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Introduction

Beneting from perfectly predictable base pairings, outstanding biocompatibility, and low cost, a considerable number of DNAbased rigid motifs^{1,2} have been developed to prepare different dimensional DNA nanostructures as biomolecular scaffolds or nanoparticle templates^{3,4} like DNA origami^{5,6} and DNA dynamic nanodevices.7,8 Remarkably, in the development of dynamic DNA nanostructural chemistry, the catalytic multiple-arm DNA junction assembly $(CMDJA)^{9,10}$ as a flexible and ideal basic building block for DNA structural nanotechnology¹¹⁻¹⁵ has attracted increasing attention. Compared to the branched DNA obtained by conventional annealing approaches with tedious temperature control,^{16,17} the isothermal CMDJA was obtained simply by relying on the base-pairing principle, in which the self-assembly can be well controlled by catalysts. Nevertheless, the CMDJA has also suffered from multiple-step reactions and weak driving forces, resulting in inherent time-consuming and suboptimal kinetic and thermodynamic properties, which

Programmable mismatch-fueled high-efficiency DNA signal amplifier†

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Herein, by introducing mismatches, a high-efficiency mismatch-fueled catalytic multiple-arm DNA junction assembly (M-CMDJA) with high-reactivity and a high-threshold is developed as a programmable DNA signal amplifier for rapid detection and ultrasensitive intracellular imaging of miRNA. Compared with traditional nucleic acid signal amplification (NASA) with a perfect complement, the M-CMDJA possesses larger kinetic and thermodynamic favorability owing to the more negative reaction standard free energy (ΔG) as driving force, resulting in much higher efficiency and rates. Once traces of the input initiator react with the mismatched substrate DNA, it could be converted into amounts of output multiple-arm DNA junctions via the M-CMDJA as the functional DNA conversion nanodevice. Impressively, the mismatchfueled catalytic four-arm DNA junction assembly (M-CFDJA) exhibits high conversion efficiency up to 1.05×10^8 in 30 min, which is almost ten times more than those of conventional methods. Therefore, the M-CMDJA could easily address the challenges of traditional methods: slow rates and low efficiency. In application, the M-CFDJA as a DNA signal amplifier was successfully used to develop a biosensing platform for rapid miRNA detection with a LOD of 6.11 aM and the ultrasensitive intracellular imaging of miRNA, providing a basis for the next-generation of versatile DNA signal amplification methods for ultimate applications in DNA nanobiotechnology, biosensing assay, and clinical diagnoses. **EDGE ARTICLE**
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further limits its deeper exploitation and wider application.¹⁸⁻²⁰ On account of these disadvantages, it is of critical significance to improve the reaction speed and efficiency of CMDJAs to advance their superiority and applicability in the areas of nucleic acid signal amplification, biomarker assay, biomolecular nanomachines, and computing.^{2,21}

Since the performance of CMDJAs that relies on the toehold strand displacement reaction (TSDR) is immensely limited by the branch migration step in the displacement domain, $22,23$ the producing rate and yield of a multiple-arm DNA junction would be influenced by the reaction equilibrium of the step of branch migration in the TSDR. Therefore, the introduction of suitable mismatches in the displacement domain of the subsequent hairpin DNA can make the corresponding ΔG more negative than the one in a traditional CMDJA consisting of completely matched reactant $DNA₁²⁴⁻²⁷$ realizing a stronger driving force accompanied with higher reactivity for improving the reaction rate and efficiency. In this study, we utilized NUPACK²⁸ to find some possibly useful mismatches in the displacement domain of the substrate hairpin DNAs and compared the thermodynamic and kinetic parameters as essential indices to pick out the optimal mismatches for a mismatch-fueled catalytic multiple-arm DNA junction assembly (M-CMDJA). As a result, we carved out a mismatchfueled catalytic four-arm DNA junction assembly (M-CFDJA) with optimal properties and used it as a high-efficiency DNA

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signal amplifier for rapid biomarker detection and ultrasensitive intracellular miRNA imaging, which helps in the development of the next generation of nucleic acid signal amplification techniques for biosensing assay and early disease diagnosis.

