Chirality-Mediated Mechanical and Structural Properties of Oligopeptide Hydrogels

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Supporting Information

ABSTRACT: The origin and the effects of homochirality in the biological world continuously stimulate numerous hypotheses and much debate. This work attempts to look at the biohomochirality issue from a different angle—the mechanical properties of the bulk biomaterial and their relation to nanoscale structures. Using a pair of oppositely charged peptides that coassemble into hydrogels, we systematically investigated the effect of chirality on the mechanical properties of these hydrogels through different combinations of syndiotactic and isotactic peptides. It was found that homochirality confers mechanical advantage, resulting in a higher elastic modulus and strain yield value. Yet, heterochirality confers kinetic advantage, resulting in faster gelation. Structurally, both homochiral and heterochiral hydrogels are made of fibers interconnected by lappet-like webs, but the homochiral peptide fibers are thicker and denser. These results highlight the possible role of biohomochirality in the evolution and/or natural selection of biomaterials.

KEYWORDS: hydrogels, mechanical properties, homochirality, heterochirality, dynamic rheometry, NMR spectroscopy, small-angle X-ray scattering, small-angle neutron scattering

INTRODUCTION

The phenomenon of homochirality, its origin and implications, is one of the most enigmatic questions in the life sciences.† From a fundamental viewpoint, of greatest interest is the core controversy of the preservation of mirror symmetry in the nonbiological world—that is, the existence of racemic mixtures of enantiomers (or heterochirality)—and the break of mirror symmetry in the biological world—that is, the existence of enantiopure systems (or homochirality).‡ Natural biomolecules are homochiral, with proteins comprised exclusively of L-amino acids, while nucleic acids and most polysaccharides contain only D-sugars. The origin of the homochirality of life is a matter of much debate, and numerous hypotheses and speculation exist in the literature.³ As to the advantages conferred by homochirality, the discussion has focused on the molecular aspects of homochirality, such as protein folding,⁴ chiroselective replication,⁵ enzyme catalysis,⁶ and DNA recognition.⁷ However, de novo design efforts have shown that introducing D-amino acids into proteins greatly expands structural motifs and even lead to enhanced stability.⁸ Such studies put the molecular advantage of homochirality into question.

In comparison, little is known about the material aspects of homochirality, that is, whether homochirality confers advantages to material properties such as mechanical strength. One might expect that such effects could be profound and lead to significant morphological differences of the resulting bulk material.⁹ In this work, we investigate whether homochirality confers mechanical advantages to soft biomaterials. Mechanical properties are important functional parameters for biomaterials.¹⁰ Specifically, we will compare the viscoelastic properties of homochiral vs heterochiral hydrogels. Hydrogels are viscoelastic soft biomaterials that have many natural (e.g., collagen) and man made (e.g., contact lenses) examples. It has been proposed that life may have originated in a hydrogel environment.¹¹ Hence, the effect of homochirality on the mechanical properties of hydrogels might have implications for the origin of life.

In organogels, three scenarios have been observed: homochirality leads to better gelling ability,⁶,¹² homochirality and heterochirality make no difference in gelling ability,¹³ and heterochirality leads to better gelling ability.¹⁴ However, the dominant situation is that homochirality leads to better gelling ability.¹⁵ In these studies of organo-gelators, the gelling ability was measured by either the minimum gelation concentration or the gelation temperature. There was no report on the mechanical properties of homochiral vs heterochiral organogels.

In contrast to organogels, where homochirality in most cases leads to better gelling ability, hydrogels appear to show a preference for heterochirality in many cases, a phenomenon called stereocomplexation.¹⁶ For example, stereocomplexes are formed between poly(D-lactic) and poly-(L-lactic) acids.
through stereospecific interlocking.16 It has been shown that the formation of the heterochiral stereocomplexes from D- and L-enantiomers is more rapid and more complete as opposed to homochiral enantiomers; and the polyethylene glycol gels formed in the presence of heterochiral stereocomplexes of the poly(lactic) acids are capable to hold more water and are more stable with higher transition temperatures than gels formed in the presence of homochiral poly(lactic) acids.26 It has been shown that hydrogels resulting from the stereocomplexation of 50/50 mixtures of D- and L-enantiomers of poly(lactic) acid with polyethylene glycol demonstrate much higher elastic moduli G′ as compared to the hydrogels where the ratio of D- and L-enantiomers was shifted toward greater homochirality (e.g., 84/16).17 Similarly, photoinduced hydrogelation of stereocomplexes constructed from poly(lactic) acid, polyethylene glycol, and methacrylate results in mechanically stronger hydrogels for the D+L mixture of poly(lactic) acid as compared to the pure L-poly(lactic) acid material (the elastic modulus G′ is ca. 2 orders of magnitude higher).18 Hydrogelation induced by poly(lactic) acid stereocomplexation persists after poly(lactic) acids are grafted to dextran.19

Such preference of heterochiral stereocomplexation is also found in other aqueous systems. For example, it was found that L-peptides stereocomplex with poly(D-lactic) acid but not with poly(L-lactic) acid.20 Also, peptate, a D-polysaccharide, stereocomplexes more efficiently with poly-(L-lysine) than with poly(D-lysine).21 In all these cases, heterochirality (D-L complexity) was preferred over homochirality (L-L or D-D complexation). In addition, heterochiral 1,3,5-cyclohexyltricarboxamide-phenyl-alanines (LLD or DDL) are excellent hydrogelators while their homochiral counterparts (LLL or DDD) do not gelate.22 However, in these studies, mechanical properties were not reported.

