Hierarchical Self-Assembly of Histidine-Functionalized Peptide Amphiphiles into Supramolecular Chiral Nanostructures

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ABSTRACT: Controlling the hierarchical organization of self-assembling peptide amphiphiles into supramolecular nanostructures opens up the possibility of developing biocompatible functional supramolecular materials for various applications. In this study, we show that the hierarchical self-assembly of histidine- (His-) functionalized PAs containing D- or L-amino acids can be controlled by both solution pH and molecular chirality of the building blocks. An increase in solution pH resulted in the structural transition of the His-functionalized chiral PA assemblies from nanosheets to completely closed nanotubes through an enhanced hydrogen-bonding capacity and π−π stacking of imidazole ring. The effects of the stereochemistry and amino acid sequence of the PA backbone on the supramolecular organization were also analyzed by CD, TEM, SAXS, and molecular dynamics simulations. In addition, an investigation of chiral mixtures revealed the differences between the hydrogen-bonding capacities and noncovalent interactions of PAs with D- and L-amino acids.

INTRODUCTION

Disordered constituents can spontaneously form stable organized patterns without any need for human intervention under equilibrium conditions in the self-assembly process. This natural process is important for the bottom-up fabrication of advanced functional nanostructures with the help of various noncovalent interactions such as van der Waals, π−π stacking, hydrogen-bonding, coordination-bonding, and electrostatic interactions. These noncovalent interactions guide the self-assembly of building blocks into supramolecular nanostructures with different morphologies including nanofibers, nanospheres, nanotubes, and nanoribbons.

Among different self-assembling molecules, peptides are interesting building blocks for the construction of supramolecular assemblies because amino acids are the building elements of peptide molecules, which provide great diversity depending on their side-chain properties. Self-assembling peptide amphiphiles (PAs) are obtained through the conjugation of hydrophobic alkyl tails to hydrophobic and polar amino acid residues. The supramolecular organization of PA assemblies is modulated through molecular design exploiting the hydrogen-bonding properties of amino acid residues, as well as their hydrophobicity and molecular chirality. The structural organization of PA nanostructures can be also controlled by different external factors such as temperature, photoirradiation, salt concentration, solvent effects, and pH, owing to the dynamic nature of the self-assembly process.

PA molecules can be functionalized with aromatic moieties including 9-fluorenylmethyloxycarbonyl (Fmoc), pyrene, and naphthalene as N-terminal capping groups or aromatic amino acids such as diphenyl groups incorporated into the PA backbone to promote self-assembly using the directionality of π−π stacking interactions along with hydrogen bonding and hydrophobicity. Furthermore, histidine-containing PA building blocks have been developed to acquire pH-sensitive control of the self-assembly behavior through the properties of the aromatic imidazole side chains. Branched PA molecules consisting of a tri-His headgroup and β-sheet-forming backbone were synthesized to develop pH-switchable injectable hydrogels as tissue scaffolds. In another example, the design of pH-sensitive oligo-His-containing PAs was presented to control the morphology of PA nanostructures at pH values between 6 and 7.5.
incorporating a terminal histidine residue was shown to self-assemble into tape structures based on bilayers, with the number of bilayers within the nanostructures and the intermolecular interdigitation depending on the concentration. The surfactant-like PA A₁₀H₆ forms fibrils and can be tagged with nanogold using nickel nitrotriacetic acid (Ni-NTA) coordination. In this study, we have designed histidine-functionalized PA building blocks containing either D- or L-amino acids (L-VVHH, D-VVHH, L-FFHH, and D-FFHH). The functionalization of the PAs with a double-His headgroup (HH) provided fine control over the supramolecular organization of the assemblies through a variety of noncovalent interactions under acidic, neutral, and basic conditions. In addition, the aromaticity differences between the β-sheet-forming regions were examined using different β-sheet-forming domains (−VVAG− and −FFAG−) in the peptide backbones. The supramolecular chirality of the PA assemblies is another important intrinsic property that can be manipulated using the chirality of the D- and L-amino acids. The effects of the amino acid chirality on the supramolecular organization were also analyzed using mixtures of different PA assemblies. In this article, we show the structural transition of the His-functionalized chiral PA assemblies from nanosheets to completely closed nanotubes with increasing pH and the effects of the molecular chirality of the building blocks on the supramolecular chiral organization.

