D-Periodic Collagen-Mimetic Microfibers

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Abstract: Self-assembling peptides have been previously designed that assemble into macroscopic membranes, nanotapes, and filaments through electrostatic interactions. However, the formation of highly ordered collagen-like fibrils, which display D-periodic features, has yet to be achieved. In this report, we describe for the first time a synthetic peptide system that self-assembles into a fibrous structure with well-defined periodicity that can be visualized by transmission electron microscopy (TEM). Specifically, we designed and synthesized a peptide that utilizes charged amino acids within the ubiquitous Xaa-Yaa-Gly triad sequence to bias the self-assembly into collagen-like homotrimeric helices that are capable of fibrillogenesis with the production of D-periodic microfibers. Potential molecular mechanisms for peptide assembly into triple-helical protomers and their subsequent organization into structurally defined, linear assemblies were explored through molecular dynamics (MD) simulations. The formation of thermodynamically stable complexes was attributed to the presence of strong electrostatic and hydrogen bond interactions at staggered positions along the linear assembly. This unexpected mimicry of native collagen structure using a relatively simple oligopeptide sequence establishes new opportunities for engineering linear assemblies with highly ordered nano- and microscale periodic features. In turn, the capacity to precisely design periodic elements into an assembly that faithfully reproduces these features over large length scales may facilitate the fabrication of ordered two- and three-dimensional fiber networks containing oriented biologically, chemically, or optically active elements.

Introduction

The oriented axial assembly of supercoiled protein motifs represents the primary structural feature associated with the load-bearing properties of native protein-based fibrils such as collagen and α-keratin.1−3 As a principal building block or “protomer”, supercoiled motifs assemble into supramolecular fibrils and macroscopic fibers through specific recognition that occurs at the interfaces between these elements, such that linear propagation occurs along the incipient fibril axis.4−6 The structural features that guide self-assembly are encoded within the sequences of the peptide subunits and include complementary packing of side chains at the interface, as well as electrostatic, hydrophobic, and hydrogen-bonding interactions.

Given the richness of the molecular-scale information programmed within proteins, it should not be surprising that synthetic materials have yet to recapitulate the self-assembly behavior of native protein-based, multistranded helical assemblies. Thus, elucidating the design principles that underlie the self-assembly of fibrillogenic proteins into structurally defined supramolecular structures is of interest not only in generating synthetic polypeptides that mimic native structural proteins but also for the design of new materials with biological, chemical, and mechanical properties that exceed those of currently available synthetic polymers.

The triple-helical domain of native collagen comprises a strict recurrence of tripeptide repeat sequence Xaa-Yaa-Gly and displays precise registration of individual amino acids in adjacent helices within a defined oligomerization state imparting structural specificity and fibril-forming properties. Proline (Pro) and (4R)-hydroxyproline (Hyp) occupy the Xaa and Yaa positions, respectively, of the triplet repeats at the highest level of statistical frequency within native collagen sequences. The high imino acid content of collagen sequences is necessary for the stabilization of its triple-helical structure, although alternative substitutions are often observed in native collagen sequences and may be necessary to fully define its biological role. We believe that
the synthesis of collagen-mimetic triple helix peptide protomers (THPs) that display the capacity to form triple helices with improved stability and that exhibit a propensity to form linear assemblies through a process of axially oriented alignment will prove to have a number of important practical applications in the design of novel biomaterials. Moreover, the capacity to mimic collagen’s ability to form linear assemblies with precisely defined periodic features would represent an important milestone in the design of ordered two- and three-dimensional fiber networks containing oriented chemically, biologically, or optically active elements.

In this report, we describe the solid-phase synthesis of a collagen-mimetic THP in which the sequence was designed to exhibit linear fibril growth upon assembly of the triple-helical structure. In addition, molecular dynamics (MD) simulations were used to assess potential molecular mechanisms for peptide assembly into triple-helical protomers and their subsequent organization into structurally defined, linear assemblies. Sequence specific peptides were generated comprised of three different Xaa-Yaa-Gly domains, including a central core of Pro-Hyp-Gly repeat sequences, including a central core of Pro-Hyp-Gly repeat sequences flanked by distinct sets of peptide repeats containing either negatively (Glu) or positively (Arg) charged amino acid residues (Figure 1). The Pro-Hyp-Gly peptide sequence forms the structurally critical hydrophobic core of the assembly, which is responsible for maintaining the thermodynamic stability of the collagen triple-helical structure. If positioned appropriately at the N- and C-terminal portions of the THP, MD simulations in association with transmission electron microscopy demonstrated that the electrostatic interactions of the charged residues (Arg and Glu) facilitated a preferential linear oligomerization of protomers within fibrils and promoted a staggered orientation between adjacent triple helices that was similar to that observed in native fibrous collagens. Likewise, MD simulation illustrated that linear oligomerization was reinforced by hydrogen bonds formed within the N- and C-terminal interface of adjacent THPs. Self-assembling peptides have been previously designed that assemble into macroscopic membranes, nanotapes, and filaments through electrostatic interactions. However, the peptide described herein is the first to yield triple-helical protomers that undergo collagen-like fibrillogenesis through complementary electrostatic interactions that are reinforced by hydrogen bond formation. Significantly, synthetic D-periodic collagen-mimetic fibers have been generated through the spontaneous self-assembly of short triple-helical oligopeptides, which heretofore has not been previously achieved.

