Nucleic acid detection using a universal electrochemical sensor for SNS differentiation

Detección de ácidos nucleicos usando un sensor electroquímico universal para la discriminación de SNS

Percy Calvo-Marzal¹, Dmitry M. Kolpashchikov¹,²,³ and Karin Y. Chumbimuni-Torres¹

¹ Department of Chemistry, University of Central Florida, 4000 Central Florida Blvd., Orlando, FL 32816
² National Center for Forensic Science, University of Central Florida, Orlando, FL 32816
³ Burnett School of Biomedical Science, University of Central Florida, Orlando, FL 32816

Recibido el 26 de diciembre del 2017, aceptado el 31 de diciembre del 2017

Abstract

Nucleic acid detection with high sensitivity and selectivity capabilities could aid in the diagnosis of various diseases such as: infections, cancer, among others. Discrimination of a mismatch or single nucleotide substitution (SNS) is challenging due to a minimum pair to pair formation double strand needed to stabilize the structure. Here we present a new approach to attain the needed stability by using additional segments strains to favor stabilization. The use of additional complementary strains facilitates the discrimination of a SNS in a cooperative way.

Keywords: Nucleic acid, Single nucleotide substitution, infection, cancer

1. Introduction

The World Health Organization (WHO) identifies infectious diseases as a major global health issue. [1-3]. This continued threat is due to the emergence of new diseases and antibiotic resistant bacteria and the rapid dissemination of these diseases due to overpopulation and globalization.[4]

To address these threats, fast, affordable, and reliable diagnostic tools are needed. Detection of single nucleotide substitutions (SNS) can differentiate between pathogenic and non-pathogenic bacteria. Pathogenic E. coli strains-O157:H7 exhibit 99% genome identity to non-pathogenic K12 strains. Nucleotide alignments reveal that 16S rRNA sequences have 9 SNS between the two strains.

State-of-the-art sensors for nucleic acid detection fail to detect these SNS, especially at ambient temperatures and when buried in the secondary (folded) structure that is characteristic of ribosomal RNA (rRNA) [5-7]. Nucleic acid amplification tests (NAATs) offer high sensitivity and rapid results [8-10] but they are prone to contamination, frequently produce false-positive results and require technical expertise.

Here, we propose to develop a point-of-care (POC) platform based on a self-assembling binary DNA approach [11, 12]. Two hybrid intermediary strands will be designed such that half of each strand is complementary to the target nucleic acid and the other half complementary to a universal DNA probe that is immobilized on a solid surface. Self-assembly of these components along with the target rRNA will result in formation of a 4J structure. A redox marker covalently attached to an intermediary strand allows formation of the 4J structure to be monitored by electrochemical detection (4J E-biosensor).

The ideal POC sensor for bacterial pathogens will exhibit a signal in the presence of pathogenic bacteria and distinguish between closely related, non-pathogenic species. Binary probes exhibit
excellent selectivity and can differentiate SNS in DNA strands [13-15]. We will use the high selectivity of binary probes to solve the challenging task of differentiating between two *E. coli* strains—O157:H7 and K12. Sequences of 16S and 23S rRNA from these two strains are 99% identical. The alignment data show that the 16S and 23S rRNA only exhibit 9 and 15 SNS between the two strains, respectively. There are 2 SNS clusters containing either two or three nucleotide variations in 23S rRNA, providing target regions to differentiate between the two strains [16].

Recent work by the PI and CoPI has demonstrated for the first time a single probe that universally recognizes DNA and RNA, differentiates between SNS at room temperature and uses an electrochemical transducer. The platform was adapted from the 4J molecular beacon approach proposed and investigated by the CoPI. This platform utilizes a hairpin DNA stem-and-loop (SL) probe attached to a gold electrode through a thiol bond (Fig. 1). Two adaptor strands, m and f, are introduced during hybridization. The m-strand is covalently bound to an electrochemical redox marker, methylene blue (MeB). The DNA-SL probe is stable in its hairpin conformation in the absence or presence of the target and hybridizes to form a 4J architecture only when the adaptor strands are present. The electrochemical signal is generated by the electron transfer (eT) between the MeB redox marker on the m-strand and the electrode’s surface.

**Figure 1: Schematic of the 4J E-biosensor for RNA/DNA detection.**

Electrochemical methods combined with highly selective recognition elements can revolutionize real-time and on-site biosensing applications towards POC. Hairpin DNA probes, which consist of a SL structure, offer high selectivity when compared to their corresponding linear probes [17], allowing for the distinction between targets that differ by a SNS [18, 19]. Kramer et al. introduced a molecular beacon approach [20], which optically monitored a hairpin DNA probe containing a fluorophore on one end and a quencher on the other end (Fig. 2A). Fan et al. [21] adapted this approach for electrochemically detection by replacing the fluorophore with a redox marker (electroactive reporter), ferrocene, while covalently attaching the probe’s other end to a gold electrode via a thiol bond (Fig. 2B). In this configuration, the electroactive species is near the electrode, facilitating a high rate of eT. Upon hybridization, the distance between the electroactive species and the electrode greatly increases, resulting in a low eT rate (signal decrease).

