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The noncovalent dimerization of a G-quadruplex/hemin DNAzyme improves its biocatalytic properties†

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While many protein enzymes exert their functions through multimerization, which improves both selectivity and activity, this has not yet been demonstrated for other naturally occurring catalysts. Here, we report a multimerization effect applied to catalytic DNAs (or DNAzymes) and demonstrate that the enzymatic efficiency of G-quadruplexes (GQs) in interaction with the hemin cofactor is remarkably enhanced by homodimerization. The resulting non-covalent dimeric GQ–DNAzyme system provides hemin with a structurally defined active site in which both the cofactor (hemin) and the oxidant (H$_2$O$_2$) are activated. This new biocatalytic system efficiently performs peroxidase- and peroxygenase-type biotransformations of a broad range of substrates, thus providing new perspectives for biotechnological application of GQs.

Introduction

Besides its function as the repository of genetic information, DNA molecules are increasingly studied for their biocatalytic properties, often referred to as DNAzymes. Catalytic DNA usually comprises a sequence in which a tertiary structure is encoded which, upon interaction with a metal cofactor, accelerates chemical reactions. An interesting tertiary structure is the G-quadruplex (GQ) that folds from guanine-rich sequences through the stacking of G-quartets, i.e., arrays of four guanines associated via Hoogsteen hydrogen bonds. A GQ exhibits enticing catalytic properties notably upon interaction with hemin (Fe(m)-protoporphyrin IX), a well-known cofactor of hemoproteins. The stacking of hemin onto an accessible G-quartet of a GQ provides a catalytic GQ/hemin system that efficiently performs peroxidase- and peroxygenase-type oxidations of different substrates in the presence of an excess of oxidants such as hydrogen peroxide (H$_2$O$_2$). However, despite dozens of examples and applications reported each year, the precise mechanism of these oxidations still remains unclear.

Hemin can interact with both accessible G-quartets of a GQ, referred to as the 3'- and 5'-end, respectively, but is more catalytically competent at the 3'-end. Various catalytic intermediates have been postulated to explain this observation, but a firm demonstration is still expected. Efforts have been invested to optimize the hemin binding site via the modification of the adjacent nucleobases or the surrounding loops of the 3'-end. However, most if not all optimizations have been attempted with monomeric GQs. We previously demonstrated that multimeric GQs (comprising several covalently linked GQ units) display better catalytic performance than the corresponding monomeric GQs. This improvement originates from the formation of a high-activity hemin binding site located at the interface between two consecutive GQ units. In these assemblies, hemin is sandwiched in between the 3'-end of a GQ unit and the 5'-end of the other GQ unit (Fig. 1A). A Schematic representation of dimeric GQ interaction with hemin: (A) a covalent multimeric GQ considered as a 3'-5' heterodimer, GQ; two non-covalent GQ assemblies that interact via their 3'-ends (3'-3' homodimeric GQ, B) and their 5'-ends (5'-5' homodimeric GQ, C). The arrows indicate the strand orientation from the 5'- to 3'-direction; only the structural core of GQs is represented for clarity.
Here, we hypothesize that providing hemin with a binding pocket formed by two 3'-ends can enhance its catalytic proficiency. To do so, we decided not to use covalent multimeric GQs (that require long sequences, i.e., being poorly efficient and having expensive syntheses, and for which only 5'-3' interfaces have been reported) but we designed 3'-3' stacked non-covalent dimeric GQs (along with 5'-5' dimers as controls), resulting from the self-association of two blunt-ended GQs with naked 3'-ends (and 5'-ends, respectively, Fig. 1B and C). In this original biocatalytic system, hemin nestles in between two 3'-ends (Fig. 1B) where it acquires a high catalytic activity that allows the resulting system to perform a broad variety of biotransformations in a highly efficient manner.

Results and discussion

Formation of a non-covalent dimeric GQ–DNAzyme system

An approach to control the dimerization of GQs is to use DNA sequences that fold into tetramolecular GQs (that is, with a single stretch of guanines comprising 3 to 5 guanines) with various 3'- and 5'-ends. The 3'-homodimeric GQ system was built with 5'-TTAG₃₃₃₃ (n = 3–5), displaying a blunt 3'-G-quartet, along with 5'-TTAG₃₃₃₃ and 5'-TTAG₃₃₃₃ (corresponding to the 5'-TTAG₃₃₃₃ sequence with a 3'-phosphorylated end) as controls since the presence of dTs (or phosphate groups) on the 3'-end is likely to preclude GQ dimerization. We experimentally established this by non-denaturing polyacrylamide gel electrophoresis (PAGE) (Fig. 2A and B); in a GQ-promoting K⁺-rich buffer (100 mM KCl), 5'-TTAG₃₃₃₃ was predominantly found as a dimer (64% versus 36% as a monomer), its 3'-homodimeric nature having been previously established by nuclear magnetic resonance (NMR) spectroscopy. Conversely, 5'-TTAG₃₃₃₃ was found as a monomer only, and both 5'-TTAG₃₃₃₃ and 5'-TTAG₃₃₃₃ were found as mixtures of monomeric and dimeric forms. We confirmed this by NMR for 5'-TTAG₃₃₃₃, 5'-TTAG₃₃₃₃ and 5'-TTAG₃₃₃₃ (Fig. S1†). The 5'-homodimeric GQ system was built using 5'-G(T₃₃₃₃)₃₃₃₃, 5'-T(G₃₃₃₃)₃₃₃₃ and 5'-T(G₃₃₃₃)₃₃₃₃. These sequences were selected since it was previously demonstrated that 5'-G(T₃₃₃₃)₃₃₃₃ self-assembles into a 5'-homodimer (PDB 2L66), 5'-T(G₃₃₃₃)₃₃₃₃ exists as a monomer only (PDB 2L7K) and 5'-T(T₃₃₃₃)₃₃₃₃ is a mixture of monomeric and dimeric forms. To further demonstrate the generality of this approach, other sequences were studied (a total of 23 sequences, Table S1†), whose GQ folding ability was assessed by circular dichroism (CD, Fig. S2†) and dimerization was assessed by PAGE (Fig. S3†).