Figure 1. Collagen-mimetic peptide, triple-helical protomer, and fibril assembly. (A) Amino acid sequence of synthetic collagen-mimetic peptide CPII indicating the distinct domain structure of collagen triads. (B) Proposed triple-helical wheel diagram of CPII, a synthetic collagen-like peptide sequence, adopting 7/2 superhelical symmetry. (C) Side view illustrating proposed interhelical electrostatic interactions driving higher order assembly of a CPII homotrimer to yield triple-helical protomers. (D) Schematic representation of CPII fibrillogenesis through axially staggered assembly between oppositely charged N- and C-terminal domains of triple-helical peptide protomers.
Results

Determination of Triple-Helical Conformation of CPII. The collagen-mimetic peptide, NH-(Pro-Arg-Gly)$_4$-(Pro-Hyp-Gly)$_4$-(Glu-Hyp-Gly)$_4$-COOH, CPII, was synthesized by a solid-phase synthetic strategy (Supporting Information). Circular dichroism (CD) spectroscopy was performed to ascertain the triple-helical character of the synthesized peptide in various buffer solutions, as well as related thermal transition curves (Figure 2A). CD spectra in various buffer solutions exhibited a positive peak at 221–222 nm, a crossover peak at 213–214 nm, and a negative peak at 197–198 nm. These spectral positions are very similar to those of native collagen and collagen-mimetic peptide assemblies and, therefore, are consistent with the triple-helical conformation through homotrimer formation.

Thermal unfolding studies were also performed to confirm the presence of triple-helical conformation of CPII in various buffer solutions (Figure 2B). Thermal denaturation demonstrated melting temperatures ($T_m$) ranging from 37 to 48 °C. The highest $T_m$ was noted in PBS, and the lowest was noted in a buffer solution of 30 mM $Na_2HPO_4$, 30 mM TES, 135 mM NaCl (ionic strength 0.225), pH 7.4. (C) Turbidity profiles for collagen-mimetic peptide CPII determined at concentrations between 0.57 and 1.42 mg/mL and temperatures between 26 and 37 °C. Absorbance was acquired at 313 nm as a function of time and turbidity presented as $A_{313} \times 2.303$.

Figure 2. Circular dichroism spectroscopy and turbidimetric analysis of CPII collagen-mimetic peptides. (A) CD spectroscopy of collagen-mimetic peptide CPII (0.5 mg/mL). Mean residue molar ellipticity $[\theta]$ 10 deg cm$^2$ dmol$^{-1}$ as a function of solvent system at 4 °C. (B) First derivative plots, $d[\theta]/dT$, of thermal denaturation profiles for CPII in varying solvent systems. Buffer 1: H$_2$O, Buffer 2: 10 mM HCl, Buffer 3: 10 mM MOPS (pH 6.85), Buffer 4: PBS (pH 7.25), Buffer 5: 10 mM MES (pH 6.0), Buffer 6: 10 mM HEPES (pH 7.0), Buffer 7: 10 mM CHES (pH 9.0), Buffer 8: 30 mM mM $Na_2HPO_4$, 30 mM TES, 135 mM NaCl (ionic strength 0.225), pH 7.4. (C) Turbidity profiles for collagen-mimetic peptide CPII determined at concentrations between 0.57 and 1.42 mg/mL and temperatures between 26 and 37 °C. Absorbance was acquired at 313 nm as a function of time and turbidity presented as $A_{313} \times 2.303$. 

\[ \text{Absorbance} = (A_{313} \times 2.303) \]

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strength 0.225), pH 7.4. The latter is similar to the reported melting temperature of ca. 40 °C for monomeric type I collagen in a similar buffer solution (Supporting Information). 10

