Aromatic—Aromatic Interactions Enhance Interfiber Contacts for Enzymatic Formation of a Spontaneously Aligned Supramolecular Hydrogel

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Supporting Information

ABSTRACT: Anisotropy or alignment is a critical feature of functional soft materials in living organisms, but it remains a challenge for spontaneously generating anisotropic gel materials. Here we report a molecular design that increases intermolecular aromatic–aromatic interactions of hydrogelators during enzymatic hydrogelation for spontaneously forming an anisotropic hydrogel. This process, relying on both aromatic-aromatic interactions and enzyme catalysis, results in spontaneously aligned supramolecular nanofibers as the matrices of a monodomain hydrogel that exhibits significant birefringence. This work, as the first example of monodomain hydrogels formed via an enzymatic reaction, illustrates a new biomimetic approach for generating aligned anisotropic soft materials.

Anisotropy or alignment is a critical feature of functional materials in living organisms or man-made systems. For example, muscle cells rely on sophisticated anisotropic protein filaments to function, and liquid crystal display is based on the electronic static alignment of a simple anisotropic molecule (e.g., 4-cyano-4'-pentylbiphenyl). Recognizing the importance of anisotropy, materials scientists are striving to create high quality anisotropic materials. While generating anisotropic materials from inorganic solids is relatively easy and successful, the introduction of anisotropy to synthetic soft organic materials, particularly hydrogels, remains less developed and a considerable challenge despite the fact that biological anisotropic soft materials with substantial complexity prevail in living organisms. Although several approaches have shown effectiveness for inducing anisotropy in gels, including the use of an electrical field, the inclusion of liquid crystals, and the utilization of mechanical stretching or shear force from flow, each process still has its limitation for creating anisotropy in hydrogels because hydrogels contain a large amount of water that usually conducts electricity, excludes common liquid crystals, and favors relaxation to the isotropic state. Thus, better approaches are needed for spontaneously generating inherently anisotropic hydrogels.

Based on the anisotropic hydrogel in nature (e.g., corneal stroma formed mainly by regularly arranged collagen fibers), one attractive approach is to use molecular self-assembly to generate anisotropic structures that immobilize water and form hydrogels. In fact, recent advances in the development of supramolecular gels make this approach particular promising because the self-assembly of hydrogelators usually affords nanofibers, which is inherently anisotropic. However, these nanofibers, being long and flexible, not only physically cross-link to form the matrices of gels but also adopt random entanglement, which results in isotropic hydrogels in most cases. Encouragingly, several exceptional examples of supramolecular hydrogels exhibit birefringence, thus offering useful hints for creating anisotropic hydrogels. For example, Stupp and co-workers demonstrated that a thermal pathway can convert isotropic solutions of peptide amphiphiles to a strongly birefringent hydrogel, in which molecules self-assemble to form large arrays of aligned nanofibers. Moreover, the addition of calcium ions into the solution of a phosphate containing peptide amphiphile results in a gel that exhibits birefringence, which coincides with the parallel alignment of the nanofibers shown in the transmission electron micrograph (TEM) of the gels. These results and other examples suggest that the enhancement of interfiber interactions should be a reliable molecular approach to induce alignment of nanofibers and thus generate anisotropy.

Although ionic forces or overlaps between alkyl chains are able to increase the interfiber interactions, ionic interactions are intrinsically isotropic and interactions between alkyl chains are relatively weak and inefficient. Therefore, we decided to explore the use of aromatic–aromatic interactions to enhance interfiber interactions that are necessary for creating anisotropic supramolecular hydrogels because (i) aromatic–aromatic interaction is not only stronger than the van der Waal’s interaction (London dispersion force) between alkyl chains but also inherently directional due to the plane-to-plane or edge-to-plane orientation; (ii) nature has extensively used aromatic–aromatic interaction as a stabilizing force for generating ordered structures in proteins; and (iii) aromatic rings have relatively compact volumes; thus, the interactions between aromatic rings lead to a more predictable and efficient self-assembly of the molecules in aqueous phase for the formation of mechanically strong or stable supramolecular hydrogels.

Besides the use of proper interfiber interactions, it is necessary to choose an appropriate pathway for molecular self-assembly. Instead of using a thermal pathway, we decided to use an enzymatic pathway because cells use enzymatic conversion to form anisotropic hydrogels and the enzymatic

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self-assembly process has emerged as a powerful method for generating ordered nanostructures and supramolecular hydrogels for a wide range of applications. For example, enzymatic self-assembly has led to supramolecular hydrogels outside cells and inside cells in vitro and in vivo, and resulted in highly stable hydrogels that are unavailable via other pathways (e.g., thermal or pH). Recently, enzyme catalyzed self-assembly has provided a route for controlled drug release and for the exploration of dynamic self-gels for a wide range of applications.