In peptide hydrogels, it has been recently reported that heterochiral hydrogels assembled from a pair of D-, L-peptides have higher elastic modulus than the parent homochiral hydrogels.23 Previously, it was reported that poly(D-lysine) and poly(L-lysine) form amyloid-like fibrils while each individual enantiomer remains a clear solution,24 although no rheological data were reported for these poly(D-lysine) + poly(L-lysine) stereocomplexes. In the case of actual amyloid fibers, somewhat contradictory observations have been reported. On the one hand, it was reported that amyloid fibers demonstrate homochiral stereospecificity in that L-peptides or proteins deposit onto preexisting L-fibers but not onto preexisting D-fibers.25 On the other hand, it was reported that inhibitors of amyloid fibers demonstrate heterochiral stereospecificity in that D-oligopeptides better inhibit the fibrilization of L-β-amylloid peptide than L-oligopeptides of the same sequence.26

This observed preference for heterochirality over homochirality in an aqueous environment raises the following question: does homochirality confer advantages to biomaterials? After all, natural protein biomaterials are homochiral. In this work, we address this question using a pair of self-repulsive but mutually attractive peptides that can co-assemble into hydrogels. Different chiral combinations are explored in a systematic fashion. Using dynamic rheometry, we found that heterochirality indeed leads to quicker gelation and higher elastic modulus in the first few hours of gelation. Afterward, homochiral gels outpace heterochiral gels and homochirality eventually leads to a much higher elastic modulus and slightly higher strain yield than heterochirality. Thus, for these peptide hydrogels, homochirality confers mechanical advantage while heterochirality confers kinetic advantage. Using a combination of NMR spectroscopy, small-angle X-ray and neutron scattering techniques (SAXS and SANS), we explored the mechanism underlying these observations.

### EXPERIMENTAL SECTION

#### Peptide Design and Synthesis

Oppositely charged undecapeptide modules (11 amino acids long) have been designed in accordance with our earlier approach27 whereby a positively charged module and a negatively charged module co-assemble into a hydrogel in phosphate-buffered saline (PBS) when mixed. The positive module contains alternating positively charged (lysine, K) and neutral (tryptophan, W; and alanine, A) amino acids, while the negative module contains alternating negatively charged (glutamate, E) and neutral (tryptophan, W; and alanine, A) amino acids. Such general design separates oppositely charged undecapeptides or different peptide chains. As a result of the electrostatic repulsions inherent within each peptide module, spontaneous hydrogelation and/or self-assembly due to slight pH, temperature, and ionic strength changes are avoided. To study the chirality effect of the hydrogels, both syndiotactic and isotactic peptides were made: two syndiotactic peptides composed of alternating D- and L-amino acids, denoted D,L-K and D,L-E, respectively; two isotactic peptides composed of all L-amino acids, denoted L-K and L-E, respectively; and two isotactic peptides composed of all D-amino acids, denoted D-K and D-E, respectively (Table 1). The N-, C- termini of each peptide were acetylated (Acetyl- ) and amidated (amide), respectively, to block terminal charges. In this work, all peptide sequences are palindromic.

All peptides were synthesized on Rink-amin MBHA resin by means of a CEM microwave synthesizer using standard solid-phase Fmoc-chemistry.28 The crude peptides were cleaved by a cocktail containing 95% trifluoroacetic acid (TFA), 2.5% trisopropylsilane, and 2.5% water volume fraction. The side chains were also deprotected during cleavage. TFA was removed by rotary evaporation under reduced pressure. The crude peptides were precipitated and washed twice by cold diethyl ether. The crude peptides were dissolved in water and lyophilized before purification.

Preparative reverse-phase HPLC (RP-LC) method was used to purify the crude peptides. In the purification of positive charged
peptides (L-K, D-K, and D,L-K), solvent A was 0.1% mass fraction HCl in water and solvent B was 0.1% mass fraction HCl in MeOH. In the purification of negatively charged peptides (L-E, D-E, and D,L-E), solvent A was 20 mM NH4HCO3 in water (pH 7.0), solvent B was 20 mM NH4HCO3 (pH 7.0) in MeOH/water (8:2, volume ratio). The chromatographic method of peptide purification was as follows: 0–40% B in 0–60 min, 40–100% B in 60–90 min with a linear gradient for each segment. The purity of each peptide was verified by analytical RPLC methods with the same solvents used for preparative RPLC.

