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# Single-step electronic detection of femtomolar DNA by target-induced strand displacement in an electrode-bound duplex

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**We report a signal-on, electronic DNA (E-DNA) sensor that is label-free and achieves a subpicomolar detection limit. The sensor, which is based on a target-induced strand displacement mechanism, is composed of a “capture probe” attached by its 5′ terminus to a gold electrode and a 5′ methylene blue-modified “signaling probe” that is complementary at both its 3′ and 5′ termini to the capture probe. In the absence of target, hybridization between the capture and signaling probes minimizes contact between the methylene blue and electrode surface, limiting the observed redox current. Target hybridization displaces the 5′ end of the signaling probe, generating a short, flexible single-stranded DNA element and producing up to a 7-fold increase in redox current. The observed signal gain is sufficient to achieve a demonstrated (not extrapolated) detection limit of 400 fM, which is among the best reported for single-step electronic DNA detection. Moreover, because sensor fabrication is straightforward, the approach appears to provide a ready alternative to the more cumbersome femtomolar electrochemical assays described to date.**

biosensors | electron transfer | gold electrode | methylene blue | signal-on

**T**he rapid, specific detection of nucleic acid sequences offers the potential for utility in both clinical and research diagnostic applications (1, 2). A wide range of electronic DNA detection schemes have been described to date (3–5), the best of which achieve limits of detection (LOD) ranging from picomolar to femtomolar. Among promising recent examples are approaches based on conductive links produced through the catalytic deposition of silver by nanoparticle-linked secondary probes (LOD 500 fM) (6), the electrocatalytic oxidation of modified bases by Os(bpy)<sub>3</sub><sup>3+</sup> (LOD 400 fM) (7), the electrochemistry of water-soluble, ferrocene-functionalized cationic polythiophenes (LOD 500 pM) (8) or ferrocene-linked triblock copolymer–DNA hybrids (LOD 100 pM) (9), the electron transfer of ferrocenyl-tethered poly(amido-amine)dendrimers in a sandwich-type enzyme-linked DNA sensor (LOD 100 pM) (10), charge transport from electroactive DNA intercalators (with magnetic sample concentration) (LOD 2 pM) (11), the chronopotentiometric detection of micrometer-long indium rod tracer in a DNA sandwich hybridization assay (with magnetic sample concentration) (LOD 2.6 pM) (12), and the anodic stripping voltammetry of silver nanoparticles deposited in a multistep reduction process initiated by a labeled secondary probe (LOD 0.1 fM) (13).

Although the detection limits of the above-described sensor technologies are often impressive, achieving them requires the addition of exogenous, label-containing secondary probes and, typically, complicated, multicomponent deposition/amplification steps. For example, although Hwang *et al.* (13) report an exceptional 0.1 fM detection limit, achieving it required a five-step assay, including an enzyme-linked secondary probe, enzymatic reduction of *p*-aminophenyl phosphate, the concomitant reductive deposition of silver, and, finally, anodic stripping voltammetry to quantify the deposited silver. In contrast to these

relatively cumbersome assays (6–13), we and others have recently described several reagentless, single-step electrochemical DNA detection methods based on immobilized, redox-tagged single-stranded DNA (14, 15) and DNA stem-loops (16–18). The latter strategy, termed E-DNA (16), is based on a redox-tagged DNA stem-loop structure that self-assembles on a gold electrode by a gold–thiol bond. The hybridization of target with the loop region induces a large conformational change in this surface-confined DNA and thus significantly affects the rate of electron transfer between the redox moiety and the electrode. The associated change in redox current produces a signal indicative of the target without the addition of exogenous reagents.

In addition to being reagentless and single-step, the E-DNA sensor has numerous advantages in terms of its applicability to real-world oligonucleotide detection. It is, for example, readily reusable, sequence-specific, and selective enough to perform even when placed directly in blood serum and in solutions contaminated with soil, foodstuffs, or other complex materials (19). For many applications, however, these advantages are offset by the modest sensitivity of such sensors: the best reported LOD for an E-DNA sensor is, at tens picomolar (16), several orders of magnitude poorer than the best multistep electronic approaches (6, 11, 13). Here, in contrast, we report an E-DNA design that couples the femtomolar detection limits of the best reagent-intensive electrochemical methods with the single-step convenience of the E-DNA-sensing platform.

