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Introduction

DNA nanotechnology has been widely used to accomplish intelligent processes in many fields including synthetic soft materials, bio-computing, sensing and bioimaging, due to their outstanding programmability, high flexibility and diverse structures.1-3 Recently, DNA nanodevices have become a hotspot in exploiting manipulation strategies in bio-processes, such as substrate channeling, cell migration, immune activation and cell identification.⁴⁻⁷ The specificity and well predictability of Watson-Crick-Franklin base pairing provides the fidelity of constructing the responsively artificial DNA nanomachines by catalytic hairpin assembly, proximity-based ligation, rolling circle amplification (RCA), hybridization chain reaction (HCR).8-11 Furthermore, convenient modification on DNA strands with various functional organic molecules greatly expanded their applications to manipulate cascade enzymatic reactions, cell surface receptors assembly and cell-cell interactions.4-6 For example, cholesterol-functionalized DNA

Controllable mitochondrial aggregation and fusion by a programmable DNA binder⁺

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DNA nanodevices have been feasibly applied for various chemo-biological applications, but their functions as precise regulators of intracellular organelles are still limited. Here, we report a synthetic DNA binder that can artificially induce mitochondrial aggregation and fusion in living cells. The rationally designed DNA binder consists of a long DNA chain, which is grafted with multiple mitochondria-targeting modules. Our results indicated that the DNA binder-induced *in situ* self-assembly of mitochondria can be used to successfully repair ROS-stressed neuron cells. Meanwhile, this DNA binder design is highly programmable. Customized molecular switches can be easily implanted to further achieve stimuli-triggered mitochondrial aggregation and fusion inside living cells. We believe this new type of DNA regulator system will become a powerful chemo-biological tool for subcellular manipulation and precision therapy.

nanostructures have been applied for cell surface engineering to realize adaptive cellular interactions.^{12–14} DNA nanomachines demonstrate precise controllability and robust operation with effective biological outputs in extracellular environment.^{15,16} However, a large number of biological processes occurs inside cells, the development of DNA-based manipulation of intracellular network still remains a big challenge.

Mitochondria are intracellular organelles playing a critical role in energy production and proliferation.¹⁷ Under oxidative stress, the dynamic fusion-fission process of mitochondria can be used to regulate metabolism: fission facilitates cell apoptosis, whereas, fusion can mitigate oxidative stress. Specifically, when abnormal level of reactive oxygen species (ROS) prompts the mutation of mitochondria genomes,18 fusion occurs between mitochondria with wild-type DNA and ones with mutant DNA to compensate for the defects.¹⁹ As an important communication process among mitochondria in the cytosol, mitochondrial fusion (mito-fusion) is regulated by membrane-anchored dynamics, of which Fzo1 heptads from two adjacent mitochondria form coiled-coil structures to tether the membrane.20 Abnormal mito-fusion can cause neurodegenerative diseases, such as Charcot-Marie-Tooth 2A and dominant optic atrophy.21 Excessive fission are closely related to tumors.^{22,23} Traditional mito-therapeutic strategy is normally based on modulating the expression of fusion proteins, which requires either delicate gene engineering process or screening of organic molecules.24,25 Recently, Wang and coworkers reported another approach for the direct induction of mitochondrial fusion based on host-guest supramolecular assembly, which coins the artificial manipulation of mito-fusion as an alternative therapeutic strategy.²⁶ Yang and coworkers developed a DNA tetrahedron machine as

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Fig. 1 Schematic illustrations and characterization of DNA binder and its intracellular operation. (A) DNA binder construction and (B) introduction of DNA binder into living cells, as well as the cellular induction of mitochondrial aggregation and fusion. (C) 10% denature polyacrylamide gel analysis of DNA binder self-assembly. Lanes 1–6 represent DNA primer, circular DNA, DNA chains, Linkage DNA, (1.0 μM) DNA binders and DNA ladder marker. (D) AFM image of DNA binders. Scale bar, 600 nm.

polyanionic barriers to successfully modulate mitochondria communications.²⁷ However, designing molecular tools that can achieve controllable assemblies of mitochondria and precise manipulation of the fusion processes remains largely underdeveloped. In this work, a new DNA nanomachine will be introduced to fill this gap.

We designed herein a DNA binder that contains multiple mitochondrial targeting modules for the controllable manipulation of mitochondrial aggregation and fusion in living cells. Such DNA binder-mediated artificial handing of mitochondria can also result in mitigated reactive oxygen species (ROS) levels and cell repairs (Fig. 1A and B). By further taking advantages of the highly programmable DNA architectures, molecular switches could be integrated into the DNA binder to achieve precise 2'-deoxyadenosine-5'-triphosphate (ATP)-controlled mitochondrial fusion. In addition, a variety of stimulitriggered process could also be rationally activated using this approach. We believe the DNA binder described in this study opens a new avenue for effective and easy manipulation of intracellular organelles and related metabolism.

