Functionalization of Quantum Rods with Oligonucleotides for Programmable Assembly with DNA Origami

Supplemental Information

Tennyson L. Doane, Rabeka Alam, Mathew M. Maye*

*Department of Chemistry, Syracuse University, Syracuse, NY 13215. a Current Address: Department of Chemistry and Biochemistry, Notre Dame University, Notre Dame Indiana, 46556, U.S.A. *mmye@syr.edu

Materials: Single stranded DNA was purchased from Integrated DNA Technologies. M13mp18 was purchased from New England Biolabs. Sodium hydroxide and chloroform were purchased from Fisher. Cadmium oxide (CdO, \(\geq 99.99\%\)), sulfur (reagent grade), selenium powder (Se, reagent grade \(\geq 99.999\%\)), 1-octadecene (ODE, 90%), trioctylphosphine oxide (TOPO, 90%), hexadecylamine (HDA, 98%), trioctylphosphine (TOP, 97%), magnesium chloride, ethyldiaminetetraacetic acid, toluene, ethidium bromide, 10x TAE buffer, 10x TBE buffer, rhodamine 6G, pentamethylethlenediamine (PEMED), and ammonium persulfate (APS) were purchased from Sigma Aldrich. GelSafe Green and GelSafe Red (Biotium) were purchased from VWR. Octadecylphosphonic acid (ODPA, 98%), n-decylphosphonic acid (DPA, 97%) and hexylphosphonic acid (HPA, 98%) were purchased from Strem Chemicals. All chemicals were used without further purification. Both Ethanol (Pharmco-AAPer) and 40% acrylamide/bisacrylamide monomer solution (Baker) were purchased and used without further purification.

Instrumentation: Optical absorption measurements were performed using either a Varian Cary 50 Bio spectrometer or a Cary 100 spectrometer, the latter of which was equipped with a multicell peltier heating accessory. The photoluminescence and anisotropy measurements were conducted with a Fluoromax4 spectrofluorometer equipped with automated excitation and emission polarizers. Quantum yield measurements were preformed on dilute samples (O.D. \(\text{Excite} < 0.1\)) and excitation of the core structure alone (\(\lambda = 500 \text{ nm}\)) versus Rhodamine 6G in ethanol as the standard.1 Additional PL were obtained using a Varian Cary Eclipse spectrometer that was equipped with a plate reader. AFM imaging was performed on a Digital Instrument Nanoscope IIIa using an E-scanner and silicon AFM tips (\(\mu\text{Masch, HG:NSC14 / AlBs}\)) that had a typical resonance frequency of \(~160 \text{ Hz}\). Imaging was performed on Mica substrates (Ted Pella) that were freshly cleaved and pre-soaked with 35 \(\mu\text{L}\) of 0.1M magnesium acetate (20–30 min) before being rinsed with 35 \(\mu\text{L}\) of 10 mM Tris Acetate buffer. Typically, a 5 \(\mu\text{L}\) sample solution was added to the surface, and diluted with 30 \(\mu\text{L}\) of 10 mM TA buffer. The substrate was subsequently allowed to sit for 15 minutes, and the solution was wicked away using filter paper and dried for 90s with argon gas. The dried samples were imaged in tapping mode. The confocal microscopy was performed on a Zeiss LSM-710 microscope. For these experiments, the 15\(\mu\text{L}\) of purified QR+origami conjugates were incubated with 1x GelSafe Green for ~10 minutes and then 2 \(\mu\text{L}\) of a 40% acrylamide-bisacrylamide solution (29:1) was added. A stock solution of 10% ammonium persulfate was prepared and 1 \(\mu\text{L}\) was added to the mixture, followed rapidly by pentamethylethlenediamine to give a 0.25% concentration, and the gel was sandwiched between two pre-cleaned microscopy cover slips.
Quantum Rod Synthesis & Phase Transfer: CdSe/CdS QRs was synthesized using previously published seeded growth approaches with modification. CdSe QR seeds was synthesized by injecting Se-TOP (0.7 mmol) into a hot mixture of CdO (0.5 mmol) dissolved in purified TOPO (8.0 mmol), phosphonic acid (0.9 mmol) and octadecene (2 ml) at 350 °C in an inert atmosphere and allowing growth of CdSe QRs to occur for 10 minutes before quenching the reaction by cooling to room temperature. CdSe QRs with /w = 2.5 ± 0.3 (l = 9.9 ± 0.9 nm, w = 2.5 ± 0.3 nm) for seeded growth of QR(615) were prepared by using DPA (0.9 mmol) and CdSe QRs with /w = 4.1 ± 0.4 (l = 19.4 ± 1.6 nm, w = 4.7 ± 0.3 nm) for seeded growth of QR(650) was prepared by using a mixture of MPA (0.1 mmol) and ODPA (0.8 mmol). Before using CdSe QRs for seeded growth of CdS QRs, CdSe QRs were purified by precipitating QRs in a combination of toluene and methanol via centrifugation two to three times. Concentration of QRs was quantified using previously reported method. For seeded growth of CdS, a solution of CdSe QR seeds (10 nmol) and S (2.0 mmol) dispersed in TOP (2 ml) was injected into a mixture of CdO (0.5 mmol) dissolved in TOPO (8.0 mmol), HPA (0.2 mmol), ODPA (0.4 mmol) and octadecene (2 ml) at 350 °C and allowing growth of QRs to occur for 10 minutes before quenching the reaction by cooling to room temperature. CdSe/CdS QRs was purified via precipitation as described above and stored in toluene. A set of TEM images and statistical analyses is shown in Fig. S1.