The LOD of the original E-DNA sensor is, in part, limited by its “signal-off” architecture (target binding reduces the redox current); the gain of signal-off sensors is limited because the target can suppress no more than 100% of the original signal. “Signal-on” sensors for which target binding increases the redox current, in contrast, have the potential for greatly improved sensitivity because, under ideal conditions (as the background signal approaches zero), the gain of such a sensor increases without limit (20). Here we describe an E-DNA sensor that, in contrast to the original E-DNA signal-off architecture, is signal-on, and thus it exhibits enhanced gain and a significantly improved detection limit.

## Results

The signal-on E-DNA sensor is based on a target-induced strand-displacement mechanism (Fig. 1). The sensor is composed of two parts. The first part is a single-stranded “capture

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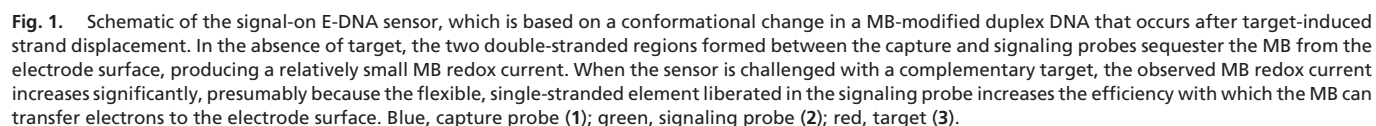
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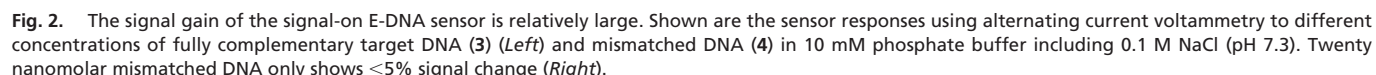
Abbreviations: E-DNA, electronic DNA; LOD, limits of detection; MB, methylene blue.

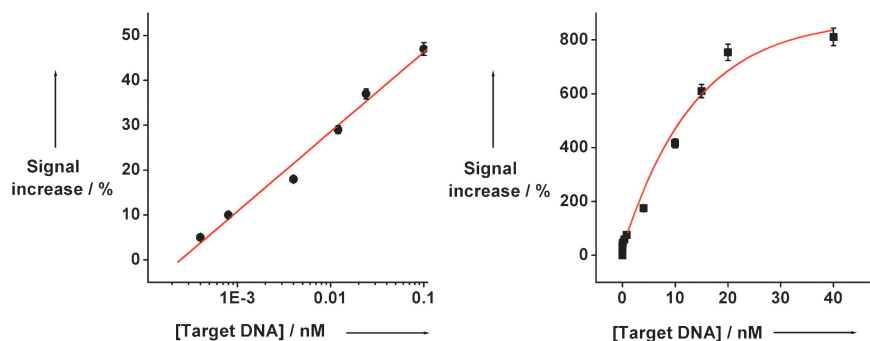
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Because of the large signal change induced by the target, the signal-on E-DNA sensor is significantly more sensitive than the





**Fig. 3.** The dynamic range of the signal-on E-DNA sensor covers target concentrations from 400 fM to  $\approx 20$  nM. The error bars represent the SD of four AC voltammetric scans conducted with a single electrode at each target DNA concentration. Multiple matched electrodes were used to collect this data set.

corresponding signal-off sensor. As a result of limited electron transfer from the signaling probe we observe only small,  $\approx 10$  nA faradaic currents in the absence of target [Fig. 2, 0 nM DNA (3)]. After target (3) binding, the faradaic current increases significantly (Fig. 2), typically reaching  $\approx 7$  times that of the original signal at saturating target concentrations (above  $\approx 20$  nM; Fig. 2 Right). The signal gain of this new sensor architecture is thus an order of magnitude greater than the  $\approx 35\%$  signal suppression observed at similar target concentrations with a signal-off E-DNA sensor (15–17). This improved signal gain leads in turn to significantly improved sensitivity; the directly measured (not extrapolated) detection limit of the current sensor architecture is 400 fM (Fig. 3 Left), and the dynamic range of the sensor spans four orders of magnitude (Fig. 3 Right). In contrast, control experiments reveal that the addition of a 5-base mismatched target (4) at a concentration of 20 nM (which would be a saturating target concentration for the perfect match) does not produce any significant signal change (Fig. 2 Right).

