

Journal of Visualized Experiments

Self-assembly of gamma-modified peptide nucleic acids into complex nanostructures in organic solvent mixtures

--Manuscript Draft--

Article Type:	Invited Methods Collection - JoVE Produced Video
Manuscript Number:	JoVE61351R2
Full Title:	Self-assembly of gamma-modified peptide nucleic acids into complex nanostructures in organic solvent mixtures
Keywords:	peptide nucleic acids, structural nanotechnology, single-stranded tiles, xeno nucleic acids self-assembly, organic solvent mixtures, PNA nanotechnology, scaffold-free self-assembly
Corresponding Author:	Rebecca Elizabeth Taylor, Ph.D. Carnegie Mellon University Pittsburgh, PA UNITED STATES
Corresponding Author's Institution:	Carnegie Mellon University
Corresponding Author E-Mail:	bex@andrew.cmu.edu
Order of Authors:	Rebecca Elizabeth Taylor, Ph.D. Sriram Kumar Ying Liu
Additional Information:	
Question	Response
Please indicate whether this article will be Standard Access or Open Access.	Standard Access (US\$2,400)
Please indicate the city, state/province, and country where this article will be filmed . Please do not use abbreviations.	Pittsburgh, PA, USA

TITLE:

Self-Assembly of Gamma-Modified Peptide Nucleic Acids into Complex Nanostructures in Organic Solvent Mixtures

AUTHORS AND AFFILIATIONS:

Sriram Kumar¹, Ying Liu¹, Rebecca E. Taylor^{1,2,3}

¹Department of Mechanical Engineering, Carnegie Mellon University, Pittsburgh, Pennsylvania

²Department of Biomedical Engineering, Carnegie Mellon University, Pittsburgh, Pennsylvania

³Department of Electrical and Computer Engineering, Carnegie Mellon University, Pittsburgh, Pennsylvania

Email addresses of authors:

Sriram Kumar (srasipur@andrew.cmu.edu)

Ying Liu (yliu5@andrew.cmu.edu)

Corresponding author:

Rebecca E. Taylor (bex@andrew.cmu.edu)

KEYWORDS:

peptide nucleic acids, structural nanotechnology, single-stranded tiles, xeno nucleic acids self-assembly, organic solvent mixtures, PNA nanotechnology, scaffold-free self-assembly

SUMMARY:

This article provides protocols for the design and self-assembly of nanostructures from gamma-modified peptide nucleic acid oligomers in organic solvent mixtures.

ABSTRACT:

Current strategies in DNA and RNA nanotechnology enable the self-assembly of a variety of nucleic acid nanostructures in aqueous or substantially hydrated media. In this article, we describe detailed protocols that enable the construction of nanofiber architectures in organic solvent mixtures through the self-assembly of uniquely addressable, single-stranded, gamma-modified peptide nucleic acid (γPNA) tiles. Each single-stranded tile (SST) is a 12-base γPNA oligomer composed of two concatenated modular domains of 6 bases each. Each domain can bind to a mutually complimentary domain present on neighboring strands using programmed complementarity to form nanofibers that can grow to microns in length. The SST motif is made of 9 total oligomers to enable the formation of 3-helix nanofibers. In contrast with analogous DNA nanostructures, which form diameter-monodisperse structures, these γPNA systems form nanofibers that bundle along their widths during self-assembly in organic solvent mixtures. Self-assembly protocols described here therefore also include a conventional surfactant, Sodium Dodecyl Sulfate (SDS), to reduce bundling effects.

INTRODUCTION:

Successful construction of numerous complex nanostructures¹⁻¹² in aqueous or substantially hydrated media made using naturally occurring nucleic acids such as DNA¹⁻¹⁰ and RNA^{11,12} has been shown in previous works. However, naturally occurring nucleic acids undergo duplex helical conformational changes or have reduced thermal stabilities in organic solvent mixtures^{13,14}.

Previously, our lab has reported a method towards the construction of 3-helix nanofibers using gamma-position modified synthetic nucleic acid mimics called gamma-peptide nucleic acids (γ PNA)¹⁵ (**Figure 1A**). The need for such a development and potential applications of the synthetic nucleic acid mimic PNA has been discussed within the field^{16,17}. We have shown, through an adaptation of the single-stranded tile (SST) strategy presented for DNA nanostructures¹⁸⁻²⁰, that 9 sequentially distinct γ PNA oligomers can be designed to form 3-helix nanofibers in select polar aprotic organic solvent mixtures such as DMSO and DMF. The γ PNA oligomers were commercially ordered with modifications of (R)-diethylene glycol (mini-PEG) at three γ -positions (1, 4 and 8 base-positions) along each 12-base oligomer based on methods published by Sahu *et al.*²¹ These gamma-modifications cause the helical pre-organization that is associated with the higher binding affinity and thermal stability of γ PNA relative to unmodified PNA.

This article is an adaptation of our reported work in which we investigate the effects of solvent solution and substitution with DNA on the formation of γ PNA-based nanostructures¹⁵. The aim of this article is to provide detailed descriptions of the design as well as detailed protocols for solvent-adapted methods that were developed for the self-assembly and characterization of γ PNA nanofiber. Thus, we first introduce the modular SST strategy, a general platform for nanostructure design using the synthetic nucleic acid mimic PNA.

The helical pitch for PNA duplexes has been reported to be 18 bases per turn in comparison to DNA duplexes, which undergo one turn per 10.5 bases (**Figure 1B**). Therefore, the domain-length of the demonstrated γ PNA SSTs was set at 6 bases to accommodate one third of a full turn or 120° of rotation to enable interaction between three triangularly arrayed helices. Also, unlike previous SST motifs, each SST contains just 2 domains, effectively creating a 1-dimensional ribbon-like structure that wraps to form a three-helix bundle (**Figure 1C**). Each 12-base γ PNA oligomer is gamma modified at the 1, 4 and 8 positions to ensure uniformly spaced distribution of mini-PEG groups across the overall SST motif. Additionally, within the motif, there are two types of oligomers: “contiguous” strands that exist on a single helix and helix-spanning “crossover” strands (**Figure 1D**). In addition, oligomers P8 and P6 are labelled with fluorescent Cy3 (green star) and biotin (yellow oval), respectively (**Figure 1D**), to enable detection of structure formation using fluorescence microscopy. Altogether, the SST motif is made of 9 total oligomers to enable the formation of 3-helix nanofibers through programmed complementarity of each individual domain to the corresponding domain on a neighboring oligomer (**Figure 1E**).

PROTOCOL:

1. γ PNA sequence design

1.1. Download the DNA Design Toolbox²² developed by the Winfree Lab at Caltech²³ into the folder containing programming scripts for designing sequences.

1.2 Within that sequence design folder, open a fourth-generation programming language compatible with the file extension “.m”, and then add the previously downloaded “DNAdesign” folder to the path using the following command:

```
>> addpath DNAdesign
```

1.3 Subsequently run the following script named “PNA3nanofiber.m” (see **Supplementary Figure 1**) using the following command:

```
>> PNA3nanofiber
```

1.4 Run this script to create variables called “thisSeqs” and “thisScore.” The variable “thisSeqs” contains the sequences of the designed oligomers, and “thisScore” is the penalty score.

1.5 Run the script multiple times to obtain the most minimal score. A sample of 20 such runs are shown in **Table 1**.

1.6 Manually confirm the desired Watson-Crick complementarity of each domain of the generated sequences for the specified SST structural motif. Sequence specifications for the 3-helix nanofiber structure are shown in **Table 2**.

1.7 Verify the following for generated sequences.

1.7.1. Avoid using four consecutive C and G bases.

1.7.2. Manually select specific sequences for N-terminal functionalization with fluorescent dye and biotin molecules to enable fluorescent microscopy studies.

1.7.3. Include a minimum of 3 mini-PEG gamma-modifications on the backbone to enable pre-organized helical conformation in the single-stranded oligomers as described by Sahu *et al.*²¹

2. Preparation of γPNA stock strands

2.1. Obtain γPNA component strands of specified sequences at 50 nmol scale synthesis with high performance liquid chromatography (HPLC) purification from a commercial manufacturer.

2.2 Resuspend each strand in deionized water to 300 μM concentrations. Store the resuspended strands in -20 °C freezer up to several months until needed for experimentation.

3. Melting curve studies of γPNA oligomer subsets

3.1. Obtain melting temperature ranges for different combinations of complementary 2-oligomer and 3-oligomer subsets by running a melt curve study in aqueous buffers such as 1x phosphate

buffered saline (PBS) or preferred polar organic solvent such as Dimethylformamide (DMF) or Dimethyl sulfoxide (DMSO) (see **Figure 2**).

NOTE: Thermal melting curves at >50% of DMF start to lose the upper baseline and show severe disturbances partly because of high absorbance of DMF at the wavelength required for the experiments. This is a noted phenomenon in the literature²⁴. It is however possible to obtain the melting curves at 5 μ M per strand concentrations in these solvent conditions.

3.1.1. Aliquot 16.7 μ L from the 300 μ M stock of each oligomer and make the final volume to a 1000 μ L either in 1x PBS or 100% (v/v) DMSO or 100% (v/v) DMF effectively obtaining 5 μ M final concentration per oligomer. Transfer this oligomer subset mixture to a 1 cm optical path, quartz cuvette.