Phase transfer of the QRs was accomplished using a histidine-mediated phase transfer route. In a typical experiment, 400 µL of 4.55 µM of the QR solution in toluene was dried to a thin film using a nitrogen stream, and then re-suspended in 200 µL of chloroform. Then 200 µL of 114 mM L-Histidine was added and the solution was vortexed, with immediate phase transfer. Next, 200 µL of 10 mM Tris-acetate buffer (pH 8.2) was added and again the solution was vortexed. The lower chloroform layer was extracted, and the solution was cleaned 3 times with excess chloroform for complete removal of organic ligands. The final aqueous layer was transferred to a 3 kDa molecular weight cutoff (MWCO, Amicon) filter and centrifuged at 6000 rpm for 5 minutes. Experimentally, repeating this step to ensure complete removal of histidine resulted in significant loss of His-QRs. The remaining solution was collected and diluted to 200 µL as a stock solution. We observed that to achieve the best stability, a reaction ratio of histidine such that the ligand:QR surface area was 40-50 / nm² was needed.

DNA Functionalization: In a typical reaction, the U-type ssDNA and it’s complementary protecting strands (U’1 and U’2 for example) were incubated in 10 mM tris acetate buffer with 100 mM NaCl at 4°C to ensure complete protection of the outermost 15 bases. This solution was mixed with the QR at a ratio of 100:1 (typical QR concentration ~0.9 µM) at 4°C. Salt aging was employed, moving from a nominal ionic strength of 30 mM to a final ionic strength of 100 mM over the period of three hours. The reaction was allowed to proceed for 48 hours to ensure maximum DNA coverage. Purification of the dsDNA coated QRs was conducted using centrifugation-induced precipitation at 13,000g for 15 minutes. Successive centrifugations after dilution with 10 mM tris acetate both decreased the overall DNA concentration and lowered the ionic strength, leading to favorable melting temperatures for dehybridizing the dsDNA. In the case of the T series, a 30% isopropanol:tris acetate buffer solution could be used to lower the melting temperature for dsDNA, and also induce better rod precipitation. In later experiments, we used ultracentrifugation to substantially increase particle pelleting for more thorough purification.
DNA Quantification: The quantification of ssDNA grafted to the QR was conducted using a modified Cy 3 methodology we have used in previous work. A 100 kDa MWCO filter was used for separating bound and unbound Cy3, with the bound Cy3 modified QRs remaining in the above solution. The filters were first pretreated with a 1% Triton 100x solution in Tris-Acetate buffer to avoid non specific absorption (Fig. S3). In this step, the filters were first loaded with 400 µL of Triton and centrifuged at 12,700 RPM for 3 minutes, with the process repeated 2 additional times. The tubes were then washed with 400 µL of 0.75M NaCl in 10 mM Tris-Acetate Buffer to remove any excess Triton X. Hybridization of the U’ – Cy 3 ssDNA to the U ssDNA coated QRs was achieved by mixing the two components at 50:1 ratio (QR ~ 100 nM) in 10 mM Tris-acetate with 100 mM NaCl at room temperature overnight and compared to a control solution of Cy 3 under identical conditions. The resulting solutions were then diluted with 200 µL of 0.75M NaCl, transferred to freshly prepared Triton-treated MWCO filters, and centrifuged at 12,700 rpm for 3 minutes with cooling. The filtered solutions (i.e. free U’-Cy3) were then transferred to a quartz cuvette and both absorbance and fluorescence measurements were recorded as a function of serial dilution to establish a calibration curve. The equation of the lines for both calibration curves were used to calculate the amount of Cy 3 in the original solution, and taking into account the dilution during centrifugation, the amount of Cy 3 that did not transfer was calculated. In later experiments, ultracentrifugation could also be used for particle pelleting and the supernatant was examined to determine unbound Cy 3.
DNA Origami Substrate Design and Synthesis. The design of the DNA Origami substrates was carried out based on the methods developed by Rothemund and Winfree. Firstly, our DNA origami “substrate” were designed with the aid of the caDNAno software package using circular ssDNA M13mp18 as the main strand. Only 7168 of the total 7249 bases were utilized (the specific base sequence for M13mp18 is available at New England Biolabs). Scheme S1a shows the specific features of the origami. It can be divided into 5 zones where three are used as inert regions (red staples) and the other two (blue staples) are used as active binding zones (Zones A and B). A column of capture strands was inserted into the middle of each capture zone, with a nominal distance of ~75 nm between the two columns. Using a dsDNA helix-helix distance of 3 nm (1 nm helix radius, 1 nm helix spacing), the capture strands are placed 6 nm apart. The large distance was chosen to prevent steric interactions and possible binding of one QR between the two zones. To develop face specific orientation on the origami, the end of the staple was designed to terminate with the 3’ facing out of the plane as previously described. Scheme S1b shows the staple schematic of the origami, with the majority of staples consisting of 32 base ssDNA. Using the knowledge that ~11 bp composes one complete turn of a dsDNA helix, attention to where the M13mp18 strand helix is facing relative to the plane of the DNA origami allows for the termination of the staple strand in an “up” or “down” orientation (Scheme 1b,
helix illustration). In both Zones A and B, strands are allowed to periodically terminate in the up face to allow the display of a capture strand. These capture strands are integrated into the substrate and leave dangling a 5b poly-T spacer, followed by the complementary sequences (i.e. $U'$). Importantly, the origami design allows for more rigorous investigations on the relation between complimentary ssDNA display (number and position) and rod capture efficiency (orientation and yield), which we are actively pursuing. A more detailed schematic and full list of staple strands are shown in Appendix S1 and S2, respectively.