Purified peptides were dissolved in PBS composed of 50 mM phosphate buffer and 100 mM sodium chloride, pH 7.4. The final concentrations of the individual peptide stock solutions were 16.0 mM, determined on the basis of the molar absorptivity of tryptophan (ε = 5 1H signals attributed to di-D,L-E) and the same receiver gain were used in all cases and the relaxation time was recorded again. To compare the signal intensity, the 1D1H spectra were recorded every hour until 8 h after mixing. Six months later, the 1D1H spectra were acquired every 30 min in the first hour for pair 1, and the same receiver gain were used in all cases and the relaxation time was recorded again. For every measurement, the monochromatic X-ray beam (λ = 0.689 Å) with a size of 0.07 mm × 0.20 mm was adjusted to pass through the centers of the capillaries. The exposure time for all samples was set to 0.2 s to avoid detector saturation and radiation damage to the samples. X-ray scattering intensities were collected using the 2D detector Pilatus 2 M (DECTRIS Ltd.). The 2D scattering images were converted into 1D scattering profiles of I(Q) vs Q in the Q-range from 0.007 Å−1 to 0.6 Å−1 by means of azimuthal averaging after solid angle correction. The resulting 1D profiles were normalized over the intensity of the transmitted X-ray beam, using the software package at the beamline 12ID-B. I(Q) is the scattering intensity of X-rays, and Q is the scattering vector amplitude which is related to the X-ray wavelength λ and the scattering angle θ by

$$Q = \frac{4\pi}{\lambda} \sin \left( \frac{\theta}{2} \right)$$

(1)

Subtraction of the solvent scattering (PBS in D2O) involved normalization based on the ratio of incident and transmitted X-ray photon counts to account for the slight differences in the thickness of different capillaries. Also additional background scattering correction was performed in accordance with the generally accepted published procedure.

SANS data were collected using the 30 m SANS instrument (NG-7) at NIST. Monochromatic neutrons at Δλ = 6 Å with a wavelength spread (Δλ/λ) of 0.14 were detected on a 64 cm × 64 cm two-dimensional detector. Data on SANS intensity were collected with a
Q-range from 0.001 Å⁻¹ to 0.4 Å⁻¹. The low-Q configuration used neutron focusing lenses and an 8 Å neutron wavelength. Scattering intensities were normalized using direct beam transmission measurements and were reduced according to published protocols. Both SAXS and SANS instruments have pinhole geometry.

The combined usage of SAXS and SANS techniques to characterize hydrogel structures has evident advantages. The high flux of the X-ray beam from the synchrotron allows one to reduce the data collection time down to 0.2 s, thus facilitating the use of SAXS to follow the gelation process in real time. In comparison, the data collection time in SANS is about 1–2 h, making it unsuited to monitor the gelation process. On the other hand, because of the different wavelength parameters (0.689 Å for SAXS vs 6 Å and 8 Å for SANS) and detector setup, the SANS instrument allows one to get down to much lower Qmin values as compared to the SAXS (0.001 Å⁻¹ for SANS vs 0.007 Å⁻¹ for SAXS). Lower Q values provide the possibility to reliably observe molecular assemblies of much greater size (up to ~2000 Å for SAXS vs up to ~500 Å for SANS).

Pair 1 of the syndiotactic peptides D,L-K and D,L-E does not gel. Instead, clusters of finite size are formed. The solution structure of pair 1 at 72 h was measured by SAXS, and the data were processed using the ATSAS software. The analysis of pairwise distance distribution functions for globular particles P(r) (eq 2) was performed using the linear regularization method of indirect Fourier-transformation using the program GNOM.

\[
P(r) = \frac{1}{2\pi^2} \int_{0}^{\infty} I(Q) \cdot (Q \cdot r) \sin(Q \cdot r) \, dQ
\]

(2)

P(r) reflects the probability that two randomly chosen points in a scattering particle are at r distance apart from each other, and P(r = 0) happens at the maximum linear dimension of the scattering particle, dmax. P(r) also provides the information about the distances between the electrons in the scattering particle, and could be calculated from the electron density distribution:

\[
P(r) = r^2 \gamma(r), \quad \gamma(r) = \left\{ \int_0^r (\rho(r + x) \, dx \right\}
\]

(3)

where \( \gamma(r) \) is the spherically (whole volume) averaged correlation function of the electron density reflecting the difference in the scattering intensity between the positions r and r+x. In the case of homogenous particles when \( \rho(r) = \rho(r+x) = \Delta \rho \), which is the difference in electron densities between the scattering particle and the solvent and is a constant

\[
P(r) = r^2 \gamma_0(r)
\]