3.1.2. Perform variable temperature UV-Vis experiments in a spectrophotometer equipped with a programmable temperature block.

3.1.3. Collect data points for the melting curves over a temperature range of 15–90 $^{\circ}$ C for both cooling (annealing) and heating (melting) cycles at a rate of 0.5–1 $^{\circ}$ C/min. Keep the samples for 10 min at 90 $^{\circ}$ C before cooling and at 15 $^{\circ}$ C before heating. Determine the melting temperature (T_m) from the peak of the first derivative of the heating curve.

3.1.4. Verify that melting temperature ranges for 2-oligomer subsets are above 35 $^{\circ}$ C in all solvent cases. Additionally, verify that the corresponding 3-oligomer subsets that contain an additional strand to their equivalent 2-oligomer subset shows a considerable increase in T_m due to increased co-operativity.

NOTE: This would verify that a single pot self-assembly of multiple oligomers can co-operatively fold into the desired nanostructure with reasonable thermal stability.

4. Self-assembly protocol for multiple distinct γ PNA oligomers

NOTE: To devise a self-assembly thermal ramp protocol for γ PNA nanostructures, slow-ramp annealing is desirable.

4.1. In the case of oligomer sequences generated, anneal the samples for 22.5 h in a thermal cycler cooling from 90 to 20 $^{\circ}$ C. Typically, melting temperature obtained for 2-oligomer and 3-oligomer γ PNA subsets lie in ranges of 40–70 $^{\circ}$ C for different solvent conditions.

4.2. Program the thermal cycler as follows: hold at 90 $^{\circ}$ C for 5 min, ramp down from 90 to 70 $^{\circ}$ C at a constant rate of 0.1 $^{\circ}$ C/min, ramp down from 70 to 40 $^{\circ}$ C at a rate of 0.1 $^{\circ}$ C/3 min, ramp down from 40 to 20 $^{\circ}$ C at a rate of 0.1 $^{\circ}$ C/min and hold at 4 $^{\circ}$ C (see **Table 3**). Samples can be stored in 4 $^{\circ}$ C for 12–24 h before characterization.

NOTE: While 2- and 3- oligomer subsets of our nanofiber system can form in 1x PBS, the full micron-scale nanofibers aggregate in 1x PBS. Therefore, solvent conditions should be optimized based on the scale and size of structure being formed as well as the type and density of gamma modifications.

4.3. For micron scale long 3-helix nanofibers, prepare anneal batch samples in 75% DMSO: H₂O (v/v), 75% DMF: H₂O (v/v), 40% 1,4-Dioxane: H₂O (v/v) based on solvent optimization studies¹⁵.

4.4. Prepare anneal batches as follows (see **Table 4**). First, prepare 10 μ L sub-stocks at 20 μ M concentrations from the 300 μ M main stocks for each oligomer by aliquoting 0.67 μ L from the main stock and making the volume to 10 μ L using deionized water.

4.5 Aliquot 1 μ L from the 20 μ M sub-stocks for each oligomer and add it to a 200 μ L PCR tube. This accounts for a total volume of 9 μ L for 9 oligomers.

4.6 Add either 30 μ L of anhydrous DMSO/DMF for the 75% DMSO and 75% DMF cases with an additional 1 μ L of deionized water to make a final volume of 40 μ L with each oligomer at 500 nM final concentrations. Add 16 μ L of 1,4-Dioxane and make the volume with deionized water to 40 μ L for the 40% Dioxane solvent condition.

4.7 Load the anneal batches on to the thermal cycler and anneal using the protocol mentioned in step 4.2.

5. Total internal reflection fluorescence (TIRF) microscopy imaging

5.1. Prepare a humidity chamber from an empty pipette tips box. Fill the box with approximately 5 mL of water to prevent drying of the sample flow channels described as follows (see **Figure 3**).

5.2. Prepare flow chamber with a microscope slide, 2 double-sided tape strips, and nitrocellulose-coated coverslip. To coat the coverslip with nitrocellulose, dip the coverslip in a beaker containing 0.1% collodion in amyl acetate and air-dry.

5.2.1. Prepare the nitrocellulose solution by making a 20-fold dilution from commercially available 2% collodion in amyl acetate with isoamyl acetate solvent.

5.3. Prepare biotinylated bovine serum albumin (Biotin-BSA) solution by weighing 1 mg of Biotin-BSA and dissolving it in 1 mL of 1x PBS. Flow 15 μ L of 1 mg/mL Biotin-BSA in 1x PBS buffer by placing the flow channel at an angle. Incubate the flow channel for 2–4 min at room temperature in a humidified chamber.

5.4. Prepare the wash buffer by dissolving 1 mg of BSA in 1 mL of 1x PBS. Wash the excess Biotin-BSA by flowing 15 μ L of the wash buffer. To passivate the surface, incubate the flow channel for 2–4 mins at room temperature in a humidified chamber.

218 5.5. Prepare the streptavidin solution by measuring 0.5 mg of streptavidin and 1 mg of BSA and
219 then dissolving using 1 mL of 1x PBS. Flow 15 μ L of 0.5 mg/mL streptavidin in 1x PBS containing
220 1 mg/mL BSA. Incubate the flow channel for 2–4 min at room temperature in a humidified
221 chamber. Flow 15 μ L of the wash buffer to wash away unbound Streptavidin.

222
223 5.6. Flow 15 μ L of previously annealed batch of γ PNA oligomers at 500 nM concentration per
224 strand in either 75% DMSO, 75% DMF or 40% 1,4-Dioxane condition. Incubate the flow channel
225 for 2–4 mins at room temperature in a humidified chamber.

226
227 5.7. Prepare 1 mM Trolox in preferred solvent conditions by diluting 10-fold from a 10 mM stock
228 of Trolox in DMSO (measure 2.5 mg of Trolox and dissolve in 1 mL of DMSO). Wash the unbound
229 nanostructures in the flow channel with 15 μ L of 1 mM Trolox having the same solvent
230 composition as the nanostructures.

231
232 NOTE: The >50% DMSO content in the flow channel would produce minor signal disturbances
233 due to the different index of refraction of the solvent condition during TIRF which is typically
234 calibrated for coverslip-water TIRF angles. This can be rectified by washing with 1 mM Trolox
235 made by diluting the 10 mM Trolox 10-fold using deionized water. This technique provides clearer
236 images but is only useful for assessing whether formation occurred, however, because it
237 produces two-phase micro-bubbles in the flow channel. As a result, nanostructures might be
238 visible at the interface of the two-phases as shown in **Figure 4D**.

239
240 5.8. Transfer the flow channel on to the slide holder and image using a fluorescence microscope
241 equipped with TIRF imaging using a 60x oil-immersion objective and a 1.5x magnifier. Scan the
242 flow channel at either 60x or 90x magnification by monitoring the Cy3 channel (see **Figure 4**).

243 244 6. Transmission electron microscopy (TEM) imaging

245
246 6.1. Weigh 0.5 g of uranyl acetate in 50 mL of distilled water to prepare a 1% aqueous uranyl
247 acetate stain solution. Filter the 1% aqueous uranyl acetate stain solution using a 0.2 μ m filter
248 attached to a syringe.

249
250 NOTE: Alternatively, 2% aqueous uranyl acetate could be purchased from a commercial
251 manufacturer.

252
253 6.2. Purchase commercially available formvar support layer coated Copper grids with 300-mesh
254 size.

255
256 NOTE: It is important to note that the formvar support layer could be dissolved away by solvents
257 like DMSO beyond 2 min. For longer sample incubation, commercially available formvar stabilized
258 with silicon monoxide on copper grids are available which allow the grid to be more hydrophilic
259 than carbon-coated grids and can withstand vigorous sample conditions and electron beam as
260 shown in **Figure 5C**.

262 6.3. Pipette 4 μ L of sample onto the grid for 15 s. Use a piece of filter paper to wick off the sample
263 by bringing the filter paper in contact with the grid from the side.

264
265 6.4. Immediately add 4 μ L of the stain solution onto the grid for 5 s.

266
267 6.5. Wick off the stain as before and hold the filter paper against the grid for 1–2 min to make
268 sure the grid is dry. Samples should be typically imaged within 1–2 h after staining. Alternatively,
269 if the grid is ensured to be completely dry, grids can be stored in a TEM grid storage box for up
270 to 3 days before imaging.

271
272 6.6. Transfer the grid to a TEM specimen holder and image using a transmission electron
273 microscope operated at 80 kV with magnification ranging from 10 K to 150K (see **Figure 5**).

274 275 **7. Different morphologies for γ PNA-DNA hybrids based on selective replacement with DNA**

276
277 7.1. Obtain DNA oligomers of specified sequences (see **Table 5**) from commercial oligonucleotide
278 manufacturers synthesized at 25 nmol scale using standard desalting. Resuspend these DNA
279 sequences using RNase-free deionized water at 20 μ M stock concentrations.

280
281 7.2. For contiguous γ PNA strand replacements with DNA, sequences D3, D5 and D9 can replace
282 strands p3, p5 and p9. Similarly, for crossover γ PNA strand replacements with DNA, sequences
283 D1, D4 and D7 can replace strands p1, p4 and p7.