The assembly of the substrates was achieved as follows: 0.5 μL of a 150μM stock of each capture strand was added to a small PCR tube (212 strands × 0.5 μL = 106 μL), and diluted to 1060 μL; 106 μL of this solution was combined with 15 μL of 52.5 nM M13mp18 to obtain a final concentration of 60 nM staple / 5 nM M13mp18. The solution was also composed of 40 mM Tris-Acetate, 10 mM magnesium acetate, and 1 mM ethylendiaminetetraacetic acid (EDTA), with a total volume of 153 μL. The mixture was then heated in a Techne TC-3000 temperature cycler according to the temperature profile previously reported. After reaching 25°C, the solution was transferred to a 100 kDa MWCO filter and ~300 μL of 0.75M NaCl in 10 mM Tris Acetate was added. The solution was centrifuged at 12,000 G for 2 min, and then refrigerated for 5-10 minutes. This process was repeated three more times (50 μL hold up volume, 350 μL of 0.75M NaCl added), and finally 350 μL of 10 mM Tris Acetate buffer was added to remove excess salt. Monitoring the filtrate via UV-Vis absorbance confirmed that this removed the majority of excess DNA (data not shown). The solution was stored in the fridge and used as required.

**QR-Origami Assembly & Gel Electrophoresis Assessment.** The loading of QR onto the origami substrates was conducted at 25°C for a period of 1-2.5 hrs in the presence of 100 mM NaCl and 10 mM tris acetate. The introduction of rigidity to a longer poly T spacer region ($U_{\text{Rigid}}$, SI Table 1) using a polyA sequence resulted in better hybridization (see SI Fig. 7). Assessment of the QR-Origami substrates was achieved via gel electrophoresis of DNA-QR conjugates using standard agarose gel electrophoresis protocols. Briefly, a 0.5% w/w percent agarose gel was prepared by dissolving 0.25 g of agarose in 50 mL of a 0.5x TBE buffer solution with subsequent heating in a conventional microwave and poured in to a VWR Mini-cast gel system with a subsequent cooling time of 40 minutes. The gel was then submerged under 0.5x TBE buffer, and each well was loaded with 20 μL of sample loaded with 15-25 v/v% glycerol. The gel was run at room temperature for 20-40 min with a constant applied voltage of 80V. In the experiment shown in Fig. 5, the samples were injected at two different well comb positions and run simultaneously; the gel was then split and one half was stained with GelSafe Red. The other half was cut into 3 slices and imaged using a Cary Eclipse equipped with a well plate reader accessory. Imaging of the stained gel segment was conducted using a 300 nm excitation source (VWR) and a digital camera (Cannon). The optimization of gel electrophoresis showed that increasing agarose concentration increased separation efficiencies at both shorter (Fig. S4a) and longer (Fig. S4b) run times; thus, a short gel electrophoresis experiment (15 minutes) with higher agarose concentration (0.75%) was used to separate excess QRs from the resulting conjugates (Fig. S4c). After centrifugation through a Freeze N’ Squeeze centrifugal filter (Bio-Rad) to extract the conjugates and concentration through a MWCO filter, photoluminescence measurements (Fig. S4c) showed that the optical properties of the rod were still intact.
**QR-Origami Assembly and Orientation Statistics.** AFM images for multiple assembly were utilized, typically encompassing 10-12 different sampled areas. Unpurified reaction mixtures were utilized to determine the extent of hybridization, and often agglomerated structures were visible but uncounted. The criteria for counting included: A) structures which could be identified individually (no structures with two or more origami in indistinguishable contact), B) origami with dimensions expected based on caDNAno predictions (~150 nm ± 15 nm) and rectangular shape, and C) raised heights ≥ QR width. All statistics reported were for assembly populations ≥ 99 origami. Orientation of the QR on the DNA origami was determined by measuring the angles of the origami and of the assembled QR relative to the horizontal axis of the image, and taking the difference of two (0-90°). QR angle was determined by finding the long z-axis given the asymmetric shape; QRs which could not be easily distinguished were excluded from orientation calculations.