Results and discussions

Fabrication and in vitro performance of DNA binders

To develop mitochondria-targetable DNA binder, a long singlestranded DNA chain was first prepared by a phi29 polymerasecatalyzed RCA reaction, which consists of 48 nt-long repeated sequences containing 24 nt (Table S1[†]) that can be recognized by multiple triphenylphosphonium (TPP)-modified linkage DNAs (TPP-LDNAs) (Fig. 1A and Scheme S1[†]). Here, the TPP cation is used to target mitochondrial and induce mitofusion.28,29 TPP-LDNA was synthesized based on a coppercatalyzed azide/alkyne cycloaddition reaction30 between an azide-modified linkage DNA and an alkynyl-functionalized TPP cation (Fig. S1-S4[†]). Native PAGE (15 wt%) showed complete formation of TPP-LDNA (Fig. S5[†]), indicating the high yield of the cycloaddition reaction (nearly 100%).31 The amounts of phi29 polymerase for DNA chain synthesis were optimized. The agarose gel electrophoresis revealed that changing amounts of phi29 polymerase led to no obvious variation on the length of DNA chains.³² Due to the low yield, there is no obvious DNA chain bands were observed at 0.05 and 0.1 U μL^{-1} of phi29 polymerase. The yield of DNA chains plateaued at 0.2 U μ L⁻¹ of phi29 polymerase by comparing the band intensities of the RCA products. Thus, 0.2 U μ L⁻¹ of phi29 polymerase was chosen as the optimized amount for RCA, and the DNA chain was obtained with major length ranging from 1500 to 2000 bases (Fig. S6A[†]). After hybridization between the DNA chain and TPP-LDNA, and purification by ultrafiltration, only the band of DNA binder that shifted toward higher base of band ruler was observed, indicating the excess TPP-LDNA was removed and the DNA binder was prepared (Fig. S6B[†]). The successful

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preparation of the DNA binder was also confirmed by 10% denature polyacrylamide gel electrophoresis and atomic force microscopy (AFM) (Fig. 1C and D). The average length of the DNA binder was measured by observing thread features in the AFM images,³³ and found to be 500 ± 150 nm (Fig. S7†). Given that 48 nt-long repeated sequences were about 16.3 nm,³⁴ 31 \pm 9 TPP-LDNA repeating units could be counted for each DNA binder. A 93% grafting ratio of TPP-LDNA on the DNA chain was further estimated by fluorescence measurements (Fig. S8†), demonstrating sufficient mito-targeting modules in the DNA binder.

We first wanted to test if the designed DNA binder can be used to induce mitochondrial aggregation and fusion in an extracellular experiment. As shown in Fig. 2A, the addition of either TPP-LDNA alone or a non-TPP-modified DNA chain/ LDNA complex (C-binder) did not change the size or aggregation status of mitochondria. While in contrast, when mitochondria were incubated with the DNA binder (binder), particle stacking with an integrated morphology could be clearly observed, suggesting a DNA binder-induced aggregation and fusion of the mitochondria. A 15-fold enlargement in the area of mitochondrial aggregates was displayed upon the treatment of DNA binder (Fig. 2B), which was in accordance with the dynamic light scattering results (Fig. 2C). The length of DNA binder was optimized³⁵ to 500 nm, and the number of interspersed bases in-between TPP-LDNAs of DNA binder was optimized to 24 nt for the efficient aggregation and fusion of mitochondria (Fig. S9[†]), which 31 of mito-targeting modules is require as the minimum number. The manipulation of extracellular mitochondrial behavior can be further visualized in solution under confocal fluorescence microscopy. The TPP-LDNA and DNA binder were hybridized with FAM-labeled oligonucleotides (FAM-O1 and FAM-O2) respectively. The fluorescence signals of these two complexes localized well with extracted mitochondria in the bright field channel of confocal images, which proved their successfully attachments on the mitochondria (Fig. S10[†]). To investigate the fusion behaviors of mitochondria, Förster resonance energy transfer (FRET) study was carried out by using fluorescent probes N-(7-nitro-2-1,3benzoxadiazol-4-yl)-1,2-dipalmitoyl-sn-glycero-3-



Fig. 2 DNA binder induced mitochondrial aggregation *in vitro*. (A) SEM images of extracted mitochondria (blank) incubated with (1.0 μ M) TPP-LDNA, (1.0 μ M) C-binder or (1.0 μ M) binder for 20 min at room temperature, respectively. Scale bar, 3 μ m. (B) The quantification of mitochondrial area distributions in the experiments as shown in the panel (A) and scored by Image J. Shown are mean \pm SD from 50 mitochondria in each case. ****P* < 0.001 (two-tailed Student's *t*-test). (C) Dynamic light scattering results of extracted mitochondria (blank) incubated with (1.0 μ M) TPP-LDNA, (1.0 μ M) C-binder or (1.0 μ M) binder for 20 min at room temperature, respectively.