The residual reduction peak observed in the absence of target could arise because of limited, long-range transfer from the MB to the electrode surface. Alternatively, a small fraction of the signaling probes might not hybridized with the 3' terminus of the capture probe, allowing direct, short-range transfer of electrons from the MB to the electrode. Finally, the capture/signaling probe double-stranded complex might be sufficiently flexible to allow the MB to collide occasionally with the electrode surface and perform short-range electron transfer. Using the Laviron equation (24) to calculate the interfacial electron transfer rates between the MB and the electrode we obtain rate constants of  $10 \text{ s}^{-1}$  and  $58 \text{ s}^{-1}$  before and after reacting with target DNA (3), respectively. The relative similarity in these rate constants (which decay exponentially with increasing distance and which could have differed by orders of magnitude) suggests that the electron transfer mechanism is similar in the both presence and absence of target, supporting the latter two hypothesized mechanisms.

The signal-on E-DNA sensor requires that the 3' terminus of the signaling probe remain physically associated with the electrode surface after target binding. To ensure that this association occurs we have designed the system to form a 15-base duplex with the 5' terminus of the immobilized capture probe. In the optimized hybridization we have used, this duplex element is stable for more than 24 h in the absence of target. The described sensor is not, however, regenerated on stringent washing (20 min in distilled, deionized water) without the reintroduction of the signaling probe; the hybridization that retains the signaling probe on the gold surface does not withstand the conditions required to disrupt target (3) binding. The sensor can be regenerated, however, with the reintroduction of a new signaling probe (by hybridization of MB-modified signaling probe for 6 h at room temperature). We speculate that alternative covalent

linkages will alleviate this requirement and lead to more readily reusable sensors.

### Discussion

The mechanism underlying the new sensor architecture is target-induced strand displacement. The low background currents produced by the initially rigid, double-stranded sensor element allow for significantly improved signal gain over earlier, signal-off E-DNA architectures (16) and provide for a 400 fM detection limit.

A reagentless, signal-on E-DNA sensor architecture was described previously in the elegant approach of Immoos *et al.* (15). In this work, which utilizes a surface-immobilized, single-stranded oligodeoxynucleotide—poly(ethylene glycol) triblock polymer, signal arises when a large conformational change is induced by the simultaneous hybridization of both the top and bottom oligonucleotide of the immobilized triblock probe with the target. This simultaneous hybridization forces a terminally linked ferrocene redox tag into proximity with the electrode surface, increasing the signaling current. The reported detection limit for the Immoos sensor (15) is, however, three orders of magnitude poorer than that reported here, presumably because the flexibility of the unbound, single-stranded triblock polymer is sufficient to allow the ferrocene to collide with the electrode surface, producing a significant background current. In the approach reported here, in contrast, the sensing DNA forms a relatively rigid double helix in the absence of target, presumably accounting for the orders of magnitude smaller background current we observe. This reduced background current ensures that the signal gain of our sensor is relatively large, thereby lowering our limit of detection to femtomolar levels.

The E-DNA sensor described here works by target-induced strand displacement, with the detection signal arising as a result of a large, binding-induced change in the probe flexibility and thus the electron-transfer distance. The observed detection limit of this simple sensor is among the best reported to date for electronic sensors. Moreover, unlike the few E-DNA detection approaches that approach or exceed this detection limit, the architecture described here is label-free and enables single-step detection. Given the combined sensitivity and simplicity of the signal-on E-DNA architecture, it appears that it may be of utility in a variety of DNA-detection applications.

### Materials and Methods

**Reagents.** Modified DNA oligonucleotides were synthesized by BioSource, Int. (Foster City, CA), purified by C18 HPLC and PAGE, and confirmed by mass spectroscopy. The sequences of these oligomers used are as follows: (1), 5'-HS-(CH<sub>2</sub>)<sub>6</sub>-GCG-AGTTAGACCGATCCCCCCTTCGTCCAGTCTTTT-3'; (2), 5'-MB-(CH<sub>2</sub>)<sub>6</sub>-GACTGGACGCCCCCCCATCGGTCTA-