**Ultracentrifugation:** Purification of the QR-origami assembly solutions could be carried out via gradient ultracentrifugation. A density gradient which has previously been shown to be successful at separating folded origami from free strands and larger aggregates was used. In a typical experiment, a ultracentrifuge tube loaded with a 15%-65% sucrose gradient (w/v) was prepared by successively adding heavier solutions from the bottom of the tube up, and the solutions were allowed to set at 4°C overnight to form a continuous gradient. A concentrated solution of QR+origami assembly solutions was prepared with a 20:1 QR excess at 100 mM NaCl concentration for 4 hours at ~14°C (to protect the DNA strands and improve hybridization) under mild stirring using a Eppendorf Thermomixer. A control solution of QRs at the same concentration was mixed as well. Ultracentrifugation was conducted at 131,000g for 30 minutes at 4°C using a Beckman Coulter Optima Max XP Ultracentrifuge. The tubes were imaged using a digital camera (Canon EOS Rebel T5 EOS 1200D) and under UV excitation. The separated regions after ultracentrifugation could be collected either via a pipette, or via a needle penetrating the tube. After collection, the solutions were dialyzed against 100 mM NaCl with 10 mM tris acetate for ~ 15 hours at 4°C with two buffer exchanges to remove excess sucrose.

**FRET Calculations and Simulations of Orientation Dependence:** The potential energy transfer between QR(615) and QR(650) was assessed using Förster resonance energy transfer (FRET) based calculations. The FRET efficiency ($E$) is given by: \[
E = \frac{1}{1 + (r/R_0)^6}
\]

where $r$ is the donor-acceptor distance and $R_0$ is defined as the Förster distance at which energy transfer is 50%, found using the following relation: \[
R_0^6 = \frac{9000 (\ln 10) \phi_D \kappa^2}{128\pi^5 N n^4} J(\lambda)
\]

with $\phi_D$ is the donor quantum yield, $\kappa^2$ is the dipole orientation factor, $N$ is Avagadro’s number, $n$ is the refractive index, and $J(\lambda)$ is the spectral overlap integral, which relates the normalized donor fluorescence ($f_D$) and acceptor absorbance extinction coefficients ($\epsilon_A$) at each corresponding wavelength, defined as:
Modeling DNA Origami Mechanical Properties: The twisting and bending of the DNA origami was computed using the CanDo software package developed by Bathe and colleagues. The default properties parameters were utilized (Table S2) with fine model resolution. The results of the modeling are summarized in SI Figure 8.

### Supporting Tables

#### Table S1: QR DNA

<table>
<thead>
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<th>Type</th>
<th>Sequence (* = Phosphorothioate bases)</th>
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<td>( U )</td>
<td>( \text{T<em>T</em>T* T<em>T</em>T TTT TTA CTC ACC TAT ATC A} )</td>
<td>16.5</td>
</tr>
<tr>
<td>( U' ),</td>
<td>GTG AGT A</td>
<td></td>
</tr>
<tr>
<td>( U' ),</td>
<td>TGA TAT AG</td>
<td></td>
</tr>
<tr>
<td>( U' )</td>
<td>TGA TAT AGG TGA GTA</td>
<td></td>
</tr>
<tr>
<td>( U_{\text{Rigid}} )</td>
<td>( \text{G<em>G</em>G* G<em>G</em>T TTT TTT TTT TTT ACT CAC CTA TAT CA} )</td>
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<tr>
<td>( T )</td>
<td>( \text{T<em>T</em>T*T TTT TTT TTC ACG ACA CAC TTT G} )</td>
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</tr>
<tr>
<td>( T' ),</td>
<td>GTC GTG A</td>
<td></td>
</tr>
<tr>
<td>( T' ),</td>
<td>CAA AGT GT</td>
<td></td>
</tr>
<tr>
<td>( T' )</td>
<td>CAA AGT GTG TCG TGA</td>
<td></td>
</tr>
</tbody>
</table>

\( T_w \) Experimentally obtained, 100 mM NaCl, pH 8.3 \([S],[S'1],[S'2]\) = 1 \( \mu \)M