phosphoethanolamine (NBD-PE) and N-(lissamine rhodamine B sulfonyl)-1,2-dipalmitoyl-sn-glycero-3-phosphoethanolamine (Rhod-PE) as the donor and acceptor.36 Two sets of mitochondria were respectively stained with NBD-PE and Rhod-PE, followed washing with PBS and mixing them together. Upon addition of 1.0 µM DNA binder into the above mixture, increased particle size was observed in the confocal images, together with FRET signals (red) under the excitation of 488 nm (Fig. S11 and S12A[†]). This result was also verified by the enhanced emission intensity at 580 nm in the fluorescence spectra (Fig. S12B⁺).^{37,38} Further disassembly of DNA binder and aggregated mitochondria via toehold-mediated strand displacement did not change the FRET results, indicating the FRET signal was generated by mitochondrial fusion. On the other hand, reduced FRET efficiency could be achieved by mixing one set of FRET pairs-labeled mitochondria with the unlabeled set39,40 under the treatment of DNA binders (Fig. S13[†]). Besides, colocalization study showed overlapping of the two sets of mitochondria signals⁴¹ were observed with increased particle size (Fig. S14[†]), demonstrating the fusion of mitochondria by DNA binders. All these above results confirmed that the DNA binder could successfully induce mitochondrial fusion *in vitro*. Both mito-targeting TPP moieties and the long DNA chain are essential for the function of DNA binder.

The performance and therapeutic results of DNA binder in living cells

Next, we studied if the DNA binder can be further used to regulate mitochondrial aggregations in living cells. After demonstrating the minimal cell toxicity of the DNA binder (Fig. S15†), these DNA nanodevices were transfected into 4T1 cells by means of liposomes (Fig. S16†). As shown in Fig. 3A, upon the introduction of 1 μ M DNA binders, the mitochondria in the cytosol of 4T1 cells (as stained by the MitoTracker) clearly changed from scattered small spheres (~0.5 μ m diameter) into elongated structures ($\geq 2 \ \mu$ m length). The percentage of cells bearing these elongated mitochondria increased from 5% to 40% (Fig. 3B), the efficacy of mitochondria elongation might be attributed to the combined results of transfection,



Fig. 3 DNA binder induced mitochondrial in living cells. (A) Confocal fluorescence images of 4T1 cells treated without (blank) and with 1.0 μ M TPP-LDNA, C-binder and binder for 24 h at 37 °C, respectively. The nucleus and mitochondria were staining with DAPI (blue) and mitochondria tracker red (red), respectively. Scale bars, 10 μ m. (B) The percentage of cells with elongated mitochondria in the experiments as shown in the panel (A). For each construct, 100 cells were scored in biological triplicate. (C) TEM images of 4T1 cells treated without (blank) and with 1.0 μ M TPP-LDNA, C-binder and binder, respectively. The red arrows represent mitochondria. Scale bar, 1 μ m. (D) The quantification of mitochondria length distributions in the experiments as shown in (C) and analyzed by ImageJ. Shown are mean \pm SD from 10 mitochondria in each case. Shown are mean \pm SD. ****P* < 0.001 (two-tailed Student's *t*-test).

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mitochondria targeting and aggregating processes. As a control, the addition of either TPP-LDNA or C-binder did not change the cellular mitochondrial size or shape (Fig. 3A and B). Transmission electron microscopy images have further evidenced the DNA binder-induced formation of fused mitochondria with increased lengths (Fig. 3C). The mitochondria in 4T1 cells remained individual form before and after treated with TPP-LDNA and C-binder. Upon addition of DNA binder, elongation of mitochondria was observed with average length of 1.8 μ m (Fig. 3D). Thus, the DNA binder can indeed be used to effectively manipulate mito-aggregation and fusion in living cells.