The reaction mechanism and the practical application of the M-CFDJA are shown in Scheme 1. First, the mechanism illustration of the catalytic formation pathway of the M-CFDJA is depicted in part A. The complementarity relationships of the segments of hairpin DNAs were specified so that it is only with the initiator target miRNA-182-5P that the hairpin DNAs are kinetically prevented from generating the four-arm DNA junctions that can be predicted to dominate at equilibrium. The first hairpin DNA $(H1)$ is opened by the initiator miRNA-182-5P from the toehold to form the duplex H1–miRNA-182- 5P and the pre-locked stem region is exposed, then the second hairpin DNA (H2) will be opened by H1's new stickyend. Similarly, the rest of the hairpin DNAs (H3 and H4) will be opened to form the corresponding multiple-arm DNA junction and the initiator will be released to participate in the next pathway. Most importantly, the optimal mismatches in

Scheme 1 (Part A) schematic illustration of the mismatch-fueled catalytic four-arm DNA junction assembly (M-CFDJA). (Part B) illustration of the mismatched domain in the substrate hairpin DNA. (Part C) illustration of the biosensor construction based on the M-CFDJA for cancer biomarker miRNA-182-5P detection. (Part D) rapid and sensitive intracellular imaging of miRNA-182-5P based on the M-CFDJA.

the neck (displacement domain) of the hairpin DNAs (part B) were selected to engineer the M-CFDJA with a fast reaction rate and high conversion efficiency. With the help of the M-CFDJA, traces of initiator can be rapidly converted into amounts of four-arm DNA junction for constructing a DNA conversion nanodevice. As a proof of concept, the evolved M-CFDJA was applied to develop a highly sensitive electrochemical biosensing platform (part C) for rapid miRNA-182-5P detection with a LOD (6.11 aM) that is nearly two orders of magnitude that of the wild CFDJA. Impressively, the M-CFDJA was further applied in rapid intracellular miRNA imaging to monitor miRNA-182-5P expression in cancer cells (part D), and it was observed to be much more sensitive than the typical FISH approach. The evolved M-CFDJA therefore showed great potential for ultimate application in biological nanobiotechnology and biomarker assay in the nascent stages of diseases. Edge Article

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Results and discussion

First, through polyacrylamide gel electrophoresis (PAGE), it was certified that the reactions in the CMDJA could successfully occur (Fig. S1†). Moreover, the three-arm and four-arm DNA junctions obtained via the CMDJA were respectively characterized using atomic force microscopy (AFM), and the results (Fig. S2†) were as expected. Next, based on the ΔG obtained from NUPACK as an evaluation index of hairpin DNA stability, we selected a suitable number of positions of mismatches following the principle that these hairpin DNAs as reactants with enough stability must possess less negative free energy to lead to a more negative reaction standard free energy as the driving force of the M-CMDJA, thus resulting in a higher amplification efficiency with low background leakage. In conclusion, one mismatch or double mismatches could be introduced into the neck of hairpin DNA (3′ -terminal) for realizing enough stability and a strong driving force, and the suitable mismatches are listed in Tables S1 and S2.† Subsequently, to verify the high efficiency of the mismatch-fueled catalytic multiple-arm DNA junction assembly (M-CMDJA), as displayed in Scheme 2 (part A), we produced a series of CMDJAs consisting of substrate hairpin DNAs with a shorter neck (A), wild hairpin DNAs (B), hairpin DNAs with one mismatch (B-M1), and hairpin DNAs with two mismatches (B-M2), respectively. And the schematic illustrations of a further two typical CMDJAs (two-arm and three-arm) are also shown in part B and part C, separately.

Efficiency comparison of annealing and self-assembly in the different M-CMDJA and CMDJAs

We first compared the yield of different multiple-arm DNA junctions by annealing and self-assembly respectively through PAGE experiments. As shown by the results in Fig. 1, with the introduction of mismatches into the hairpin DNA substrates, the corresponding yields of the multiple-arm DNA junction formed by either self-assembly (lanes 1–4) or annealing (lanes 5 and 6) were all dramatically improved (two-arm, Fig. 1A and B;

Scheme 2 Schematic illustration of (part A) the different CMDJAs consisting of: (a) hairpin DNA with a shorter neck (A), (b) wild hairpin DNA (B), (c) hairpin DNA with one mismatched base (B-M1), (d) hairpin DNA with two mismatched bases (B-M2), (part B) the catalytic two-arm DNA junction assembly (CTDJA, two-arm), and (part C) the catalytic three-arm DNA junction assembly (CTDJA, three-arm).

three-arm, Fig. 1C and D; four-arm, Fig. 1E and F), demonstrating that the introduction of mismatches into hairpin DNA substrates can not only enhance the thermodynamic tendency of the self-assembly pathway, but also solve the problem of the free-energy landscapes with kinetic traps²⁹ that cannot be addressed via conventional annealing methods.