**Turbidity Analysis of Peptide Assembly in Solution.** As an initial measure of fibrillogenesis, turbidity profiles were determined for CPII at varying peptide concentrations and solution temperatures in 30 mM TES, 30 mM phosphate, and 135 mM NaCl after initial “thermal annealing” by heating the peptide solution to 75 °C for 40 min followed by cooling to room temperature (Figure 2C). Similar to collagen fibrillogenesis, growth phases were consistent with a first-order rate process that required a minimum peptide concentration to trigger peptide self-assembly. However, a number of features of CPII aggregation were distinct from collagen fibrillogenesis, including shorter lag phases and a decline in turbidity after an initial plateau. The latter phenomenon is consistent with sedimentation of CPII peptide aggregates, which, unlike monomeric collagen, do not form a physical network or gel. In addition, we observed that the growth rate for CPII self-assembly decreased with increasing temperature. Leikin et al. 11 and others 10,12 have suggested that fibrillogenesis is favored as the temperature approaches the \( T_m \) of collagen due to the release of structured water, increasing mobility of side chains, and loosening of the triple helix, all of which may promote structural interactions that drive collagen self-assembly. However, Glu-Hyp-Gly guest triplets may decrease the propensity toward triple-helix formation in (Pro-Hyp-Gly) containing host peptides. 13 Thus, given the relatively small size of CPII compared to collagen, local unfolding of the negatively charged C-terminal domain with increasing temperature may limit the thermodynamic stability of self-assembled structures. As a consequence, although the natural logarithm of the rate constants was linearly dependent on 1/T, obeying the Arrhenius relationship, the calculated activation energy for the growth phase of CPII was negative (−74 kJ mol\(^{-1}\)).

**TEM Analysis of CPII Fibril and Microfiber Formation.** CPII self-assembly, initiated as described for turbidity studies, was examined at varying time intervals by transmission electron microscopy. In the absence of thermal annealing, nonbanded fibers with average lengths of 3.0–4.0 μm and diameters of 12–15 nm were observed (Figure 3).

However, after thermally annealing solutions of CPII, fiber growth proceeded within several hours by initial formation of smooth fibrils that were hundreds of nanometers in length and tens of nanometers in diameter (Figure 4). These fibrils displayed tapered tips similar to the tactoidal ends of native collagen fibers from which continued fiber growth is thought to occur.
Samples examined after longer incubation periods demonstrated the formation of uniform micron-length fibers that were approximately 70 nm in diameter with well-defined transversely banded structure akin to the D-periodicity of collagen (Figure 5). Analysis of high magnification images revealed that the length of D-periodic gap and overlap regions was approximately 18 nm. Of note, fibril formation did not occur in the absence of salt (e.g., water or 10 mM HCl) and at high phosphate concentrations (i.e., 60 mM Na$_2$HPO$_4$), only nonbanded, poorly formed linear aggregates were noted.

Molecular Simulations of CPII Triple-Helical Protomers and Higher Order Assemblies. Molecular dynamics simulations were performed to examine potential structural features that contribute to the assembly of CPII THPs, as well as to the formation of axially oriented staggered protomer–protomer assemblies. A system of individual CPII peptides, in parallel or antiparallel orientation, reached an energy minimum after 5 ns in a NpT dynamics simulation at 23 °C and 1 atm (Figure 6 and Supporting Information). A thermodynamically stable, close-packed, triple-helical protomer was observed for both systems, and the presence of a significant number of hydrogen bonds contributed to their thermodynamic stability. Although the distance matrix and the number of hydrogen bonds between the peptides were similar for both THP systems, we observed a significant difference in the distribution of electrostatic potential with strong positive and negative potentials noted at the termini of THPs produced from parallel oriented CPII peptides. Significantly, subsequent simulations of this system yielded a linear assembly of head-to-tail, axially oriented protomers that was promoted by the presence of substantial electrostatic and hydrogen bonding interactions, which were much more favored than THPs generated from antiparallel oriented CPII peptides.

Additional simulations were conducted to investigate whether these linear peptide assemblies display a propensity for further fibrillogenesis-like growth. Specifically, we simulated the interactions of three linear complexes, each produced from four THPs that were initially assembled from three CPII peptides in parallel orientation. After a 10 ns NpT molecular dynamics simulation at 23 °C and 1 atm, an energy minimum was achieved with formation of thermodynamically stable staggered complexes attributed to strong electrostatic effects at the N- and C-termini of THPs and hydrogen bond interactions between THPs arising from parallel oriented peptides. Indeed, in a manner similar to collagen self-assembly, these linear fibril-like structures displayed staggered alternating high and low charge density domains. In summary, both the experimental data and the results of molecular simulations suggest that staggered complex formation of higher order THP multimers gives rise to the observed D-periodic collagen-mimetic microfibers (Figure 7).

Discussion

The association of three peptide chains that interact to form triple-helical protomers initiates collagen supramolecular structures. Native protomers have at least one triple-helical collagenous domain and two noncollagenous domains of variable sequence and size that are positioned at the N- and C-termini. Distinct supramolecular structures are formed by the oligomerization of protomers through interactions that involve the formation of end-to-end connections, lateral associations, and supercoiling of helices. However, the critical protomer–protomer recognition steps that result in assembly of collagen into a fibril or fiber network remain poorly defined.