We next asked whether the DNA binder could be potentially used as a therapeutic tool in neurodegenerative diseases, which are known to be closely associated with the abnormal mitochondrial fusion process. Human neuroblastoma SH-SY5Y cells were selected as a model disease cell line. 1-Methyl-4-phenyl-1,2,3,6-tetrahydropyridine (MPTP) was used to interrupt the mitochondrial metabolism of SH-SY5Y cells by increasing the intracellular ROS and Ca²⁺ levels (Fig. 4A, B and S17†).^{42,43} While after the addition of 2–8 μ M DNA binders, the cellular generation of ROS was gradually decreased to that of the blank sample. Meanwhile, an obvious increase in the SH-SY5Y cell viability (from 55 to 87%) was shown after adding the DNA binders (Fig. 4C). However, addition of C-binder construct (non-TPP-modified DNA chain/LDNA complex) in MPTP treated cells showed negligible change on the ROS level and cell viability, which indicated the DNA binder construct (without mito-targeting modules) did not contribute to the oxidative stress mitigation. Similar therapeutical results could be achieved by using glutathione (GSH), an antioxidant, as the positive control. The above therapeutic process was also verified by the intracellular Ca²⁺ levels measurements and propidium iodide staining⁴⁴ experiments (Fig. S17 and S18†).

Confocal fluorescence microscopy was also utilized for the visual evaluation of subcellular status upon different treatment. As shown in Fig. 4D, after adding MPTP, stress-induced mito-fission resulted in obviously more mitochondrial fragmentations as compared to the blank sample.²⁶ While the introduction of DNA binders in these MPTP-pretreated cells can instead result in distinct mitochondrial aggregations and elongations.



Fig. 4 Therapeutic performance of DNA binder in living cells. (A) Schematic illustrations of blank, MPTP, MPTP/C-binder and MPTP/binder. (B) ROS generation and (C) cell viability of different experimental conditions. Blank, SH-SY5Y cells; MPTP, 1 mM MPTP-pretreated SH-SY5Y cells; MPTP/C-binder, 1 mM MPTP-pretreated SH-SY5Y cells with 8 μ M C-binder; MPTP/x-binder, 1 mM MPTP-pretreated SH-SY5Y cells with $x \mu$ M DNA binder, x = 2, 4, 8; GSH, 5 mM GSH with 1 mM MPTP-pretreated SH-SY5Y cells for 24 h at 37 °C. Shown are mean \pm SD (n = 3). **P < 0.01, ***P < 0.001 (two-tailed Student's t-test). (D) Confocal fluorescence images of SH-SY5Ycells (blank), 1 mM MPTP-pretreated SH-SY5Y cells with or without the addition of (1.0 μ M) DNA binders for 24 h at 37 °C. The nucleus and mitochondria were staining with DAPI (blue) and mitochondria tracker red (red), respectively. Scale bars, 10 μ m. (E) The percentage of cells with elongated mitochondria in the experiments as shown in the panel (D). For each construct, 100 cells were scored in biological triplicate.

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The percentage of cells bearing these elongated mitochondria increased from 2.86% to 38.1% upon addition of DNA binders (Fig. 4E). Therefore, the DNA binder could be considered as an artificial cell repair tool for mitochondria-related therapy. The easy modification of oligonucleotides enables the DNA binder system to be functionalized with cell-targeting and stimuli-recognition modules for improved performance.

The controlled manipulation of mitochondria aggregation and fusion by DNA binder in living cells

The DNA binder system was further studied on engineering molecular switch to achieve stimuli-triggered mitochondrial

fusion. Since ATP is overexpressed in cancer cells and can be used as a trigger for controlling anticancer therapy,^{45,46} we chose an ATP-recognizing aptamer as a proof-of-concept (Fig. 5A). This 27 nt-long ATP aptamer was first divided into two parts, named as SA1 and SA2,^{47,48} and labelled with a TAMRA dye and a BHQ2 quencher, respectively (Table S1†). Due to the fluorescence resonance energy transfer,⁴⁹ the fluorescence of TAMRA-SA1 could be efficiently quenched by the nearby BHQ2 upon the formation of an ATP/SA1/SA2 complex (Fig. S19A†). By integrating repeated SA2 sequences into the long RCA chain, a responsive DNA chain (R-DNA chain) was generated (Table S1†). Further fluorescence experiment verified that ATP could be used to selectively induce the self-assembly of R-DNA chain



Fig. 5 ATP-triggered mitochondrial aggregation by programmable DNA binder. (A) Schematic illustrations of ATP-controlled R-binder response for mitochondrial aggregation and fusion. (B) SEM images of extracted mitochondria (blank) or those incubated with the (1.0 μ M) R-binder in the absence (R-binder) or presence of (1.0 μ M) ATP (R-binder/ATP), as well as a non-ATP aptamer-based NR-binder (1.0 μ M) in the presence of (4.0 μ M) ATP (NR-binder/ATP) for 20 min at 37 °C. Scale bar, 3 μ m. (C) The quantification of mitochondrial area distributions in the experiments as shown in the panel (b) and scored by ImageJ. Shown are mean \pm SD from 50 mitochondria in each case. ****P* < 0.001 (two-tailed Student's *t*-test). (D) Confocal fluorescence images of 4T1 cells with (R-binder) or without (blank) incubation with 1.0 μ M R-binder (without oligomycin) for 24 h at 37 °C. The nucleus and mitochondria in the experiments as shown in the panel (b). For each construct, 100 cells were scored in biological triplicate. Shown are mean \pm SD. ****P* < 0.001 (two-tailed Student's *t*-test).

and TAMRA-SA1, and as a result, to develop an ATP-triggered responsive DNA binder (R-binder) (Fig. S19B[†]).