#### Table S2: Modeling Parameters for DNA Origami Mechanical Properties used in CanDo simulations

<table>
<thead>
<tr>
<th>Property</th>
<th>Value</th>
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<td>Axial Rise per base Pair</td>
<td>0.34 pN</td>
</tr>
<tr>
<td>Helix Diameter</td>
<td>2.25 nm</td>
</tr>
<tr>
<td>Crossover Spacing</td>
<td>10.5 bp</td>
</tr>
<tr>
<td>Axial Stiffness</td>
<td>1100 pN</td>
</tr>
<tr>
<td>Bending Stiffness</td>
<td>230 pN m^2</td>
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<tr>
<td>Torsional Stiffness</td>
<td>460 pN m^2</td>
</tr>
<tr>
<td>Nick Stiffness Factor</td>
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Supporting Figures

Figure S1. TEM images (i) of QR(615) (a) and QR(650) (b) with statistical analysis (ii-iv). The QR(615) have an average $l/w = 6.8 \pm 0.8$ ($l = 32.3 \pm 2.9$ nm, $w = 4.7 \pm 0.3$ nm) and the QR(650) have an average $l/w = 8.8 \pm 1.4$ ($l = 50.1 \pm 6.1$ nm, $w = 5.7 \pm 0.4$ nm).
Figure S2. (a) Normalized optical absorption (solid lines) and emission (dotted lines) spectra at 500 nm excitation for QR(615) and QR(650) CdSe/CdS QR. Insets show full absorption spectra. The quantum yields ($\phi$) of the CdSe/CdS QRs in toluene and after DNA functionalization for QR(615) were 0.3 and 0.17, respectively, while the corresponding values for QR(650) were 0.2 and 0.07. (b) Fluorescence anisotropy measurements of the as-synthesized QR(615) and QR(650) showing excitation dependent polarized emission.

Figure S3. Comparison of the absorbance and fluorescence (a and b) of remaining Cy3 after hybridization to unprotected and protected U-ssDNA coated QR 615Ls relative to a control solution (shown in figure 1d). A dilution calibration curve was used to assess how much Cy3-U’ did not hybridize. Hybridization conditions: 50:1 Cy-3 to QR concentration (QR = ~80nM) with100 mM NaCl in 10 mM NaCl incubated overnight at 25°C. Subsequent analysis revealed a nearly two fold increase in the amount of DNA when using protected strands (c).
Figure S4. (a-d) Additional AFM images of the DNA origami on Mica.
Figure S5. Additional AFM images of $U_{\text{rigid-QR}(615)}$-origami conjugates. Statistical analysis found that 70% of origami contained at least one rod, with 29% having two per origami ($N=148$) after 2.5 hr incubation.
Figure S6. Representative large scan range (4x4 μm) AFM images of unpurified $U_{\text{rigid}}$-QR(615)-origami conjugates used for assessing assembly yields and orientation control. Unpurified products contain significant number of aggregates in addition to free-standing origami. Only individual origami were counted for assembly yields and relative orientations.
Figure S7. AFM images (517 x 517 nm) in height (a and c), phase (b and d) mode for two different QR(615)-origami revealing variations in orientations of the QR.

Figure S8. Statistical analysis of AFM images for the number of QRs per origami for different DNA configurations and times.
Figure S9. Detailed orientation analysis of (a) all assembled rods \((n = 142)\) and (b) dual-functionalized origami rods \((n = 32)\) as a function of angle relative to the main axis of the rectangular origami. Roughly 90\% of dual modified species contained at least one rod >60°, and nearly 40\% had both rods correctly oriented. The average height of the origami was found to be 2.6 ± 0.4 nm and the raised height of the assembled QRs were found to be 11.6 ± 3.8 nm.

