5 mM), the CD increases to positive values of about 150 mdeg during slow cooling experiments, whereas for fast cooling conditions, the CD decreases to negative values of about –100 mdeg, and then remains constant. Considering that disassembly is rather improbable below \( T_r \), we believe that this plateau is related to the coexistence of single fibrils along with some superstructures, which add a positive contribution to the CD. This is also in agreement with the intensity of the negative CD signal, which is not as high as expected from the experiment performed at 0.1 mM.

To probe further the existence and nature of such hierarchical self-assembly pathways, and to assign their absolute configuration, we performed AFM imaging on enantiopure (S)-TATA solutions after either slow or fast cooling. AFM micrographs of self-assemblies, obtained from a diluted 0.1 mM solution, reveal the sole presence of single fibers with a P helicity that can slightly coagulate by lateral aggregation (Figures 5A,B). In a striking difference, for high concentrations (1 mM), only large M superhelical bundles were observed by AFM (Figure S6B,C, Supporting Information). For medium concentrations of 0.5 mM (Figures 5C–F and Figure S6D,E in the Supporting Information), the observations vary depending on the cooling regimes. Interestingly, for the fast cooling process, single P-helices with some coagulation between them formed (Figure 5C). For slow cooling, larger bundles appeared with a global M-superhelix arrangement but internally composed of individual P-helices (Figure 5 D–F). Such an important morphological difference is consistent with the various spectroscopic responses observed by CD. In addition, DFT calculations combined with VCD experiments (Figure 4C), are in agreement with the presence of a bisignate (+/–) peak for a P conforma-

To explain these experimental data we propose that, although following a similar primary nucleation step to lead to short monomolecular stacks (Figure 6, steps 1—3, grey circles), (S)-TATA primary fibrils can then give rise to different hierarchical structures depending on the concentration and cooling rates between two identical temperatures (50 to 20 °C). At low concentrations, independently of the cooling rate, the growth of single fibrils with a P helicity and coagulation of a small number of these fibrils is the dominant process which is characterized by a negative CD signal and a \( –/+/+ \) bisignate VCD signal (Figure 6, steps 4 and 5, blue circles). At high concentrations, independently of the cooling rate, an additional level of self-assembly takes place from the short monomolecular stacks (with a P helicity) that consists of the formation of polyfibrillar nuclei that further enter in a superhelical growth regime. The positive CD signal and \( +/– \) bisignate VCD signals observed result from these highly twisted bundles of supramolecular fibers with a M helicity (Figure 6, steps 4’ and 5’, red circles). Finally, at medium concentrations, the cooling rate can drive the system towards one of the other self-assembly pathways. Very importantly, and as shown by the memory effect in step 3 (for instance well seen in Figure 4D, fast cooling at 1 mM), the primary aggregates have identical columnar packing for both cooling regimes (i.e.; P-helices). Conversely, beyond the primary polymerization process, the hierarchy of these supramolecular structures can be modified by the cooling rate of the environment, which influences the secondary nucleation step.
Figure 5. (A, B) AFM height image of (S)-TATA self-assemblies from a 0.1 mM solution in toluene and (B) of a single fibril with a P-helicity, (inset shows its height profile); (C–F) AFM height image of (S)-TATA self-assemblies from 0.5 mM solutions in toluene. (C) Fast cooling regime (10 Kmin⁻¹). (D–F) Slow cooling regime (1 Kmin⁻¹). In SEL, which represent magnified regions of SD, the presence of only single P-helices in larger M-superhelices (i.e. P-helix ⊂ M-superhelix) is identified (see for instance the regions pointed by the arrows, and an extended micrograph in the Supporting Information).

Figure 6. Schematic representation of the two main self-assembly pathways for (S)-TATA molecules depending on the cooling rate at a typical concentration of 0.5 mM in toluene. The mechanism of primary nucleation is shared by both pathways up to the formation of P-helical fibrils (3), but then differs to evolve towards a primary growth regime of P-helical fibrils (for fast cooling) or toward a secondary nucleation and growth regime of M-superhelical fibers (i.e., P-helix ⊂ M-superhelix; for slow cooling).
Conclusions

We have discussed the hierarchical self-assembly mechanism of a series of TATA molecules based on a combination of spectroscopic techniques together with microscopy experiments and DFT calculations. We have first proposed that TATAs self-assemble according to a cooperative nucleation and growth polymerization process. The presence of a strong thermal hysteresis between the assembly and disassembly processes is related to the existence of a kinetic barrier during the aggregation process, thus offering the possibility to delay the nucleation process and to reach sigmoidal growth for temperatures comprised between \( T_n \) and \( T_c \). This behavior relates to the particular propeller shape of the TATA molecules. It induces strong entropic and conformational contributions to the self-assembly process which relates to 1) the breakage of the intramolecular rotation at the nitrogen center of the \( C_2 \)-symmetric TATA, and to 2) the heterochiral mismatch of their \( \Delta \) and \( \Delta' \) conformers in the stacking of the columnar polymers. We have also unraveled the presence of divergent chiral self-assembly pathways, which can take place at a similar concentration and range of temperature but for different cooling rates of the system. Kinetic analysis of the CD melting curves, interpretation of VCD bisignate bands, as well as AFM imaging suggest together that the difference of chirality is related to the formation of structures of higher hierarchy (i.e., superhelices) which readily form upon slow cooling. Such a large structural change obtained from such a small thermal effect is of particular fundamental interest for gaining knowledge and control over complex self-assembly processes. This unprecedented observation of a hierarchical bifurcation is possible because the system shows a peculiar novelty compared to the other systems published so far in the literature, as the kinetic product (single P-helix) is thermally stable for a long time without conversion to the thermodynamic product (P-helix\( \subset \)M-superhelix), so that it can be properly isolated and characterized. The reason of this stability relates to the very low kinetic lability of TATA molecules in the primary stacks, as well as to their locked conformation as P propellers in the single helix (as shown by the strong memory effect deciphered here). This feature is also of potential practical interest for fine-tuning the physical properties of functional materials of these systems.[9]

Overall, these results highlight that pathway complexity in supramolecular polymerization can be also encountered in systems that produce similar primary nucleation steps and primary growths, but which diverge in their secondary or higher organizational order (and such as it is found in nature, for instance with the secondary nucleation involved in the formation of \( \beta \)-amyloids).[62] Therefore, they enrich the understanding of supramolecular polymerization mechanisms towards stepwise hierarchical structures that involve increasing levels of spatial and temporal organizations, as encountered in complex living systems.

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Conflict of interest

The authors declare no conflict of interest.

Keywords: hierarchical self-assembly · organogelators · pathway complexity · supramolecular polymers