(4)

where \( \gamma_0(r) \) is the normalized correlation function or so-called characteristic function.\(^{36}\) defined only by the geometry of the scattering particles, \( \gamma_0(r) \) = 1, for \( r = 0 \); and \( \gamma_0(r) \) = 0, for \( r > d_{\text{mac}} \). The radius of gyration of the scattering globular particle, \( R_g \) is derived from the second moment of P(r) as

\[
R_g^2 = \frac{\int_0^{d_{\text{mac}}} P(r) r^2 \, dr}{\int_0^{d_{\text{mac}}} P(r) \, dr}
\]

(5)

\( R_g \) is the root-mean-square distance of all unit-volume elements from the center of gravity of the scattering particle, and in the case of X-rays, the distribution of the mass is defined by the electron density distribution within the scattering particle. A simulated annealing algorithm was used to restore low resolution 3D structures of D,L-K + D,L-E clusters in solution built from densely packed dummy atoms implemented in the DAMMIN program.\(^{37}\) To build the most probable and reliable 3D model, multiple DAMMIN shape solutions (at least 20 runs) were aligned with respect to their principal axes of inertia followed by the structural discrepancy minimization using SUPCOMB program\(^{40}\) and averaging by means of the DAMAVER routine.\(^{41}\)

ATSAS software\(^{36,42}\) also has been used to estimate the zero-angle scattering intensity I(0) from the Guinier analysis of ln I(0) vs Q² plots. Since I(0) characterizes the mass of the scattering particles, it was used to monitor the gelation process of all hydrogels over time. At the same time, the fibers formed by pairs 2–5 have one dimension (length) much greater than the other two (cross-section). The length of the fibers exceeded the upper detection limits of our techniques (~500 Å for SAXS and ~2000 Å for SANS). Hence we analyzed the scattering data of pairs 2–5 in terms of the cross-sectional dimensions of the scattering particles using the standard approach of multiplying I(0) by Q. This approach essentially removes information about the length of the scattering particles. A simulated annealing algorithm, analogous to that used for restoring the 3D shape for pair 1, was used to restore the 2D cross sections for pairs 2–5. Here, we model the 2D cross sections of the fibers formed by pairs 2–5 using the algorithm in a purpose-written program that is described elsewhere.\(^{43}\) In SANS data processing, the dummy atoms were arranged on a flat grid of 20 × 20 close-packed dummy atoms, each 3 Å in diameter. In SANS data processing, where bigger 2D cross-sectional patterns were modeled, the grid has 20 × 50 close-packed dummy atoms, each 3 Å in diameter to model dimensions of about 200 Å, and 30 × 150 close-packed dummy atoms, each 5 Å in diameter to model dimensions of about 800 Å. This allows one to model pictorial cross-sectional slices of the hydrogels showing the fibers and how they are connected into the fibrous network. The program calculated the pair distance distribution function, \( P(r) \), for the model cross-section composed of the dummy atoms. \( P(r) \) is the distribution of distances between area elements in the cross-section, weighted by the scattering density at each radial distance, \( r \). The optimization procedure is in general described elsewhere.\(^{32,43}\) After optimization, the radius of gyration of the cross-section \( R_g \) in Å, the maximum cross-sectional dimension \( d_{\text{mac}} \) in Å, the cross-section area \( S_c \) in Å², and the zero-angle scattering intensity I(0), which is proportional to the mass per unit length of the fiber (in arbitrary unit per Å), were determined from \( P(r) \). \( d_{\text{mac}} \) is the r value at which \( P(r) \) goes to 0. The zero-th and the second moments of \( P(r) \) yield I(0) and \( R_g \) values, respectively. \( R_g \) is the contrast-weighted mean distance of all area elements from the center of scattering density. The program also outputs the model cross sections as atomic coordinate files in the Protein Data Bank format which allowed their pictorial presentation. On the basis of these coordinates and the grid dimensions, the area of the cross-section, \( S_c \), can be calculated. Corrections for scaling and incoherent background were applied to the model scattering profile so it could be compared directly with experimental scattering data.\(^{35}\)

### RESULTS AND DISCUSSION

Of the five peptide pairs, pair 1 is made of syndiotactic peptides; pairs 2 and 3 are made of homochiral peptides of opposite chiralities, and pairs 4 and 5 are made of homochiral peptides of the same chirality (Table 1). Pairs 2 and 3 are mirror images of each other while pairs 4 and 5 are mirror images of each other. Figure 1 shows the photographs of the five pairs in NMR tubes; pair 1 does not gel and remains a clear solution; pairs 2 and 3 form opaque gels; pairs 4 and 5 form translucent gels. From visual observation, gelation is instantaneous for pairs 2 and 3 but much more gradual for pairs 4 and 5. As it has been mentioned in the Experimental Section, the electrostatic repulsion prevents the hydrogelation and/or self-assembly of the individual peptide modules. This is also confirmed by the narrow ¹H NMR signals in the spectra of all individual peptide modules as well as by the absence of their scattering in the control SAXS experiments (see Supporting Information).

