Nanoengineering a single-molecule mechanical switch using DNA self-assembly

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Abstract

The ability to manipulate and observe single biological molecules has led to both fundamental scientific discoveries and new methods in nanoscale engineering. A common challenge in many single-molecule experiments is reliably linking molecules to surfaces, and identifying their interactions. We have met this challenge by nanoengineering a novel DNA-based linker that behaves as a force-activated switch, providing a molecular signature that can eliminate errant data arising from non-specific and multiple interactions. By integrating a receptor and ligand into a single piece of DNA using DNA self-assembly, a single tether can be positively identified by force–extension behavior, and receptor–ligand unbinding easily identified by a sudden increase in tether length. Additionally, under proper conditions the exact same pair of molecules can be repeatedly bound and unbound. Our approach is simple, versatile and modular, and can be easily implemented using standard commercial reagents and laboratory equipment. In addition to improving the reliability and accuracy of force measurements, this single-molecule mechanical switch paves the way for high-throughput serial measurements, single-molecule on-rate studies, and investigations of population heterogeneity.

Online supplementary data available from stacks.iop.org/Nano/22/494005/mmedia

(Some figures in this article are in colour only in the electronic version)

1. Introduction

The ability to precisely manipulate individual molecules has led to stunning new discoveries in physics, biology, and medicine [1, 2], as well as powerful new methods in nanoscale engineering. For example, single-molecule force measurements have revealed the basic mechanical properties of nucleic acids [3], the dynamics and functioning of molecular motors [4, 5], and the role of hydrodynamic forces in the circulatory system in regulating enzymatic activity [6]. In addition, these measurements have yielded fundamental insights into the dynamical strength of molecular interactions [7], which have led to the development of creative new tools for nanoscale assembly [8].

Mechanical forces can be applied to individual molecules using a broad range of tools, including optical traps, magnetic tweezers, mechanical cantilevers, and the centrifuge force microscope [9, 10] (figure 1). Yet a common requirement of these methods is that single-molecule constructs must be specifically tethered between two surfaces (e.g. beads, cover slips or cantilevers) to enable their manipulation and detection. This leads to one of the major challenges in single-molecule experimentation—verifying that exactly one molecular tether is being pulled, and distinguishing this tether from non-specific...
Figure 1. Single-molecule manipulation techniques include those of (a) optical tweezers, (b) magnetic tweezers, (c) the atomic force microscope, and (d) the centrifuge force microscope.

Figure 2. Single-molecule linking geometries: a cartoon showing receptor–ligand unbinding using a variety of linking strategies including (a) direct contact/short linker, (b) a long linker, and (c) a looped linker.

and unintended interactions that may occur (e.g. surface–surface interactions, formation of multiple bonds). The success and reliability of single-molecule experiments depends upon the creation of reliable, verifiable and robust linking techniques. This is particularly important for bond rupture studies (e.g. characterizing the strength of molecular adhesion bonds [11], DNA base pairing [12], and cell adhesion and signaling [13]), as the dissociation between two molecules can be difficult to positively identify due to the lack of an obvious mechanical signature.

We have met this experimental challenge with a new single-molecule attachment technique that facilitates reliable and accurate single-molecule force measurements. Using DNA self-assembly techniques, we have nanoengineered a unique linker that behaves as a force-activated single-molecule switch. This switch changes conformation under force to signify bond rupture, providing an identifiable molecular signature that eliminates the possibility of accidentally measuring non-specific, multiple and unknown interactions. Furthermore, this construct enables the same pair of interacting molecules to be brought back together following rupture, opening the way to high-throughput serial measurements, single-molecule on-rate studies, and studies of population heterogeneity. Our approach is simple, versatile and modular, and can be easily implemented using standard commercial reagents and laboratory equipment. We elaborate on this approach in section 1.1, and provide an overview of standard linker geometries for single-molecule manipulation experiments.

1.1. Overview of linker geometries in single-molecule experiments

A variety of different linker strategies have been employed in single-molecule experiments. These include different chemical attachment strategies (e.g. SMCC, click chemistry, EDC, etc), surface preparations (e.g. silanization methods), and linkage materials (e.g. DNA, PEG) [14–18]. Most of these approaches can be classified into a few basic categories based on their geometry (figure 2). In this subsection, we review these categories, and show how appropriately designed linkages can improve the efficiency and reliability of single-molecule rupture measurements.