Kinetic performance comparison of the M-CMDJAs and CMDJAs under different conditions

To study the kinetic performance of the developed M-CMDJA accurately, we adopted real-time fluorescence analysis to characterize the corresponding reaction progress of the M-CMDJA and the CMDJA (Fig. 2). Through the fluorophore (FAM) and quencher (BHQ1) labeled on the ends of the hairpin DNA substrate (Table $S1\dagger$), the corresponding fluorescence response increased as the multiple-arm DNA junction was formed. As shown by the results in Fig. 2 (Fig. 2A, two-arm; Fig. 2C, threearm; Fig. 2E, four-arm), in the presence of miRNA-182-5P, the fluorescence response of the M-CMDJA (Scheme 2, part A, B-M1 and B-M2) increased much faster to a higher value beyond those of the wild CMDJA (Scheme 2, part A, A and B), manifesting the obviously improved kinetic and thermodynamical performance of the M-CMDJA. In order to obtain a comparison of the accurate kinetic parameters of these control groups, the reaction rate was obtained based on the value of fluorescence response intensity versus the time. The results in Fig. 2B (two-arm), Fig. 2D (three-arm), and Fig. 2F (four-arm) show that not only were the initial reaction rates of the M-CMDJA obviously faster

Fig. 1 Non-denaturing PAGE characterization of different M-CMDJAs and CMDJAs: (A) two-arm, (C) three-arm, and (E) four-arm (lane 1, B; lane 2, B-M1; lane 3, B-M2; lane 4, A (lane 1–4: self-assembly with 100 nM target miRNA-182-5P); lane 5, B; lane 6, B-M1; lane 7, B-M2, lane 8, A (lane 5–8: annealing)), and the corresponding gray scale intensity of (B) two-arm, (D) three-arm, and (F) four-arm (a, lane 1 and lane 5; b, lane 2 and lane 6; c, lane 3 and lane 7; d, lane 4 and lane 8) (all the hairpin DNA substrates were 1μ M, PAGE 8%, 50 mA, 25 min).

than those of the wild CMDJA (Table 1), but the instantaneous reaction rates of the M-CMDJA in the reaction process were also improved to some extent. Subsequently, through the relationship of fluorescence response and fluorophore concentration depicted in Fig. $57, \dagger$ the specific average reaction rates (the ratio between the resultant concentration and the reaction time) of the M-CMDJA were all faster than those of the wild CMDJA (Table 1) (the concentration of multiple-arm DNA junction assembled after 5200 s was computed based on Fig. 2). Impressively, compared with other comparison groups (twoarm DNA junction and three-arm DNA junction), the mismatch-fueled catalytic four-arm DNA junction assembly (M-CFDJA) with two mismatched bases possesses the best kinetic performance (Table 1, initial reaction rate: 274.31 a.u. s^{-1} , average reaction rate: 1.97×10^{-10} M s⁻¹). To explain this phenomenon, based on the conclusion that a stronger thermodynamic driving force from a more negative ΔG allows acceleration of the TSDR verified by previous works, $30-32$ we speculate two reasons as follows: (1) though the four-arm system requires more reaction steps that could dramatically decrease the reaction rate, its much stronger thermodynamic driving force beyond those of the two-arm system and three-arm system (Table S3†) offsets the kinetic decay, resulting in a just slightly faster reaction of the four-arm system (Table 1); (2) the

Fig. 2 Real-time fluorescence responses of different M-CMDJAs and CMDJAs ((A) two-arm, (C) three-arm, and (E) four-arm) with and without the target miRNA-182-5P (1 nM) and the reaction rate of different M-CMDJA and CMDJAs ((B) two-arm, (D) three-arm, and (F) four-arm) in response to 1 nM target miRNA-182-5P (the meanings of A, B, B-M1, and B-M2 are shown in Scheme 2; EX WL: 492 nm, EM WL: 525 nm, gain: 75, time: 5200 s, interval: 15 s).

target miRNA-182-5p as catalyst in the four-arm system could be recycled more easily to further accelerate the reaction of the four-arm system.