## EXPERIMENTAL SECTION

### Materials

Fmoc and tert-butycarboxyl- (Boc-) protected amino acids, Rink Amide 4-methylbenzhydrylamine (MBHA) resin, and 2-(1H-benzotriazol-1-yl)-1,1,3,3-tetramethyluronium hexafluorophosphate (HBTU) were purchased from Novabiochem. All other solvents were purchased from Sigma-Aldrich.

### Synthesis of Peptide Amphiphile by Solid-Phase Peptide Synthesis

All peptides were synthesized using standard Fmoc synthesis. All reactions were performed with 2 equiv of Fmoc-protected amino acid, N,N-diisopropylethylamine (DIPEA) for 2 h. Removal of the Fmoc protecting group was performed using a mixture of triisopropyl silane (TIS), and H₂O in a ratio of 95:2.5:2.5 for 2 h. Excess TFA was removed by rotary evaporation. The remaining peptide was triturated with ice-cold diethyl ether, and the resulting white precipitate was freeze-dried. All peptides were purified by preparative high-performance liquid chromatography (prep-HPLC), and positive peptides were treated with 1 mM HCl.

### Liquid Chromatography-Mass Spectrometry (LC-MS)

For structural and chemical analysis of the peptides, an Agilent Technologies 6530 Accurate-Mass quadrupole time-of-flight (Q-TOF) liquid chromatography-mass spectrometry (LC-MS) system with a Zorbax SB-C8 column was used. The concentration of the sample for LC-MS measurement was 0.5 mg/mL. The solvents were water (0.1% formic acid) and acetonitrile (ACN) (0.1% formic acid). The LC-MS run for each sample was 30 min starting with 2% ACN and 98% H₂O for 5 min. Then, the gradient of ACN reached 100% until 20 min. Finally, the ACN concentration was decreased to 2% for the last 5 min. The solvent flow rate was 0.65 mL/min, and a sample volume of 5 μL was injected.

### Determination of Critical Aggregation Concentrations (CACs)

The critical aggregation concentrations (CACs) of the PAs were determined using hydrophobic Nile Red (9-diethylamino-5-benzo[x]phenoxazinone). Nile red exhibits an emission blue shift upon inclusion in a hydrophobic environment. The primary stock solution (1.256 mM) of Nile red was prepared in ethanol (0.4 mg of Nile red/1 mL of ethanol), and the stock solution was diluted to 78.125 μM using ethanol (the total volume of the final Nile Red solution was 1.5 mL). A 6.4 μL volume of Nile Red in ethanol (78.125 μM) was added to PA solutions (996.8 μL) at different concentrations, and the mixtures were vortexed. The samples were then stood overnight at room temperature. In the spectrometer, the excitation wavelength was 550 nm, and the emission spectra were collected between 580 to 750 nm using a Cary Eclipse fluorescence spectrophotometer.

### Circular Dichroism (CD)

A Jasco J-815 circular dichroism (CD) spectrophotometer was used for CD analysis. PA solutions [1% (w/v)] were prepared in water. The samples were gelled by the addition of 5 μL of 0.1 M NaOH (for pH 7.4) or 2.5 μL of 1 M NaOH (for pH 10) and incubated overnight. The PA samples prepared at pH 4.5 formed gels after overnight incubation without the addition of any NaOH. After the gelation, the samples were diluted to a 0.5 mM concentration using water. Spectra were measured from 300 to 190 nm with a 0.1 data pitch, a 1 nm bandwidth, and a 4 s data acquisition time. All samples were measured in a 1 mm quartz cell. Averages of three measurements were used, and sensitivity was selected as the standard.

### Transmission Electron Microscopy (TEM)

Imaging of the peptides was achieved by transmission electron microscopy (TEM) (FEI, Tecnai G2 F30) at 100 kV. For PA nanofiber staining, uranyl acetate solution in water (2 wt %) was used. The 2 mM stock solution was diluted 40 times. Diluted samples were placed on a lacey-carbon-coated copper grid. Ten milliliters of diluted sample solution was dropped on a grid and allowed to stand for 8 min. The excess was removed by pipet. Then, 20 mL of 2 wt % uranyl acetate solution was put on a parafilm sheet. The grid was placed on the top of the drop with its top side down and held there for 5 min. Stained grids were dried in the fume hood at room temperature overnight.

