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Catalytic activity of dual-hemin labelled oligonucleotide: conformational dependence and fluorescent DNA sensing††

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The conformation-dependent peroxidase activity of dual-hemin labeled oligonucleotide was identified and conveniently utilized to design a sensitive homogenous fluorescent method for DNA sensing.

The sensitive, selective and fast detection of sequence-specific DNA fragment has attracted considerable interest in various fields such as cancer diagnosis and food safety.1 Great efforts have been devoted to this area by designing and optimizing DNA probes labelled with different signal molecules. Fluorescein or rhodamine derivatives labelled probes such as molecular beacon have been widely used in fluorescent DNA sensing.2 Electrochemical indicators such as ferrocene and methylene blue have also been used to label DNA probes for the development of electrochemical DNA biosensors.3 Typically, both the fluorescent and electrochemical DNA biosensors transduce the target DNA hybridization event into the structure change of the probe, and then output detectable signal directly through the labelled molecule in a “one target to one signal molecule” manner.

In order to improve the sensitivity, different signal amplification strategies have been designed to transduce one target recognition event to multiple signal molecules. One of the most feasible transduction pathways is to incorporate multiple signal probes in the detection system by target DNA cycle strategy such as endonuclease cycle,4 exonuclease cycle5 or strand-displacement amplification.6 Another convenient and promising avenue is to introduce an enzymatic cycle for producing the multiple signal molecules, in which the enzyme or catalyst can be labelled to a DNA probe16,7 or formed upon the target DNA recognition.8 This work used small hemin molecules as the biomimetic peroxidase to label the two ends of oligonucleotide and design a target DNA regulated enzymatic cycle by labelling two hemin molecules to two ends of oligonucleotide. The dual-hemin labeled DNA probe was synthesized via amide reaction (Scheme 1A) and characterized with mass spectroscopy (Fig. S1-S3, ESI†). This probe could spontaneously form an intramolecular dimer of hemin, which form a circular structure and led to a low peroxidase activity of regulated by its adsorption on some carbon nanomaterials such as graphene oxide (GO), which is achieved by assembling the hemin labelled single-stranded DNA (ssDNA) on GO.9 However, the quenching properties of GO2 on the fluorescence of catalytically generated dityramine and the adsorption of some formed probe/target duplex on GO surface greatly decreases the DNA detection sensitivity.9 Therefore, it is highly desired to seek other ways to effectively regulate the catalytic activity of hemin.

Due to the weak solubility, hemin can usually self-aggregate into catalytically inactive dimer in aqueous solution.10 Grafting hemin to DNA can significantly improve its solubility, and thus dissociate the hemin aggregate to monomer to recover its peroxidase activity.11 This work proposed a “DNA switch” to regulate the aggregation of hemin and dissociation of hemin dimer by labelling two hemin molecules to two ends of oligonucleotide. The dual-hemin labeled DNA probe was synthesized via amide reaction (Scheme 1A) and characterized with mass spectroscopy (Fig. S1-S3, ESI†). This probe could spontaneously form an intramolecular dimer of hemin, which form a circular structure and led to a low peroxidase activity of

Scheme 1. Schematic illustration of (A) chemical structure of dual-hemin labelled probe, (B) target DNA regulated catalytic activity of probe, and (C) homogenous fluorescence strategy for DNA sensing.
hemin (Scheme 1B). Upon the hybridization of the probe with complementary target DNA, the intramolecular hemin dimer dissociated into highly active hemin monomers. Thus the hybridization recognition could be used to introduce an enzymatic cycle for producing multiple signal molecules. Using the oxidation reaction of non-fluorescent tyramine by hydrogen peroxide (H$_2$O$_2$) as a model, the enzymatic catalysis produced fluorescent dityramine (Scheme S1, ESI†). The reaction possessed quick dynamics and could be completed within 10 min, leading to a relatively fast fluorescence method for target DNA detection (Scheme 1C). This method was highly sensitive and convenient, showing a promising application in practice.

To investigate the structure change of the probe upon its recognition to target DNA, the UV-visible absorption spectra of the probe and the formed probe/target duplex were firstly measured. The absorption spectrum of probe/target duplex, which was formed by mixing 1 µM probe and 1 µM target DNA to incubate for 10 min, showed a characteristic peak of hemin monomer at 402 nm (Fig. 1A, curve a), while the absorption spectrum of the probe showed a blue-shifted, broadened and hypochromic peak at 378 nm (Fig. 1A, curve b). This could be attributed to the formation of a face-to-face dimer of hemin, which was benefited from the flexible structure of ssDNA probe. Similarly, the absorption spectrum of a control stem-loop structured reference probe (rP1) bearing a conclusive intramolecular hemin dimer also showed a broadened hemin peak at 378 nm (Fig. S4, ESI†), which further confirmed the formation of the circular structure and hemin dimer in the probe.

Another reference probe (rP2) was designed to examine the dissociation of face-to-face dimer by replacing one hemin molecule labelled on the 5′ end of the probe with a fluorescent reporter X-rhodamine (ROX) (Scheme S2, ESI†). In the absence of target DNA, the fluorescence of ROX on rP2 was greatly quenched by hemin after reaction with 10 nM probe (blue), which indicated the formation of hemin-ROX heterodimer. The quenching efficiency of photoinduced electron transfer was very sensitive to the distance between hemin and ROX. After rP2 hybridized with target DNA, the fluorescence of ROX was apparently recovered (Fig. 1B, curve b), which demonstrated the separation of ROX and hemin and significantly increased distance between hemin and ROX due to the formation of rigid probe/target duplex structure. Thus the spontaneously formed intramolecular dimer could be dissociated by hybridizing probe with target DNA.

