A DNA Origami-Based Chiral Plasmonic Sensing Device

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ABSTRACT: Accurate and reliable biosensing is crucial for environmental monitoring, food safety, and diagnostics. Spatially reconfigurable DNA origami nanostructures are excellent candidates for the generation of custom sensing probes. Here we present a nanoscale biosensing device that combines the accuracy and precision of the DNA origami nanofabrication technique, unique optical responses of chiral plasmonic assemblies, and high affinity and selectivity of aptamers. This combination enables selective and sensitive detection of targets even in strongly absorbing fluids. We expect that the presented sensing scheme can be adapted to a wide range of analytes and tailored to specific needs.

KEYWORDS: DNA origami, gold nanorods, chiral plasmonics, aptamers, biosensing

Various materials and biochemical entities have properties that are potentially useful for biorecognition and/or transduction in sensing applications.1 The difficulty often lies in the integration of those properties for the realization of efficient and reliable biosensors. This is especially challenging for nanoscale sensing devices. Herein, we describe design, fabrication, and characterization of a nanoscale sensing device that combines the beneficial biosensing properties of DNA origami technique, chiral plasmonics, and aptamers.

DNA nanotechnology, which utilizes DNA as a construction material for the assembly of nanostructures,2,3 is finding increasing use in biosensing applications.4 Recently, the DNA origami technique has emerged as a powerful method for fabrication of complex nanostructures that can serve as templates for assembly of various functional components with nanometer precision.5–10 In addition, reconfigurable DNA origami structures have enabled realization of nanoscale devices with functionalities tailored for the characterization of bimolecular interactions,11–13 biosensing,14–17 and drug delivery.18

Metal nanostructures and their plasmon resonances have been extensively used as optical transducers for sensing. Utility of plasmonics traditionally has stemmed from the ability to generate strong electromagnetic field enhancement.19 Alternatively, the coupling of plasmonic excitations in metal nanoparticles and its sensitivity to interparticle distances can be utilized for sensing.20,21 Among coupled plasmonic systems, reconfigurable chiral plasmonic assemblies enabled by DNA nanotechnology22 are particularly promising candidates for sensing applications because their structural configurations can be readily correlated with the optical responses, e.g., circular dichroism (CD).23 Optical detection based on CD spectros-

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nanorods (AuNRs), 55 nm × 23 nm, are assembled on top of the DNA origami to enable optical detection of the chiral configuration with CD spectroscopy. Detailed description of the DNA origami design and AuNRs assembly is provided in the Supporting Information.

The operation principle of the nanosensor is illustrated in Figure 1B, C. The state of the aptamer-based lock changes upon target binding. The DNA origami structure transduces the state of the lock (closed/open) into two spatial configurations of the AuNRs dimer (chiral/relaxed) with distinct plasmonic CD responses. To demonstrate the versatility of our system, we designed two types of locks. The double stranded (ds)-lock consisted of an aptamer and a complementary strand (cs) and undergoes closed to open transition upon analyte binding (Figure 1B). The split aptamer (sp)-lock consisted of partial aptamer strands and closes upon target binding (Figure 1C). In both cases, we used adenosine as a model analyte and adapted its corresponding DNA aptamer.

Agarose gel electrophoresis, transmission electron microscopy (TEM), and CD spectroscopy were used to characterize the assembly of the DNA origami-AuNRs structures. The structures assembled with high yields (Figure 2A and Figure S1 and S2) and exhibited distinct CD responses in chiral and relaxed states (Figure 2B). The relaxed state had a slight RH CD response preference due to structural imperfections.

First, we tested the performance of nanosensors with ds-locks (Figure 1B). The polyacrylamide gel electrophoresis (Figure S4) showed the shortest length of cs to form a stable hybrid with the aptamer is 12 nt (cs12) and that this hybridization was efficiently broken by adenosine as the adenosine competed against the cs to bind with the aptamer (Figure S5). The sequences of the aptamer and the cs12 were incorporated into the two origami bundles as ds12-lock (Figure S6A) and the origami was in RH chiral state when the ds12-lock was closed. The agarose gel electrophoresis showed that the origami with the ds12-lock shifted faster confirming the closed configuration (Figure S7). After the attachment of AuNRs, the DNA origami-AuNRs constructs with the ds12-lock (ds12-nanosensor) exhibited strong RH CD response. Addition of adenosine resulted in a concentration dependent decrease of the CD signal, indicating the equilibrium shift from the RH chiral state to the relaxed state (Figure 3A). The lowest concentration of adenosine to induce clear change in CD spectra was 300 μM. The highest measured adenosine

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**Figure 1.** Design and operation principle. (A) Schematics of the DNA origami-based chiral plasmonic nanosensor. The DNA aptamer-based molecular lock is employed as a biorecognition element incorporated into reconfigurable DNA origami structure, which hosts two gold nanorods (AuNRs). Two AuNRs constitute a three-dimensional (3D) chiral plasmonic object with circular dichroism optical response that is dependent on the angle between the rods. (B,C) Operation principle of the nanosensor with different aptamer-based molecular locks. (B) Double-stranded (ds)-lock opens upon target binding, and the 3D spatial configuration of the origami-AuNR construct changes from chiral to relaxed. (C) Split aptamer (sp)-lock closes upon target binding, and the spatial configuration of origami-AuNR constructs changes from relaxed to chiral.

**Figure 2.** DNA origami-AuNR structures and their chiroptical responses. (A) TEM image of the DNA origami-AuNR structures. The structures showed typical binding preference to the TEM grid, i.e., parallel origami bundles and rods. Scale bar, 100 nm. (B) CD responses in RH chiral (red line) and relaxed (blue line) states.