### Small-Angle X-ray Scattering (SAXS)

Small-angle X-ray scattering (SAXS) experiments were performed on beamline BM29 at the European Synchrotron Radiation Facility (ESRF), Grenoble, France. Solutions were loaded into the 96-well plate of a European Molecular Biology Laboratory (EMBL) BioSAXS robot and then injected by an automated sample exchanger into a quartz capillary (1.8-mm internal diameter) in the X-ray beam. The quartz capillary was enclosed in a vacuum chamber to avoid parasitic scattering. After the sample had been injected into the capillary and had reached the X-ray beam, the flow was stopped during SAXS data acquisition. Beamline BM29 was operated with an X-ray wavelength of λ = 1.03 Å (12 keV). The images were captured using a PILATUS 1 M detector, and data processing was performed using the dedicated beamline software ISPYB (Information System for Protein Crystallography: Beamlines) (BM29). Because of the strong anisotropy of the observed SAXS patterns for some samples (discussed below), in some cases, remasking and rebinning of the data was necessary to generate satisfactory radially averaged intensity profiles.

### Computational Experiments

The dynamics of PA nanostructures were investigated by all-atom explicit-solvent molecular dynamics (MD) simulations. The PA nanostructures were modeled as cylindrical nanofibers, and these structures were used as initial structures for the simulations. Each PA nanofiber system was built using 19 layers composed of 12 PA molecules each, and each system contained 228 PA molecules. The starting configuration for each PA nanofiber was based on the results of a previous study, which suggested that 19 layers with 12 PAs in each layer configuration gives rise to the most stable configuration for PAs having a similar length consistent with experimental results. Each layer was built by placing 12 PAs with 30° angles between them. Adjacent layers were assembled with a 5 Å separation and 15° angle rotation (Figure S8). The PA nanofibers were solvated with water molecules modeled using TIP3, and Na⁺ and Cl⁻ ions were added to reach a salt concentration of 0.15 M. The resulting simulation-system boxes comprised approximately 120000 atoms. MD simulations of the PA nanofibers were performed using the NAMD program (version 2.9) with the CHARMM force field [5051].
Prior to production simulations, simulation systems were minimized with 1000 minimization steps. Then, 50-ns production simulations were carried out for each PA nanofiber system at 1 atm pressure and 310 K temperature. Electrostatic interactions were calculated using the particle-mesh Ewald method with a grid spacing. The cutoff for van der Waals interactions was taken as 12 Å, with a switching function after 10 Å. Simulation trajectories were integrated with a time step of 2 fs, with all interactions calculated at every time step. Atomic coordinates were collected every 10 ps. The analyses were applied to the last 10 ns of each trajectory. Hydrogen bonds were calculated using the program CPPTRAJ. Calculations of nonbonded interaction energies for the simulation trajectories were carried out using the VMD program.

**Sample Preparation for Chiral Mixtures.** For the preparation of chiral mixtures, the PA powders were initially dissolved in hexafluoroisopropanol (HFIP), which is a strong H-bond-donating solvent, to prevent individual self-assembly of the PA molecules prior to chiral mixing. The PA solutions (with D or L chirality) prepared in HFIP were mixed at 100%, 75%, 50%, 25%, and 0% molecular ratios. Then, HFIP was removed under a vacuum, and the chiral mixture powders were resuspended in water at a PA concentration of 0.5 mM. The pH was adjusted using NaOH solution.

## RESULTS AND DISCUSSION

**Design, Synthesis, and Self-Assembly of the Histidine-Functionalized PA Building Blocks.** Amino acids exhibit a great diversity in their side chains including charge, hydrophobicity, polarity, and aromaticity. Histidine (His) contains an aromatic imidazole side chain that plays an important role in protein–protein interfaces and catalysis because of its capacity for multiple interactions with other molecules through cation–π (when His is protonated, His+), π–π stacking, hydrogen–π, hydrogen-bonding, and coordination interactions with metallic cations. PAs consisting of a hydrophobic alkyl tail and aliphatic or aromatic amino acid domains were functionalized at the N-terminus with double-histidine head groups (Figure 1). The integration of two histidines within the PAs resulted in self-assembling building blocks with multiple interaction capabilities at different pH values. Two different β-sheet-forming domains, namely, −VVAG− and −FFAG−, were also conjugated to the molecules so that hydrogen-bonding capacities of aliphatic and aromatic side chains could be studied. In addition to the differences in amino acid residues, the PA molecules were built with either L or D chirality to reveal the effects of molecular chirality on the supramolecular packing and to understand the impacts of the chiral differences on the self-assembly behavior (Figure 1). All molecules were synthesized according to the solid-phase peptide synthesis method, and the purities of the molecules were determined by LC-MS (Figures S1−S4).