**NMR Monitoring of the Assembly Process.** The above visual observations were confirmed by NMR spectroscopy. Figure 2 shows the ¹H NMR spectra for pair 1 (6 months after mixing) and pairs 2–5 (17 h after mixing). For pair 1, the peptide ¹H signals are still very sharp after 6 months, consistent with no gelation. For pairs 2–5, hardly any peptide ¹H signals are left, consistent with extensive gelation because gelled
peptides are no longer detectable by solution NMR due to their extremely short $T_2$ values.\textsuperscript{30} Figure 3 shows the gelation process monitored by NMR through the $^1$H signal from the $\varepsilon$-H of the lysine side chains. The $^1$H signal intensity of pair 1 did not change for 6 months, again consistent with no gelation. Of the four gelling pairs, the two heterochiral pairs 2 and 3 gelled much faster than the two homochiral pairs 4 and 5. However, the eventual extent of gelation, measured by the $^1$H signal intensity from nonincorporated peptides, is almost the same for all four gelling pairs after 18 h of gelation (ca. 98%). Unfortunately, the repetitiveness of the amino acid sequences in pair 1 makes it impossible to study the structure of its assemblies using multidimensional NMR. Therefore, to confirm that pair 1 forms clusters of finite size, the diffusion coefficient, $D$, of the peptides was measured, with a small molecule TFA as the reference point. Figure 4 shows the diffusion coefficient data in the mixture and in each parent peptide solution. The total peptide concentration in all three solutions is 16 mM. Keep in mind that, at pH 7.4, $D,L$-$K$ carries six positive charges, $D,L$-$E$ carries six negative charges, and TFA carries one negative charge. Hence, in the $D,L$-$K$ solution, it is highly likely that the positively charged peptide associates with several copies of the
negatively charged TFA. But such peptide-TFA association is unlikely in the D,L-E solution as both the peptide and TFA are negatively charged. This difference in peptide-TFA association explains why both peptide and TFA have smaller diffusion coefficients in the D,L-K solution than that in the D,L-E solution. In the mixture, the (D,L-K + TFA) complex is replaced by the (D,L-K + D,L-E) complex. The replacement of TFA by D,L-E can be attributed to three factors: (i) D,L-E might interact with D,L-K more strongly because it has multiple carboxylic groups, hydrophobic groups, and H-bond donors/acceptors while TFA has only one of each; (ii) D,L-E (8 mM) is in great excess of TFA (trace amount); (iii) gelation is kinetically much less reversible than TFA binding. Consistent with such replacement, the diffusion of the (D,L-E + D,L-K) complex is slower than both D,L-E and (D,L-K + TFA) as shown in Figure 4. In contrast, diffusion of TFA lies between that of the two parent solutions, indicating that there might still be a portion of TFA bound to the (D,L-E + D,L-K) complex.