Moreover, it is worth mentioning that the CMDJA not only hardly exhibited improved kinetic performance but also suffered from an extremely obvious background leakage when reducing the matched bases in the neck of the substrate hairpin DNAs (Scheme 2, part A, A) to make the ΔG more negative for obtaining a stronger driving force, which was also verified by the PAGE experiments shown in Fig. S6,† further demonstrating the superiority of introducing suitable mismatches in the M-

CMDJA that can successfully improve the performance and decrease the background leakage simultaneously.

Conversion efficiency comparison of the M-CMDJAs and CMDJAs using fluorescence

To further explore the conversion efficiency of the M-CMDJA and wild CMDJA, the relationships between fluorescence response value and concentration of the initiator miRNA-182-5P were studied respectively, and the results are shown in Fig. 3 (Fig. 3A–D, two-arm; Fig. 3E–H, three-arm; Fig. 3I–L, four-arm). When the concentration of miRNA-182-5P increased, the fluorescence intensity of these comparison groups all increased obviously and showed good linear relationships with the logarithm concentration. Among these results, the M-CFDJA showed the most sensitive response to miRNA-182-5P with the regression equation for the value of fluorescence response ν s. c of Fi = 51389.09 + 3588.69 lg c (from 100 fM to 10 nM, $R =$ -0.9977) (Fi: the fluorescence intensity, c: the miRNA concentration). Then, the accurate conversion efficiency (N) at different concentrations of initiator miRNA-182-5P in fluorescence was calculated (Table S4†) based on Fig. S7.† Impressively, the M-CFDJA obtained the highest conversion efficiency $(N_{\text{M-CFDIA-F}};$ 1.96×10^6), almost five times that of the wild CFDJA ($N_{\text{CFDJA-F}}$) $4.15\times 10^5)$ and at least four times that of the other comparison groups (two-arm and three-arm, Table 2), suggesting the outstanding thermodynamic performance of the M-CFDJA and that mismatch introduction could improve the thermodynamic performance of the CMDJA. These results reported above demonstrate that the developed M-CMDJA indeed excelled either with a faster reaction rate or a higher conversion efficiency compared to the wild CMDJA, precisely due to the introduction of mismatches in the substrate hairpin DNA that could obviously improve the driving force, promoting the whole hybridization reaction. Edge Article. Computer article. Published on 26 September 2022. Downloaded the Creative Common and Common and Creative Co

Conversion efficiency comparison of the M-CMDJAs and CMDJAs in electrochemistry

In order to apply the M-CMDJA to develop an ultrasensitive electrochemical biosensing platform, we explored the response of a biosensing platform based on different CMDJAs and M-CMDJAs to miRNA-182-5P at different concentrations, respectively (Fig. 4). Next, the corresponding conversion efficiency

Fig. 3 Fluorescence responses of the (A) wild CTDJA (two-arm) to the target miRNA-182-5P ((a) 10 pM, (b) 100 pM, (c) 1 nM, (d) 10 nM, (e) 100 nM, (f) and 500 nM), (C) M-CTDJA (two-arm) with two mismatches to the target miRNA-182-5P ((a) 5 pM, (b) 10 pM, (c) 100 pM, (d) 1 nM, (e) 10 nM, and (f) 100 nM), (E) wild CTDJA (three-arm) to the target miRNA-182-5P ((a) 1 pM, (b) 10 pM, (c) 100 pM, (d) 1 nM, (e) 10 nM, and (f) 100 nM), (G) M-CTDJA (three-arm) with two mismatches to the target miRNA-182-5P ((a) 500 fM, (b) 1 pM, (c) 10 pM, (d) 100 pM, (e) 1 nM, and (f) 10 nM), (I) wild CFDJA (four-arm) to the target miRNA-182-5P ((a) 500 fM, (b) 1 pM, (c) 10 pM, (d) 100 pM, (e) 1 nM, and (f) 10 nM), and (K) M-CFDJA (four-arm) with two mismatches to the target miRNA-182-5P ((a) 100 fM, (b) 1 pM, (c) 10 pM, (d) 100 pM, (e) 1 nM, and (f) 5 nM). Corresponding calibration plot for the fluorescence intensity vs. lg c: (B) wild CTDJA (two-arm), (D) M-CTDJA (two-arm) with two mismatches, (F) wild CTDJA (three-arm), (H) M-CTDJA (three-arm) with two mismatches, (J) wild CFDJA (four-arm), and (L) M-CFDJA (four-arm) with two mismatches (c represents the concentration of miRNA-182-5P, unit of c was M, error bars, SD, $n = 3$; EX WL: 492 nm, EM WL: 520–700 nm, gain: 75).