Recently, there have been several reports of collagen-like peptides that self-associate to higher molecular weight linear structures through the presence of chemical moieties at the N- and C-termini that promote the formation of cysteine knots,\(^\text{15}\) facilitate native chemical ligation,\(^\text{16}\) or lead to noncovalent aromatic-stacking interactions.\(^\text{17}\) Thus far, none of these systems have yielded D-periodic microfibers that are characteristic of collagen. In this report, we designed and synthesized a peptide, CPII, that utilizes charged amino acids within the ubiquitous Xaa-Yaa-Gly triad sequence to bias the self-assembly into collagen-like homotrimeric helices that are capable of fibrillogenesis with the production of D-periodic microfibers.


Glutamic acid and arginine are the most common charged amino acids found in collagen, and investigations of synthetic peptides have demonstrated that relatively stable triple helices can be produced when these residues are found in the Xaa and Yaa positions of the triplet, respectively. Differences can be produced when these residues are found in the Xaa and Yaa positions of the triplet, respectively. However, differences in triple-helix stability have been noted among a range of evaluated tripeptide repeats that have been introduced into guest sites in collagen-mimetic peptide sequences. These studies have demonstrated that highly stable motifs such as Pro-Hyp-Gly can stabilize the collagen structure such that Glu-Hyp-Gly repetitive triplets, which have a weaker propensity for triple-helix formation, can be incorporated into the collagen sequence. In the sequence of CPII, (Pro-Hyp-Gly)$_4$ formed a critical hydrophobic core, which facilitated the capacity of N- and C-terminal (Pro-Arg-Gly)$_4$ and (Glu-Hyp-Gly)$_4$ domains to reinforce the preferred linear oligomerization within a fibril and promote staggered orientation between adjacent fibrils in the proposed structure of the collagen-mimetic fiber. Of note, CPII shows a high propensity for self-association following a nucleation-growth mechanism even at lower concentrations (<1.0 mg/mL) and neutral pH. This observation contrasts sharply with the self-assembly behavior of the (Pro-Hyp-Gly)$_{10}$ homotrimer, in which TEM studies indicated the presence of nonspecific aggregates only at much higher concentrations (ca. 7 mg/mL).

The significance of our electrostatically driven approach lies in the observation that self-assembly of this protomer produced D-periodic microfibers that resemble native fibrous collagen. D-periodicity in collagen has its origins in differential intermolecular interactions that occur between protomers, which are mediated either through electrostatic interactions between oppositely charged side chains or through hydrophobic interactions. These interactions are maximized when each protomer is displaced axially with respect to its nearest neighbor. In a parallel molecular assembly, the repeated expression of this staggered association generates a D-periodic structure. Each D-period is made up of two zones, one containing higher charge density than the other. This staggered structure of unequal molecular charge density accounts for the characteristic banding pattern of fibrils stained with uranyl salts. Uranyl acetate is only weakly dissociated in aqueous solution and exists as anionic and cationic complexes that bind to charged side chains. The repeating darkly staining transverse bands are thought to be a result of preferential penetration of electron-dense uranyl ions into regions of lowest packing density. The D-periodicity of the synthetic collagen-mimetic microfibers was approximately 18 nm, which, although considerably shorter than the 67 nm D-period observed for native collagen, is longer than one might have anticipated given the 10 nm length of the designed protomer. The banding distance observed for CPII assemblies suggests that the “functional” unit for fiber formation may involve oligomers consisting of multiple, laterally associated protomers, such that their staggered assembly defines an arrangement in which overlap zones create periodic regions of varying peptide packing and charge density. Notably, synthetic fibers were initially observed to exhibit pointed tips similar to the tectoidal ends of native collagen fibers from which continued fiber growth is thought to occur. Prokop et al. and others have suggested that this distinct morphological feature is consistent with a model for tip growth for native collagen based on the helical packing of aligned and parallel D-staggered molecules. Given this similarity, such an assembly process may be operative in our system.

In summary, our experimental studies show that D-periodic collagen-mimetic fibrils and microfibers, which are very similar to those formed in vivo, can be obtained through the linear assembly of a small collagen-mimetic peptide driven through electrostatic interactions. Mechanistically, molecular modeling suggests that staggered axial growth appears to be the preferred higher order structure for CPII peptides due to strong electrostatic effects and hydrogen bond formation between parallel oriented peptides assembled as triple-helical protomers. We believe that the synthesis of collagen-mimetic THPs that exhibit a propensity to form linear assemblies through a process of axially oriented alignment will prove to have a number of important practical applications in the design of novel biomaterials. Moreover, the capacity to mimic collagen’s ability to form linear assemblies with precisely defined periodic features represents an important milestone in the design of ordered materials for diverse applications in medicine and bionanotechnology.
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Supporting Information Available: All experimental procedures as well as HPLC purification and mass spectroscopic data for CPII and additional results for molecular dynamics simulations. This material is available free of charge via the Internet at http://pubs.acs.org.