As shown in Fig. 5B, C, and S20,† in the presence of ATP, Rbinder can induce an efficient in vitro aggregation of mitochondria with increased particle sizes and integrated morphology. As a control, R-binder itself (no ATP) resulted in no morphological changes. These results have been further validated in living 4T1 cells, as the formation of elongated mitochondria can be triggered by intracellular ATP (Fig. 5D and E). The efficacy (30.1%) of mitochondria elongation by R-binder might be attributed to the combined results of transfection, in situ self-assembly, mitochondria targeting and aggregating processes. To further confirm that the ATP-triggered mitochondrial aggregation and fusion, we treated the cells with oligomycin, a commonly chemical inhibitor that is known to suppress the expression of ATP in cells.48 As expected, after adding oligomycin, no obviously morphological changes were observed in cells (Fig. S21⁺). As another control, by replacing the SA1 with a non-ATP aptamer sequence (CSA1), no mitochondrial aggregation was observed both in vitro and in living cells (Fig. 5B, C, and S21[†]). Taken together, these results demonstrated that aggregation and fusion of mitochondria were triggered by the ATP. This proof-of-concept could be extended to further control the mitochondria metabolism by programmable DNA binder, such as fusion triggered by damage-related marker cytochrome c⁵⁰ and terminated by stand displacement.⁵¹

Conclusion

In sum, we report here a programmable DNA-based molecular tool for the manipulation of mitochondrial aggregation and fusion in living cells. The designed DNA binder could efficiently gather mitochondria together based on multivalent mitotargeting modules. The DNA binder may potentially even be used as an alternative strategy for cell therapy. By redesigning a stimuli-responsive DNA binder, we demonstrated that ATP could be used as a switch to achieve regulatable mitochondrial aggregation. Further conjugation of cell targeting moieties, such as folate and aptamers, could reduce the nonspecific therapeutical effects of the system.⁵² The highly editable and predictable nature of Watson–Crick–Franklin base pairing endows the DNA binder as a new versatile platform for regulating intracellular metabolism, which also further extended the functions of existing DNA nanodevices.

Methods

Reagents

All the chemicals were purchased from Sigma unless otherwise noted. Commercial reagents were used without further purification. 4',6-Diamidino-2-phenylindole (DAPI) and SYBR gold were purchased from invitrogen (Carlsbad, CA, USA). *N*-(7-Nitro-2-1,3-benzoxadiazol-4-yl)-1,2-dipalmitoyl-*sn*-glycero-3-

phosphoethanolamine (NBD-PE) and *N*-(lissamine rhodamine B sulfonyl)-1,2-dipalmitoyl-*sn*-glycero-3-phosphoethanolamine (Rhod-PE) were obtained from Macklin (Shanghai, China). Mitochondrial extraction kit was obtained from Abbkine

Scientific Co., Ltd (USA). Mitochondrial tracker red (MitoRed) purchased from KeyGEN Biotech Co., Ltd (Nanjing, China). Phi29 DNA polymerase, T4 DNA ligase, exonuclease I, exonuclease III and dNTPs were purchased from New England Biolabs Ltd (USA). DNA oligonucleotides were synthesized and purified by Sangon Biotech Co., Ltd (Shanghai, China). The detailed sequences have been listed in Table S1.† All aqueous solutions were prepared using ultrapure water (18.2 M Ω cm, Milli-Q, Millipore).

Apparatus

The concentrations of nucleic acids were all measured using a NanoDrop one UV-vis spectrophotometer. The gel electrophoresis was performed on a Tanon EPS-300 electrophoresis analyser (Tanon Science & Technology Company, China) and imaged on Bio-rad ChemDoc XRS (Bio-Rad, USA). All the in vitro fluorescence measurements were measured on an F-7000 spectrometer (HITACHI, Japan). All the intracellular images were taken by a Nikon A1& SIM-S&STORM super resolution microscope (Tokyo, Japan). ROS levels and cell viabilities were measured with a Safire microplate analyzer (Molecular Devices, America). AFM imaging was performed under ambient conditions with a dimension icon atomic force microscopy (Bruker, Germany). Dynamic lights scattering (DLS) was observed on a 90 plus/BI-MAS equipment (Brookhaven, USA). Scanning electron microscopy (SEM) images were recorded using a JEOL JSM-7800F field emission scanning electron microscope. MALDI-TOF MS was carried out by ABSCIEX MALDI TOF-TOF 4800 plus.

