Single-step multivalent capture assay for nucleic acid detection with dual-affinity regulation using mutation inhibition and allosteric activation†

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The rational modulation of receptor affinity through distal-site mutation and allosteric control is valuable in biosensor design to tune the useful dynamic range. Our ability to programatically engineer dual-affinity regulation into diverse affinities of target binding and activities of hybridization chain reaction, however, remains limited. By programmable engineering of the switching equilibria of the recognition hairpin using distal-site mutation inhibition and allosteric activation, we obtained a set of receptors varying significantly in affinities of target binding and activities of the hybridization chain reaction. For the first time, we developed an electrocatalytic biosensor for nucleic acid detection with a tunable dynamic range based on a conformational switch triggered bidirectional hybridization chain reaction and blocker assisted multivalent binding. This designable biosensor thus enables single-step incubation, diverse affinities of target binding, diverse efficiencies of signal amplification and diverse single nucleotide discrimination for quantitative analyses of nucleic acids of various lengths in serum, which holds great potential as a compelling platform suitable for liquid biopsy.

Introduction

Mutation and allosteric control are two main strategies for biomolecules regulating their affinities in many naturally occurring biology processes.1–5 The underlying mechanisms that regulate molecular binding affinities are increasingly providing inspiration in biosensors to tune, extend and narrow the useful detection range. Mutation and allosteric control are often employed in biosensors to regulate diverse affinities of receptors responding to their target molecules, which efficiently transfer a given input to an altered output.6–10 Biomimetic sensors that employ mutation or allosteric regulation have a broad control of the dynamic range, but most of them have low sensitivity of signal transduction.11,12 The biosensors based on precise valence control of enzymes have high enzymatic amplification for signal transduction. However, they have a limited adjustable dynamic range, which is only dependent on the number of linked enzymes.13,14 The dual-affinity regulation of receptors with diverse affinities of target binding and activities of the hybridization chain reaction are ideally suited to both extend the dynamic range and amplify signal transduction. This unique property of dual-affinity regulation meets the desirable demand of biosensors with a tunable dynamic range for the analysis of biomarkers in clinical samples with different concentrations. For example, the levels of the overall cell-free DNA expression profile vary from the aM level to the pM level in peripheral blood samples in both healthy controls and cancer patients.15,16

The detection of cell-free DNA expression profiles at the genetic level enables broad liquid biopsy approaches in infection diagnostics,17 early childhood wheezing,18 fetal chromosome abnormalities,19 cancer management,20–25 etc. As cf-DNA is more chemically and bioenvironmentally stable than RNA, cf-DNA is particularly informative, suitable as a molecular biomarker.26,27 Sensitive, robust and specific detection of DNA plays an important role in efficient diagnosis, prognostic analysis of cancer metastasis and cancer therapies. However, sensitive and single nucleotide resolution detection of DNA remains challenging because of the low abundance of targets in peripheral blood.

In the past few decades, some popular and potent technologies using signal amplification strategies have been developed to increase the nucleic acid detection sensitivity. Typically, DNA nanostructure based signal amplification methods mainly include the rolling circle reaction (RCA),28–34 hybridization chain reaction (HCR),35–44 entropy-driven catalysis (EDC),45–48 catalyzed hairpin assembly (CHA),49–54 etc. The hybridization chain reaction is even widely employed in biosensors to amplify the signal. Fan and coworkers had integrated the hybridization chain reaction amplification and tetrahedral DNA nanostructure (with outstanding hybridization conditions) to build an ultrasensitive
nucleic acid detection platform. However, it suffers from extensive sample processing with multiple steps of incubation and washes, which restrain its applications in on-site, real-time analytical devices suitable for point of care (POC) detection.

Existing hybridization chain reaction amplified biosensors have several disadvantages. First, the efficiency of the hybridization chain reaction that occurred at interfaces is low due to the limited mass transfer of the hairpins to the surface. Second, the hairpin probes used in the hybridization chain reaction are not universal due to the sequences of the hairpins varied with the change of the initiator sequence. Third, the stabilities of the hairpins that will affect the signal to noise (S/N) ratio are greatly dependent on the target sequence. Fourth, although ultrasensitive nucleic acid detection performance has been obtained by using hybridization chain reaction amplification, multiple processing steps are always required. The cancer cell capture and analysis efficiency of the hybridization chain reaction (mbHCR), which can produce long chain products with multiple branched arms for multivalent capture and multiple biotins linked with SA-HRP for further signal amplification. A blocker was employed to prevent the binding of unreacted hairpins with the probes on the electrode. Thus, the target binding, signal amplification and multivalent binding can be finished in single-step incubation. Furthermore, by employing the allosteric activation strategy, we could sensitively detect nucleic acids of diverse lengths with excellent selectivity using a universal signal reporter in diluted serum.