284
285 7.3. Aliquot 1 μ L from the 20 μ M sub-stocks for each γ PNA or DNA oligomer and add it to a 200
286 μ L PCR tube as in step 2.7. Add 30 μ L of anhydrous DMSO and 1 μ L of RNase-free deionized water
287 to make the final volume to 40 μ L (see **Table 6**).

288
289 7.4. Load the anneal batches on to the thermal cycler and anneal using the protocol mentioned
290 in step 4.2.

291
292 7.5. Characterize γ PNA-DNA hybrid nanostructures using the TIRF protocol following steps in
293 section 5 or using TEM imaging protocol mentioned in section 6 (see **Figure 6**).

294 295 **8. Different morphologies for γ PNA nanofibers in varying concentrations of SDS**

296
297 8.1. Prepare a 20% (wt/v) SDS main stock by measuring 20 mg of SDS and dissolving it in 100 μ L
298 of deionized water.

299
300 8.2. Prepare a 6% (wt/v) SDS sub-stock by aliquoting 3 μ L from the 20% SDS stock and making the
301 volume to 10 μ L using deionized water.

302
303 8.3. Anneal the γ PNA oligomers in final concentrations of 5.25 mM and 17.5 mM SDS as follows.
304 Aliquot 1 μ L from the 20 μ M sub-stocks for each γ PNA oligomer and add it to a 200 μ L PCR tube
305 as in step 2.7. Add 30 μ L of anhydrous DMSO and 1 μ L of 6% and 20% SDS to make the final

volume to 40 μ L to achieve SDS final concentrations of 5.25 mM and 17.5 mM, respectively (see **Table 7**).

8.4. Load the anneal batches of varying SDS concentrations on to the thermal cycler and anneal using the protocol mentioned in step 4.2.

8.5. Characterize γ PNA nanostructures in the presence of SDS using the TIRF protocol following steps in section 5 or using TEM imaging protocol mentioned in section 6 (see **Figure 7**).

REPRESENTATIVE RESULTS:

The protocols discussed in the sections above describe the design of an adapted SST motif from DNA nanofibers for the robust generation of self-assembled nanofibers structures using multiple, distinct γ PNA oligomers. This section describes the interpretation of data obtained from the successful recreation of the protocols described.

Following the protocol described in section 5 for TIRF imaging of samples of γ PNA oligomers annealed in 75% DMSO: H₂O (v/v) most readily provides evidence of well-organized architectures under microscopic observations as shown in **Figure 4A**. 75% DMF: H₂O (v/v) solvent condition results in spicule-shaped or needle-like nanostructures (**Figure 4B**) and, in our experience, the 40% 1,4 dioxane: H₂O (v/v) condition shows sparse decoration of filamentous nanostructures when viewed using TIRF microscopy (**Figure 4C**). Furthermore, the samples of γ PNA nanotubes formed in 75% DMSO: H₂O (v/v) demonstrate bundling of nanofibers at high magnification or nanoscopic resolutions during TEM imaging following steps mentioned in section 6 (**Figure 5**). Quantitative analyses of the width of the nanostructures showed a median width of 16.3 nm with maximum values beyond 80 nm¹⁵.

For devising a scheme towards generating γ PNA-DNA hybrid nanostructures in organic solvent mixtures to enable further functionalization using DNA, it is important to consider the oligomeric position in the SST motif being replaced with isosequential DNA oligomers. Adapting steps mentioned in section 7 for the case of the nanotube construct described here, isosequential DNA oligomers that replace the contiguous γ PNA oligomers forms straight filamentous structures whereas replacement of the crossover γ PNA oligomers forms stellate structures when viewed using TIRF and TEM imaging (**Figure 6**). Quantitative analyses of the width of the nanostructures that were replaced with contiguous DNA oligomers showed median widths around 19 nm¹⁵.

Finally, upon adapting steps from section 8, γ PNA nanofibers adopt thinner morphologies consistent with reduced bundling in the presence of SDS concentrations below its critical micelle concentrations (CMC, 8.2 mM). γ PNA nanofibers also adopt highly networked morphologies at high SDS concentrations in comparison to its CMC when viewed using TIRF imaging (**Figure 7**). TEM imaging of γ PNA oligomers self-assembled in the presence of SDS indicate that the 5.25 mM SDS condition indicate the most substantial reductions in structural bundling with widths ranging 8–12 nm as shown in (**Figure 7C**).

FIGURE AND TABLE LEGENDS:

Figure 1: Peptide nucleic acid oligomers as building blocks for complex nanostructures. (A) Chemical structures of DNA (PNA), PNA and MP-containing γ PNA units. (Figure has been reprinted with permission from Ref¹⁹.) (B) PNA-PNA helices are less twisted than DNA-DNA helices, having 18 bases per turn instead of 10.5. (C) This three-helix structural single stranded tile (SST) motif consists of 9 distinct 12-base-long γ PNA oligomers, each having two six-base domains. (D) Within the structure there are two types of oligomers: “contiguous” strands that exist on a single helix and helix-spanning “crossover” strands. (E) This 18-base-long structural motif can polymerize to form micron-scale filaments. Panels B, C and E have been modified with permission from Ref¹⁵.

Figure 2: Representative melt curves of select γ PNA oligomers in different solvent conditions. 2-oligomer (p4, p5) subset showing 6-base domains can bind with reasonable thermal stability (black curve) in 1x PBS. 3- oligomer (p4, p5, p6) subset shows increased thermal stability due to cooperativity in 1x PBS (blue curve) and shows little to no instability with respect to T_m when solvent is changed to organic solvents like DMF (red dotted curve). Figure has been modified with permission from Ref¹⁵.

Figure 3: Flow chart for fluidic flow channel for sample TIRF imaging. A step-by-step workflow indicating steps involved in sample preparation for TIRF imaging of γ PNA nanofibers after flow channel assembly.

Figure 4: TIRF microscopy imaging of γ PNA nanofiber self-assembled in different solvent conditions. γ PNA nanofibers visualized using TIRF microscopy (5 μ m scale bar) while monitoring the Cy3 channel when γ PNA oligomers are self-assembled in (A) 75% DMSO: water (v/v), (B) 75% DMF: water (v/v) and (C) 40% Dioxane: water (v/v) solvent conditions. (D) When γ PNA nanofibers self-assembled in 75% DMSO: water (v/v) bound to the flow channel is washed with 1mM Trolox in water, two-phase microbubbles of DMSO-water form with nanostructures aligning along the interface of the microbubble. Figure has been modified with permission from Ref¹⁵.

Figure 5: TEM imaging of γ PNA nanofibers self-assembled in 75% DMSO: water (v/v) using formvar support layer copper 300 mesh grids. (A) TEM images of γ PNA nanofibers visualized under low magnification (15x). (B) TEM images of γ PNA nanofibers visualized under high magnification (150x) shows bundling of nanotubes along the width at nanoscopic resolutions. (C) TEM images of γ PNA nanofibers visualized under low magnification (10x) using a Formvar-Silicon Monoxide support layer on a Copper grid allows imaging under vigorous specimen conditions. Figure has been modified with permission from Ref¹⁵.

Figure 6: Different morphologies for γ PNA-DNA hybrids based on selective replacement with DNA. (A) TIRF (5 μ m scale bar) and (C) TEM images (60x magnification) of self-assembly of γ PNA-DNA hybrids upon replacing contiguous γ PNA oligomers (p3, p5, p9) with isosequential DNA oligomers (D3, D5 D9) show nanotubes adopting a straight filament morphology. (B) TIRF (5 μ m scale bar) and (D) TEM images (25x magnification) of self-assembly of γ PNA-DNA hybrids upon replacing crossover γ PNA oligomers (p1, p4, p7) with isosequential DNA oligomers (D1, D4 D7)

show nanofibers adopting a stellate morphology. Figure has been modified with permission from Ref¹⁵.

Figure 7: Different morphologies for γ PNA nanofibers in varying concentrations of SDS. TIRF images (5 μ m scale bar) of self-assembly of γ PNA nanofibers in the presence of SDS at concentrations of (A) 5.25 mM and (B) 17.5 mM. Self-assembly in the presence of SDS concentrations less than the CMC (8.2 mM) shows thinner morphologies from TIRF images based on fluorescence intensity. (C) This is verified by TEM images (100x magnification) of the system where median width for nanofibers lies in the range of 8–12 nm. At concentrations significantly above the CMC, γ PNA nanotubes appear to form higher order assemblies through highly networked nanotube structures. Figure has been modified with permission from Ref¹⁵.

Table 1: 20 sample algorithmic results for potential optimization of γ PNA sequence design. 20 sample **algorithmic** results with sequence outputs in column 1 and their corresponding score in column 2. Repeated iterations of the script are performed to obtain the most minimal score.

Table 2: γ PNA sequence design results for nanofibers. Individual oligomer sequences were generated as indicated in this table. Underlined bases indicate the gamma-position modifications with mini-PEG. Table has been modified with permission from Ref¹⁵.

Table 3: Anneal ramp protocol for thermal cycler. Table has been modified with permission from Ref¹⁵.

Table 4: Protocol for preparing anneal batches of γ PNA oligomer in different solvent conditions.

Table 5: Isosequential DNA sequences as replacement oligomers to γ PNA. Table has been modified with permission from Ref¹⁵.