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METHODS

Materials

Fmoc-protected amino acids, Fmoc-Gly, Fmoc-Arg(PMC)-OH (PMC: 2,2,5,7,8-pentamethylchroman-6-sulfonyl group), Fmoc-Glu(Ot-Bu)-OH (t-Bu: tert-butyl protecting group), Fmoc-Pro-OH and Fmoc-Hyp(t-Bu)-OH and Wang resin (substitution level 0.66 µmol/g) were purchased from Novabiochem. Activating agent DCC/HOBt, dichloromethane (anhydrous CH₂Cl₂), N-methylpyrrolidone (NMP), and dimethylformamide (DMF) were purchased from Applied Biosystems (Foster City, CA), while 20% (v/v) piperidine in DMF was obtained from Advanced Chemtech (Louisville, KY). N-ethyl-diisopropylamine (DIEA), 1,2-dithioethane, trifluoroacetic acid (TFA), and thioanisole were obtained from Fluka Chemicals (Seelze, Germany), acetic anhydride from Fisher Scientific (Waltham, MA), and diethyl ether and phenol purchased from Aldrich (St. Louis, MO). N-[tris(hydroxymethyl)methyl]-2-aminoethanesulfonic acid (TES) and all other reagents for buffer solutions were purchased from Aldrich.

Peptide Synthesis

The collagen-mimetic peptide, NH₂-(Pro-Arg-Gly)₄-(Pro-Hyp-Gly)₄-(Glu-Hyp-Gly)₄-COOH, CPII, was synthesized by a solid phase synthetic strategy using Applied Biosystems Model 433A automated peptide synthesizer using standard Fmoc chemistry. Stepwise chain elongation of the peptide was monitored throughout by high-pressure liquid chromatography (HPLC) and matrix-assisted laser desorption/ionization time-of-flight (MALDI-TOF) mass spectrometry. For Fmoc solid phase peptide synthesis, Wang resins were used and stepwise elongation of the amino acids was accomplished by activating Fmoc amino acids using DCC/HOBt coupling reagent. The couplings were performed in NMP as a solvent. The complete cleavage of the Fmoc group during individual coupling steps was carried out by base-induced deprotection using 20% (v/v) piperidine in DMF. The synthesis was run in 0.25 mmol scale with iterative coupling conditions involving 10-fold excess of Fmoc-protected reacting amino acids.
Conditional double coupling of unreacted amino groups with acetic anhydride ($\text{Ac}_2\text{O}$) was used in coupling steps involving Hyp, Pro and Arg, in order to obtain high yield of the desired product and minimize the formation of side products. On completion of the final amino acid coupling (proline) in the sequence, removal of side-chain protecting groups was achieved using acid-mediated cleavage with trifluoroacetic acid (TFA). The N-terminus of the peptide was not capped with acetic anhydride prior to its cleavage from the resin.

Peptides were cleaved off the resin at 25°C using 10 mL ‘cocktail’ of thioanisole (1.25 mL)/ethanedithiol (0.625 mL)/phenol (1.875 g)/water (1.25 mL) diluted in 25 mL of TFA (100%) over a period of 150 min. This procedure afforded a collagen-like peptide with a free COOH and amino group at the N and C terminus, respectively. The crude peptide was precipitated overnight with cold diethyl ether, filtered through a sintered funnel, and extracted over 20 min with 30% aqueous acetic acid and washed five times with distilled water (10 mL). Lyophilization yielded 570 mg of a white cotton-like material.

**HPLC Purification**

The crude peptide was dissolved in formic acid/TFA/trifluoroethanol/water (1:1:4, v/v) and purified by preparative reversed phase (RP) HPLC on a Zorbax SB C18 silica column (22 x 250 mm, 5 mm particle size, 300 Å pore size, Agilent) equilibrated at room temperature in 0.1% aqueous TFA. The peptide was eluted using a linear gradient of acetonitrile (AcN) in 0.08% aqueous TFA at 25 °C at a flow rate of 5 mL/min. Specifically, the gradient was created using two solvents. Solvent A consisting of 0.1% TFA in H$_2$O and solvent B, which was acetonitrile (AcN) in 0.08% aqueous TFA (v/v, 0.08% TFA). The gradient conditions were 3% solvent B for 10 min, 3-15% solvent B for 35 min, 15-65% solvent B for 110 min, 65-100% solvent B for 120 min, 100% solvent B for 130 min. The column effluent was monitored at 235 nm and the eluate was manually collected and stored at room temperature prior to further analysis.
The purity and the integrity of the HPLC fractions were screened by MALDI-TOF (Voyager-DE STR instrument (PerSeptive Biosystems, Framingham, MA) and those fractions containing the desired product were pooled. Rechromatography of some of the enriched fractions, which were lyophilized prior to their HPLC purification, was carried out using the same above mentioned conditions to obtain the final material in high purity. Subsequent lyophilization of the repurified fractions furnished CPII (320 mg) in 98% purity as demonstrated by MALDI-TOF MS and analytical HPLC.