The hierarchal self-assembly of L-VVHH, D-VVHH, L-FFHH, and D-FFHH is controlled by the hydrophobic alkyl tail, the peptide backbones with different hydrogen-bonding abilities, and the hydrophilic double-His headgroup. The pH under self-
assembly conditions was determined according to the pKₐ value of the imidazole ring in the His residues, which is approximately 6.5. Below this pH (e.g., in slightly acidic conditions, pH 4.5), the side chains of His are protonated, and hydrophilic His residues solubilize the PA molecules in water. The solvophobic interactions enhanced by hydrogen bonding between the β-sheet-forming residues and cation−π interactions of the His residues facilitate the supramolecular organization of the PAs under acidic conditions. At physiological pH, the imidazole ring can be in either the neutral or positively charged form, and the molecules are able to self-assemble into supramolecular nanostructures with the help of cation−π hydrogen bonding, and π−π interactions. The self-assembly is also assisted under basic conditions because the neutral imidazole ring can behave as both a hydrogen donor and a hydrogen acceptor and can form extensive hydrogen bonding and π−π stacking arrays. To reveal the aggregation behavior of the PAs, the critical aggregation concentrations (CACs) of the molecules were first studied using a Nile Red assay. The CACs of the PAs were determined as being higher than approximately 1 μM in water. Hence, all structural studies were carried out above this concentration to obtain histidine-functionalized PA assemblies under slightly acidic (pH 4.5), neutral (pH 7.4), and basic (pH 10) conditions.

**Structural Characterization of the PA Assemblies.** The secondary structures of the l-VVHH, D-VVHH, l-FFHH, and D-FFHH assemblies were characterized using circular dichroism (CD). The typical β-sheet secondary-structure organization shows a characteristic CD spectrum with maximum and minimum peaks at about 195 and 216 nm, respectively. At pH 4.5, the CD spectrum of l-VVHH revealed a twisted β-sheet secondary structure with red-shifted maximum and minimum peaks at about 205 and 224 nm, respectively, due to the slight distortion from perfect β-sheets. In addition, the CD spectrum of d-VVHH showed mirror-image symmetry with respect to that of l-VVHH because of chiral inversion (Figure 2a). The position of the positively charged
aromatic His residues in the L- and D-VVHH assemblies could change hydrogen-bonding energies between the residues and prevent the formation of rigid β-sheets at pH 4.5. Although the CD spectrum of L-VVHH was similar to that of the L-FFHH assembly (Figure 2d), a decrease in CD signal was observed because of differences in the solubilities of the PA assemblies resulting from the enhanced aromaticity of L-FFHH compared to L-VVHH at pH 4.5. On the other hand, mirror-image symmetry was not observed in the CD spectrum of D-FFHH compared to that of L-FFHH (Figure 2d). The CD signal depends on the concentration of the assemblies, and slight differences in the concentrations of the molecules could result in the non-mirror-image symmetry due to experimental errors.

At pH 7.4, the L- and D-VVHH assemblies preserved their twisted β-sheet organization with mirror-image symmetry (Figure 2b). However, the secondary structure of the L- and D-FFHH assemblies changed dramatically from twisted β-sheet to superhelical peptide assemblies. The CD spectrum of L-FFHH exhibited a maximum at 195 nm and two minima at about 207 and 217 nm, indicating helical organization of the

Figure 3. TEM images of the supramolecular L-VVHH, D-VVHH, L-FFHH, and D-FFHH nanostructures at different pH values.
The reason for this change is attributed to enhanced π−π stacking of neutral −HH− and −FF− residues of the PAs at pH 7.4. The potential face-to-face or face-to-edge orientation of the aromatic groups of L- and D-FFHH molecules through the PA assemblies resulted in helical organization under these conditions. Moreover, the observation of the two additional peaks at about 225 and 240 nm also pointed to an increase of the π−π interactions between the L- and D-FFHH molecules at pH 7.4. On the other hand, this phenomenon was not observed in the CD spectra of the L- and D-VVHH assemblies because −VV− residues lack π−π stacking capabilities and have different hydrogen-bonding properties at pH 7.4. This difference prevented the possible π−π stacking of the His residues and made the twisted β-sheet organization favorable for the L- and D-VVHH assemblies at pH 7.4 (Figure 2b).