To address the above-mentioned question, we designed two pairs of precursor/hydrogelator (i.e., 1a/1b and 2a/2b) differing only in one phenylalanine (Phe) residue and examined their enzymatic hydrogelation by using polarized optical microscopy, transmission electron microscopy (TEM), and rheology. Our study shows that the addition of an enzyme (e.g., alkaline phosphatase (ALP)) into the solution of 1a affords an aligned monodomain supramolecular hydrogel of 1b, which consists of aligned nanofibers (Figure 1). Having one less phenylalanine residue, 2b forms a nonbirefringent hydrogel that consists of a network of randomly oriented nanofibers. In addition, the hydrogel of 1b (or 2b) formed by the change of pH exhibits little birefringence and consists of few aligned nanofibers. Thus, this work, as the first example of monodomain supramolecular hydrogels formed by an enzymatic reaction, illustrates a new, effective approach for generating anisotropic supramolecular soft materials, which ultimately may lead to the formation of aligned nanostructures directly in a cellular environment.

Figure 1B shows the detailed molecular design of the two pairs of precursor/hydrogelator. The precursor 1a, as a pentapeptidic derivative, consists of a (naphthalene-2-ly)acetyl group (Nap), three phenylalanine residues (Phe-Phe-Phe), a lysine (Lys), and a tyrosine phosphate moiety (−O−P−Tyr). The precursor 2a, as an analog of 1a, consists of Nap, Phe-Phe, Lys, and −O−P−Tyr. Nap enhances intermolecular aromatic−aromatic interactions, the phenylalanine residues (Phe), besides providing aromatic−aromatic interactions, constitute the peptide backbone with Lys to provide the donors and acceptors for intermolecular hydrogen bonds. Since ALP catalytically dephosphorylates −O−P−Tyr to Tyr to form the hydrogelators, 1b and 2b, the phosphate group on Tyr acts as a trigger for enzymatic hydrogelation. The hydrogelator 1b, bearing three phenylalanine residues, likely has stronger intermolecular aromatic−aromatic interactions than 2b (bearing two phenylalanine residues) does. After the molecular design, we used solid phase peptide synthesis (SPPS) to synthesize 1a, 2a, 1b, and 2b (Scheme S1).

As a soluble precursor, 1a (or 2a) (4 mg) dissolves in distilled water (0.5 mL) to a final concentration of 0.8 wt % and pH = 7.4. After the addition of ALP (2 U/mL), the solution of 1a (or 2a) transforms to a transparent hydrogel (Figure S1, here denoted as Gel1b (or Gel2b)) within 2 h because the precursor 1a (or 2a) becomes the corresponding hydrogelator, 1b (or 2b). Gel1b, being placed between the cross polarizers, exhibits little birefringence when the vial is at 90° and 0° with polarizers (Figure 2A) but displays significant birefringence when the vial is at 45° with the polarizers (Figure 2B). This result indicates that the nanofibers of 1b in Gel1b have largely uniform alignment to result in a significant birefringence. In contrast, Gel1b hardly shows any birefringence in either of the orientations (90° and 0° in Figure 2C and 45° in Figure 2D). This result indicates that the nanofibers of 2b in Gel2b likely have random orientations. To further verify the alignment of nanofibers in Gel1b, we carried out the enzymatic gelation of Gel1b in two capillaries with a diameter of 0.3 mm. As shown in Figure 2E, F, Gel1b barely exhibits birefringence when the capillaries are at 90° and 0° with the polarizers (Figure 2E) but displays bright birefringence when the capillaries are at 45° with the polarizers (Figure 2F). In addition, light extinguishes at the crosspoint of two capillaries containing Gel1b (Figure 2G), confirming uniform alignments of the nanofibers of 1b in Gel1b. Although the change of the solution pH of 1b (or 2b) also results in a hydrogel (Figure S2A (or S3A), denoted as Gel′(1b) (or Gel′(2b))) at the concentration of 0.8 wt % and pH of 7.4, Gel′(1b) shows sporadic, random, and orientation-independent birefringence (Figure S2B and S2C), and Gel′(2b) hardly shows...
any birefringence (Figure S3B and S3C). These results indicate enzymatic hydrogelation as a feasible and effective pathway to generate an aligned, monodomain supramolecular hydrogel.