**Rheological Characterization of Heterochiral and Homochiral Materials.** Time-sweep rheological monitoring of the gelation process was entirely consistent with visual observations and NMR measurements. No gelation was observed for the syndiotactic pair 1, where the detected values of elastic ($G'$) and viscous ($G''$) moduli were very low (see Supporting Information). In contrast, the two heterochiral pairs 2 and 3 showed fast gelation with $G'$ reaching plateau within 5−10 h (∼5 kPa) with very slight growth afterward. Gelation of the two homochiral pairs 4 and 5 was significantly slower with $G'$ of both gels reaching plateau values around 48 h (∼90 kPa, Figure 5A). From Figure 5B, it can be seen clearly that the heterochiral pairs gelled faster initially but were outpaced by the homochiral pairs around 4−4.5 h. All heterochiral and homochiral hydrogels appear to be fairly stiff materials with the elastic modulus $G'$ significantly higher than the viscous modulus $G''$ (phase angle $\delta = \arctan (G''/G') \sim 4−7^\circ$, see Supporting Information). The frequency spectra for all four pairs point to the formation of very stable materials (Figure 5).
Structural Analysis of Heterochiral and Homochiral Aggregates Using SAXS and SANS. As it has been already mentioned above, the structure of pair 1 assemblies cannot be obtained from NMR spectroscopy because of the sequence repetitiveness of the peptides; therefore, we resort to SAXS to investigate its solution structure. The SAXS scattering profile for pair 1 at 72 h after mixing demonstrates a scattering intensity significantly higher than the individual peptides, consistent with the formation of larger molecular aggregates (Figure 6A). To gain further structural insight into the oligomers formed by pair 1, we resort to Guinier analysis. Often, the analysis of the Guinier region is capable to corroborate the formation of the aggregates of finite size, while the nonlinearity of Guinier plots typically suggests the presence of very different large assemblies, sometimes even of the size beyond the detection limit of SAXS. In the case of pair 1, Guinier plots for rod-like particles (in $Q(I(Q) - Q^2$; bottom inset in Figure 6A) are linear ($Q \sim 0.13-0.22 \, \text{Å}^{-1}$) and show the characteristic upturn pointing to the formation of elongated aggregates of finite length. Here, the scattering data are analyzed only from the viewpoint of the cross-sectional dimensions of the scattering particles since the multiplication of $I(Q)$ by $Q$ essentially removes the length data of the scattering particle. The Guinier plot ln $I(Q)$ vs $Q^2$ (characterizing the particles of the arbitrary shape, or sometimes called globular particles) shown in the top inset in Figure 6A is also linear ($Q \sim 0.02-0.05 \, \text{Å}^{-1}$), and this suggests that pair 1 assemblies into finite aggregates with fairly similar dimensional characteristics. The indirect Fourier transform of the experimental scattering data $I(Q)$ vs $Q$ using GNM results in pairwise distance distribution function $P(r)$ characteristic for elongated assemblies with $R_g = 13.4 \pm 0.4 \, \text{Å}$ and $d_{max} = 55 \, \text{Å}$ (Figure 6B). Of note, we are fully aware that despite such dimensional similarities of the assemblies, aggregation of two oppositely charged peptides could result in differently shaped particles. Still, to get a pictorial understanding of possible 3D shapes of the aggregates, we used the ab initio low resolution shape reconstruction program DAMMIN. This program which is based on the simulated annealing algorithm uses the dummy atom model to reconstruct the shape of the particle solution by minimization of the differences between the model scattering and the experimental X-ray scattering data $I(Q)$ vs $Q$. A series of more than 20 separate runs of DAMMIN were performed resulting in a set of different 3D shapes which, of course, are not unique and are not necessarily absolutely identical with each other. These resulting different 3D models were superimposed using the best-matching alignment program SUPCOMB. This program starts from the inertia-axis alignment of our modeled 3D objects; such alignment is then refined by minimization of the normalized structural discrepancy (NSD) parameter. The NSD value is a quantitative indicator of the structural similarity of the aligning models. NSD = 0, for identical structures, and NSD > 1, for systemically different structures. All of our more than 20 shapes have an NSD ~ 0.4, which speaks in favor of their structural similarity. Their average low resolution 3D model (inset in Figure 6B), is not unique, but gives a general illustration of the possible average shape of the pair 1 aggregate in solution of finite size. The formation of oligomers of finite size from pair 1 is also in a good agreement with NMR results which show pair 1 has a sharp $^1$H spectrum (Figure 2A) but reduced diffusion coefficients (Figure 4).
Contrary to pair 1, the mixing of all other four pairs resulted in the formation of fibrous hydrogel networks. The growth of this fibrous network is evidenced by the growth in SAXS scattering intensity $I(Q)$ vs $Q$ within 72 h of the gelation monitoring as well as from the changes in the zero-angle scattering intensity $I(0)$ from the Guinier analysis of $\ln I(Q)$ vs $Q^2$ plot (Figure 7).

Fully consistent with NMR and rheological data, the homochiral pairs initially have lower $I(Q)$ and $I(0)$ values than the heterochiral pairs, which are indicative of slower initial fiber growth for the homochiral pairs. However, the growth of homochiral fibers soon outpaces that of the heterochiral fibers and the homochiral pairs have higher $I(Q)$ and $I(0)$ values than the heterochiral pairs after a few hours. At 72 h, the zero-angle scattering intensity $I(0)$ from pairs 4 and 5 is about 2 times that of pairs 2 and 3, and about 70 times that of pair 1 (Figure 6A and Figure 7(Center)). From these data, it is clear that stronger gel but slower gelation of the homochiral pairs is related to the morphology of the peptide fibers.

As to the shape and dimension of the peptide fibers, the linearity of the Guinier plots for rod-like particles (see the insets in all panels in Figure 7) points to the formation of elongated asymmetrical assemblies in all gelling pairs, indicative of fiber formation from the very beginning of SAXS monitoring. The length of these fibers is beyond the maximum resolved size of our SAXS setup, which is $\sim$500 Å. However, the cross-section of the fibers is within the detection limit of SAXS; therefore, we modeled the cross sections of the peptide fibers that best fit the scattering data and tracked the changes in these cross sections over time. The cross-section pairwise distance distribution functions, $P_c(r)$, as well as the cross-section shape are shown in Figure 8 for all four pairs. From $P_c(r)$ and from the modeled cross-sectional shapes, $R_c$, $d_{max}$, $S_c$, and $\rho$, all four cross-sectional parameters, $d_{max}$, $R_c$, $S_c$, and $\rho$, behave similarly to the elastic modulus $G'$: within the first few hours, the homochiral values are lower than the heterochiral values; but eventually, the homochiral values are much higher than the heterochiral values. This feature provides a structural explanation to the observed stronger gel but slower gelation associated with homochirality: the heterochiral fibers are formed quicker than the homochiral fibers, but eventually the homochiral fibers are thicker than the heterochiral fibers.