**Figure 3.** Ds12-nanosensor. (A) Relative CD responses at different concentrations of adenosine. Concentrations are given in mM. (B) CD responses of the nanosensor in the presence of adenosine (2 mM) or guanosine (saturated solution, 2.5 mM).
concentration was 10 mM due to the limited solubility. The typical response time of the ds12-nanosensor was in the range of tens of minutes (Figure S11) and the limit of detection (LOD) was ~270 μM (see the Supporting Information). To demonstrate that the CD change was induced by the specific interaction between the aptamer-based lock and the adenosine, we constructed the structures in which the aptamer strand in the lock was replaced by a scramble strand that formed the same 12 base pairs (bp) with the cs12 sequence but lacked the ability to bind adenosine. No adenosine response was observed when the aptamer sequence was scrambled or omitted (Figure S8). The ds12-nanosensor also responded to the adenosine triphosphate (ATP), which is another target of the adenosine aptamer (Figure S9). No decrease in the CD signal was observed in the presence of the guanosine (Figure 3B), which confirms the selectivity of our sensing device.

To test whether the ds-lock can be used to detect adenosine at concentrations below 300 μM, we shortened the hybridization length between the aptamer and the cs from 12 bp to 9 bp by substituting the cs12 with the cs9 sequence (ds9-lock, Table S2). Fewer base pairs of the aptamer-cs hybrid are expected to benefit the competing binding between the adenosine and the aptamer. Although free cs9 and aptamer were not expected to form a stable hybrid at room temperature the origami-AuNRs constructs with the ds9-lock (ds9-nanosensor) exhibited strong CD signals, possibly due to the enhanced local concentration effect after the incorporation into the origami structures. The ds9-nanosensor exhibited clear CD changes at the adenosine concentration of 30 μM (Figure 4A), 1 order of magnitude smaller than the ds12-nanosensor. The LOD of ds9-nanosensor was ~20 μM. Furthermore, the ds9-nanosensor responded to the addition of adenosine within 1 min (Figure S11), much faster than ds12-nanosensor. Although shortening the hybridization length of the ds-lock improved the detection limit and speed, the correlation between the CD signal and the adenosine concentration became less steep (Figure 4D).

Next, we tested the performance of the nanosensor with the split aptamer (sp)-molecular lock (Figure 1C), which consisted of two partial aptamers (apt1, apt2) that do not interact with each other but can form an apt1-apt2-adenosine complex in the presence of the adenosine. We inserted the apt1 and apt2 on the two bundles of the origami as shown in the Figure S6B. The complex formation of apt1-apt2-adenosine enabled the origami to possess a LH chiral configuration. The CD spectra of the origami-AuNRs with the sp-lock (sp-nanosensor) in adenosine showed that the higher the adenosine concentration, the higher the LH CD signal generated (Figure 4B). Of note, although the LH CD spectra became clear only after the adenosine concentration reached 250 μM, the RH CD signal of the relaxed state exhibited clear decrease in CD at 50 μM of adenosine. The high adenosine concentration was required to compensate the constructs’ intrinsic tendency toward the RH configuration in the relaxed state. The LOD of sp-nanosensor was ~65 μM. When the sp-lock was inserted on the opposite side of the origami bundle to close the construct as the RH structure, the signal started to increase at 25 μM (Figure S10).

The DNA origami technique provides the ability to combine both types of locks in one device. We used the ds9-lock responsible for RH chiral to relaxed transition and sp-lock responsible for relaxed to LH chiral transition to test the performance of the dual lock system (ds9-sp-nanosensor). The addition of adenosine is expected to open the ds9-lock to drive the equilibrium from the RH toward the relaxed state and close the sp-lock to shift the equilibrium from the relaxed to the LH state. The ds9-sp-nanosensor not only showed clear change in CD already at 30 μM of adenosine (Figure 4C), similar to ds9-lock based system, but also exhibited steeper CD signal–adenosine concentration dependence (Figure 4D) compared to the nanosensor with only ds9-lock. The LOD of ds9-sp-nanosensor was ~25 μM. In addition, the response time of the double lock-nanosensor was in the range of several minutes (Figure S11).

Finally, we tested our devices in absorbing fluids. The CD responses of the nanosensor were almost identical at optical densities of environment up to OD ≈ 3 (Figure 5). At ODs above 3, the CD response became too noisy for reliable measurement.
In summary, we developed a nanoscale biosensing device that combines the benefits of the DNA origami technique, chiral plasmonics and aptamers. One of the main advantages of such combination is the high level of programmability. Aptamer based biorecognition elements can be easily incorporated into DNA origami structures at desired locations. While the availability of split aptamers is limited, the double stranded lock strategy can be adapted to a wide variety of aptamers and targets. Dynamic DNA origami structures can be optimized to undergo desired spatial reconfiguration upon target binding and the spectral position of optical responses can be tuned by the size and material composition of the nanorods. In addition, the nanosensor enables optical detection in environments with strong optical extinction, which would simplify preparation procedures of biologically relevant samples. Our results demonstrate a promising route toward the development of the aptamer-based sensing platforms utilizing optical responses of chiral plasmonic nanostructures.

ASSOCIATED CONTENT

\* Supporting Information
The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acsami.8b19153.

Materials, experimental details, gel electrophoresis images, TEM images, and absorption and CD spectra (PDF)

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A.K. and Y.H. conceived the research and designed the experiments. K.N. synthesized the AuNRs. K.N. performed the gel electrophoresis experiments, and performed the CD measurements. A.K. and A.N. performed TEM characterization. A.K. and Y.H. wrote the manuscript with the contributions of all authors. All authors have given approval to the final version of the manuscript.

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Notes
The authors declare no competing financial interest.

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