Figure S10. Normalized fluorescence spectra of extracted Origami+dye gel segments for control experiments probing the binding of zone-A and Zone-B, \(T\)-Cy3+Origami (i) \((\lambda_{\text{ex}} = 500\) nm), \(U\)-Atto 647N+Origami (iii) \((\lambda_{\text{ex}} = 590\) nm), and for origami containing both \(T\)-Cy3 and \(U\)-Atto647N (iii), \((\lambda_{\text{ex}} = 500\) nm & 590 nm).
Figure S11. (a) Comparison of the migration of stained origami, QR 650, and QR615-origami reaction mixtures in gels of different agarose concentrations in 0.5X TBE with 80V running voltage for 15 minutes. Increasing the running time to 40 minutes (b) shows a greater separation between free QR and conjugates. (c) Due to fluorescence loss at long running times, however, a short running time experiment was used; subsequent Freeze ‘n Squeeze extraction was utilized to confirm retention of QR optical absorption (dotted line) and fluorescence (solid line, λ\text{Exc} = 400 nm) of the excess QR615 and the conjugates relative to the original solution. Electrophoresis conditions: 0.75% agarose, 0.5X TBE, 80V, 15 minutes.

Figure S12. Images of the static origami as designed in caDNAno (a) and after CanDo modeling as viewed from the y, x, and z axes (b,c, and d, respectively). Significant bending is observed at the ends of the origami which likely displaces the capture strands from the idealized “row” anticipated in the static structure.
Figure S13. Representative (a) 2% uranyl acetate stained TEM and (b) AFM images showing the degree of flexibility of the DNA origami, in agreement with CanDo modeling agreements. Red arrows indicate high deformed structures while white arrows show flat origami. Deposition conditions were in 12 and 100 mM Mg$^{2+}$ concentration for TEM and AFM, respectively.
Appendix S1. Enlarged Origami Schematics