To quantify the birefringence in the hydrogels, we used polarized microscopy to image the hydrogels made of different samples (i.e., 1b and 2b) and by different pathways (i.e., enzyme or pH). Birefringence is a material property that derives from molecular alignment, which results in a fast and a slow axis in the material. The polarization of light changes due to a differential phase shift between two orthogonal polarized components, one of which, being parallel to the fast axis, travels faster than the other that is parallel to the slow axis. Retardance ($\Delta n$) signifies the differential phase shift between two wavefronts of the orthogonal components that have traveled through a birefringent material and is typically measured as a distance in the unit of nm (Figure 1A). Polarized microscopy with the OpenPolScope method24 measures the retardance of every resolved point of specimen and converts the results of computation into a color gradient of the retardance images. Thus, we used PolScope to obtain retardance images of Gel$_{1b}$, Gel$_{2b}$, Gel$_{1b'}$, Gel$_{2b'}$, and a solution of 1a or 2a. As shown in Figure 3A, the thin film of Gel$_{1b}$ contains largely uniform domains (with the sizes of hundreds of micrometers) that exhibit a retardance of approximately 35 nm or larger. Gel$_{1b'}$, however, exhibits mostly multiple domains that have retardance between 0 and 25 nm (Figure 3C). As shown in Figure 3B and 3D, Gel$_{2b}$ and Gel$_{2b'}$, under PolScope, give a retardance value of almost 0 nm, confirming that Gel$_{2b}$ and Gel$_{2b'}$ are nonbirefringent. As expected, a solution of 1a or 2a at the concentration of 0.8 wt % and pH of 7.4 are nonbirefringent, as proven again by PolScope imaging (Figure S4). These results further support that the combination of strong aromatic–aromatic interactions and enzymatic hydrogelation facilitates the formation of anisotropic supramolecular hydrogels.

Transmission electron microscopy (TEM) also confirms the orientation orders of the nanofibers in Gel$_{1b}$. As shown in Figure 4A, the nanofibers in Gel$_{1b}$ have diameters ($d$) of $8 \pm 2$ nm and form exceptionally long, aligned bundles that stretch over many micrometers (Figure S5), suggesting strong interfiber interactions. In contrast, TEM shows that Gel$_{1b'}$ contains a large amount of randomly entangled nanofibers ($d = 8 \pm 2$ nm) that form a typical network (Figure 4B) as those found in conventional supramolecular hydrogels. A TEM image of Gel$_{1b'}$ indicates a mixture of parallel aligned nanofibers and entangled nanofibers ($d = 8 \pm 2$ nm) (Figure S6A and S6B). Gel$_{1b'}$, however, contains entangled nanofibers ($d = 8 \pm 2$ nm) to form a network, exhibiting the same morphology as that of Gel$_{1b}$ (Figure S6C and S6D). The agreement between the TEM images and PolScope images confirms that the alignment of the nanofibers formed during enzymatic hydrogelation is responsible for birefringence of Gel$_{1b}$.

To further verify the results in TEM images, we also compared the rheological properties of the hydrogels. Figure 5 shows the strain and frequency dependence of dynamic storage moduli ($G'\prime$) and loss moduli ($G''\prime$) of all four gels are larger than those of their loss moduli ($G''\prime$), indicating that all the samples behave as viscoelastic gel materials. The values of $G'\prime$ of the hydrogels change little during the frequency sweep (from 0.1 to 200 rad/s) (Figure S7), suggesting that the matrices of those gels have good tolerance to external shear force. In agreement with TEM data, Gel$_{2b'}$ which consists of entangled nanofibers (more cross-linking), shows stronger mechanical strength than Gel$_{1b}$ which contains parallel aligned nanofibers (less cross-linking). On the other hand, when both of them have similar nanofiber networks (Figure S6), Gel$_{1b}$ exhibits a slightly higher storage moduli than Gel$_{2b'}$, agreeing with that hydrogelator 1b has one more phenylalanine residue to provide stronger aromatic–aromatic interactions than 2b does.

In conclusion, this communication describes a rational design that utilizes aromatic–aromatic interactions to promote interfiber contacts between nanofibers and to drive the alignment of nanofibers for producing inherently anisotropic supramolecular hydrogels via enzyme catalyzed molecular self-assembly. Although direct evidence of aromatic–aromatic interfiber interaction remains to be established, the crystal structure of the self-assembly motif (Nap-Phe-Phe)$_2$22 in 1 strongly supports the notion of aromatic–aromatic interfiber
interactions. In addition, intermolecular aromatic–aromatic interaction from the overlapping of phenyl and/or naphthyl groups, which are responsible for forming single molecular width nanofibers, should favor interfiber interactions. The alignment of the nanofibers likely stems from the stereochemical cooperation between phosphatases and the precursors/hydrogelators because the use of the enantiomer of 1a for enzymatic hydrogelation hardly results in the alignment of fibers (Figure S7). Since the hydrogel directly prepared with 1b by a change in pH exhibits little birefringence and consists of few aligned nanofibers, enzymatic conversion is indispensable for the alignment of the nanofibers. The increase of the concentration of enzyme speeds up the gelation process, but shows little influence on birefringence or alignment. Our work, for the first time, illustrates PolScope imaging is a useful and effective method to study anisotropy of supramolecular hydrogels. By establishing enzyme catalysis as a new pathway to complement other processes for generating inherently anisotropic hydrogels, this work may lead to the production of aligned nanostructures in biological systems or living organisms.

**ASSOCIATED CONTENT**

* Supporting Information

The details of the synthesis, NMR spectra and LC-MS data for all compounds, and rheological data. This material is available free of charge via the Internet at http://pubs.acs.org.

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Notes

The authors declare no competing financial interest.

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