(a) Direct contact/short linkers. First, we consider the simplest strategy of direct molecular coating onto a surface (figure 2(a)), which is typically accomplished via adsorption or conjugate chemistry with short linkers (e.g. SMCC, EDAC). This approach has been commonly used, particularly in early single-molecule experiments [19, 9]. Some advantages of this approach are that it is typically simple, fast and inexpensive, particularly for the case of physical adsorption of molecules and cells to a surface. However, the first major problem that arises from this geometry is the introduction of unwanted interactions with the surfaces. Since the sample molecules are typically located within nanometers of the surfaces, non-specific surface–surface and molecule–surface interactions (e.g. van der Waals and depletion forces) may significantly affect the measured force [20]. Furthermore, any impurities or contamination in the sample may result in unwanted and unidentifiable adhesion between the surfaces. To deal with these difficulties, careful characterization and minimization of
non-specific interactions are necessary. Reduction of non-specific adhesion can be accomplished by blocking agents such as BSA, casein, Pluronics F127, and Tween, but they are often difficult to eliminate completely. For bond rupture measurements, the second difficulty with this geometry is the lack of a specific mechanical signature for the molecule of interest. Thus, detachment events arising from both multiple attachments and non-specific attachments are difficult to distinguish from the single-molecule interaction of interest. Minimization of multiple attachments can be accomplished by diluting the active molecules on the surfaces, and by controlling the distance, touch time and touch force between the functionalized surfaces during the experiment. While careful control experiments and the use of statistical techniques to estimate the number of single bonds can help to mitigate difficulties arising from direct contact geometry [21], we will see that the use of longer linkers can improve the reliability, accuracy, and ease of analysis of the resulting single-molecule data.

(b) Long linkers. A variety of linking strategies have been developed for keeping the molecules of interest far from the surfaces to which they are attached (figure 2(b)). This can be accomplished using long polymer linkers such as PEG [17] or DNA [16], or with more exotic materials such as the M13 filamentous bacteriophage [22]. Long linkers can serve the double purpose of both eliminating close range non-specific interactions and facilitating the positive identification of each tether as a single molecule; the required linker length depends upon the resolution of the force-probe instrument and the compliance of the molecule of interest. When used on one side, a sufficiently long linker can already eliminate surface–surface non-specific interaction, as well as allowing positive identification of a single interaction by using the known force–extension behavior of the tether. When used on both sides as depicted in figure 2(b), the linkers can additionally eliminate non-specific surface–molecule interactions. We note that long linkers between the force probe and the molecule of interest may complicate the force–loading history and effective energy landscape, but this can generally be accounted for [23, 15].

(c) Looped linkers. More recently, single-molecule experiments have been performed using a looped linker geometry, in which the interacting molecules/sites of interest are connected to each other by a molecular tether [24, 25]. This looped linker geometry as shown in figure 2(c) has two primary advantages. First, it provides an additional signature for bond rupture events, as this molecular transition will create a well-defined increase in the tether length. Unlike for the non-looped geometries, unbinding of the receptor–ligand pair cannot be confused with accidental unbinding from the anchoring ends. Second, the looped linker allows for repeated testing of the exact same pair of molecules, provided the conditions for reforming the bond are favorable. This opens up new possibilities for studying population heterogeneity in a population of molecules—a highly useful, but difficult to realize, benefit of single-molecule experiments. This looped linker concept was recently introduced as ReaLiSM (receptor and ligand in a single molecule) [24], where the A1 domain of the von Willebrand factor was linked by an amino acid chain to platelet GPIbα, and the two interacting protein domains were repeatedly bound and unbound. While highly effective, the ReaLiSM construct as presented may be difficult for many labs to create, requiring expertise in protein engineering and purification.

Here we present a versatile DNA-based alternative constructed using DNA origami methods [26, 27]. By mixing a long piece of single-stranded DNA with a carefully designed soup of short DNA oligomers, we have constructed looped single-molecule linkers with an integrated receptor–ligand pair via DNA self-assembly (figure 3).