Analytical HPLC of peptides was carried out using microbore RP-HPLC consisting of a Model 140A solvent delivery system and a Model 1000S diode array UV detector (Applied Biosystems). The separation was achieved using a Zorbax SB-C18 column (3m C-18 silica column 4.6 X 50 mm, 120A mBore) equilibrated at 25°C in 0.1% TFA and developed at a flow rate of 90 mL/min using a linear gradient of acetonitrile. The column eluate was monitored at 214 nm and eluted at 30 °C using linear gradient of acetonitrile in 0.1% TFA (Figure 1S).

**Figure 1S.** Reverse phase analytical HPLC profile of purified CPII.  
*Conditions*: Zorbax SB-C18 column (3 µ C-18 silica column 4.6 X 50 mm, 120A mBore), 30 °C, 90 ml/min, linear gradient of acetonitrile in 0.1% TFA. Rt = 10.05 min. Solvent A: 0.1% TFA in H₂O, Solvent B: 80% AcN (v/v, 0.08% TFA).
Mass Spectroscopy Analysis

MALDI-TOF MS analysis was performed using a Model Reflex III mass spectrometer (Bruker, Billarica, MA) equipped with a 337-nm nitrogen laser or MALDI-TOF Voyager program module and/or Voyager-DE STR BioSpectrometry Workstation MALDI-TOF mass spectrometer (Perseptive Biosystems, Inc., Framingham, MA, USA). α-Cyano-4-hydroxycinnamic acid (MALDI-Quality, Hewlett-Packard, Palo Alto, CA) was used as the matrix. The HPLC fractions (0.3 µL) were spotted on a target site of a sample grid, followed by addition of 0.3 µL of matrix (saturated α-cyano-4-hydroxycinnamic acid) dissolved in ethanol/water (1:1, v/v). The sample matrix was allowed to dry at room temperature for 5 min, and each sample was desorbed with 100 laser shots, each giving a spectrum. Measurements were performed at an acceleration voltage of 20 kV and spectra averaged to give the final spectrum. The instrument was operated in the reflector mode and was calibrated using the external standard peptides. MALDI-TOF mass spectrum (m/z): calcd, 3525.71; obsd, 3524.91 (monoisotopic mass) (Figure 2S).

Figure 2S. MALDI-TOF mass spectroscopy purified CPII. Matrix: α-cyano-4-hydroxycinnamic acid. calcd, 3525.71; obsd, 3524.91 (monoisotopic mass).
Circular Dichroism Spectroscopy

CD spectra were obtained on a JASCO J-810 CD spectropolarimeter (Jasco Inc., Easton, MD) equipped with a PFD-425S Peltier temperature control unit. Peptide solutions (0.5 mg/mL, 142 µM) were prepared in various buffer solutions and allowed to equilibrate at 4 °C for 24 h and spectra acquired from 5 °C to 25 °C, averaging 10 scans between 190–260 nm at 2 nm intervals and a 2 nm bandwidth. The scan rate was 20 nm min⁻¹ with a 4 second response time. Spectra were baseline corrected before converting to mean residue ellipticities [θ]. CD melting curves were generated by monitoring the absorbance at 222 nm, while the temperature was increased 2 °C/min from 20 to 100 °C with an equilibration time of 1 min at each temperature. The Tm values were determined from the first derivative d[θ]/dT of the CD melting curve (Table 1S).

Table 1S. Solvent dependant melting temperatures (Tm) for triple helical collagen-mimetic peptide CPII

<table>
<thead>
<tr>
<th>Buffer System</th>
<th>Tm (°C)</th>
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</thead>
<tbody>
<tr>
<td>Buffer 1 (H₂O)</td>
<td>44.4</td>
</tr>
<tr>
<td>Buffer 2 (10 mM HCl, pH 3.5)</td>
<td>38.9</td>
</tr>
<tr>
<td>Buffer 3 (10 mM MOPS, pH 6.85)</td>
<td>38.6</td>
</tr>
<tr>
<td>Buffer 4 (PBS, pH 7.25)</td>
<td>48.8</td>
</tr>
<tr>
<td>Buffer 7 (10 mM CHES, pH 9.0)</td>
<td>48.4</td>
</tr>
<tr>
<td>Buffer 8* (30 mM TES, pH 7.4)</td>
<td>37.4</td>
</tr>
</tbody>
</table>