Synthesis of alkynyl TPP cation (but-3-yn-1yltriphenylphosphonium)

Triphenylphosphine (520 mg, 2 mmol) and 4-bromobutyne (400 mg, 3 mmol) was dissolved in acetonitrile (20 mL). The reaction mixture was heated to 80 °C for 72 h under nitrogen. The solvent was removed at room temperature followed by addition of benzene (80 mL). The resulting mixture was cooled to -20 °C for 0.5 h and the product was filtered off as white solid (512 mg, 82% yield). ¹H NMR (500 MHz, DMSO-*d*6) δ : 8.27–7.24 (m, 15H), 3.93 (dt, J = 13.1, 7.5 Hz, 2H), 3.02 (d, J = 2.7 Hz, 1H), 2.57 (dtd, J = 12.9, 7.6, 2.7 Hz, 2H). ¹³C NMR (126 MHz, DMSO-*d*6) δ : 135.55, 135.52, 134.28, 134.20, 130.75, 130.65, 118.72, 118.04, 81.60, 81.46, 74.47, 20.50, 20.09, 12.51, 12.49. ESI-MS: C₂₂H₂₀P⁺ [M]⁺ calculated, 315.12; found, 315.03. The spectroscopic data agreed with previous reports.⁵³

Synthesis of TPP-LDNA

The TPP-LDNA was synthesized by adding 5 mM (5 μ L) alkynyl TPP cation to 100 μ M (200 μ L) of linkage DNA (or SA1). The mixture was diluted with 250 μ L DMSO before addition of 40 μ L (50 mM) ascorbic acid. The cycloaddition reaction between azide group of linkage DNA (or SA1) and alkynyl group of TPP cation was initiated by adding 10 mM (50 μ L) CuSO₄ stock solution. The reaction was incubated for 24 hours and the resulting TPP-LDNA was then purified by gel filtration.⁵⁴ MALDI-TOF MS: LDNA [M + K]⁺ calculated, 10 944.3; found, 10 951.7.

TPP-LDNA $[M + K]^+$ calculated, 11 259.3; found, 11 266.8. MALDI-TOF MS results confirmed the successful synthesis of TPP-LDNA (Fig. S4⁺).

Preparation of circular DNA (circ-DNA)

For the synthesis of circ-DNA1, 4.2 μ L of 100 μ M phosphorylated linear DNA1 (linear DNA-12 or linear DNA-36) and ligation DNA1 were mixed at a ratio of 1 : 1. After the annealing procedure, 1 μ L of T4 DNA ligase (400 U μ L⁻¹), 2 μ L of 10× T4 DNA buffer and 8.6 μ L of ultrapure water were added for 16 h incubation at 25 °C. Then, the mixture was heated at 65 °C for 10 min to inactivate T4 DNA ligase. After reaction, 4 μ L of exonuclease I (20 U μ L⁻¹) and 4 μ L of exonuclease III (100 U μ L⁻¹) were added, and the mixture was incubated at 37 °C for 8 h to degrade ligation DNA1. Afterward, the mixture was incubated at 80 °C for 15 min to denature the exonuclease I and exonuclease III. The circ-DNA1 (circ-DNA12 or circ-DNA36) was generated and stored at 4 °C before use. The circ-DNA2 was obtained by using phosphorylated linear DNA2 and ligation DNA2 following the same approach.

Preparation of DNA chain

The DNA chain was synthesized by rolling circle amplification (RCA).55 10 μL of 3 μM circ-DNA1 (or circ-DNA2) and 0.5 μL of 100 µM DNA primer1 (or DNA primer2) were mixed and annealed at 95 °C for 4 min, then gradually cooled to 25 °C at a rate of 1 °C min. The mixture was incubated with phi29 DNA polymerase (0.2 U μ L⁻¹), bovine serum albumin (BSA) (0.4 μ g μL^{-1}) and deoxynucleotide triphosphates (dNTPs) (0.1 mM) at 37 °C for 5 min in 150 μ L of 1× Phi29 reaction buffer. The RCA reaction was terminated through heated at 65 °C for 10 min, then purified by 30 kDa MWCO ultrafiltration device (Millipore) to obtain the long DNA chain (R-DNA chain). DNA chain has repeating strand segments with sequences complementary to circ-DNA. The 36 nt and 12 nt interspersed bases DNA chains were respectively obtained by using circ-DNA36 and DNA primer1, and circ-DNA12 and DNA primer12 following the same approach. The concentration of the repeat unit in DNA chain was obtained by measuring ultraviolet absorbance at 260 nm and used as the concentration of DNA chain.⁵⁶ Briefly, DNA chain with n repeats had an approximate extinction coefficient of 471500*n*, where 471 500 L (mole⁻¹ cm⁻¹) is the ε_{260} of linear RCA repeats unit. Therefore, the molar concentration of RCA repeats could be calculated using the absorbance data and 1 cm length, which was used as the concentration of DNA chain for a given experiment.