2. Materials and methods

2.1. Linker design and construction

Two different kinds of linkers were designed and assembled based on techniques outlined from previous DNA origami work [26, 27]. Both linkers incorporate functional ‘sticky’ ends (we used double biotin on both ends) which act as anchors for single-molecule experiments, as well as two functional sites near the middle of the linker to form the loop. One set of constructs forms the loop by hybridizing a single ‘bridge’ oligo across two distinct locations, while the other set used two separate oligos each functionalized with digoxigenin or anti-digoxigenin, which can bind to each other as a receptor–ligand pair to form the loop.

To make these linkers, M13mp18 single-stranded DNA (New England Biolabs) was first linearized by hybridizing a 40-nucleotide oligo to form a double-stranded region and then cleaving this region with BtsCI restriction enzyme (New
The linearized single-stranded DNA was then mixed with complimentary oligos (Bioneer, Inc.) and subjected to a temperature ramp from 90 to 20°C with a 1°C min\(^{-1}\) ramp in a PCR machine (Bio-Rad) to allow the oligos to anneal properly. For the linker with the bridge oligo formed loop, 121 oligos excluding the bridge oligo were added in tenfold molar excess, with the bridge oligo added in equimolar concentration to the scaffold strand. For the receptor–ligand loop construct, 120 oligos excluding the antibody oligo were in tenfold molar excess, which was added in equimolar concentration and subjected to a temperature ramp from 40 to 10°C with a 0.5°C min\(^{-1}\) ramp after the other 120 were linked.

Detailed protocols and the full sequences of all oligos are available in the supplemental information (available at stacks.iop.org/Nano/22/494005/mmedia). All oligos were purchased from Bioneer, Inc., with the exception of the 5′ double-biotin oligo (Integrated DNA Technologies), the digoxigenin oligos (Integrated DNA Technologies), and a few plain oligos ordered with next-day service (Invitrogen).

### 2.2. DNA–protein conjugation

A 3′ thiol modified oligo was reduced and linked to monoclonal and polyclonal anti-digoxigenin (Roche Applied Science) using sulfo-SMCC (Pierce) and the accompanying protocol. The NHS group on the SMCC was first linked to free amines on the antibody (at 1 mg ml\(^{-1}\)) with a 30 min reaction at room temperature using 20-fold molar excess of SMCC in PBS at pH 7.4. At the same time, the thiol oligo was deprotected and reduced by incubating in 50 mM TCEP (Pierce) for 20 min and then cleaned using a PCR clean up kit (Qiagen). Following the first SMCC reaction, excess SMCC was removed with a Zeba desalting column (Pierce), pre-equilibrated with PBS buffer. The activated protein was then mixed with the reduced thiol oligo in a 1:1 molar ratio for 30 min at RT.

Conjugation was verified by visualization on a 4–20% polyacrylamide gel (Bio-Rad) run in 1× TBE buffer at 150 V for 40 min, where a shift from the protein linkage was readily apparent (see supplemental figure 3 available at stacks.iop.org/Nano/22/494005/mmedia). Typically, 5–50% of the oligos were conjugated to protein, and purification of the protein–DNA conjugate was accomplished by excising the gel band and using an electro-elution kit with the accompanying protocol (Gerard biotech).

### 2.3. Single-molecule force spectroscopy

The final unpurified linkers with double-biotin ends were incubated with streptavidin polystyrene beads (Corpuscular) for 15 min, and then injected into a chamber with PBS buffer for use in the optical trap. The optical trap setup consists of a single stationary trap and a piezo-controlled micropipette integrated into an inverted light microscope (Nikon). The setup is functionally identical to previously described instruments [6, 28], but with 160× overall magnification instead of 400×. High-speed video microscopy is used to measure bead positions in 1D with a resolution of ~4 nm at ~2 kHz. The optical trap is calibrated using the blur-corrected power spectrum fit [29], with additional calibration information provided by the dsDNA overstretching transition [30].

Single-molecule force measurements are performed by bringing linker functionalized beads held in the optical trap into contact with streptavidin-coated beads held in the micropipette to form molecular tethers. Tension in each tether is applied by moving the bead in the micropipette, and quantified by measuring the displacement of the bead in the optical trap. The observed distance between the beads gives a measure of the tether length.