*Buffer 8: 30 mM Na₂HPO₄, 30 mM TES, 135 mM NaCl (ionic strength 0.225), pH 7.4

Turbidity Measurements

The turbidity profiles for native collagen Type I and collagen-mimetic peptide CPII were acquired by monitoring the optical density at 313 nm as a function of time using a Cary 50 UV-vis spectrophotometer with attached Peltier temperature controller. All turbidity measurements were performed in a sealed cell to minimize evaporative losses and repeated a minimum of three times. For comparative analysis, turbidimetric analyses were determined for collagen
Type I at concentrations and temperatures between 0.2 - 1.0 mg/mL and 26 - 37 °C, respectively. The isolation and purification of monomeric collagen type I from rat tail tendon followed protocols detailed elsewhere. Fibril formation was initiated using a "cold-start procedure" by mixing 0.5 mL of collagen dissolved in 10 mM HCl with 0.5 mL of buffer solution to yield a final buffer composition of 30 mM TES, 30 mM Na$_2$HPO$_4$·7H$_2$O, and 135 mM NaCl (ionic strength 0.225), pH 7.4. Both collagen and buffer solutions were maintained at 4 °C prior to mixing and, thereafter, placed in the sample holder maintained at the experimental temperature (Figure 3S).

Collagen-mimetic peptide fibril forming solutions were prepared by adding 100 µL of buffer to 600 µL of peptide solution in 10 mM HCl. Both solutions were maintained at 4 °C prior to mixing. The final peptide concentration ranged between 0.5 – 1.4 mg/mL in a final buffer composition of 30 mM TES, 30 mM Na$_2$HPO$_4$·7H$_2$O, 135 mM NaCl, pH 7.4. The peptide solution was heated to 75 °C for 40 min and then cooled to room temperature, after which the cuvette was placed in the UV-vis sample holder maintained at the target temperature for turbidity studies.

The turbidity curves were analyzed as described by Silver and Birk. Specifically, turbidity ($A_{313} \times 2.303$) was plotted as a function of time in minutes to define the lag ($t_{lag}$) and plateau times ($t_p$) of the assembly process. Data obtained from turbidity-time assays at different peptide or collagen concentrations were used to calculate the apparent rate constants (k) for the lag and growth phases of self-assembly by regression of plots of $c/t_{lag}$ and $c/(t_p - t_{lag})$ versus concentration, respectively. The activation energy ($E_a$) for each phase was determined using the Arrhenius equation, $k = A e^{-E_a/RT}$, where $E_a$ can be from a plot of ln k versus 1/T.
Figure 3S. Turbidity profiles for collagen Type I. Turbidimetric analyses were determined at concentrations between 0.2 - 1.0 mg/mL and temperatures between 26 °C and 37 °C. Absorbance was acquired at 313 nm as a function of time and turbidity presented as $A_{313} \times 2.303$. Characteristic sigmoidal plots with a lag region, a growth region, and a plateau were observed for Type I collagen with an enhancement in the rate and magnitude of self-assembly noted with increasing collagen concentration. The calculated $E_a$ was 206 kJ.mol$^{-1}$ for the lag phase and 196 kJ.mol$^{-1}$ for the growth phase, which is similar to that reported by Silver et al. ³

Electron Microscopy

A fibril forming solution was prepared, as described above, with a final concentration of 0.71 mg/mL of CPII in 30 mM TES, 30 mM Na$_2$HPO$_4$·7H$_2$O, and 135 mM NaCl at pH 7.4. The peptide solution was heated to 75 °C for 40 min and then gradually cooled to room temperature, after which the sample was stored at 23 °C. After pre-defined incubation periods, 5 µL of
peptide solution was placed on 200 mesh Formvar-carbon film-coated copper grids, excess solution was removed with filter paper, and the sample stained with 0.5% uranyl acetate for 10 sec and then washed twice in water for 10 sec. The preparation was examined in a JEOL 1210 transmission electron microscope at 90 kV accelerating voltage.