Preparation of DNA binder

 $25 \ \mu L$ of $10 \ \mu M$ TPP modified linkage DNA (TPP-LDNA) and $300 \ \mu L$ of DNA chain were mixed together for 2 h at $37 \ ^{\circ}C$ to synthesize the DNA binder. The prepared DNA binder was purified by 30 kDa MWCO ultrafiltration device (Millipore) for three times. The concentration of TPP-LDNA on the DNA chain was used as the concentration of DNA binder. Given that the excess amount of TPP-LDNAs were used, all the binding sites of DNA chain were considered to be occupied by TPP-LDNA.

Therefore, the concentration of DNA binder is considered equal to the concentration of RCA repeats. As controls, the Cbinder was obtained by using linkage DNA and DNA chain following the same approach.

To measure the amounts of linkage DNA in each DNA binder, 5-carboxyfluorescein (FAM)-labelled linkage DNA (FAM-LDNA) was used as components to synthesize a fluorescent DNA binder (F-binder). From the fluorescence intensities of FAM in DNA binder and the standard calibration curves for FAM-LDNA, its amounts could be obtained.

The responsive DNA binder (R-binder) that was able to activate with ATP was constructed by using TPP modified SA1 (TPP-SA1) and R-DNA chain. As controls, no responsive DNA binder (NR-binder) that was unable to activate with ATP was constructed by using TPP modified negative control SA1 (TPP-CSA1) and R-DNA chain.

To demonstrate the response of the split ATP aptamer to ATP, 2 μ M tetramethylrhodamine labeled SA1 (TAMRA-SA1) or tetramethylrhodamine labeled CSA1 (TAMRA-CSA1), and 2 μ M black hole quencher (BHQ2) labeled SA2 (BHQ-SA2) with various concentrations of ATP (CTP, GTP or UTP) were used and incubated for 1 hour at 37 °C. Fluorescence measurements were performed with excitation at 550 nm and emission spectra were collected in the range of 570–650 nm. The slit width was set to be 2 nm for both excitation and emission.

To characterize the ATP activated R-binder response, 2 μ M TAMRA-SA1 and 2 μ M R-DNA chain with 4 mM ATP were used and incubated at 37 °C for 1 hour. Afterward, the solution was purified by ultrafiltration (30 000 MW cut off membrane, Millipore) for three times. The fluorescence measurement of concentrate and filtrate at the third ultrafiltration were performed with excitation at 550 nm and emission spectra were collected in the range of 570–650 nm. All the data were plotted using the Origin software.

Polyacrylamide gel electrophoresis analysis

10% native polyacrylamide gel was prepared using 1× TBE buffer. The loading samples were prepared by mixing 7.5 μ L DNA samples and 1.5 μ L 6× loading buffer, and placed for 3 min before injected into polyacrylamide gel. The gel electrophoresis was run at 100 V for 60 min in 1× TBE buffer, stained with 1× SYBR gold, and scanned with a molecular imager Gel Doc XR.

AFM imaging

 $6 \ \mu L$ DNA binder was deposited on freshly cleaved mica surface and incubated for 10 min. After the solvent water was absorbed with filter paper, 20 μL ultrapure water was added to wash the mica for three times. The mica was then dried with a nitrogen flow and scanned in tapping mode.

Cell culture

The SH-SY5Y and 4T1 cell lines (Procell Life Science & Technology, Wuhan, China) were cultured in Dulbecco's modified Eagle's medium (DMEM) supplemented with 10% fetal bovine serum (FBS), 100 μ g mL⁻¹ streptomycin and 100 U per mL penicillin-streptomycin at 37 °C in a humidified incubator

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containing 5% CO_2 and 95% air. Short tandem repeats (STR) profiling and mycoplasma testing were conducted for each cell line before use. Cell numbers were determined with a Petroff-Hausser cell counter (USA).

3-(4,5-Dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide (MTT) assay

MTT assays were performed to investigate DNA binder biocompatibility. SH-SY5Y and 4T1 cell lines were respectively seeded within replicate 96-well plates at a density of 1×10^6 cells per well. Plates were maintained at 37 °C for 24 h. After discarding the medium, the cells were washed twice with PBS and incubated with serial concentrations of the DNA binder transfected with Lipofectamine® 2000 transfection reagent. Cells incubated with only the PBS was served as control. After 24 hour-incubation, the cells were washed twice with PBS buffer and 100 µL of a fresh medium containing 10 µL of MTT (5 mg mL⁻¹) was added and incubated for 4 h. After removing the remaining MTT solution, 100 µL of dimethylsulphoxide was added to dissolve the formazan crystals precipitates. After shaking the plate for 10 min, the optical density at a wavelength of 490 nm was measured with a Safire microplate analyzer.