### 3. Results and discussion

We successfully created looped single-molecule linkers via DNA self-assembly. Two different kinds of linker constructs were generated and tested: (i) linkers looped by a short complementary strand of DNA to study the kinetics of DNA base pairing, and (ii) linkers looped by a receptor–ligand pair to study protein–protein interactions. As detailed below, the proper assembly and functionality of these linkers was verified using gel-shift assays and optical trap measurements. We demonstrated their effectiveness for single-molecule force spectroscopy by measuring the kinetics of bond rupture for both DNA hybridization and an antibody–antigen interactions, and by showing how the molecular signature of a looped tether can be used to improve the accuracy of the data.

#### 3.1. Verification of the linker assembly

We first tested the linkers looped by a single DNA oligo bridge, as they served as a good model system for testing and optimizing linker assembly, independent of protein-coupling efficiency. For these oligo bridge constructs, we made two different loop lengths: 2580 base pairs and 600 base pairs. Additionally, we varied the length of the bridge oligo on one side to be 30 bp, 20 bp, 15 bp, and 10 bp, while the other side was maintained at 30 bp. The formations of both long and short loops were easily distinguishable from those of unlooped products by a gel shift due to slower migration on a 0.7% agarose gel (figure 4), for both the 30 bp and 20 bp bridge constructs. We were unable to observe any looped construct in the gel when using the 15 bp or 10 bp bridge oligo, presumably due to the harsh conditions of electrophoresis (e.g. low salt, high temperature, high voltage). Confirmation that the shifted gel band was indeed the looped construct was accomplished by cutting the construct with a single-cut enzyme in the loop region (figure 4)—the shifted band (looped DNA) was largely unaffected by the enzyme, while the lower band (unlooped DNA) was completely digested into two separate pieces.

Next, these products (with double-biotin ends) were verified directly in the optical trap by pulling them end to end with linear ramps of force (figure 4). Unlooped linkers show characteristic DNA force–extension behavior with typical contour lengths of 2000–2300 nm, consistent with the number of DNA bases within the construct. The looped linkers initially start with a shorter contour length, then exhibit a sudden increase to this full contour length when the DNA...
bridge ruptures under the application of mechanical stress. We measured an average increase in contour length of 884 nm and 208 nm for the long and short loops, respectively (figure 4 inset), which is within a few nanometers of the expected length changes of 877 and 204 nm predicted from the worm-like chain polymer model using a contour length of 0.34 nm per base pair [31, 3]. As can be seen in figure 4, after bridge rupture both curves roughly follow the curve for the unlooped linker. As the linker was stressed above 65 pN, the DNA overstretching transition could be observed, which served as an additional mechanical signature for identifying single-molecule tethers. However, pulling the molecule through this transition always resulted in detachment of the linker from the functionalized beads, presumably due to the force-induced melting of the biotinylated anchor oligos off of the ssDNA scaffold. While this effectively limits the use of this linker to measurements below about 65 pN, this could likely be overcome by covalently cross-linking the DNA linker or by using much longer anchoring oligos. Regarding the observed length of the linkers, we note that a distribution of lengths is expected from multiple-tether measurements, even if every tether is identical on a molecular level. Since the bead in the pipette is rotationally constrained, tethers may be held at different angles, causing the measured distance between the beads to differ from the molecular tether length.

Receptor–ligand looped linkers were also created in order to measure the force-dependent kinetics of an antibody–antigen interaction. Oligos coupled to digoxigenin and to its antibody were assembled to form linkers with a loop length of 600 base pairs. The verification of these constructs was conducted in the same way as for the DNA bridge looped linkers, using both gel electrophoresis shift assays and single-molecule pulling experiments. We note that while the polyclonal looped constructs could be readily seen in a gel as a distinct shifted band (identical to the DNA bridge looped construct in figure 4), the monoclonal constructs could not. This is likely due to the much lower affinity between digoxigenin and its monoclonal antibody, and is consistent with the lack of a band for the 10 bp and 15 bp DNA bridge constructs. However, both the monoclonal and polyclonal constructs could be observed in the optical trap, and exhibited force–extension curves that matched those of the 600 bp oligo bridge construct.