**Molecular Dynamics Simulations**

A total of five molecular dynamics (MD) simulations were performed with GROMACS version 3.3.1 in order to examine potential structural features that contribute to the assembly of CPII THPs, as well as to the formation of axially oriented staggered protomer-protomer assemblies (Table 2S). In brief, simulations were performed on a local Infiniband 2 dual core Xeon 5140 cluster with 64 processors using (i) steepest descent minimization; (ii) 20 ps of constant number-pressure-temperature (NpT) simulations at -223 °C and 1 atm to equilibrate water with mobile water molecules, but with peptides restrained; (iii) 10 ps of constant number-volume-temperature (NVT) simulations at -223, -173, -123, -73, -23, and 23 °C with SHAKE constraints and 2 fs time steps; and (iv) 5 to 10 ns of NpT simulation at 23 °C and 1 atm to equilibrate the system. The Parrinello-Rahman pressure coupling algorithm was used to isotropically scale pressure with a time constant of 2 ps, while temperature was controlled using the Nosé-Hoover temperature coupling algorithm with a time constant of 0.5 ps. SHAKE constraints were used on all H-heavy atom bonds to permit a dynamic time step of 2 fs. Electrostatic interactions were calculated by the Particle Mesh Ewald (PME) method. In all cases, a periodic box was utilized with more than 1 nm of solvent between the peptide surface and box boundaries to reduce potential artifacts arising from periodicity. The box was filled with simple point charge (SPC) water molecules. For the data analysis, electrostatic potentials were calculated using APBS software. The extent of hydrogen bond formation was calculated with tools in GROMACS 3.3.1.
Table 2S: MD Simulations for CPII Peptide and Protomer Self-Assembly

<table>
<thead>
<tr>
<th>System</th>
<th>Components</th>
<th>Component orientation</th>
</tr>
</thead>
<tbody>
<tr>
<td>1: 3Peptide-Parallel</td>
<td>3 peptides</td>
<td>Three parallel oriented peptides</td>
</tr>
<tr>
<td>2: 3Peptide-Antiparallel</td>
<td>3 peptides</td>
<td>Three anti-parallel oriented peptides</td>
</tr>
<tr>
<td>3: 3Trihelix-Parallel</td>
<td>3 triple helical protomers</td>
<td>Assembly of three THPs, each formed from parallel oriented peptides</td>
</tr>
<tr>
<td>4: 3Trihelix-Antiparallel</td>
<td>3 triple helical protomers</td>
<td>Assembly of three THPs, each formed from anti-parallel oriented peptides</td>
</tr>
<tr>
<td>5: 3Complex-Parallel</td>
<td>3 complexes</td>
<td>Assembly of three complexes, each formed from four THPs produced from parallel oriented peptides</td>
</tr>
</tbody>
</table>

The initial structure for the CPII peptide was obtained by mutating the known X-ray crystal structure of [(Pro-Pro-Gly)₁₀]₃ (Protein ID: 1K6F)¹⁰ using the mutagenesis tools in Pymol¹¹ followed by energy minimization. Molecular dynamics simulations were initially performed to examine the formation of a triple helix from CPII peptides either in parallel or anti-parallel orientation at a separation distance of 2 nm (Figure 4S: Case 1, 2). Water between the three peptides was manually removed to reduce computational time for the peptides to repel water from the assembly.¹² Although the removal of water might influence the ability of the three CPII peptides to search for other conformations, the focus of this simulation was to investigate the structural features leading to THP assembly. CPII protomer-protomer interactions were subsequently investigated by simulating THPs produced from peptides assembled in either parallel or anti-parallel orientation, solvated in water, and separated by 2 nm (Figure 4S: Case 3, 4). A second form of protomer-protomer interaction was also examined by simulating three linear complexes, each produced from four THPs that were initially assembled from three CPII peptides in parallel orientation. The three linear complexes were then solvated in water, separated by 2 nm, and positioned in a parallel array (Figure 4S: Case 5).
**Figure 4S.** Case 1: Three CPII peptides oriented in parallel and solvated in water. Case 2: Three CPII peptides oriented in anti-parallel and solvated in water. Case 3: Three triple helical protomers from parallel oriented CPII peptides solvated in water. Case 4: Three triple helical protomers from anti-parallel oriented CPII peptides solvated in water. Case 5: Three complexes each formed from four triple helical protomers initially produced from parallel oriented CPII peptides solvated in the water. Peptides are represented as residue type in VDW style, for residue type, blue: Arg; red: Glu; green: Gly; cyan: Hyp; white: Pro; red dot: water.

**Figure 5S.** System total energy normalized by the number of water molecules. (A) Assembly of three peptides to form triple helical promoter (Cases 1 and 2). (B) Assembly of triple helical promoters (Cases 3 and 4). (C) Assembly of complexes formed from triple helical protomers (Case 5).
**Figure 6S.** Assembly results for different simulation cases. 

- **(A)** Three peptides oriented in parallel (Case 1). 
- **(B)** Three peptides oriented in anti-parallel (Case 2). 
- **(C)** Three triple helical protomers from parallel oriented CPII peptides (Case 3). 
- **(D)** Three triple helical protomers from anti-parallel oriented CPII peptides (Case 4). 
- **(E)** Three complexes each formed from four triple helical protomers produced from parallel oriented CPII peptides (Case 5).

The peptide was represented as residue type with VWD style separately, for residue type, blue: Arg; red: Glu; green: Gly; cyan: Hyp; white: Pro. The images were made with VMD software support.
REFERENCES