**Figure 2: A) Schematic diagram of molecular beacon (MB) and B) E-DNA biosensor.**

The seminal work of Fan et al. [21] opened the door to SNS differentiation using hairpin DNA-SL probes with electrochemical transducers. Although SNS differentiation was possible using this approach, the electrochemical sensor operated as a signal OFF upon hybridization. The primary drawbacks of the signal OFF sensing platform include **i)** false positive responses caused by interactions other than the target binding [21, 24] and **ii)** limited signal since the original signal current can only be suppressed up to 100%, upon hybridization with the target.

These limitations could be addressed by designing a signal ON sensing platform. Different approaches were proposed for signal ON, including the use of DNA-PEG-DNA triblock macromolecule [25], a target-induced strand displacement mechanism [26] and sensors based on hairpin DNA-SL probes [27-30]. However, these signal ON sensing platforms produce a nonzero background signal in the absence of the target. Since the redox marker could be in sufficient proximity to the electrode. In contrast, the proposed 4J architecture integrated onto an electrode displays zero background signal, since the hairpin DNA-SL probe does not contain a redox marker. Instead, the electroactive species is covalently bound to the end of one adaptor strand. The eT is observed only upon hybridization to form the 4J architecture, offering a signal ON sensing platform with zero background signal.

Targets that form stable secondary structures are unlikely to hybridize with the DNA-SL probe because
The proposed fundamental study of the 4J system approach is necessary to engineer a POC device for differentiation of SNS in folded and unfolded nucleic acids. Such a device could present valuable applications in environmental, food safety, clinical diagnostics, and national defense [32-34]. There is a particular interest to detect rRNA to aid in diagnosing bacterial infections. The WHO report that about five million people die annually from lower respiratory infections, diarrheal diseases and tuberculosis [1]. A variety of techniques for rRNA detection have been reported, such as northern blotting [35-38], PCR [39-43], and fluorescence [44-46]. However, these methods are prone to contamination, require skilled personnel, and lack the portability necessary for POC diagnostics [47]. Electrochemical methods have been recently explored as an alternative approach [24,30, 48-51] since they have greater potential for on-site testing while offering fast, simple, portable and inexpensive detection [52-54].

![Figure 3: Schematic representation of hairpin DNA-SL probe hybridize to folded targets. A) Duplex hybrid and B) 4J system.](image)

2. Results

Sequence descriptions can be found in Table 1.

<table>
<thead>
<tr>
<th>Synthetic oligonucleotide</th>
<th>Abbreviation</th>
<th>Sequence</th>
</tr>
</thead>
<tbody>
<tr>
<td>DNA stem loop probe</td>
<td>DNA-SL probe</td>
<td>5'-S-S-(CH₂)₆-TTT-TTT-TTCG-GTT-AAC-ATA-CAA-TAG-ATC-GCG-3'</td>
</tr>
<tr>
<td>Adaptor strand m</td>
<td>m-GCG-MeB</td>
<td>5'-CAA-ACA-CCA-T-TAT-GTT-AAC-GCG-MeB-3'</td>
</tr>
<tr>
<td></td>
<td>m-3T-MeB</td>
<td>5'-CAA-ACA-CCA-T-TAT-GTT-AAC-3'</td>
</tr>
<tr>
<td></td>
<td>m-10T-MeB</td>
<td>5'-CAA-ACA-CCA-T-TAT-GTT-ATAA-3'</td>
</tr>
<tr>
<td>Adaptor strand f</td>
<td>f1</td>
<td>5'-GAT-CTA-TTG-TGT-CAA-ACT-CCA-3'</td>
</tr>
<tr>
<td></td>
<td>f2</td>
<td>5'-GAT-CTA-TTG-ATG-CTG-ATC-3'</td>
</tr>
<tr>
<td>Target</td>
<td>miRNA-122</td>
<td>5'-UGG-AGU-GUG-ACA-AUG-GUG-UUU-G-3'</td>
</tr>
<tr>
<td></td>
<td>T-DNA</td>
<td>5'-CTG-GAT-AGC-GAT-ATG-TGT-TTT-3'</td>
</tr>
<tr>
<td>Single nucleotide</td>
<td>SNS</td>
<td>5'-UGG-AGU-GUG-ACA-AUG-GUC-UUU-G-3'</td>
</tr>
</tbody>
</table>