using the electrochemistry method (Table S5†) was also computed based on the relationship of the electrochemical signal response with the concentration of multiple-arm DNA junction (Fig. S8†). Significantly, the M-CFDJA (four-arm) obtained the highest conversion efficiency $(N_{\text{M-CFDJA-E}}$: $1.05 \times 10^8)$ almost eight times that of the wild CFDJA $(N_{\rm CFDJA\text{-}E};\, 1.38\times 10^7,$ Table 3).

Moreover, the conversion efficiency of the M-CFDJA obtained using the electrochemistry method was nearly two orders of magnitude that obtained using the fluorescence method (N_M) $_{CFDJA-F}$: 1.96 \times 10⁶, Table 2), which was because of the high sensitivity of the electrochemistry method and the probably fewer interferences of the biphasic electrode operation.³³ These results reported above confirm that the M-CMDJA has obviously surpassed the wild CMDJA in both kinetic and thermodynamic performance, enabling it to be further used in electrochemical biosensing platform construction for ultrasensitive miRNA detection.

Table 2 Comparison of the conversion efficiency between the M-CMDJAs and CMDJAs in fluorescence

Method	Wild CMDJA	M-CMDJA
Two-arm	$1.44 - 1.95 \times 10^4$	$8.26 - 4.37 \times 10^4$
Three-arm	$7.69 - 1.96 \times 10^5$	8.79×10^{1} to 5.00×10^{5}
Four-arm	8.08×10^1 to 4.15×10^5	1.92×10^2 to 1.96×10^6

Additionally, we chose some conventional isothermal amplication methods as control groups and compared their corresponding performance. As the results show in Table 4, the M-CFDJA possesses the lowest reaction time, and widest range and highest conversion efficiency even at a moderate temperature, breaking the bottleneck of long reaction times and low conversion efficiencies in conventional NASA.

Application evaluation of the M-CFDJA in cancer biomarker miRNA detection

As a practical application, the M-CFDJA was used as an efficient DNA signal amplifier to develop an ultrasensitive electrochemical biosensing platform for rapid miRNA-182-5P detection. Firstly, the successful construction of the biosensing platform was verified through CV (Fig. S3A and S4†), EIS (Fig. $S3B[†]$), and CC experiments (Fig. $S5[†]$). Then, after the reaction time was optimized (Fig. S9†), the prepared electrochemical biosensing platform displayed a sensitive response to miRNA-182-5P, as shown in Fig. 4K and L, and the corresponding regression equation is $I = 0.3016$ lg $c + 5.3702$ (c_{mixNA} $_{182\text{-}5P}$: 0.5 fM-1 nM, $R = 0.9993$. The corresponding detection limit was as found to be low as 6.11 aM for miRNA-182-5P, which is 88.4-fold lower than that of the electrochemical biosensor constructed with the wild CFDJA (0.54 fM) (Table S6†), showing the excellent sensitivity of the biosensor constructed based on the M-CFDJA. Moreover, as shown in Table 5, the proposed sensing platform showed quite a wide linear range and a much lower detection limit compared to those of other biosensors applied for nucleic acid detection, which was ascribed to the versatile nature of the M-CFDJA with hyper conversion efficiency. Chemical Science

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Moreover, the biosensing platform constructed using the evolved M-CFDJA also showed outstanding reproducibility (Fig. S10A†), selectivity (Fig. S10B†), and stability (Fig. S10C†). Importantly, as shown in Fig. S10D,† the feasibility of the prepared sensing platform for the detection of miRNA-182-5P in practical samples was studied using total RNA extracted from the human cancer cells 22RV1, A549, and HeLa. The obtained results (Fig. $S10D\dagger$) were in line with the previous reports,^{49,50} suggesting the superiority of the M-CFDJA in ultrasensitive biosensor development for rapid miRNA detection from cancer cells.