Under basic conditions, complete neutralization of the His residues resulted in increase of the π−π interactions and extensive hydrogen bonding for all groups of the PA assemblies. Although aromatic interactions were not observed in the CD spectra of the L- and D-VVHH assemblies at pH 7.4, the planar orientation of the neutral His residues under basic conditions without the need for extra aromatic residues led to the formation of π−π stacking interactions within the L- and D-VVHH assemblies. Two additional peaks at about 225 and 240 nm in the CD spectra of the L- and D-VVHH assemblies indicated an enhanced aromaticity with twisted β-sheet secondary-structure organization (Figure 2c). On the other hand, the L- and D-FFHH assemblies showed helical organization with an increase in the CD signals at about 225 and 240 nm, pointing to enhanced aromaticity (Figure 2f).

All PA structures were imaged by TEM at pH 4.5, 7.4, and 10 (Figure 3). At pH 4.5, sheetlike nanostructures were observed in the TEM images of all PAs with L or D chirality. Increasing pH resulted in the formation of twisted nanostructures instead of sheetlike arrangements because of the reduced charge on the imidazole ring in the His residues enhancing the packing of the molecules into ordered supramolecular assemblies. Under basic conditions, the imidazole groups were completely neutralized, and π−π interactions dominated the supramolecular organization. Hence, the L-VVHH and D-VVHH PA molecules formed well-organized nanotubes because of the multilayer molecular packing of the building blocks. Although well-organized nanotube formation could not be observed in the TEM images of the L-FFHH and D-FFHH PA molecules at pH 10 because of the sample preparation, atomic force microscopy (AFM) images of the L-FFHH and D-FFHH assemblies clearly showed nanotube organization at pH 10 (Figure S7).

Prior to the nanotube formation at pH 10, the PAs initially could form sheetlike nanostructures because of the positive charges on the His residues at pH 4.5. The charge screening resulting from the increase in pH promoted the π−π interactions and resulted in the formation of twisted PA nanostructures at pH 7.4 (Figure 3). Whereas twisting was observed in the TEM images of all molecules at this pH, the L- and D-FFHH assemblies showed more well-ordered twisting behavior than the L- and D-VVHH nanostructures. This mechanism was also analyzed by AFM imaging, and the nanotube formation at pH 10 was observed in the AFM images of all PA assemblies (Figures S6 and S7).

SAXS analysis provides information on nanostructure shape and dimensions in solution. The scattering intensity profiles of...
we performed all-atom explicit-solvent molecular dynamics simulations of L-VVHH, D-VVHH, L-FFHH, and D-FFHH assemblies. H-bond analysis for the simulations of L-VVHH and L-FFHH PAs under different pH conditions are shown in Figure 5a. As the pH increases, the H-bond-forming ability of both L-VVHH and L-FFHH increases. Similar behavior was also observed for D-VVHH and D-FFHH assemblies (Figure S9). The change in hydrogen-bonding capacity of the PAs under different pH conditions could stem from the interactions between histidine residues. At pH 4, all of the histidine residues are protonated, which results in high repulsion between histidine side-chain atoms (Figure 5b). At pH 6.5, where the histidine residues are partially charged, repulsive interactions between charged side chains are balanced by attractive interactions such as π–π, cation–π, and hydrogen-bond interactions. At pH 10, the interactions between histidine side chains become attractive, because all side chains are in neutral form.

Moreover, H-bond analyses also showed that FFHH PAs contain fewer interpeptide hydrogen bonds compared to VVHH. The smaller numbers of H-bonds in FFHH PAs could give rise to an increase in the twisting of FFHH PA nanostructures compared to the twisting of VVHH PAs. To elucidate this behavior, we calculated nonbonded interactions between phenylalanine and valine side chains (Figure S10). Phenylalanine side chains in L-FFHH PAs have more attractive interactions than valine side chains in L-VVHH PAs. However, phenylalanine side chains have high repulsive van der Waals interactions, whereas valine side chains have attractive van der Waals interactions (Figure S10). Overall, phenylalanine side chains have repulsive nonbonded interactions, and valine side chains have attractive nonbonded interactions. Thus, steric hindrance between phenylalanine residues could play a significant role in determining the morphology of PA nanostructures.