However, the exact homochiral-heterochiral crossing point of $R_c$, $d_{max}$, $S_c$, and $\rho$ (Figure 9) are 2–3 h earlier than that of $G'$ (Figure 5B vs Figure 9). This suggests that structural features at a scale larger than the upper detection limit of SAXS ($\sim$500 Å)

Figure 8. SAXS monitoring of the gelation process. Left and right columns show the time evolution of the 2D average cross-section of the peptide fibers. Four central panels show the consistent growth in the corresponding pairwise distance distribution function of the cross-section, $P_c(r)$, over time (in all model calculations the goodness of fit (its variance from) with respect to experimental data reflected by $\chi^2 \sim 0.5$–0.8). Red: pair 2; green: pair 3; orange: pair 4; blue: pair 5.
also contribute to the mechanical properties of these hydrogels. To explore structural features at a larger scale, we resort to SANS, which has an upper detection limit of \( \sim 2000 \text{ Å} \). SANS data were collected after 96 h of gelation. In full agreement with SAXS results, SANS scattering profiles \( I(Q) \) vs \( Q \) (Figure 10A) demonstrate evident distinction between heterochiral (2 and 3) and homochiral (4 and 5) hydrogels. As expected, the two heterochiral pairs 2 and 3 are very similar to each other, and the same is true for the two homochiral pairs 4 and 5 (Figure 10A). As seen from the linearity of Guinier plots for rod-like particles \( \ln I(Q) \) vs \( Q^2 \) (inset in Figure 10A), both heterochiral and homochiral hydrogels are made of elongated asymmetrical fibers. Interesting structural feature of both homochiral and heterochiral hydrogels follows from the Guinier plots for flat particles \( \ln Q^2 I(Q) \) vs \( Q^2 \) (Figure 10B). Here, the distinct upturn in the region \( Q > 0.06 \text{ Å}^{-1} \) (see, the region at \( Q^2 > 0.0036 \text{ Å}^{-2} \)) suggests the presence of flat structural elements in all hydrogels in addition to the network formed by elongated fibers. Indeed, based on the SANS data in the Q-range starting from \( Q_{\text{min}} \sim 0.005 \text{ Å}^{-1} \) (inaccessible to our SAXS setup) and the grid of \( 30 \times 150 \) dummy atoms (5 Å each), the modeling of average 2D cross-section of the hydrogel fibers reveals such flat webs attached to the fibers. As seen from the 2D shapes (Figure 11A) and their corresponding pair distance distribution functions (Figure 11B), the cross sections of both heterochiral and homochiral hydrogel networks includes the fibers per se with the attached lappet-like webs growing from the side surface of the fibers. The fiber cross-section dimensions obtained from SANS are identical to those obtained from SAXS (compare Figure 11 vs Figure 8 at 72 h).

One could reasonably suggest that the hydrogel networks are formed by the individual fibers interconnected with each other by flat, lappet-like webs. In an attempt to model the greater cross-sectional slice of such interconnected hydrogel networks, we used the SANS data from the lowest Q-range starting from \( Q_{\text{min}} \sim 0.005 \text{ Å}^{-1} \) and the grid of \( 30 \times 150 \) dummy atoms (5 Å each). The resulting pictorial structures (Figure 12) vividly demonstrate that homochiral networks are formed by thicker fibers interconnected by thicker webs than the heterochiral networks, and this translates into higher mechanical strength for the homochiral networks. At present, it is not clear, however, why some of the peptide modules do not incorporate into the fibers, and form the web-like connections instead. One possible explanation could be that the residual surface charges of the fibers serve as nuclei for lateral growth resulting in flat, interconnecting structures. To the same extent, since the length of the fiber, as it has been mentioned above, could not be reliably determined from our SAX(S)N data, it is hard to estimate the relationship between the fiber length and the longitudinal extension of the webs illustrated by the schematic cartoon in Figure 12C.

Of note, the combined application of NMR, dynamic rheometry, and small-angle scattering techniques to monitor the gelation process show a discrepancy with regard to the time point of the completion of gelation (cf. Figure 3, Figure 5A, and Figure 9). For NMR, which traces the decay of the free peptide concentration as a result of its incorporation into the hydrogel fiber, this amounts to about 14 h (Figure 3). Dynamic rheometry, on the other hand, shows that the mechanical properties of these hydrogels evolve over a longer period of time.
strength ($G'$) of the hydrogels do not change significantly after about 30 h and reaches plateau at about 48 h (Figure 5A). As for SAX(N)S, the dimensional parameters of the fiber cross-section appear to grow up to 72 h and remain unchanged between within the interval 72–96 h (Figure 8 and Figure 11).

On the basis of these observations, one might suggest that after the incorporation of the peptides into the fiber, accompanied by the growth of the fibrous network, further restructuring of the fibers per se do not contribute significantly to the mechanical properties of bulk material. Yet, it is quite possible that in addition to the individual fiber parameters, other factors also contribute to the mechanical properties of the hydrogels. One of such factor, for example, could be the persistence length of the fibers, $L_p$, see, for example, MacKintosh theory, and the issue of the determination of $L_p$ in hydrogel fibers from the SAX(N)S data is worthy of future careful study.