Measurement of the ROS generation

Human neuroblastoma SH-SY5Y cells were seeded in a 6-well plate at a density of 50 000 cells per well for 24 h at 37 °C. Then, incubated with 1 mM MPTP in the absence and in the presence of 8 μ M C-binder, 2, 4 and 8 μ M of DNA binders transfected with Lipofectamine® 2000 transfection reagent (Thermo Fisher Scientific), as well as 5 mM GSH for another 24 h. After washing with PBS for three times, the cells were stained with 2',7'-dichlorodihydrofluorescein diacetate (DCFH-DA), or 10 μ g mL⁻¹ propidium iodide and 5 mg mL⁻¹ of DAPI, and measured by a Safire microplate Analyzer or Nikon A1& SIM-S&STORM super resolution microscope, respectively.

Measurement of Ca²⁺ levels

Intracellular Ca²⁺ levels were measured according to the instruction of Fluo-4AM calcium kit from Beyotime (Jiangsu, China). SH-SY5Y cell lines were seeded within replicate 96-well plates at a density of 1×10^6 cells per well. Plates were maintained at 37 °C for 24 h. After discarding the medium, the cells were washed twice with PBS and incubated with 1 mM MPTP in the absence and in the presence of 8 μ M C-binder, 2 μ M, 4 μ M or 8 μ M of DNA binders transfected with Lipofectamine 2000 transfection reagent (Thermo Fisher Scientific), or 5 mM GSH for another 24 h. The cells were washed three times and stained with 5 μ M Fluo-4AM at 37 °C in the dark for 30 min. Then, the cells in the wells were washed three times with PBS and the level of fluorescence was acquired at $\lambda_{ex/em} = 488/516$ nm with Safire microplate analyzer (Molecular Devices, America).⁵⁷

Cellular imaging and data analysis

 1×10^4 4T1 cells were seeded in a confocal dish for 24 h at 37 $^\circ$ C, then transfected, respectively, with 1 μM binder, TPP-LDNA,

C-binder, or F-binder using a Lipofectamine® 2000 transfection reagent (Thermo Fisher Scientific). 24 h after transfection, the cells were washed twice with Dulbecco's phosphate-buffered saline (DPBS) and stained with 200 nM of mitochondrial tracker red and 5 mg mL⁻¹ of DAPI for 15 min for imaging.

To demonstrate the potential capability of DNA binder for molecular mitochondrial aggregation/fusion, 1×10^4 human neuroblastoma SH-SY5Ycells were seeded in a confocal dish for 24 h at 37 °C, then incubated with 1 mM MPTP in the absence and in the presence of 1 μ M DNA binder transfected with Lipofectamine® 2000 transfection reagent (Thermo Fisher Scientific) for another 24 h. Afterward, the cells were washed twice with DPBS and stained with 200 nM of mitochondrial tracker red, as well as 5 mg mL⁻¹ of DAPI for 15 min for imaging.

For the ATP controlled mitochondrial aggregation/fusion experiment, 1×10^4 4T1 cells were seeded in a confocal dish for 24 h at 37 °C, then transfected, respectively, with 1 µM TPP-SA1 and R-DNA chain, TPP-CSA1 and R-DNA chain, or TPP-SA1, R-DNA chain and 10 µM oligomycin using a Lipofectamine® 2000 transfection reagent (Thermo Fisher Scientific). After incubation for 24 h, the cells were washed with DPBS and stained with 200 nM of mitochondrial tracker red and 5 mg mL⁻¹ of DAPI for imaging.

All the fluorescence images were collected with Nikon A1& SIM-S&STORM super resolution microscope. FAM, mitochondrial tracker red and DAPI were excited with 488 nm, 640 nm and 405 nm lasers, respectively. A $100 \times$ oil immersion objective was used for imaging cells. Image analysis was performed with a NiS-elements AR analysis software. Data analysis was done using the Origin and GraphPad Prism software.

Author contributions

Longyi Zhu: conceptualization, investigation, writing – original draft. Yiting Shen: investigation. Shengyuan Deng: data curation. Ying Wan: conceptualization, resources. Jun Luo: resources. Yan Su: data curation. Mingxu You: methodology. Chunhai Fan: resources, supervision. Kewei Ren: conceptualization, writing – original draft, supervision.

Conflicts of interest

The authors report no declarations of interest.

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