3.2. Demonstration of single-molecule force spectroscopy

We tested the dynamic strength of DNA hybridization with the optical trap by repeatedly applying linear force ramps to the DNA bridge constructs to determine the distribution of the rupture force. The molecular signature of the looped linker serves as a powerful filtering method to distinguish the rupture of the DNA bridge from non-specific, unknown and multiple interactions. This is illustrated in figure 5 (left) for the rupture force of the 20 bp bridge, where positive identification of the correct rupture transition (using the change in tether length, overall tether length, and overstretching of the linker) enabled the removal of erroneous data that accounted for 57% (34/60) of the measured events. In the resulting data, we measured a mean rupture force of 52 pN with a standard deviation of 6 pN at a nominal loading rate of 100 pN nm$^{-1}$ (this was a combination of experiments with a mean loading rate of 98 pN nm$^{-1}$ and a standard deviation of 35 pN nm$^{-1}$). This agrees within error with the expected force of 39 ± 15 pN for the mechanical shearing of DNA (based upon their empirical formula) [12].

When testing the other DNA bridge lengths, we observed fewer rupture events for the 30 bp bridge, as the biotin–streptavidin bonds anchoring the linker would often rupture first. In addition, we also saw evidence of the 15 bp and 10 bp bridges in single-molecule pulling experiments, despite not seeing these constructs with the gel-shift assay. While the formation of these loops should be energetically favorable even with the additional entropic cost of closing the loop [32], it is possible that the conditions of electrophoresis lower the stability of these constructs leading to their absence in the gel assays.

As another demonstration, we measured the force-dependent unbinding kinetics of digoxigenin with its antibody in the optical trap (figure 5 (right)). By making repeated measurements of bond rupture under a constant force, we found a characteristic lifetime of 1.3 s (with a 95% confidence band of 0.9–2.0 s) at a force of 49 ± 2 pN for the polyclonal antibody, using maximum likelihood estimation with an exponential decay model. We found this interaction to be relatively strong, in agreement with other single-molecule measurements that used it as a molecular anchor [22].
Figure 5. Single-molecule force spectroscopy results for the rupture of (left) DNA hybridization and (right) antibody–antigen interactions. Left: rupture force histogram for shearing a 20 bp DNA segment, demonstrating the filtering of erroneous data via the looped linker molecular signature. Right: survival trajectory for digoxigenin against its antibody under constant force, with results of maximum likelihood estimation superimposed.

Figure 6. Trajectory demonstrating repeated rupture and formation of a single receptor–ligand pair (digoxigenin with its monoclonal antibody): (left) force versus time and (right) force versus extension traces for repeated cycles of force application and release. Bond rupture events are observable by a sudden drop in force and an increase in tether length, as demarcated by red arrows. Rebinding/bond formation during a low force clamp can be observed by subsequent bond rupture under the application of force.

Note that without the looped linker the high bond strength of this interaction can make rupture measurements difficult, as it can be difficult to distinguish the rupture of digoxigenin–antibody from the failure of molecular anchors in the absence of an additional molecular signature.

Not only were we able to measure bond rupture, but single-molecule bond formation could also be observed. In many cases with the digoxigenin–antibody construct, we were able to reform the complex after dissociation by bringing the beads closer together and waiting for a short time (figure 6). However, we were not able to reform the oligo bridge after rupture under similar conditions, suggesting that the formation of secondary structure or the extra time to diffusively align the two strands slowed the rebinding kinetics.