Origami Left
Capture Zone A
| 1   | GACAGATGAACCTTCATCAAGA                        |
| 2   | GTAATCTTGACTCAACTTTAAT                       |
| 3   | CATTGTGAATTAAACCCTCTGTT                      |
| 4   | TACCAGACGACCCTTTAATTG                       |
| 5   | TCTTTTGATAAAATGCCTGAGT                      |
| 6   | AATGTGTAAGTATCGCACTCCA                      |
| 7   | GCCAGCCTTTCACCCTGTATCA                      |
| 8   | TAAAGGGAGCCGCCGATTTAGAGCTTTGCGTCA          |
| 9   | AAGGCGGAAAGGCACCGCTTC                      |
| 10  | CAGGCGAACCAGACTGACCAACTTTGAAAGAG           |
| 11  | ATTCAGCATAAGCTGCTGACGTTAGTACAGAC           |
| 12  | TTTATTTGAGATGTTTATAAAATGCAAGACCGGT         |
| 13  | AAAATAAGAGCAACACTATCAACCTTTAGCGA          |
| 14  | TGGTGCTCAGGATAGAGTATAGATAAAAACCA           |
| 15  | AGGGCTCATAATTATTTATGCAAGAGGTCATTT         |
| 16  | TGGTGTAATCGGCTCAGGAAGAAAGAGATTCAA         |
| 17  | AAGGCGGATCTCAGGAGGTGTTGGAATAAGGTGTA       |
| 18  | TCACCTACCCAATCAACGTGAGTATGAATT            |
| 19  | GGGCAGAACCTGCTCATATTATATTACGGAGCA         |
| 20  | TAGTGCGAGAGGCTTTGCAAGCACAACCTCAA          |
| 21  | CAGGGATGGCTCAGGCTTAAATTTTTAGAA           |
| 22  | CCCTGAGAAAGGCACGAGACGGAGGACGGAC           |
| 23  | ACACCCGGAACCGCACAAGAGACGTGGACT            |
| 24  | CCAAAACGGGAAAGCCGCCGAGCGTTGGCAGAAAGAGAGCTCCACTA |
| 25  | GCCAGTACCCGCCACCCCTCACCGGAACGAGGCGACGAGCTCAATCAT |
| 26  | GAACACCAAAAGCTGCTCATTTAGTATAAGTATA       |
| 27  | AAGGACCAGTCAGGAGCTGCGAGAAACACCA          |
| 28  | CGGAAAAGAAGTTTTTGCCAGACATAACGCCAA        |
| 29  | ATAAATTGCTGAAATATAATGAGCGAACAGAC         |
| 30  | GTTTAGTCAAATACCCATACCTTTACACCAAGG         |
| 31  | TTAAGCGCCATTCCGCCATTCTGCTCATCTGCA        |
| 32  | CCTAGGAACCAGACCCCTCAGTGCGCTGAGA          |
| 33  | GGGTTCAGTGAATAAGAAGACCCGCCAGCTTGC       |
| 34  | CCTGAGGAAAGAAAATCTACGGTCACAACTAATG       |
| 35  | CAGATAGGGGTATAGTAAATTAATTACGAGC         |
| 36  | TTCAACTGTAGCTCAACATGTCCGGGAGAAGCC      |
| 37  | TTATATATGATATTTAACCCGTCGGCGCATCG        |