Results and discussion

The design of the single-step electrocatalytic assay for the analysis of nucleic acids is depicted in Fig. 1. Our re-engineered multibranched and bidirectional hybridization chain reaction contains two monomer hairpins: recognition hairpin (rH) with a single-stranded extension at the 5' termination and signal hairpin (sH-biotin). The sequences of the two hairpins used in this work followed the sequences designed by Pierce and coworkers except that a section of the sequence was inserted in the rH for target recognition. In the absence of the target, the rH and sH do not hybridize each other because the hairpins are protected by long stems, and therefore the signal output is suppressed. Binding of a specific target DNA sequence to the loop of rH breaks the stem and triggers the hybridization chain reaction in both directions by the two hairpins into the multibranched double helix. Long mbHCR products are produced for efficient electrochemical signal amplification and multivalent capture. In order to obtain target binding, the hybridization chain reaction and multivalent capture at single-step incubation, the mbHCR products are pretreated with SA-HRP and the blocker. With optimized design, the blocker is employed to hybridize with unreacted rH and thereby it helps to prevent the capture of excess rH on the gold electrode surface. However, the blocker cannot hybridize easily with the rH in the mbHCR products due to the low equilibrium stability. It is important to note that the addition of active-2 for enhancing rH binding affinity will not disrupt the hybridization of rH with the blocker. This assumption was estimated via NUPACK at 25 °C (Fig. S1†).

The useful dynamic range of the DNA binding switch and hybridization chain reaction are controlled by using distal-site mutation inhibition and allosteric activation, in which target binding affinity and hybridization chain reaction activity are controlled by either destabilizing or stabilizing the receptor (Fig. 1, top). To create this situation, we tuned the switching equilibria of rH to alter its dual-affinity by reducing or extending the length of the stem in rH. The free energy of rH was
estimated via NUPACK at 25 °C (Fig. S2†). It decreased from
–26.07 kcal mol⁻¹ (rH, original hairpin with 18 base pairs of
stems) to –30.63 kcal mol⁻¹ (rH-v1, 21 base pairs of stems) and
–35.41 kcal mol⁻¹ (rH-v2, 24 base pairs of stems) indicating the
progressive reduction of the dual-affinity of the receptors
through mutation inhibition. When adding allosteric effector
activator 1 or activator 2 (see sequences listed in Table S1†)
to help open the hairpin structure, the equilibrium probability of
the stem in rH decreased suggesting the easier opening of rH-a1
and rH-a2 by target DNA than that of original rH.

To demonstrate the feasibility of tuning the target binding
affinity and hybridization chain reaction activity of mbHCR, gel
electrophoresis analysis was first performed. The product of
receptor hybridized with the target increased with the reduced
switching equilibrium constant of rH (Fig. 2A). The product of
mbHCR was also apparently related to the stability of rH. The
molecular weight of the multibranched nicked double helix
became larger by decreasing the stability of the rH (Fig. 2B),
which demonstrated the negative and positive modulation of
the target binding affinity and hybridization chain reaction
activity though mutation inhibition and allosteric activation.

We next characterized the kinetics of the dual-affinity
controlled mbHCR through monitoring the fluorescence intensity
change initiated by 0 or 100 nM target DNA. By employing
mutation inhibition and allosteric activation to modulate the
target binding affinity and hybridization chain reaction activity
of the receptors, the kinetic rate of fluorescence intensity increased
with the increasement of dual-affinity (Fig. 3A–E). We further
demonstrated the bidirectional initiation of mbHCR in the
following experiments. 5’ termination (obtain 3’ trigger) and 3’
termination (obtain 3’ trigger) were employed to block one
direction initiator. Then we monitored the real time fluorescence
intensity of the hybridization chain reaction induced by the same
amount of 3’ trigger and 5’ trigger (Fig. 3F and G). We observed
a relatively low rate constant for the 5’ directional HCR because
the loop-based displacement in the 5’ directional HCR was
harder (much higher energy barrier to open the hairpin) than
the toehold-based displacement in the 3’ directional HCR. The AGE
analysis further proved the lower efficiency of the 5’ directional
HCR than the 3’ directional HCR (Fig. 3H).

