ABSTRACT: We describe the co-assembly of two different building units: collagen-mimetic peptides and DNA origami. Two peptides CP and sCP are designed with a sequence comprising a central block (Pro-Hyp-Gly) and two positively charged domains (Pro-Arg-Gly) at both N- and C-termini. Co-assembly of peptides and DNA origami two-layer (TL) nanosheets affords the formation of one-dimensional nanowires with repeating periodicity of ~10 nm. Structural analyses suggest a face-to-face stacking of DNA nanosheets with peptides aligned perpendicularly to the sheet surfaces. We demonstrate the potential of selective peptide-DNA association between face-to-face and edge-to-edge packing by tailoring the size of DNA nanostructures. This study presents an attractive strategy to create hybrid biomolecular assemblies from peptide- and DNA-based building blocks that takes advantage of the intrinsic chemical and physical properties of the respective components to encode structural and, potentially, functional complexity within readily accessible biomimetic materials.

Recent years have witnessed substantial advancement in molecular self-assembly with programmable biomolecules. One of the ultimate goals is to rationally design molecular structures that rival the complexity and functionality of naturally occurring biomolecular assemblies. Success in this endeavor critically relies on the development of new strategies for the self-assembly of hybrid structures that consist of multiple types of biomolecules. Biology has demonstrated that co-assembly of different structural units can lead to structurally complex supramolecular assemblies that display exquisitely tailored function. The assembly of infectious tobacco mosaic virus from self-association of coat proteins on an RNA template represents a notable example (Figure 1a). These native nucleoprotein assemblies have inspired efforts to create synthetic protein-DNA biomaterials that could hypothetically recapitulate the structural and functional complexity of viruses. Several approaches have been described that employ different mechanisms to control the protein-DNA interface, which include electrostatic charge complementation, covalent conjugation, and sequence-specific molecular recognition.

Herein, we demonstrate that DNA nanotechnology, which has exhibited great capacity for the rational design of complex structures across multiple length-scales and holds promise for engineering hybrid biomaterials, can be employed to enable co-assembly of larger ordered peptide-DNA nanostructures (Figure 1b).

In contrast to both naturally occurring nucleic-acid-protein co-assemblies and previous human-made nucleic-acid-peptide co-assembly, in which the nucleic acids generally serve as assembly templates without forming complex structures, our approach allows us to use pre-assembled, rationally designed DNA nanostructures to produce more structurally sophisticated peptide-DNA hybrid assemblies. We believe that our study will provide new information for the design and construction of peptide-DNA co-assemblies using tailored DNA nanostructures that will ultimately promote greater control over the structural and functional complexity of artificial hybrid structures.

Previously, we described a peptide CP that self-assembled into nanoscale sheets. It has a sequential triblock architecture.
comprising positively, neutral, and negatively charged (Xaa-Yaa- Gly) triads\(^{9}\) (Figure 2a) that self-associated into collagen-like triple helices. Electrostatic attractions between oppositely charged residues on adjacent peptides\(^8,10\) drove the assembly of the triple helices into peptide sheets. The greater length of the positively charged (Pro-Arg-Gly) domain, versus the negatively charged (Glu-Hyp-Gly) domain, ensured that the extra (Pro-Arg-Gly) triads protruded from sheet surfaces.\(^8\) We hypothesized that those arginine-rich overhangs might be recruited for interacting with negatively charged DNA-based materials. We observed that two-layer (TL) DNA origami\(^{6c}\) nanosheets (Figure S1; dimension \(\sim50\ nm \times \sim50\ nm \times 5\ nm,\) measured by TEM) could deposit onto the surface of pre-assembled CP\(^{+}\) sheets (Figure S2), although the arrangement of the TL nanosheets on the CP\(^{+}\) nanosheets was sporadic and not ordered.