Table 6: Protocol for preparing anneal batches of γ PNA-DNA hybrid nanostructures in 75% DMSO: H₂O (v/v) through replacement of contiguous or crossover γ PNA oligomers with isosequential DNA oligomers.

Table 7: Protocol for preparing anneal batches of γ PNA nanostructures in 75% DMSO: H₂O (v/v) in the presence of SDS concentrations below and above CMC.

Supplementary Figure 1: Programming script PNA3nanofiber.m for designing oligomer sequences.

DISCUSSION:

This article focuses on adapting and improving existing nucleic acid nanotechnology protocols towards organic solvent mixtures. The methods described here focus on modifications and troubleshooting within a defined experimental space of select polar aprotic organic solvents. There is yet unexplored potential for other established nucleic acid nanotechnology protocols to be adapted within this space. This could improve potential applications through integration in

other fields such as polymer and peptide synthesis which typically are performed in similar organic solvents^{25,26}. Additionally, we focus here on critical steps to be observed while practicing the above-mentioned protocols.

During the preparation of sample for self-assembly, it is important to keep the volume percentages of water at 25% (v/v) for DMSO and DMF conditions. Accordingly, it is important to recognize that the organic solvent used for self-assembly should be from anhydrous stocks as well. The nanofiber structures are hydrophobic and aggregate in increased water content.

Unlike scaffolded DNA origami approaches that can create structures of discrete lengths, the current SST motif design for γ PNA nanofibers and other previously established DNA SST nanotubes does not yield structures of discrete lengths. Nanotubes polymerize achieving a range of multi-micron lengths (up to 11 μ m). For the same reason, yield associated with structure formation cannot be quantified.

However, due to the uncharged peptide backbone of γ PNA, there are no dependencies on counter ion balance in relation to structure formation. Imaging buffers, therefore, do not need to include cations like Mg^{2+} typically required for the stabilization of nanostructures made from naturally occurring nucleic acids.

Lastly, γ PNA nanostructure bundling and aggregation in different solvent conditions are also dependent on the individual oligomer concentrations introduced during self-assembly. Concentrations ranging from 1 μ M and higher of individual oligomers increase the propensity for bundling and aggregation and affect clear imaging of nanostructures either during TIRF or TEM imaging.

ACKNOWLEDGMENTS:

This work was supported in part by National Science Foundation grant 1739308 and by the Air Force Office of Science Research grant number FA9550-18-1-0199. γ PNA sequences were a generous gift from Dr. Tumul Srivastava of TruCode Gene Repair, Inc. We would like to thank Dr. Erik Winfree and Dr. Rizal Hariadi for their helpful conversations on DNA Design Toolbox MATLAB code. We would also like to thank Joseph Suhan, Mara Sullivan and the Center for Biological Imaging for their assistance in the collection of TEM data.

DISCLOSURES:

The authors declare no competing financial interests.

REFERENCES:

1. Fu, T. J., Seeman, N. C. DNA double-crossover molecules. *Biochemistry*. **32**, (13), 3211-3220 (1993).
2. Winfree, E., Liu, F., Wenzler, L. A., Seeman, N. C. Design and self-assembly of two-dimensional DNA crystals. *Nature*. **394**, (6693), 539-544 (1998).
3. Shih, W., Quispe, J., Joyce, G. A 1.7-kilobase single-stranded DNA that folds into a nanoscale octahedron. *Nature*. **427**, (6975), 618-621 (2004).

482 4. Rothemund, P. W. K. Folding DNA to create nanoscale shapes and patterns. *Nature*. **440**,
483 (7082), 297-302 (2006).

484 5. He, Y. et al. Hierarchical self-assembly of DNA into symmetric supramolecular polyhedra.
485 *Nature*. **452**, (7184), 198-201 (2008).

486 6. Douglas, S. M., Dietz, H., Liedl, T., Högberg, B., Graf, F., Shih, W. M. Self-assembly of DNA into
487 nanoscale three-dimensional shapes. *Nature*. **459**, (7245), 414-418 (2009).

488 7. Andersen, E. S. et al. Self-assembly of a nanoscale DNA box with a controllable lid. *Nature*. **459**,
489 (7243), 73-76 (2009).

490 8. Dietz, H., Douglas, S. M., Shih, W. M. Folding DNA into twisted and curved nanoscale shapes.
491 *Science*. **325**, (5941), 725-730 (2009).

492 9. Han, D. et al. DNA origami with complex curvatures in three-dimensional space. *Science*. **332**,
493 (6024), 342-346 (2011).

494 10. Liu, Y., Kumar, S., Taylor, R. E. Mix-and-match nanobiosensor design: Logical and spa421
495 tial programming of biosensors using self-assembled DNA nanostructures. *WIREs Nanomedicine*
496 *Nanobiotechnology*. **10**, e1518 (2018).

497 11. Chworos, A. et al. Building programmable jigsaw puzzles with RNA. *Science*. **306**, (5704), 2068-
498 2072 (2004).

499 12. Delebecque, C. J., Lindner, A. B., Silver, P. A., Aldaye, F. A. Organization of intracellular
500 reactions with rationally designed RNA assemblies. *Science*. **333**, (6041), 470-474 (2011).

501 13. Wang, X., Lim, H. J., Son, A. Characterization of denaturation and renaturation of DNA for
502 DNA hybridization. *Environmental health and toxicology*. **29**, e2014007 (2014).

503 14. Bonner, G., Klibanov, A.M. Structural stability of DNA in nonaqueous solvents. *Biotechnology*
504 *and Bioengineering*. **68**, 339 (2000).

505 15. Kumar, S., Pearse, A., Liu, Y., Taylor, R.E. Modular self-assembly of gamma-modified peptide
506 nucleic acids in organic solvent mixtures. *Nature communications*. (accepted, 2020)

507 16. Barluenga, S., Winssinger, N. PNA as a biosupramolecular tag for programmable assemblies
508 and reactions. *Accounts of chemical research*. **48**, 1319-31 (2015).

509 17. Berger, O., Gazit, E. Molecular self-assembly using peptide nucleic acids. *Peptide Science*. **108**,
510 e22930 (2017).

511 18. Rothemund, P.W. et al. Design and characterization of programmable DNA nanotubes.
512 *Journal of American Chemical Society*. **126**, 16344–16352 (2004).

513 19. Yin, P. et al. Programming DNA tube circumferences. *Science*. **321**, 824–826 (2008).

514 20. Yang, Y. et al. Self-assembly of DNA rings from scaffold-free DNA tiles. *Nano Letters*. **13**,
515 1862–1866 (2013).

516 21. Sahu, B. et al. Synthesis and characterization of conformationally preorganized, (R)-
517 diethylene glycol-containing -peptide nucleic acids with superior hybridization properties and
518 water solubility. *Journal of Organic Chemistry*. **76**, 5614–5627 (2011).

519 22. DNA Sequence Design Tools http://www.dna.caltech.edu/DNA_Sequence_Design_Tools/
520 (2020)

521 23. Dirks, R.M., Lin, M., Winfree, E., Pierce N.A. Paradigms for computational nucleic acid design.
522 *Nucleic Acids Research*. **32** (4), 1392–403 (2004).

523 24. Sen, A., Nielsen, P. E. On the stability of peptide nucleic acid duplexes in the presence of
524 organic solvents. *Nucleic Acids Research*. **35**, 3367–3374 (2007).

- 525 25. Yang, Z., Williams, D., Russell, A.J. Synthesis of protein-containing polymers in organic
526 solvents. *Biotechnology and Bioengineering*. **45**, 10–17 (1995).
- 527 26. Coin, I., Beyermann, M., Bienert, M. Solid-phase peptide synthesis: from standard procedures
528 to the synthesis of difficult sequences. *Nature Protocols*. **2**, 3247 (2007)