An analysis of H-bond properties depending on the molecular chirality of the PA molecules was also performed under acidic (pH 4), slightly neutral (pH 6.5), and basic conditions (pH 10) (Figure S9). L-PA molecules tend to form higher numbers of H-bonds compared to D-PA molecules depending on pH increase (Figures S9 and S11). This observation could be related to the differences of the conformations of D- and L-amino acids in the peptide backbone, which can affect noncovalent interactions between the molecules. In fact, several studies showed that the D-form of
amino acids could alter β-sheet-forming ability of PA nanofibers depending on the amino acid chirality of the molecules.71

Self-Assembly of the Chiral Mixtures. To systematically analyze the effects of molecular chirality on the self-assembly behavior of the PAs, chiral mixtures consisting of both D-VVHH and L-VVHH at different molar ratios were prepared at pH 10 according to the protocol given in detail in the Experimental Section. The CD spectra of 100% L-VVHH and 100% D-VVHH showed a nearly symmetric Cotton effect. This reflects the opposite handedness of packing in the nanostructures at pH 10. According to secondary-structure analysis of the chiral mixtures in Figure 6, an increase in the D-VVHH amount in the L-VVHH solution decreased the β-sheet CD signal because of packing constraints. A further increment in the D-VVHH amount led to a β-sheet signal that was mirror-symmetric with respect to that of L-VVHH (Figure 6). Racemic mixtures not only showed nearly zero CD signal but also formed twisted nanostructures rather than nanotubes (Figure 7). SAXS data from selected mixtures studied (at pH 10) can be fitted to a model for bilayers consistent with the twisted nanotape structures observed by TEM. The SAXS data along with model form-factor fits are shown in Figure 8. The fit parameters are listed in Table S3.

Simulation of the Self-Assembly of the Chiral Mixtures. In addition to simulation studies of individual L-VVHH, D-VVHH, L-FFHH, and D-FFHH assemblies, the noncovalent interactions between the chiral mixtures were also analyzed through molecular dynamics simulations performed for 50 ns. H-bond analysis of these simulations suggests that the pure L-PA systems have more hydrogen bonds than mixtures of L-PA and D-PAs (Figure S11). As the amount of D-PA in L-PA systems increases, the hydrogen-bonding capacity decreases. However, when the system is composed of only D-PAs, then the hydrogen bonding tends to increase. Pure L-PA and D-PA nanostructures tend to have higher hydrogen-bonding abilities than nanostructures composed of racemic mixtures of these PAs.

CONCLUSIONS
In summary, the hierarchical assemblies of the His-functionalyzed PAs and their chiral mixtures were characterized using different techniques and molecular dynamics simulations to reveal supramolecular organization of the PA assemblies and their chiral mixtures. It was shown that the supramolecular organization of His-functionalyzed PAs is controlled by the solution pH and the molecular chirality of the building blocks. Whereas self-assembly of the PAs into nanosheets is driven by solvophobic interactions under acidic conditions, the deprotonation of the His residues due to a pH increase changes the self-assembly behavior and results in the formation of stable twisted nanostructures at neutral pH. Moreover, an increase in solution basicity facilitates the transition from twisted nanostructures into closed nanotubes through an enhanced hydrogen-bonding capacity and π–π stacking of the imidazole ring at pH 10.

ASSOCIATED CONTENT

Supporting Information
The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.langmuir.7b01266.

LC-MS and CAC results; supporting AFM, TEM, and SEM images of the PA assemblies; details of SAXS data fitting with estimated structural parameters; and results of simulation studies (PDF)
Acknowledgments

This work was partially supported by grants from TUBITAK (114Z728), TUBA-GBIP, and FP7-Marie Curie IRG. M.H.K. and G.C. are supported by TUBITAK-BIDEB 2210-C and 2211-C fellowships. The simulations reported in this work were performed at TUBITAK ULAKBIM, High Performance and Grid Computing Center (TRUBA resources). We thank Mr. M. Guler for help in TEM imaging. I.W.H. is grateful to EPSRC Grid Computing Center (TRUBA resources). We thank Mr. Macit Mumcuoglu, D.; Sardan Ekiz, M.; Gunay, G.; Tekinay, T.; and Guler, M. O. for assistance with the measurements, and award of bioSAXS beamtime on BM29 (ref MX-1769), ESRF, Grenoble, France (U.K.) for Grant EP/L020599/1. We thank the ESRF for the support. Jeröme Kieff, of the National Cancer Institute for technical assistance with the measurements.

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Notes

The authors declare no competing financial interest.

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