We proposed that a new design with weakened attraction between triple helices might allow the TL nanosheets to assemble into more ordered structures in the presence of the peptide (Figure 1b). Two new collagen-mimetic peptides CP\(^{++}\) and sCP\(^{++}\) were designed and synthesized to include three positively charged (Pro-Arg-Gly) triads in both the N- and C-blocks (Figure 2b,d, top). Despite the shorter central core of the sCP\(^{++}\) (5 (Pro-Hyp-Gly) triad repeats, compared to the 7 repeats for CP\(^{++}\)), the circular dichroism (CD) spectropolarimetry of both CP\(^{++}\) and sCP\(^{++}\) in Tris buffer (5 mM, pH 8.0) displayed a characteristic signature of collagen-like triple helix, comprising a maximum peak at 224 nm and a minimum peak at 197 nm (Figure S3a). Thermal denaturation experiments indicated melting transition temperatures (\(T_m\)) of 63 and 52 °C for CP\(^{++}\) and sCP\(^{++}\) (Figure S3b). The decrease in \(T_m\) value observed for sCP\(^{++}\) versus CP\(^{++}\) can be attributed to the shorter central core having fewer canonical (Pro-Hyp-Gly) triads in peptide sCP\(^{++}\).\(^{9b}\) Compared to similar sequences of (Pro-Hyp-Gly), (\(T_m = 41 °C,\))\(^{11} \) CP\(^{++}\) afforded more stable triple helices. Indeed, the (Pro-Arg-Gly) triad sequence in host-guest systems of collagen-mimetic peptides showed a comparable thermostability to the canonical triad (Pro-Hyp-Gly).\(^{12}\) However, the electronic repulsion between arginines in adjacent helices prevented the triple helices of both peptides from self-assembling to higher-order structures, as has been observed previously for charge complementary collagen-mimetic peptide sequences.\(^{8,9}\)

TEM investigation demonstrated that co-assembly of CP\(^{++}\) and TL nanosheets (molar ratio of CP\(^{++}\):TL = 1600:1 and TL at 20 nM) led to the formation of one-dimensional (1D) banded nanowires (Figure 2b). The width of nanowires was measured to be \(47 \pm 3\ nm,\) which is close to the dimension of TL cross section (\(\sim50\ nm \times \sim50\ nm\)) (Figure S1). A high peptide/TL sheet molar ratio (at least 1600:1) was required to maintain stable co-assembly. With a peptide concentration at 32 \(\muM,\) and peptide-to-DNA sheet ratio at least 1600:1, the axial length of the nanowires depended on the amount of TL nanosheets. With lower concentrations of TL nanosheets (0.35 nM), shorter assemblies were observed (Figure S4). Within nanowires, thickness of light bands (4.8 ± 0.7 nm) matched the thickness of TL nanosheets. The spacing between adjacent light bands (dark region in the TEM images) was measured to be 5.9 ± 0.6 nm (Figure 2c), which is on the order of the theoretical length of the central block of CP\(^{++}\) triple helix (6.0 nm, 0.286 nm rise/residue\(^{12} \times \sim21\ residues). Similar 1D nanostructures were observed from sCP\(^{++}\)-TL co-assembly incubated at 4 °C.

Figure 2. Nanostructures from TL nanosheets and collagen-like peptides. (a) Sequence of CP\(^{+}\) peptide. (b) Sequence of CP\(^{++}\) peptide (top) and TEM images of nanowires (bottom) assembled with TL and CP\(^{+}\). (c) Histograms of the distances of TL-layer plus CP\(^{+}\) peptide-layer (red, \(x + y\) in (g)), and the distances of CP\(^{+}\) peptide-layer (green, \(y\) in (g)). (d) SAXS/WAXS profile for CP\(^{+}\)-TL nanowires. (e) Sequence of sCP\(^{++}\) peptide (top) and TEM images of nanowires (bottom) assembled with TL nanosheets and sCP\(^{++}\); (f) Histograms of the distances of TL-layer plus sCP\(^{++}\) peptide-layer (red, \(x + y\) in (g)), and the distances of sCP\(^{++}\) peptide-layer (green, \(y\) in (g)). (g) Proposed co-assembly structure. Scale bars, 50 nm.
(Figure 2e, bottom), but with shorter interband distance (3.8 ± 0.9 nm) (Figure 2f), which is on the order of the theoretical length of the central block of sCP++ triple helix (4.3 nm, 0.286 nm rise/residue × 15 residues). However, the sCP++-TL co-assembly generated nanowires of noticeably shorter length (46 ± 19 nm) compared to those observed for CP++-TL (133 ± 92 nm), presumably due to the reduced thermal stability of the sCP++ triple helix (Figure S5). Significant aggregation occurred between nanowires, given the presence of much more positively charged peptides and negatively charged TL sheets in nanowires (Figure S4). The aggregation may be mitigated by designing new TL nanosheets carrying neutral charge on the edges.