4. Conclusion

We have presented a simple and effective method for making functional looped linkers using DNA self-assembly, which can increase the accuracy and reliability of single-molecule force measurements. The method is versatile enough to be useful for a wide range of molecular interactions, and simple enough to be made by researchers of diverse backgrounds without significant investment of time or money (see the appendix). We demonstrated this functionality by constructing and testing two different looped linkers, designed for studying the dynamic strength of DNA base pairing and receptor–ligand interactions. In addition, we showed how the molecular signature provided by this ‘DNA mechanical switch’ enables the removal of erroneous data that can arise from non-specific, unknown, and multiple interactions. Not only is this construct useful for traditional bond rupture measurements and force spectroscopy, but it also enables the same pair of interacting molecules to be brought back together following rupture, opening the way toward high-throughput serial measurements, single-molecule on-rate studies, and studies of population heterogeneity.
The goal of this work was to develop a looped single-molecule linker, with an easily identifiable molecular ‘signature’, that is both versatile and easy to implement. While the DNA origami construction of the linker may sound intimidating due to the sheer quantity of oligos used, it is rather simple in practice. We recommend dividing the ssDNA scaffold into fixed and variable regions. The variable regions are those that may be changed in the future to have different labels, while the fixed regions will only have unlabelled oligos. Then all of the ‘fixed’ oligos can be ordered pre-mixed to save time and eliminate the possibility of mixing error (Bioneer was accommodating in this regard). In our case, we designated 12 variable oligos each spaced apart by 9–10 fixed oligos, and ordered the 109 fixed oligos pre-mixed in a single tube. The variable oligos gave us the flexibility to choose different oligo labels and to choose different loop lengths for our construct, without significant protocol modification.

The choice of oligo length should also be considered. We chose 60 nucleotide oligos to minimize the number of oligos needed (companies vary in maximum oligo size), but in hindsight the pre-mixing of the fixed oligos makes the number of oligos less important. The main differences to consider with shorter oligos will be differences in annealing behavior, requisite protocol modifications, and price. Regarding annealing, longer oligos have higher melting temperatures, which should increase the annealing speed in a decreasing temperature ramp. On the other hand, longer oligos tend to have more secondary structure which can sometimes impede proper annealing, especially at lower temperatures. Regarding protocol modification, oligos that are 30 bp or shorter can be washed out with PCR clean up kits, which would necessitate the use of a different cleaning method in our SMCC conjugation. Regarding price, the yield of shorter oligos can be significantly higher, making them more cost effective.

It is worth considering cost in more detail, as it could be a deciding factor in the use of these linkers. The cost of oligos continues to decrease, and many companies offer discounts for large oligo orders or those ordered in plates instead of individual tubes. As an example, our construct cost under $1000 with unmodified oligos. Modified oligos will increase the cost depending on the modification, from tens to hundreds of dollars per oligo. However, this initial investment can produce several liters of product at the nanomolar concentrations that are typically required for single-molecule experiments. Surprisingly, the most expensive ingredient on a molar basis when using short unpurified and unmodified oligos is the M13 scaffold [26].

To generate linkers of different lengths, alternative ssDNA scaffolds could be used. For example, ssDNA can be generated using asymmetric PCR, or obtained by strand-separating standard PCR products. Product from rolling-circle amplification could also be used for the generation of periodic linkers. An alternative method for constructing looped linkers that we have been exploring is ligating together the products of multiple PCR reactions.

Lastly, we will discuss the time, equipment, and expertise requirements for making these linkers, as these were primary considerations for us in starting this project. By following the protocols in the methods section (described in detail in the supplemental information available at stacks.iop.org/Nano/22/494005/mmedia), the linkers can be completely constructed and verified in a single working day, and this time could probably be reduced with further optimization. The protein conjugation procedure takes about 2–3 h, with an additional 2 h for purification. The oligo annealing process (excluding protein-conjugated oligos) takes 70 min, and can be done in parallel with the protein conjugation step. Annealing the protein-conjugated oligo(s) takes another 1 h, and final verification of the looped linker on a gel takes an additional 1–2 h. Overall, it can be done in 6–8 h from start to finish. The equipment requirements are minimal, and the protocols can be carried out with only modest pipetting skills. The only major equipment needs are for a centrifuge, and horizontal and vertical electrophoresis systems (the vertical one was used for protein–oligo purification, but we have seen gel shifts in agarose as well). We used a PCR machine for the annealing protocol, but this is not strictly necessary and could be replaced with a hot water bath cooling slowly to room temperature. All reagents and kits used are commercially available.

These linkers are intended to be versatile for studying a variety of molecules, and in fact much of the protocols will remain the same even as specific molecules on the linker are changed. The modularity of the linker makes it simple to use different proteins by just swapping one functionalized oligo for another. If the linker is constructed with only the protein-conjugated oligos omitted, this stock solution of partially constructed linker can then be mixed with the specific protein oligos to get the final construct. The only additional step required will be to repeat the chemical conjugation of oligo to protein for each specific case. In our example linkers, we used SMCC, but there are a variety of other bioconjugation techniques that could be used instead. In the case of antibodies, it is also possible to use antibody binding proteins such as protein G to simplify the process of swapping in different antibodies.