Having validated the facile dual-affinity regulation of mbHCR,
we next attempted to detect nucleic acids through an

![Fig. 2](image-url)  
**Fig. 2** (A) Analysis by AGE of reactions of receptors with the target. Lane 1, 1 μM rH-v2 + 0.2 μM target-24; lane 2, 1 μM rH-v1 + 0.2 μM target-24; lane 3, 1 μM rH + 0.2 μM target-24; lane 4, 1 μM rH-a1 + 0.2 μM target-24; lane 5, 1 μM rH-a2 + 0.2 μM target-24; lane 6, 50 bp DNA ladder markers. (B) Analysis by AGE of the mutation inhibition and allosteric activation controlled multibranched hybridization chain reaction. Lane 1, 1 μM rH-v2/sH; lane 2, 1 μM rH-v2/shH and 0.2 μM target-24; lane 3, 1 μM rH-v1/shH. Lane 4, 1 μM rH-v1/shH and 0.2 μM target-24. Lane 5, 1 μM rH/shH. Lane 6, 1 μM rH/shH and 0.2 μM target-24. Lane 7, 1 μM rH-a1/shH. Lane 8, 1 μM rH-a1/shH and 0.2 μM target-24. Lane 9, 1 μM rH-a2/shH. Lane 10, 1 μM rH-a2/shH and 0.2 μM target-24. Lane 11, 50 bp DNA ladder markers.

![Fig. 3](image-url)  
**Fig. 3** Kinetics of mbHCR with dual-affinity regulation. 0 or 100 nM target-24 was added into the system at t = 0. Fluorescence intensity of each sample was monitored within 10 s of mixing by pipetting. Here, [H] = [F-sh-Q] = 50 nM. The control trace (light gray) and reaction trace (green) show that mbHCR was initiated with 0 or 100 nM target-24. The kinetics of mbHCR was hindered by mutation inhibition (A, B) or accelerated by adding allosteric effectors (D, E). (F, G) kinetics of the hybridization chain reaction initiated in two directions. Same amount of 3’ trigger (F) or 5’ trigger (G) was added into the system at t = 0. Increasing traces of fluorescence indicated that the loop-toehold initiated hybridization chain reaction is slower than the end-toehold initiated hybridization chain reaction. (H) Analysis by AGE of the two directional hybridization chain reactions. Lane 1, 50 bp DNA ladder markers; lane 2, 3 μM 3’ trigger (3 μM rH + 3 μM target-24 + 3 μM 5’ termination); lane 3, 3 μM 5’ trigger (3 μM rH + 3 μM target-24 + 3 μM 3’ termination); lane 4, 3’mbHCR (1 μM rH + 1 μM sh-biotin + 0.2 μM 3’ trigger); lane 5, 5’mbHCR (1 μM rH + 1 μM sh-biotin + 0.2 μM 5’ trigger).
electrochemical method. Our previous work has proved that the multivalent hybridization capture efficiency on the surface is higher than monomer hybridization. However, a limitation existing here is that the unreacted monomer hairpin can’t be removed easily from mbHCR products. In order to obtain the DNA analysis with single-step incubation and wash, we first employed the gap hybridization strategy to diminish the binding efficiency of monomer hybridization on the surface. The monomer hybridization is destabilized because a 10 nt gap is reserved between the capture probe and rH stem. Tetrahedron DNA nanostructures (Fig. S3† validates the successful self-assembly of TDN) with a capturing probe on the vertex were employed to capture the target induced mbHCR products and generate the electrocatalytic i-t current of the enzymatic reaction. We investigated the influence of the capture reaction temperature and the length of single-stranded extension in rH. We found that the reactions were taking place well at 37 °C. However, the influence of the length of single-stranded extension in rH was enhanced with the increased length of single-stranded extension in rH (Fig. S5†). However, the influence of the 10 nt gap generated instability of hybridization will be weakened by increasing the extension length of rH, which means that the competitive hybridization of excess rH with the capture probe will dramatically disrupt the detection of target nucleic acids at low concentrations. The sensitivity of target-24 analysis using rH-target-24 (4-10) was better than that using rH-target-24 (4-12) (data not shown).