Total internal reflection fluorescence microscope (TIRFM) imaging of miRNAs in living cancer cells

In order to verify the outstanding performance of the M-CFDJA for imaging of cancer biomarker miRNA in living cells, we then applied the system to image and monitor miRNA-182-5P in living cells using the TIRFM. 22RV1 cells (high expression of miRNA-182-5P), A549 cells (high expression of miRNA-182-5P), and HeLa cells (low expression of miRNA-182-5P) were chosen to investigate the capability of the M-CFDJA-mediated intracellular miRNA-182-5P imaging strategy compared with the wild CFDJA and conventional fluorescence in situ hybridization (FISH) method. As shown by the results in Fig. 5, the M-CFDJA displayed quite a strong fluorescence response in

Fig. 4 SWV signal responses of the biosensor based on the (A) wild CTDJA (two-arm) to the target miRNA-182-5p ((a) 50 fM, (b) 100 fM, (c) 1 pM, (d) 10 pM, (e) 100 pM, and (f) 1 nM), (C) M-CTDJA (two-arm) with two mismatches to the target miRNA-182-5P ((a) 0.6 fM, (b) 2.0 fM, (c) 10 fM, (d) 100 fM, (e) 1.0 pM, (f) 10 pM, and (g) 100 pM), (E) wild CTDJA (three-arm) to the target miRNA-182-5P ((a) 20 fM, (b) 100 fM, (c) 1 pM, (d) 10 pM, (e) 100 pM, (f) 1 nM, and (g) 10 nM), (G) M-CTDJA (three-arm) with two mismatches to the target miRNA-182-5P ((a) 50 fM, (b) 100 fM, (c) 1 pM, (d) 10 pM, (e) 100 pM, (f) 1 nM, and (g) 10 nM), (I) wild CFDJA (four-arm) to the target miRNA-182-5P ((a) 1 fM, (b) 10 fM, (c) 100 fM, (d) 1 pM, (e) 10 pM, (f) 100 pM, and (g) 1 nM), and (K) M-CFDJA (four-arm) with two mismatches to the target miRNA-182-5P ((a) 0.5 fM, (b) 1.0 fM, (c) 10 fM, (d) 100 fM, (e) 1.0 pM, (f) 10 pM, (g) 100 pM, and (h) 1 nM); calibration plot for the peak current vs. lg c based on the (B) wild CTDJA, (D) M-CTDJA (two-arm) with two mismatches, (F) wild CTDJA (three-arm), (H) M-CTDJA (three-arm) with two mismatches, (J) wild CFDJA (four-arm), and (L) M-CFDJA (four-arm) with two mismatches (c represents the concentration of miRNA-182-5P, error bars, SD, $n = 3$; red dashed, slope; green dashed, LOD).

Table 5 Comparison of the biosensor constructed using the M-

Method Mismatched Wild 6.14×10^2 to 1.88×10^6 9.09×10^{1} to 4.75×10^{6} Two-arm 6.65×10^{1} to 1.39×10^{6} 9.04×10^1 to 5.93 \times 10 ⁶ Three-arm 8.38×10^2 to 1.05×10^8 5.78×10^2 to 1.38×10^7 Four-arm		

CFDJA with other methods for miRNA detection

Analytical methods	Linear range	Detection limit	Ref.
Fluorescence	1 pM to 10 nM	0.35 pM	44
Electrochemiluminescence	10 fM to 0.1 nM	6.6 fM	45
Chronocoulometry	2.0 fM to 1.0 nM	2.0 fM	46
Electrochemical	0.14 nM to 10 nM	40 pM	47
Electrochemical	1 pM to 1 nM	10 pM	48
Electrochemical	0.5 fM to 1 nM	6.11 aM	This work

Table 4 Comparison of the proposed M-CFDJA with some conventional isothermal amplification methods

22RV1 cells and A549 cells, whereas nearly no fluorescence response (FAM, green) of the FISH system and a weak fluorescence response of the CMDJA in 22RV1 cells and A549 cells could be observed, respectively. Considering these results, it can be concluded that the M-CFDJA system could convert miRNA in living cells into reaction products with higher concentration in comparison with the traditional FISH method and wild CMDJA method, suggesting that the M-CFDJA can still achieve excellent performance for ultrasensitive miRNA detection in living cells.