References


Nanoengineering a single-molecule mechanical switch using DNA self-assembly: Supplemental

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This supplemental section provides detailed information for the construction of looped single-molecule linkers using DNA self-assembly. Following a description of the DNA oligos that we used to construct this linker, a protocol for linker construction is presented.

1. Oligo details

The full sequence of all the oligos used are shown in Figure 1 and are based on the M13 sequence given by New England Biolabs and used for previous DNA origami work \([1]\). The oligos are stored at 100 \(\mu\)M at \(-20^\circ\)C, and all mixing of oligos is done at these stock concentrations unless noted. We will refer to the numbering throughout this section as we explain the construction of the various linkers.

2. Detailed protocol for looped linker construction

2.1. \textit{Step 1: ssDNA linearization}

(i) Mix the following in a clean PCR tube:

- 5 \(\mu\)L M13mp18 ssDNA (NEB product N4040S 0.25mg/mL or 100nM)
- 2.5 \(\mu\)L 10x buffer 4 (NEB)
- 0.5 \(\mu\)L 100 \(\mu\)M cut-site oligo (oligo 000A)
- 16.5 \(\mu\)L water

(ii) Briefly bring to 95\(^\circ\)C (30 seconds), lower to 50\(^\circ\)C, and add 1\(\mu\)L BtsCI enzyme.

(iii) Incubate for 1 hour at 50\(^\circ\)C.

(iv) Heat deactivate the BtsCI enzyme by incubating at 95\(^\circ\)C for 1 minute.

Note: To assay the linearization efficiency, add 1.21 \(\mu\)L of a mixture of all numbered oligos to 5 \(\mu\)L of linearized ssDNA and construct the dsDNA piece by heating to 90\(^\circ\)C.
Figure 1. Sequences of all oligos used. The numbered oligos cover the entire sequence of the M13 scaffold strand, and the lettered strands are functionalized or special purpose oligos. The bridge oligos are labeled red for regions that bind M13 bases 1950-1980, green for regions that bind starting at M13 base 2580, and blue for regions that bind starting at M13 base 4560.
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Figure 2. Looped linker construction using DNA origami: Circular single-stranded DNA is enzymatically cleaved at a single site and mixed with over 100 oligonucleotides to self assemble into a looped linker held together by a single “bridge” or “staple” oligo.

and cooling to 20 °C at 1 °C/minute or slower. The product can either be run on a 0.7% agarose gel to separate circular from linear, or the strand can be cut with a single cut enzyme (we added 1 µ AfeI) to make sure most or all of the ssDNA was linearized.

2.2. Step 2: Conjugating protein to oligo

The protocol we used for conjugating protein to oligo was loosely based on the Pierce protocol provided with the sulfo-SMCC.

(i) **Deprotect the SH oligos** by mixing the following in a clean PCR tube

- 5µL 0.5M TCEP (Pierce)
- 40µL water
- 5µL oligo 033A at 100 µM

Let the mixture sit for at least 30 minutes at room temperature

(ii) **Make 10mM Sulfo-SMCC solution**

- add 20µL DMSO to 2 mg Sulfo-SMCC and pipette up and down to mix well
- dilute into 450µL of PBS
- Use immediately

(iii) **Activate the protein**

- suspend protein in PBS at 1mg/ml concentration (use desalting column or centrifuge filter if necessary to concentrate or change buffer)
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- use 20x molar excess of Sulfo-SMCC for 1mg/ml protein (e.g. 1mg/ml Antibody - 20x: 20μL protein @ 6.7μM + 0.27μL Sulfo-SMCC @ 10mM).
- React for 30 minutes at room temperature.