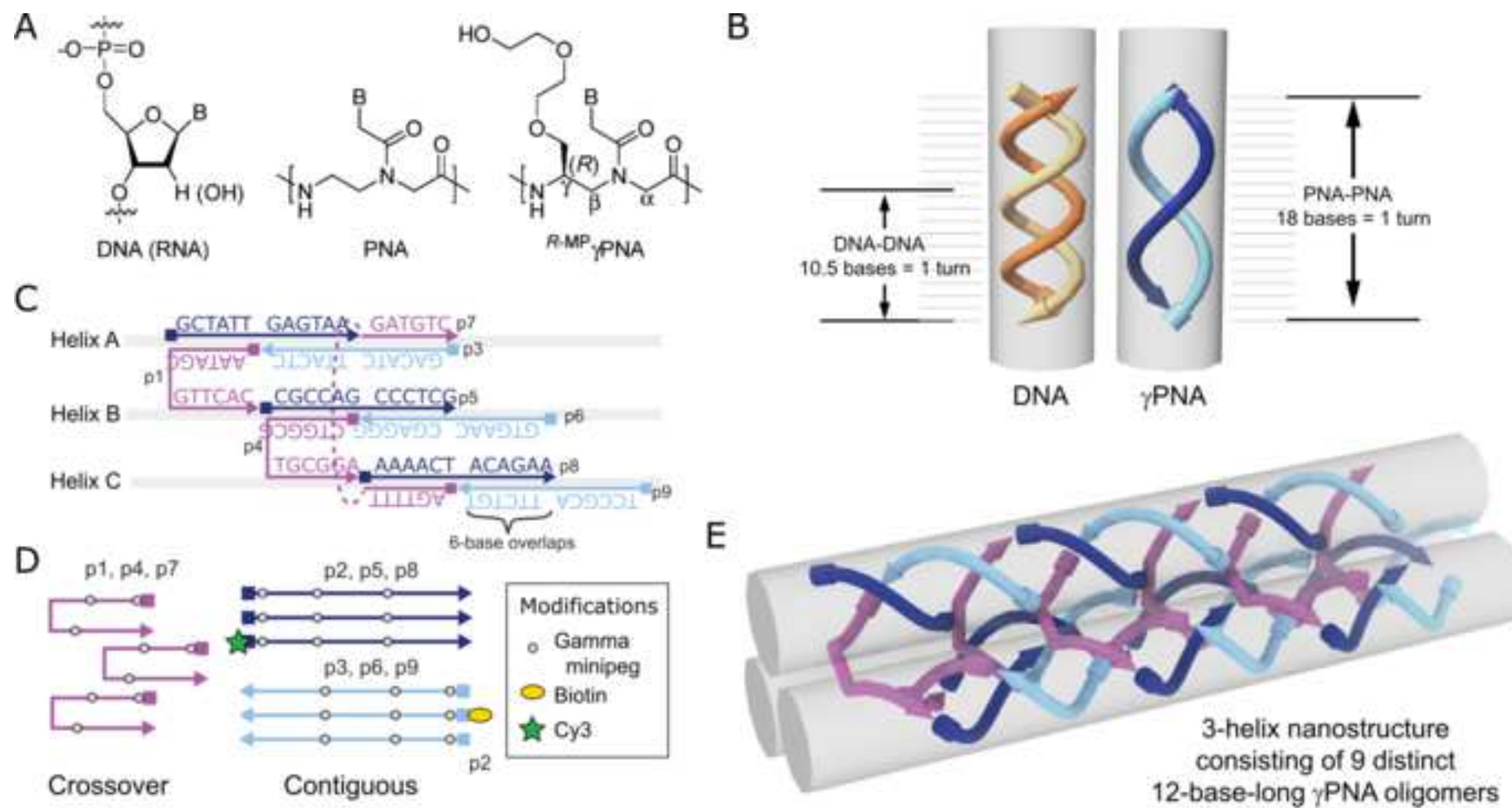
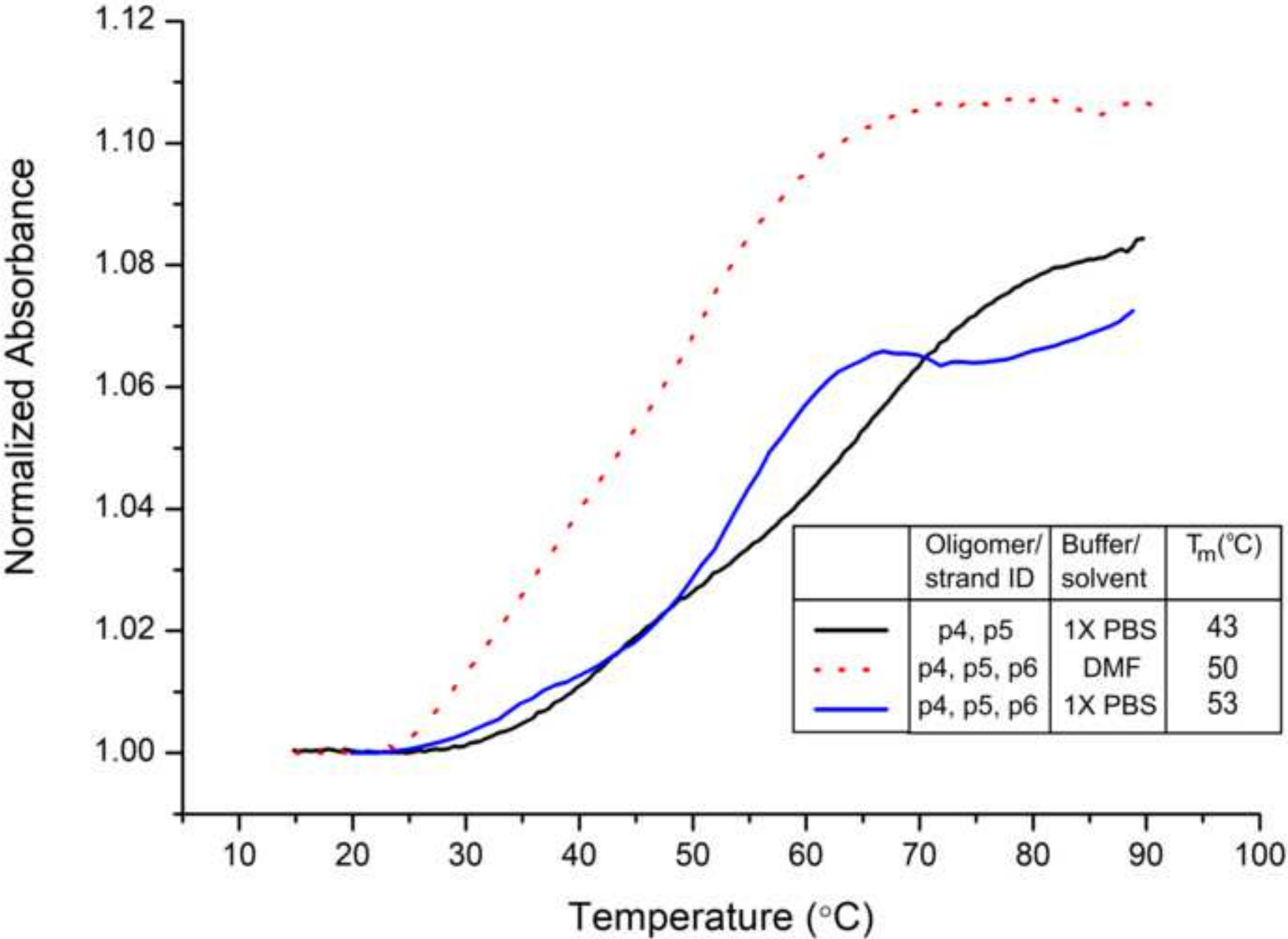
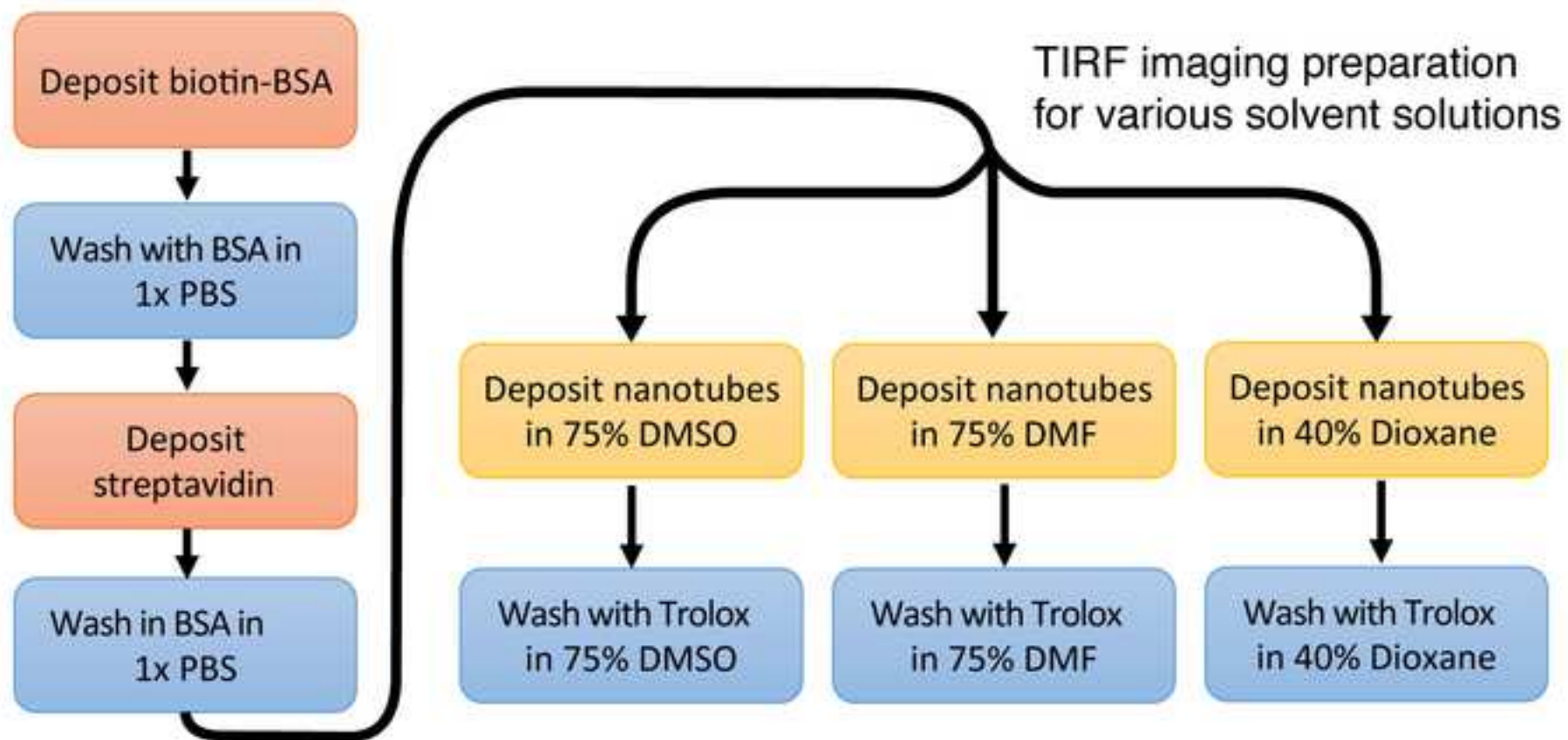