### Table 1: Nucleic acid sequences used for 4J E-biosensor design and analysis.

**Computational modeling** was used to design the sequence of the m intermediary adaptor strand by studying different nucleic sequences. The m adaptor strand (m-GCG-MeB) was designed to anchor the MeB near the surface of the gold electrode after 4J formation. Results obtained from computational modeling showed that m-GCG-MeB will hybridize with the DNA-SL probe in the absence of the f adaptor strand and the target. Consequently, a false positive electrochemical signal was observed (Fig. 5A-b) due to the energetically favored hybridization between the complementary base pairs and the DNA-SL probe. The revised sequence of the m adaptor strand replaced guanine-cytosine-guanine (GCG) with thymine-thymine-thymine (TTT) at the 3' end to eliminate the unwanted interaction between the m adaptor strand and the DNA-SL probe. Thus, a false positive electrochemical response in the absence of the target was not observed (Fig. 4B-b), indicating that m-3T-MeB did not bind to the DNA-SL probe by itself.
The concentration of the DNA-SL probe during immobilization on the electrode’s surface was optimized and analyzed upon hybridization. Fig. 5A shows that the current density maximized at a probe concentration of 0.1 μM, resulting in a surface density of 2.72x10^{11} molecules/cm^2. Furthermore, the concentrations of m and f1 adaptor strands were varied simultaneously in equimolar concentrations (0.10, 0.25, 0.50, 0.75, 1.0 μM). As seen in Fig. 5B, the current density increases as the concentrations of adaptor strands (m-10T-MeB and f1) increase, reaching a maximum at 0.75 μM.

**Sensor selectivity** was studied with a target containing a SNS (see sequences in Table 1). As observed, the current density was negligible (Fig. 6 left-c) when compared to the fully matched target at the same concentration (50 nM; Fig. 6 left-b). Furthermore, even when the target containing a SNS was used in four-fold excess (200 nM) the current density remained negligible (Fig. 6 left-d), thus reflecting the capability of the 4J E-biosensor to detect a fully matched target even in excess amount of a single base mismatched nucleic acid, a property important in practice.\(^{55}\)

It is also worth notice to highlight that the m strand by itself along with the target does not hybridize as zero signal is observed as shown in Fig. 6 right (blue line).

**Sensor regeneration.** It is known that DNA-based sensors comprised of a linear probe and a fully matched target can be regenerated by the addition of external stress, such as heating the duplex hybrid to its respective melting temperature, or by changing pH to drastic conditions\(^{50}\). In contrast, Lubin et al.\(^{58}\) reported the regeneration of a signal OFF DNA-SL probe sensor by rinsing with water for 30 s. This is most likely due to the DNA-SL probe configuration, which thermodynamically favors a hairpin configuration vs. the duplex hybrid. Similarly, the 4J structure has been shown to dissociate at room temperature after rinsing with water. This dissociation indicates a unique instability of the 4J complex in the absence of ions, most likely Mg\(^{2+}\), which are known to provide stability to DNA 4J structures.\(^{59}\) Thus, the 4J E-biosensor was reused up to 7 times after the original hybridization with over 97% recovery (Fig. 7).
Sensor response. Since the 4J E-biosensor was able to withstand regeneration, calibration curves were performed using the same sensor (Fig. 8). Follow each hybridization event, the sensor was rinsed for 30 s. with water and used for the next target concentration. The current density increased with increasing concentrations of miRNA-122 from 5 to 75 nM as shown in Figure 9A. The linear dynamic range (LDR) was from 5 to 50 nM and the response became nonlinear beyond the upper level, indicating sensor saturation. The limit of detection (LOD) was calculated to be 3.2 nM.

Figure 8: A) Calibration curve for miRNA-122 and B) T-DNA. Inset: SWVs for 1, 5, 15, 25, 30, 40, 50, and 75 nM, in HB from 0 to -0.5 V.

The target-binding arms of the adaptor strands m and f can be changed to tailor the sensor to each new target sequence using one optimized DNA-SL probe, providing a unique universal sensing platform. The sensor was incubated with a target DNA (T-DNA) sequence using the same DNA-SL probe to investigate the ability to detect other nucleic acids along with adaptor strands m-10T-MeB and f2 (sequences in Table 1). As seen in Fig. 9B, the current density increased with increasing concentrations of T-DNA from 5 to 75 nM. The LDR was from 5 to 50 nM and the response became nonlinear beyond the upper level. The LOD was calculated to be 500 pM.

3. Conclusions

Here it was demonstrated that the 4J E-biosensor was successfully used for RNA and T-DNA detection using the same DNA-SL as a universal probe. Regeneration of the sensor was possible with simple rinse with water. Multiple analytes can be detected with one single probe, increasing the potenability for commercialization.

Acknowledgements

The authors acknowledge NSF-nanobiosensing grant number #1706802 and Florida Health Department grant # 7ZK05, the College of Sciences and the Department of Chemistry at the University of Central Florida for financial support of this research. DMK was supported by NSF CCF grants #1117205 and 1423219 and NIH R15AI10388001A1.

References


[29] Wu, Y.; Lai, R. Y. Effects of DNA Probe and Target Flexibility on the Performance of a


E-mails: karin.chumbimunitrorres@ucf.edu, dmitry.kolpashchikov@ucf.edu