To overcome the above limitations, we then employed the blocker to hybridize prior with excess rH prohibiting its capture on the surface. The hybridization cases of diverse lengths of the blocker with the mbHCR product and rH were estimated via NUPACK at 25 °C (Fig. S6 and S7†). The hybridization probabilities of the blocker with the mbHCR product and rH are both enhanced with the increased length of the blocker (Fig. 4, top). The detection performance influenced by the length of the blocker was then investigated. We found that a 12 nt blocker was optimum to produce the best S/N (Fig. 4, bottom). This is because the 14 nt blocker binding to mbHCR products with high efficiency will prevent the multivalent capture. The 10 nt blocker has the lowest probability binding to mbHCR products. However, the 10 nt extension in rH-target-24 (4-6) has relatively lower efficiency of multivalent capture when compared with 12 nt extension in rH-target-24 (6-6). The results also show that the background signal produced at 25 °C was lower than that produced at 37 °C (37 °C used in Fig. S5† and 25 °C used in Fig. 4).

Our previous work also demonstrated that the multivalent capture obtained nearly saturated hybridization efficiency in 15 minutes at room temperature. A multivalent binding reaction was carried out within 30 minutes in this work. As predicted, when we used five rH with different target binding affinities and hybridization chain reaction activities for nucleic acid analysis, the detection limits shifted in intervals by almost one order of magnitude. With the increase in the dual-affinity of rH, the detection limit was enhanced from 5 nM (rH-v2) to 10 fM (rH-a2) (Fig. 5). With mutation inhibition and allosteric activation, the detection sensitivity of the constructed electrocatalytic biosensor could be tuned about 500 000-fold. The hybridization chain reaction performed in solution and the multivalent capture performed on the surface are both efficient. Thus the nucleic acid analysis performances (detection sensitivity and detection speed) are outstanding (Table S2†). Further optimization (such as optimizing the concentration of the activator) can probably lead to even higher sensitivity.

The developed single-step electrocatalytic assay using rH-a2 with high sensitivity could also be adapted for the analysis of nucleic acids of diverse lengths in diluted serum. The detection performance in serum with or without dilution is shown in Fig. S8.† We obtained an optimized S/N in 75% serum and 25%
2× SPSC buffer. The reduced signal observed in serum is possibly due to the nonspecific interaction of the added biomolecules with target DNA and the decrease of salt concentration in serum, which will decrease the efficiency of target binding and the hybridization chain reaction. The sequences of rH and sH used in this work were designed by Pierce and coworkers, excepting a section of recognition sequence was inserted in the rH for target binding. Fig. 6A–C show the excellent performance for the analysis of the DNA targets of three diverse lengths with detection limits of 50 fm (target-DNA-24), 50 fm (target-DNA-21), and 100 fm (target-DNA-18), respectively. The AGE analysis, typical CV curves and i–t curves all validated that the mbHCR triggered by target-DNA-18 was less efficient than that triggered by target-DNA-24 and target-DNA-21 (Fig. S9–S11†). Five samples (S1 to S5) containing different amounts of nucleic acids (lines indicate the known concentrations) were measured. The results (Fig. 6D) show an impressive validation of the good accuracy of our single-step electrocatalytic assay in 75% serum and 25% 2× SPSC buffer.

The sequence-specificity of our assay with single mismatch targets in various locations was considered finally. Clearly, perfectly matched targets could be easily differentiated from other mismatches at the same concentration of 10 nM in buffer by using rH-a2 as the receptor, revealing a good single nucleotide resolution (Fig. S12†). The location of single-base mismatches did not affect significantly the signal output. It is easy to distinguish a perfectly matched target from five single-base mismatched targets at different locations (Fig. S13†). Besides, the selectivity can be increased by using rH+v2 as the receptor because a higher energy barrier has to be broken to open the recognition hairpin (Fig. S14†). These results successfully demonstrate the high potential of the electrocatalytic assay as a single nucleotide resolution biosensor for nucleic acid detection in clinical samples.

Conclusions
In summary, a novel single-step electrocatalytic strategy, called blocker assisted mbHCR, has been proposed. This method enables the detection of nucleic acids with a tunable detection range by programmable modulation of the target binding affinity and hybridization chain reaction activity using mutation inhibition and allosteric activation. The dual-affinity regulation demonstrated here both extended the dynamic range and amplified the signal transduction. The blocker assisted single-step strategy and dual-affinity regulation can be applied in other solid-phase biosensors. Besides, the detection cost can be reduced as the signal hairpin is universal. The selectivity of single nucleotide polymorphism also can be tuned through dual-affinity regulation. The analysis of DNA of multiple lengths in serum with high precision revealed the high potential of the developed strategy as a reliable and sensitive electrocatalytic biosensor for the detection of nucleic acids in complex matrices.

Conflicts of interest
There are no conflicts to declare.

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