(iv) **Wash TCEP from oligos** just before Sulfo-SMCC reaction is finished
- use Qiagen PCR clean up kit, following the protocol except eluting into 50μL PBS for 10μM final concentration (note: using a Tris buffer will interfere due to the presence of amines)

(v) **Wash the Sulfo-SMCC from the activated protein**
- pre-equilibrate a Zeba desalt column (Pierce) with PBS
- do one or two passes of the protein through the column to eliminate Sulfo-SMCC
- resuspend at 5μM concentration

(vi) **Mix the activated protein with the deprotected oligos**
- Use 1:1 molar ratio or excess of protein (e.g. 1:1 - 2μL oligo @ 10μM + 4μL antibody @ 5μM)
- React for 30 minutes at room temperature

2.3. **Step 3: Purifying protein conjugated oligo**

Once the protein has been conjugated to the oligos, it may be necessary to purify the conjugated oligo from unreacted byproducts depending on the yield of the reaction. We were typically able to get 5-50% yields (see Figure 2), and even with a 50% yield we had trouble getting the loop to form with the digoxigenin antibody complex without purification. This is probably due to unconjugated oligos competing with the conjugated ones and due to excess protein reacting with dig-labeled oligos on the DNA construct.

To purify, we ran the same product in a 4-20% polyacrylamide gel (Bio-rad) enough to separate the conjugated and unconjugated oligos (see Figure S2), typically 150V for 40 minutes in 1x TBE buffer. Then we stained the gel for 10-15 minutes in a 1x solution of Sybr Gold (Invitrogen) and used a razor blade to cut out the relevant band(s). Once the gel slices were cut, we used an electroelution kit and supplied protocol to extract the conjugated oligos from the gel. Briefly, we put the slices in a midi sized electroelution tube (Gerard biotech) with 600μL of buffer, and ran them in a horizontal electrophoresis at 150V for an additional hour. We estimated the final concentration based on the known amount of oligo put into each gel lane, the conjugation yield, and the amount of dilution in the electroelution step.

2.4. **Step 4: Assembly of DNA linkers**

We have made several different linkers over the course of this study. All of these utilize the same basic protocol which was adapted from other DNA origami work [1] without further optimization. This involves mixing the oligos with the M13 ssDNA (typically in
a 10x molar excess) and subjecting the mixture to a temperature ramp from 90°C to 20°C at 1°C/minute. This protocol may be slower than necessary, as it was based on folding complex 2D shapes rather than simply making a linear piece of dsDNA as we are doing here. We used a thermal cycler to apply the temperature ramp, but heating a water bath and letting it cool to room temperature over the same time should provide the similar results.

The following mixtures of oligos will be used in the construction of various linkers:

- **plain linear**: mix of all numbered oligos
- **linear with double biotin ends**: mix of all numbered oligos except substitute 001A for 001 and 121A for 121
- **short loop with double biotin ends**: same mixture as linear with double biotin ends, but do not include 033 or 044 (or use truncated versions of 033 and 044 to ensure all bases of M13 are paired).
long loop with double biotin ends: same mixture as linear with double biotin ends, but do not include 033 or 077 (or use truncated versions of 033 and 077 to ensure all bases of M13 are paired).

To make the desired construct, mix the following in a clean PCR tube and apply the temperature ramp to anneal oligos:

- 5µL of linearized ssDNA from section 2.1
- 1.2µL of one of the above mixtures for 10:1 oligo:DNA ratio
- For looped constructs, additionally add:
  (i) 1µL of 1000x dilute digoxigenin oligo 044A for dig-antibody construct
  (ii) 1µL 1000x dilute bridge oligo 033-044B for short loop with 50bp oligo bridge
  (iii) 1µL 1000x dilute bridge oligo 033-077B for long loop with 50bp oligo bridge

The temperature ramp completes the protocol, except for the dig-antibody construct. For the dig-antibody looped construct, add an additional 1µL of purified antibody conjugated oligo at an approximate concentration of 100nM once the first temperature ramp is complete. Note that it if a larger volume of lower concentration oligo is used, it is important to maintain the proper buffer conditions for hybridization. Additional concentrated buffer may need to be added in this case. To anneal this oligo, we used a temperature ramp from 40°C to 10°C at 0.5°C/minute, though we did not exhaustively test simpler or shorter protocols. For the looped constructs, we typically got a 50% looping yield following this protocol. This looped product can be excised and purified from a gel as we did with the antibody conjugated oligo, but we did not find this purification step to be necessary for this study.

References