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Fig. 5 TIRFM images of miRNA-182-5P in different cells of type 22RV1, A549, and HeLa based on different methods: (A) FISH, (B) CFDJA, and (C) M-CFDJA. The cell nucleus is shown in blue, and the fluorophore FAM corresponding to miRNA-182-5P is in green (concentrations of all the hairpin DNAs were 1μ M, Hoechst was used for nuclear staining, cells were at the same passage number).

Conclusions

In conclusion, unlike the conventional NASA, the mismatchfueled catalytic four-arm DNA junction assembly (M-CFDJA) we proposed realizes an improved driving force by introducing suitable mismatches in reactant DNA, which lead to stronger kinetic and thermodynamic trends realizing fast reaction rates and outstanding conversion efficiency. Based on these advantages, the M-CFDJA is successfully used as an efficient DNA signal amplifier in an electrochemical biosensor for the rapid and highly sensitive detection of miRNA-21 and the highly sensitive intracellular imaging of miRNA, providing an ingenious approach to explore potential strategies for ultimate application in biosensing detection and clinical diagnoses. Moreover, the M-CFDJA provides novel insight into the investigation of high-efficiency nucleic acid signal ampli fication methods for biological research and DNA nanobiotechnology.

In comparison with other catalytic DNA circuit systems, like HCR,⁵¹ CHA,⁵² and EFTRA,⁵³ the M-CFDJA we proposed could facilely generate much more stable and programmable DNA nanostructure units with faster reaction rates and higher amplification efficiency, from which not only a higher sensitivity of a biosensor based on the M-CFDJA can be realized, but also the obtained DNA nanostructure unit possesses significant potential for application in other areas of DNA nanodevices⁵⁴ and biochemical research.² Nevertheless, the M-CFDJA system still suffers from an inherently low driving force, which largely depends on the random collisions in Brownian motion and the difference between the reaction standard free energy before and after reaction³² (in comparison with the enzyme-driven strategies NASBA,⁵⁵ HRCA,⁵⁶ and LAMP⁵⁷), limiting further improvement of its amplification performance. Additionally, when used in live cell analysis, the M-CFDJA system can only be delivered into cells via complicated liposome-based transfection, from

which the limited quantity of transfected probes may restrict the sensitivity.⁵⁸⁻⁶⁰

In order to address these limitations, some possible methods as follows may be adopted in the future: (1) introducing some energy sources as an additional driving force, such as chemical fuels,^{61,62} magnetic fields,⁶³ acoustic fields,⁶⁴ electric fields,⁶⁵ and light,⁶⁶ which may endow the M-CFDJA system with strong thermodynamical and dynamical favorability; (2) choosing alternative measures to avoid the complicated liposome-based transfection step in the delivery of nucleic acid probes into live cells, like constructing functional DNA nanostructures^{67,68} and combining DNA with biocompatible materials,69,70 which have much better cell internalization efficiencies via macropinocytosis and caveolae-mediated endocytosis pathways. In this way, it may be possible to further improve the comprehensive performance of the M-CFDJA for ultimate application in biochemical and medical fields.

Data availability

Research data are not shared.

Author contributions

Xiao-Long Zhang and Sha-Sha Li conceived the project. Xiao-Long Zhang and Sha-Sha Li designed the sequences, performed the experiments and analyzed the data. Ruo Yuan supervised the project. Xiao-Long Zhang wrote the paper. Xiao-Long Zhang, Ruo Yuan, and Ya-Qin Chai commented and revised the paper. Sha-Sha Li, Wei-Wei Liu, and Ling-Qi Kong assisted in revising the paper.

Conflicts of interest

There are no conflicts to declare.

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