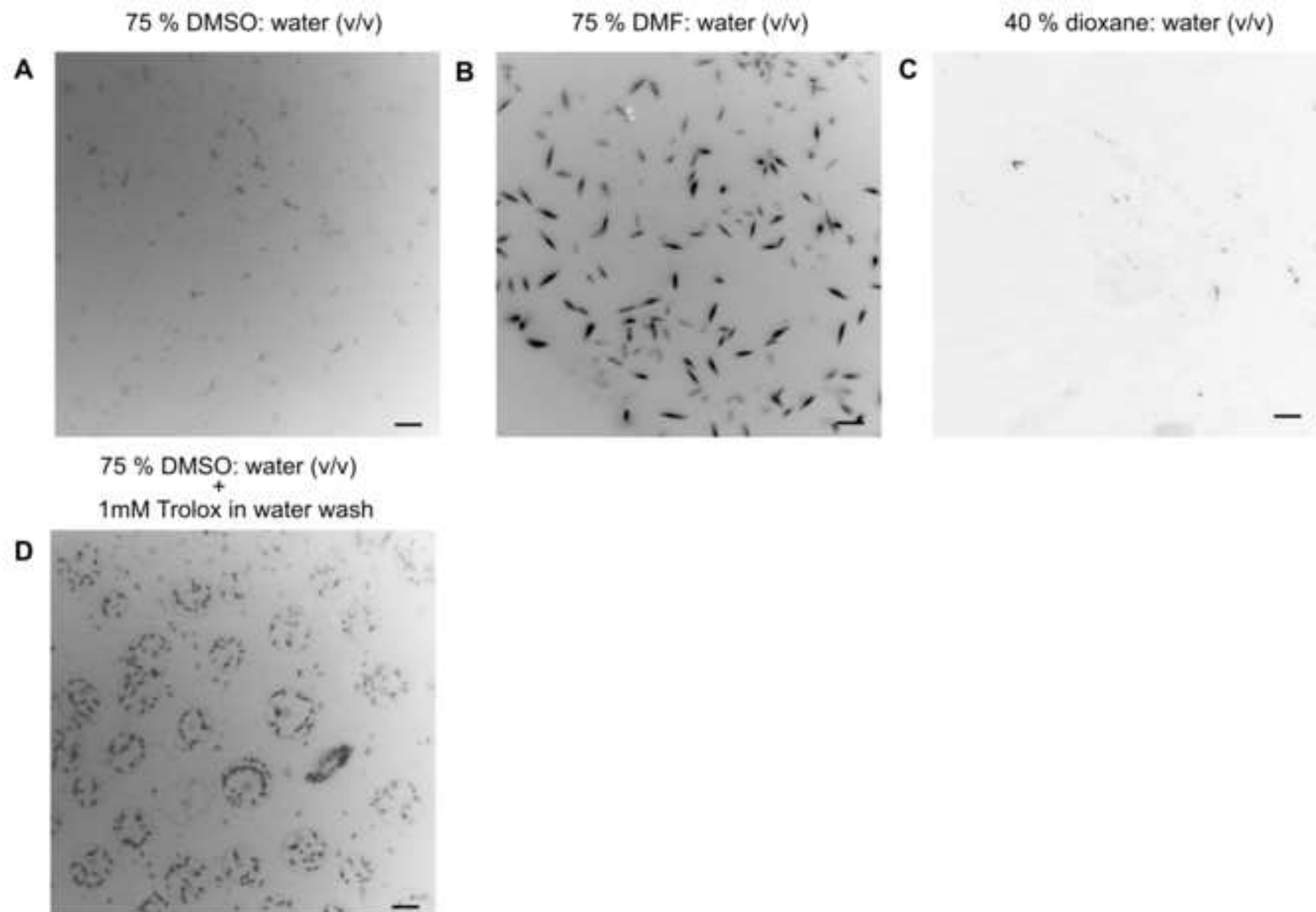
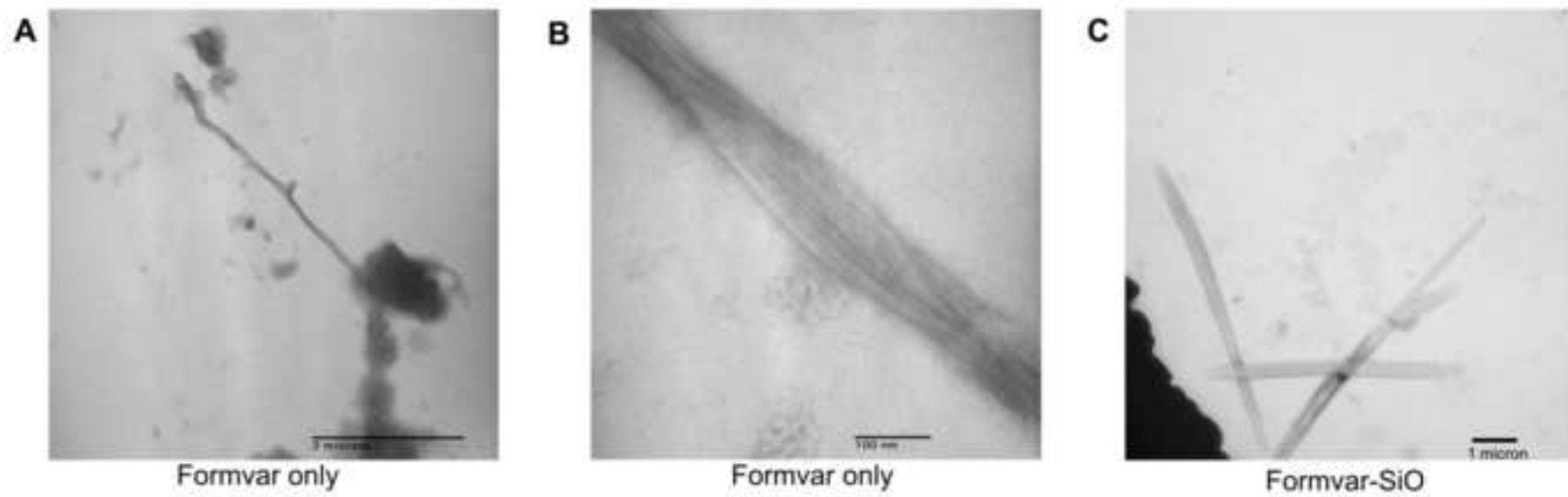