The CP++-TL nanowires proved more robust toward structural analysis and were further investigated using synchrotron SAXS/WAXS (Figures 2d and S6). A Bragg reflection was observed in the scattering profile, indicating a high order of internal structure. The peak at q = 0.061 ± 0.004 Å⁻¹ corresponds to a d-spacing of 10.3 ± 0.7 nm (Figures 2d and S6). We assigned it to the periodic center-to-center distance of adjacent light bands (equals to x + y in Figure 2g), which agrees with the axial spacing 10.6 ± 1.0 nm derived from TEM measurements (Figure 2c). Scherrer analysis of the diffraction peak width provided an average length of 109 nm for the nanowires if we assumed that the peak broadening was mainly caused by the finite nanowires length (Figure S6). A structural model was proposed that TL nanosheets (light band in TEM) are separated by peptides (dark band in TEM) and adopt a face-to-face stacking configuration (Figures 1b and 2g). Since the intersheet distances corresponded to the length of the peptides’ central blocks, rather than the total length of peptides, the peptide helices likely aligned perpendicularly to the surface of the TL nanosheets, with the positively charged, nonordered (Pro-Arg-Gly)₃ domains associating with the negatively charged surface of TL nanosheets (Figure 2g). Additionally, hydrogen bonds could also form between the guanidinium group of arginine and the phosphate backbone or nucleobases of the nucleic acids.¹⁵

Interestingly, the TL nanosheets did not stack edge-by-edge (5 nm in thickness), suggesting that TL’s edges could not provide a thermodynamically stable interaction with the CP++ or sCP++ (at extended state, the (Pro-Arg-Gly)₃ domain has a radius of 3.0 nm (0.33 nm contour length/residue × 9 residues)). We designed a DNA-brick⁶b 2H × 8H (cross section 5 nm × 20 nm) nanoribbon and a 4H × 12H (cross section 10 nm × 30 nm) nanoribbon to further verify this observation and demonstrate that the phenomenon can be used to regulate the peptide-DNA co-assembly. The 2H × 8H nanoribbon had the same thickness as TL nanosheet, while the 4H × 12H nanoribbon was twice as thick as TL nanosheet. Both nanoribbons assembled into two-dimensional (2D) arrays, in the presence of CP++ (Figure 3). Nonetheless, similar to the TL nanosheet, 2H × 8H ribbons assembled via CP++ interaction only with the large faces (8H) (Figure 3a), while 4H × 12H ribbons assembled via CP++ interaction with both the large face (12H) and thin edge (4H) (Figure 3b), in agreement with the co-assembly results of CP++ and TL nanosheets. Notably, extensive edge-to-edge stacking was not observed by using ribbons compared to the use of 2D nanosheet. It is presumably due to the self-twisting property of such long structures that hindered the perfect alignment between adjacent ribbons (Figure 3a left, b left). Moreover, most co-assemblies using 4Hx12H ribbons adopted both face-to-face and edge-to-edge stacking within same structures.

Significant research effort is being directed currently toward the creation of hybrid biomaterials containing a combination of peptide/protein and nucleic acid components.³⁻⁵,¹⁷ Here, we report a facile approach to co-assemble 2D DNA nanostructures and collagen-mimetic peptides into 1D hybrid materials. With DNA nanosheets, the peptides directed a face-to-face stacking of DNA nanosheets and formed nanowires. The separations between DNA nanosheets could be controlled through the length of the central block in peptide triple helices.
We attribute the periodic structure in co-assemblies to the alignment of peptide helices, which we propose is perpendicular to the faces of DNA nanosheets. Moreover, we showed that patterns of peptide-DNA interactions could be modulated by tailoring the design of DNA nanostructures. These types of hybrid biomaterials may have an advantage for the creation of functional devices in that the distinct structures and chemical properties of the two classes of biomolecules could be combined potentially in a synergistic manner.

## ASSOCIATED CONTENT

### Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/jacs.7b08087.

Experimental details, DNA sequences, and characterization of peptide-DNA nanostructures (PDF)

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