| 38  | TAACCCAGGCTGCACACTGTTTTGTCTCCAGTTT     |
| 39  | GGAACCAAGGGAAAGAGACCAAGAGGGCGGCTAGGAGATAGGCTTGG |
| 40  | ATAAAGAACCACCCACCCTCACGATCAGAAATCCCGCACCTGCTCCATGTTTA |
| 41  | ACCAGAAAAGGGCGACATTCAGTACCCAGCGG    |
CCACATTTAATAAAAAACGAACACTATCATATGTGTT
GCGTTATGTGTAGACTGGATAATTTAGGAATA
TTTTGTNTAAATATGCAAACACTACTTTCAAAATTC
GTAGATTCTAGCTGAAAAATCCCTCTGTAATAC
AGTGTGGGAAAGGCGATCGGTGTCACGTTGTT
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GCTCAACCGATTGAGGGAGGACAAATCAGAAG
AAATACCGGAACAACATTATTAGATTCTAC
TTGAGGGCTTCAAATACGCCCAGGAGGAGCCCG
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GAAGGACCCTGCAACAGTGCCAGGTCAGTATTA
ACACCTTCTCTGTGAAATTTGCGCGGCGGGGAG
AGGCGAAGAGTGTTTTTATAAATGACTGAGGCCACCGATAATTGAATCG
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AACAACACGTTAAAACACAGAAATGAAATAACCTTG
GAGGCCGCTGAGCCAGCGTCAGTATTTTCGCG
GCCAATTATCCGCTCACAATTAAAACAGAGGT
AGCATAGAAAGGAACAAACTAAAAAGCCAGAATG
GAAGGTTCATCGGCTAATTTCGAGAATGATCTCT
TACCGGCCAGTTACAAAATAAGACGAGCGACA
ATAAAAGAAAATCCAGACCGTGTATCAATATATG
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CATTACAATAATGAAAATCTAAACCACCAGCAG
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CTGCAGTAAAAGAGTTGCTCATTACGCAAATTAACCTCCAGTGCG
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TTGGCAATTTTTTCAGGTGACTTGATTTCA
CAAACCTAGCTTTGCCATCTCTCAATAAATAAG
AGCAATTCCCAAATCAAGAATGACCAGACA
AAAGGAATAAGATAAAACACCGAAATTACCTTT
TTAACAGATGAAATATACAGAAATTTCTTTTGC
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CATTTCGAATCATATAACTAGTAAATAAGAGA
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ATAGCCCCTCAATATACGGGTGCAAACTC
GCGTTAATGAGTGACGTTAATCGCGGAACCTG
GCTGGGCTCCAAAAAGGAGCCACAGGAGGTG
DNA Origami Capture Strands

201 TTGTCAAGGTAATATTTGACCGGAGAAGGATTA TTTT TGATATAGGTGAGTA
202 TAGAAAACAGGGGACATTACCATACGATTATTTATT TTTTT TGATATAGGTGAGTA
203 ATAAAAATCGTCATAAATATTTTCATCGTAG TTTTT TGATATAGGTGAGTA
204 GTTGTAGTTTCATTCCATATGACATCAAAGG TTTTT TGATATAGGTGAGTA
205 GGAATTGCAATTTCATATAAGCTA4
Supporting References