In summary, the time course and the morphology of the peptide aggregates in the five pairs provide a structural explanation to the differences in their material properties. The syndiotactic pair 1 forms finite size aggregates (hetero-dimer) with no fibrous network. Macroscopically, this pair remains a clear solution. Both heterochiral pairs and

Figure 10. SANS data collected after 96 h of gelation. (A) Scattering profiles $I(Q)$ vs $Q$ of all four gelling pairs. Guinier plots for rod-like particles, $Q \times I(Q)$ vs $Q^2$, are shown in the inset, and the linearity in this region (for $Q$ from 0.007 to 0.014 Å$^{-1}$) indicates the formation of elongated fibers in all hydrogels. (B) Guinier plots for flat particles, $Q^2 \times I(Q)$ vs $Q^2$, and the linearity in this region (for $Q$ from 0.08 to 0.14 Å$^{-1}$ shown by two arrows in the figure) indicates the formation of flat, lappet-like webs interconnecting the peptide fibers in all hydrogels. Statistical error bars correspond to one standard deviation and represent error in the scattering intensity estimation. Red: pair 2; green: pair 3; orange: pair 4; blue: pair 5.

Figure 11. Analysis of the SANS data. (A) 2D average cross-section of individual peptide fibers, including the flat, lappet-like webs interconnecting the peptide fibers. (B) Corresponding cross-section pairwise distance distribution functions, $P_c(r)$, for respective homochiral and heterochiral fibers shown in (A) (in all model calculations $\chi^2 \sim 1.0–1.3$). Red: pair 2; green: pair 3; orange: pair 4; blue: pair 5.

Figure 12. Pictorial presentation of the 3D slice of the hydrogels under study showing the cross sections of the individual fibers interconnected with flat, "lappet-like" webs shown in gray. Reconstruction from the 2D cross-shape restored from SANS data with the low $Q_{\text{min}}$ values ($<0.003–0.005$ Å$^{-1}$): (A) for the heterochiral hydrogel pair 2 ($D$-K$+$L-E) and (B) for the homochiral hydrogel pair 5 ($L$-K$+$L-E). Cross-sectional total size of the slice in both cases is 170 Å × 500 Å, in all model calculations $\chi^2 \sim 1.0–1.3$. (C) shows a pictorial cartoon illustrating the schematic fibrous network organization.
homochiral pairs form fibrous networks, and the fibers are interconnected by flat lappet-like webs. Macroscopically these pairs exist as solid-like hydrogels. As shown by SAXS analysis, the heterochiral pairs quickly form fibers, but these fibers show limited growth after a few hours. In contrast, the homochiral pairs have slow initial fibrillization, but the fibers grow steadily for up to 72 h. Eventually, the homochiral fibers outgrow the heterochiral fibers, resulting in a network made of thicker and denser fibers, which leads to higher $G$ and $\gamma_{\text{fibril}}$.

As to why the syndiotactic pair 1 forms oligomers of finite size, and, among the isotactic pairs, why the homochiral pairs 4 and 5 form thicker fibers than the heterochiral pairs 2 and 4, the NMR, SAXS, and SANS analyses conducted in this work cannot provide a definitive answer. One possible explanation could be that, similar to amyloid-$\beta$-proteins $\beta\text{-40}$ and $\beta\text{-42}$, some peptide pairs form “open” oligomers that are prone to grow while some peptide pairs form “closed” oligomers that are difficult to grow. Such “openness” or “closedness” could be defined by the degree of electrostatic charge compensation in the aggregates as well as by the availability of probable $\beta$-sheet links and/or hydrophobic contacts—all stemming from the structural differences between L- and D-enantiomers. Our more detailed research in this area is underway.

■ CONCLUSIONS

For hydrogels co-assembled from a pair of self-repulsive but mutually attractive oppositely charged oligopeptides, chirality is shown to be an influential factor in determining the rate of gelation as well as the mechanical properties of the resulting biomaterial. Homochirality is associated with stronger gels but slower gelation. In other words, homochirality confers mechanical advantage but heterochirality confers kinetic advantage to this class of biomaterials. Structurally, homochiral peptide pairs form networks made of thicker and denser fibers while heterochiral peptides pairs form networks more quickly. The observed mechanical advantage posed by homochirality provides another angle to assess its role in biology. The generality of this observation and its relevance to the origin of life awaits further investigation.

■ ASSOCIATED CONTENT

1 Supporting Information

Analytical HPLC and ESI-MS spectra of the peptides; the photo of the in-house designed simple humidifier for sealed-cell rheometer; detailed rheological data on all peptide systems, NMR spectra, and SAXS data on the individual peptides and model fits to the experimental SANS and SAXS data. This material is available free of charge via the Internet at http://pubs.acs.org.

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Notes
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