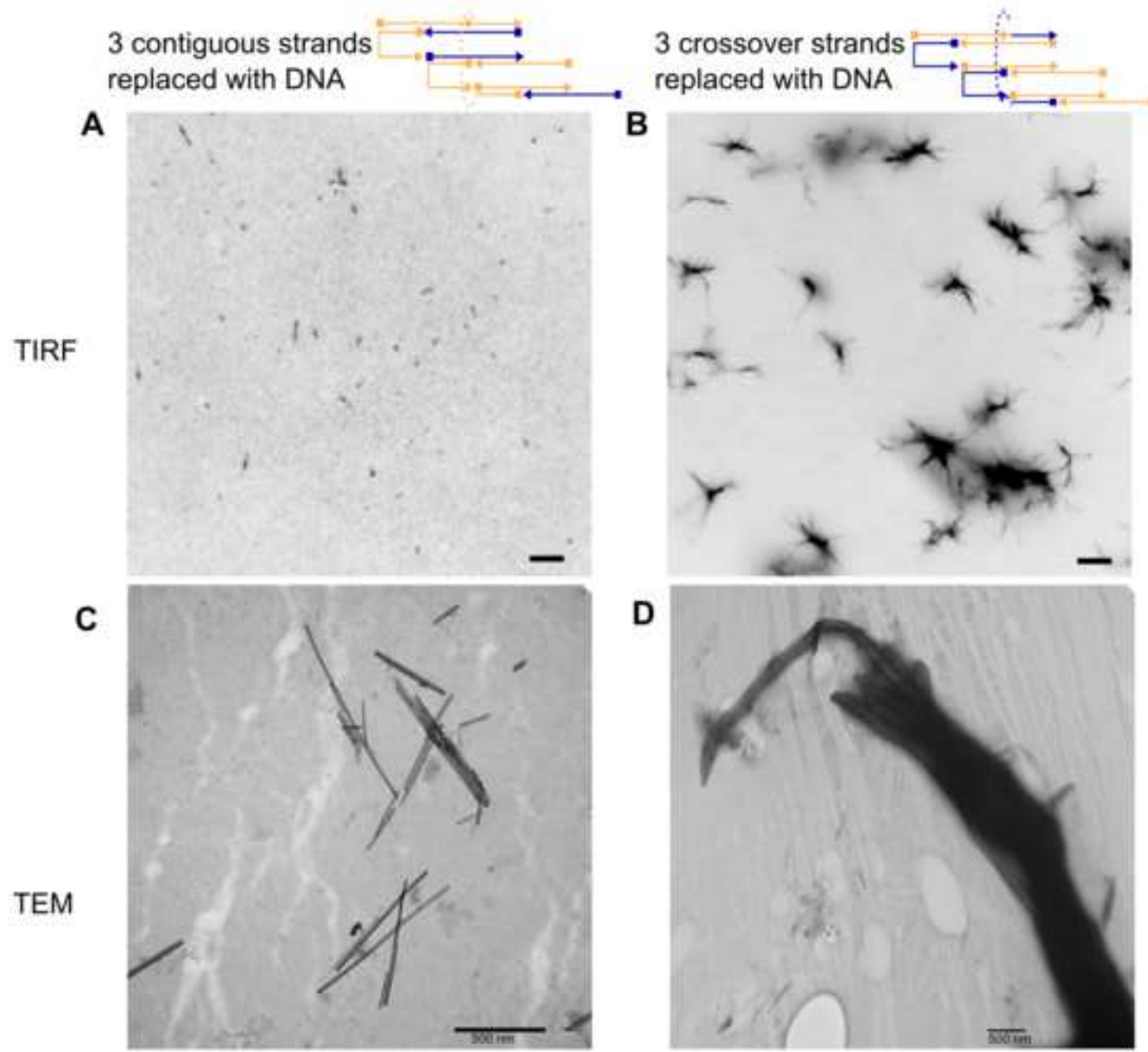
Figure 2

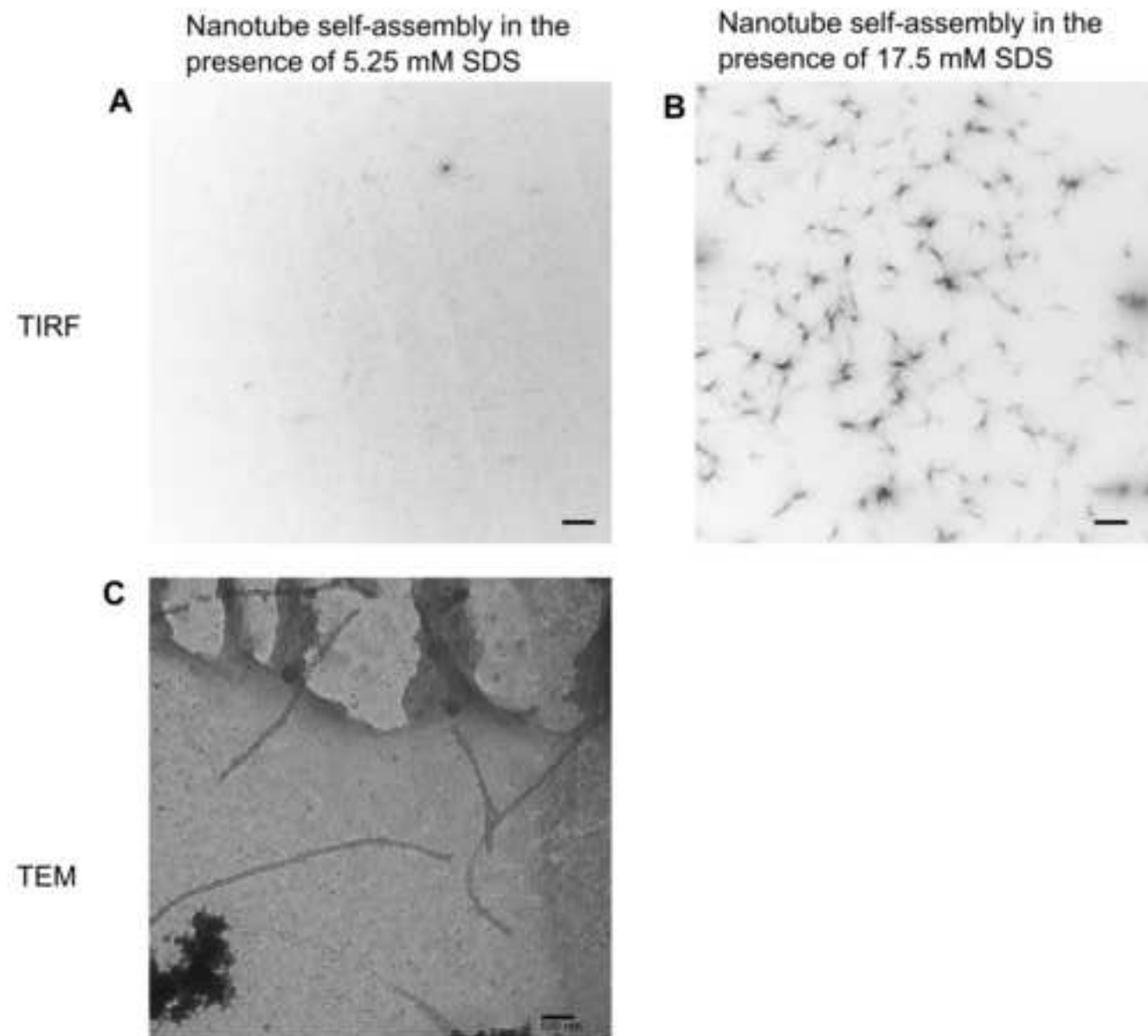












MATLAB script string output "thisSeq"
'acttcgctaaaa cgaagtgaggaa cagtatttcctc aacgagaacacc ctcgttgccgcg ttttaggcgggc ggattgatactg caatcccatagc g
'ccctcccgaccg ggagggttcagg cacttgccctgaa tggcgttgctat acgccattccaa cggcgttgga gagatacaagt tatctcatttac at
'tcgcaggagaca ctgcgaggataa aacggcttatcc ccacttgccccg aagtggacgatt tgtctcaatcgt aaatgtgccgtt acattttacaa c
'gagccccctact gggctcttgata tttcgctatcaa tccacgaccgcc cgtggacaataa agtaggttattg acagatgcgaaa atctgttccttt gg
'cgggagaaccac ctcccgtttagg ctgaccctaaa gcttacactcgc gtaagccgatga gtggtttcatcg gcaatagtcaag tattgcctgct g
'aatgagccgtgc ctcatcttatct tagggcagataa ctttcgcccaa cgaaagcagtc gcacggggactg caatacgcccta gtattggaggaa t
'gatggaagcccc tccatgcctgt ccagagacaggc aagtcaagtagg tgacttttattg ggggctcaataa gcaacgctctgg cgttgctcggtg c
'tagccccccagt gggctatctcgt ttcaggacgaga ctgcggtgtcg cgcaagagtatt actgggaatact gatttacctgaa taaatcaaaggc c
'taggcaacagac tgcctaccactc tttggagagtgg tagcccctgcgg gggctattcatc gtctgtgatgaa ttcccgccaac cgggaaacctta c
'aatgtgtctatc cacattcccttc cgccaagaaggg gtgctgttatgc cagcacgactaa gatagattagtc cctcggttggcg ccgaggtcaaac g
'ttggcgttatcc cgcaaagtctt ggacgaaagcat accgagtgaaca ctcggtggggtc ggataagacccc tctacatcgcc ttagagtgcct t
'ctgtaaggcttc ttacaggtgctt cacgcaaagcac ttcgggatggag cccgaacaatct gagaccagattg gcggactgcgtg gtccgcctattt c
'actccaatctat tggagttttgtt ctaccaacaaa cgctcaagtaat tgagcgaacggc atagatgccgtt ctgcgggggtag ccgcaggtcttc at
'ggttgttccacg acaacctgaagc ttatgtgcttca ttgccccgagcg gggcaatcttc cgtggagaaaga ctactgacataa cagtagacgatg c
'gatttgctgtct caaatcccttct agcgtagaagg cataaaaccag tttatgcctcac agacaggtgagg ctacggaaacgt ccgtagtcgtcc ct
'gaaggctaatg acctcatcctg gcagttcaggat ttggtagcggag taccaaaaatcg cattagcgattt acgacaaactgc tgtcgtaagtgg c
'tgtagtggctt actacacccttg ggacatcaaggg ttcggcaggcgg gccgaacgctaa agaccattagcg acgagtatgtcc actcgtattttg c
'tggaaacctgta gttccattcgca atcacctgcgaa ccacacttactg gtgtggtcggct tacaagagccga ggcgttggtgat aacgccctatct ca
'agaggctgtatt acctctgtgagt cggtttactcac actttccagcaa gaaagtccatcc aatacgggatgg tagcccaaaccg gggctaaggcga t
'cggcaaacctatg ttgccgatttct acttacagaaat cgtgtcctaaag gacacgggatgg catagtccatcc ttaggcgtaagt gcctcacagcga c

Penalty Score "thisScore"
0.08
0.0544
0.0688
0.0528
0.0656
0.0592
0.0688
0.0656
0.0736
0.0768
0.0576
0.072
0.064
0.0688
0.0688
0.0576
0.0624
0.0656
0.0688
0.0704