The fluorescence of dityramine was used to trace the peroxidase activity change of the probe upon its recognition to target DNA. The fluorescent spectra were recorded after mixing 0.7 mM tyramine and 2.0 mM H$_2$O$_2$ with DNA or the mixture of target and probe to initiate the catalytic reaction for 10 min. In the absence of probe, the fluorescent spectra of the mixtures of 0.7 mM tyramine and 2.0 mM H$_2$O$_2$ without with 10 nM target DNA showed negligible fluorescent peak at 410 nm (Fig. 2A, black and red), which could be attributed to the slow oxidation of tyramine by H$_2$O$_2$ to produce a small quantity of dityramine. In presence of 10 nM probe, the fluorescent peak at 410 nm slightly increased due to the weak catalytic activity of hemin dimer (Fig. 2A, blue), which produced more dityramine. However, after 0.7 mM tyramine and 2.0 mM H$_2$O$_2$ were mixed with the mixture of 10 nM probe and 10 nM target DNA for 10 min, the fluorescent peak of dityramine sharply increased (Fig. 2A, pink), indicating the high peroxidase activity of the probe due to the dissociation of hemin dimer after the hybridization of probe with target DNA. The conformation-dependent peroxidase activity could be used for fluorescent detection of target DNA.

As mentioned above, the hemin dimer showed weak catalytic activity, thus the fluorescent spectrum showed slightly increased fluorescent signal over time (Fig. 2B). With the increasing amount of target DNA from 2.5 to 10 nM in 10 nM probe, the fluorescent signal of the mixture of 0.7 mM tyramine and 2.0 mM H$_2$O$_2$ with the hybridization product at the same reaction time increased obviously (Fig. 2B). At 10 nM target DNA, the time-dependent curve reached a plateau at 10 min, indicating a relatively fast enzymatic reaction kinetics. Thus this work selected 10 min as the optimal reaction time. The ratio of fluorescent signals in the presence of 10 nM probe/target duplex or probe was used to optimize the detection conditions (Fig. S5, ESI†). At 2.0 mM H$_2$O$_2$, the ratio increased with the increasing tyramine concentration and reached the maximum value at 0.7 mM. Higher concentration of tyramine led to high fluorescent background and decreased ratio. Thus 0.7 mM tyramine was used as the optimal condition, at which the maximum ratio occurred at the H$_2$O$_2$ concentration of 2.0 mM. The ion strength and pH of reaction buffer greatly influenced both the hybridization and enzymatic reaction rate. High NaCl concentrations decreased the fluorescent signal due to the inhibited peroxidase activity of hemin. The optimal NaCl concentration and pH were 100 mM and 7.4, respectively (Fig. S6, ESI†).

The peroxidase activity change of hemin upon the recognition of the designed probe to target DNA to dissociate the hemin dimer offered a method for homogeneous fluorescent detection of target DNA. Theoretically, one target DNA could recover the peroxidase activity of two hemin molecules labelled on the same probe for catalyzing the oxidation reaction of tyramine to form fluorescent dityramine. Thus “one target” could lead to the formation of “multiple signal molecules”, which provided a novel signal amplification strategy. The FL intensity was proportional to target DNA concentration ranging from 0.1 to 10 nM ($R^2 = 0.993$) (Fig. 3A). The detection limit was estimated at 3σ to be 19 pM, which was 10 times lower than that using GO to regulate the peroxidase activity of ssDNA-hemin probe, and also lower than some fluorescent (1 nM) chemiluminescent (1 nM) and colorimetric (0.2 µM) DNA detection methods based on G-quadruplex/hemin DNAzyme. Compared with DNAzyme, the preparation of dual-hemin labelled probe needs a labelling step. However, it is a ready-to-use probe, while hemin-DNAzyme must be labelled to recognition unit for...
specific detection (ESI†). The specificity of this protocol was demonstrated to be acceptable for discriminating the single- (smT) and three-base mismatched target (tmT) from the complementary target (Fig. 3B). In addition, 10 or 100-fold concentration of non-complementary target (ncT) showed much lower FL intensity increment than target DNA, indicating the excellent selectivity for DNA detection.

This work designed a dual-hemin labelled oligonucleotide probe as a novel “DNA switch” with regulative peroxidase activity, which was achieved by the hybridization of the probe with target DNA. Using fluorescent oxidation product of tyramine as tracing molecule, the intramolecular hemin dimer with low peroxidase activity ensured the catalytic activity of hemin to produce fluorescent dityramine. Using fluorescent oxidation product of tyramine as tracing molecule, the intramolecular hemin dimer with low peroxidase activity ensured the catalytic activity of hemin to produce fluorescent dityramine. Based on the conformation-dependent peroxidase activity, the strategy of “one target” to “multiple signal molecules” and a sensitive, specific and homogeneous fluorescent DNA sensing method was proposed. This method can be easily extended to detect different target DNA by simply changing the base sequence of the labelled small molecule catalyst on oligonucleotide probe.

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Notes and references

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Graphical Abstract:

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