γPNA Sequence ID	Sequence	Domain 1 complement
p1	N-AATAGCGTTCAC-C	p2 domain 1
p2	N-GCTATTGAGTAA-C	p1 domain 1
p3	N-GACATCTTACTC-C	p7 domain 2
p4	N-CTGGCGTGCGGA-C	p5 domain 1
p5	N-CGCCAGCCCTCG-C	p4 domain 1
p6-Biotin	N-Biotin-GTGAACCGAGGG-C	p1 domain 2
p7	N-AGTITTGATGTC-C	p8-Cy3 domain 1
p8-Cy3	N-Cy3-AA AA CTACAGAA-C	p7 domain 1
p9	N-TCCGCATTCTGT-C	p4 domain 2

Domain 2 complement
p6-Biotin domain 1
p3 domain 2
p2 domain 2
p9 domain 1
p6-Biotin domain 2
p5 domain 2
p3 domain 1
p9 domain 2
p8-Cy3 domain 2

Temperature Range	Ramp rate
90 °C	Hold for 3 minutes
90-80 °C	0.1°C/minute
80-70 °C	0.1°C/minute
70-60 °C	0.1°C/3 minutes
60-50 °C	0.1°C/3 minutes
50-40 °C	0.1°C/3 minutes
40-30 °C	0.1°C/minute
30-20 °C	0.1°C/minute
4 °C	Hold indefinite

	Stock concentration	75% DMSO: water (v/v) anneal sample
γPNA oligomer (x9)	20 μM	9 μL (1 μL x 9)
DMSO	-	30 μL
DMF	-	-
1,4-dioxane	-	-
deionized water	-	1 μL
Total volume	-	40 μL

75% DMF: water (v/v) anneal sample	40% dioxane: water (v/v) anneal sample	Final Concentration
9 μ L (1 μ L x 9)	9 μ L (1 μ L x 9)	500 nM
-	-	75 % (v/v)
30 μ L	-	75 % (v/v)
-	16 μ L	40 % (v/v)
1 μ L	15 μ L	25 or 60% (v/v)
40 μ L	40 μ L	-

DNA Sequence ID	Sequence
D1	5'-AATAGCGTTCAC-3'
D3	5'-GACATCTTACTC-3'
D4	5'-CTGGCGTGCGGA-3'
D5	5'-CGCCAGCCCTCG-3'
D7	5'-AGTTTTGATGTC-3'
D9	5'-TCCGCATTCTGT-3'

	Stock concentration	DNA Contiguous strand replacements
γPNA oligomer (x6)	20 μM	6 μL (1 μL x 6)
DNA oligomer (x3)	20 μM	3 μL (1 μL x 3)
DMSO	-	30 μL
deionized water	-	1 μL
Total volume	-	40 μL

DNA Crossover strand replacements	Final Concentration
6 μL (1 μL x 6)	500 nM
3 μL (1 μL x 3)	500 nM
30 μL	75 % (v/v)
1 μL	25 % (v/v)
40 μL	-

	Stock concentration	SDS concentrations below CMC
γPNA oligomer (x9)	20 μM	9 μL (1 μL x 9)
6% SDS	6% (wt/v)	1 μL
20% SDS	20% (wt/v)	-
DMSO	-	30 μL
deionized water	-	-
Total volume	-	40 μL

SDS concentrations above CMC	Final Concentration
9 μL (1 μL x 9)	500 nM
-	5.25 mM
1 μL	17.5 mM
30 μL	75 % (v/v)
-	25 % (v/v)
40 μL	-

Name of Material/ Equipment	Company
γPNA strands/oligomers	TruCode Gene Repair Inc.
UV-Vis Spectrophotometer	Agilent
Quartz cuvettes	Starna
Thermal cycler	Bio Rad
0.2 mL PCR tubes	VWR
Anhydrous DMF	VWR
Anhydrous DMSO	VWR
Anhydrous 1,4-Dioxane	Fisher Scientific
10X Phosphate Buffered Saline (PBS)	VWR
Microscope slides	VWR
Glass cover slips	VWR
2% Collodion in Amyl Acetate	Sigma-Aldrich
Isoamyl Acetate	VWR
Biotinylated Bovine Serum Albumin (Biotin-BSA)	Sigma-Aldrich
Bovine Serum Albumin (BSA)	Sigma-Aldrich
Streptavidin	Sigma-Aldrich
Trolox	Sigma-Aldrich
Total Internal Reflection Fluorescence microscope	Nikon
Transmission Electron Microscope	Joel
Tweezers	Dumont
Uranyl Acetate	Electron Microscopy Sciences

Formvar, 300 mesh, Copper grids	Ted Pella Inc.
Formvar-Silicon monoxide Type A, 300 mesh, Copper grids	Ted Pella Inc.
DNA oligomers/strands	IDT
Sodium Dodecyl Sulphate (SDS)	VWR

Catalog Number	Comments/Description
	Section 2.1
Varian Cary 300	Section 3.1.2
29-Q-10	Section 3.1.1
C1000 touch	Section 4.1
53509-304	Section 4.5
EM-DX1727-6	Section 4.6
EM-MX1457-6	Section 4.6
AC615121000	Section 4.6
75800-994	Section 3.1.1
89085-399	Section 5.2
48382-126	Section 5.2
9817	Section 5.2
200001-180	Section 5.2
A8549	Section 5.3
A2153	Section 5.4
189730	Section 5.5
238813	Section 5.7
Nikon Ti2-E	Section 5.8
JEM 1011	Section 6.6
0203-N5AC-PO	Section 6.3
22400	Section 6.1

1701-F

1829

97064-860

Section 6.2

Section 6.2

Section 7.1

Section 8.1



Department of Mechanical Engineering
Carnegie Mellon University
5000 Forbes Avenue
Pittsburgh, Pennsylvania 15213-3890
Phone: (412) 268-2500
Fax: (412) 268-3348

May 31, 2020

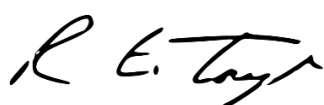
To the Editors for Journal of Visualized Experiments,

In this resubmission of manuscript **61351_R1**, this manuscript titled “Self-Assembly of Gamma-Modified Peptide Nucleic Acids into Complex Nanostructures In Organic Solvent Mixtures” has been revised to address the editor comments and questions. We have listed our point-by-point revisions within this document. We have also uploaded two documents – one with highlighted tracked changes and one with the revised document.

We would like to thank our editors for their insightful comments. Our point-by-point response to the editorial requests (with indicated lines in the tracked changes document) are described below:

1. We have now edited and replaced commercial sounding software MATLAB with a more generic term in section 1 and all other occurrences (lines 87-98,427-428).
2. We have moved our hyperlink to the reference section (ref 22) and used superscript citations.
3. We have removed all second person pronouns from all sections (line 93, section 1.2).
4. We have verified the uploaded PNA3nanofiber.m file and checked that the text references the same file (section 1.3).
5. We have also edited out the image of the algorithm and uploaded it as Supplementary Figure 1 as suggested (section 1.3).
6. We have also defined all acronyms with their expanded terms as suggested by our editor in sections 2 and 3 (lines 125-126, 135-136).
7. We have also described concentrations as our editors have suggested in the correct format in section 3.1.1.
8. We have now rearranged statements as suggested by our editor for sections 5.3, 5.4, 5.5, 5.6 and 5.7. We have also added descriptions to specific statements to include concentrations, incubation related to flow channels, sample flow throughs to enable readers to understand our procedure wherever necessary.
9. We have also added a supplementary figure 1 caption in lines 451-452.

Sincerely,



Rebecca E. Taylor, Ph.D.

Assistant Professor of Mechanical Engineering, and, by courtesy,

of Biomedical Engineering and Electrical and Computer Engineering
Carnegie Mellon University
Email: bex@andrew.cmu.edu

```

function [thisSeqs,thisScore] = PNA3nanofiber()

clear all
close all

%-----
% SST strands
% 4 helices x 3 strands per "section" = 12 strands
% each is 12-nt long

SSTseq = repmat('NNNNNNNNNNNNN ',1,9);

% SSTconstr = [a b c d e];
% a = strand id number
% b = base number - start complementarity region (from 5' or N-terminal end of strand)
% c = strand id number
% d = base number - end of complementarity region (from 3' or C-terminal end of strand)
% e = number of bases that should be complementary

%Visualization of 3-helix nanofiber structural motif
% 0 5 10 15 20 25 30
%S0N% \-07->[---02--->\-07->[---02--->\-16->
%S0C%    /-01-]<---03---]
%S1N%    \-01->[---05--->
%S1C%        /-04-]<---06---]
%S2N%        \-04->[---08--->
%S2C%            /-07-]<---09---]
%S0N% \-07->[---02--->\-07->[---02--->\-16->

SSTconstr=[
    2 1 1 6 6; 2 7 3 12 6; 7 7 3 6 6;
    5 1 4 6 6; 5 7 6 12 6; 1 7 6 6 6;
    8 1 7 6 6; 8 7 9 12 6; 4 7 9 6 6
];
[St,wc,eq] = constraints(SSTseq,SSTconstr,[]);
rS=randbase(St);
% CONSTRAIN INITIAL CONDITION
S=constrain(rS,wc,eq)

% OPTIMIZE
load Bad6mer
global bestS bad6mer
bad6mer=make_bad6mer;

[thisSeqs,thisScore] = optimize(S,St,wc,eq, 'score_spurious');
thisSeqs
thisScore

```



Click here to access/download
Supplemental Coding Files
PNA3nanofiber.m