Next, the influence of the presence of hemin was studied by both PAGE and CD (Fig. 2B and S2†); hemin was found not to disturb GQ dimerization and was even able to promote and stabilize the GQ dimer. As an example, the proportion of the dimeric form of 5'-TTAG₃₃₃₃ increases from 64 to 76% (Fig. 2B) and the melting temperature (assessed by CD-melting experiment) increases by 13 °C (Fig. 2C) after the addition of hemin. This strong GQ dimer/hemin interaction thus opened the way towards the use of these new catalytic systems for performing peroxidation-like reactions. The oxidation of 2,2’-azino-bis[3-ethylbenzothiazoline-6-sulfonic acid] (ABTS) by H₂O₂ was used as a model reaction, given that it is a firmly established model reaction and it provides a readily monitorable output (appearance of a characteristic absorbance at λ = 414 nm as a function of the time) promoted by H₂O₂ (0.6 mM) in the presence of GQs (0.25 mM) and hemin (0.5 mM) in 40 mM HEPEs buffer (pH 7.5) or 100 mM KCl and 0.05% (v/v) Triton X-100 (Fig. S4†). (B) Catalytic oxidation of ABTS (2 mM, absorbance at λ = 414 nm as a function of the time) promoted by H₂O₂ (0.6 mM) in the presence of GQs (0.25 mM) and hemin (0.5 mM) in 40 mM HEPEs buffer (pH 7.5), optimized by pH-dependent catalytic reactions, Fig. S6† with 100 mM KCl and 0.05% (v/v) Triton X-100, at 25 °C. The initial reaction velocities (Vₒ) of oxidations catalyzed by 5'-TTAG₃₃₃₃, 5'-TTAG₃₃₃₃, 5'-TTAG₃₃₃₃ and 5'-TTAG₃₃₃₃ are Vₒ = 11 054, 5333, 1624 and 1054 nM s⁻¹, respectively.
as nucleobases that disrupt GQ dimerization: again the presence of DAs had drastic consequences on the catalytic activity of the 3'-homodimers, with 5'-TTAG\_3\_T\_T\_T\_G\_T\_G being far more efficient than 5'-TTAG\_A\_T\_T\_T\_G\_T\_G and 5'-TTAG\_A\_A\_T\_T\_T\_G\_T\_G (V\_0 = 288, 47 and 18 nM s\(^{-1}\), respectively, Fig. 3B), while it barely affected those of the 5'-homodimers (V\_0 = 140, 185 and 187 nM s\(^{-1}\) for 5'-(G\_T\_T\_T\_G\_T\_G\_G\_T, 5'-A(G\_T)\_G\_T\_T\_G\_T\_G and 5'-AA(G\_T)\_G\_T\_T\_T\_G\_T\_G, respectively, Fig. 3C).

**Potassium promotes GQ dimerization and DNAzyme activity**

We then decided to further investigate GQ dimerization via \(^1\)H-NMR. The analysis of 5'-TTAG\_3\_T\_T\_T\_G\_T\_G was performed in the presence of increasing amounts of KCl (from 0 to 300 mM, Fig. 4A, B and SI\_A\). The \(^1\)H-NMR signals between 10 and 12 (the so called imino protons) are characteristic of the guanines embedded in G-quartets. Our results confirmed first that the GQ was not folded without K\(^+\); they also showed that monomeric GQs are favoured at low-concentration of K\(^+\) and that an increase in K\(^+\) concentration favoured dimeric GQs. This was further demonstrated by size exclusion chromatography (SEC)-HPLC (Fig. 4C\_79 and PAGE experiments (Fig. 4D). A comparative quantitative analysis of these two techniques showed remarkable agreement in the percentage of dimeric GQs promoted by K\(^+\) (Fig. 4E), with >75% of the dimeric form in the presence of 300 mM KCl. We also verified the behaviour of both 5'-TTAG\_T\_T\_T\_G\_T\_G and 5'-TTAG\_T\_T\_T\_G\_T\_G under similar conditions: as expected, the monomeric form was found to be predominant for both sequences even at 300 mM KCl concentration (Fig. S1B and C\_7). This K\(^+\)-promoted dimerization had also strong consequences in terms of biocatalysis. Indeed, varying the KCl concentration from 0 to 500 mM strongly impacted the catalytic performance of 5'-TTAG\_T\_T\_G\_T\_G but not that of 5'-TTAG\_T\_T\_T\_G\_T\_G and 5'-TTAG\_T\_T\_T\_G\_T\_G in the presence of 300 mM KCl, 5'-TTAG\_T\_T\_G\_T\_G was found to be 41- and 10-fold more catalytically active than 5'-TTAG\_T\_T\_T\_G\_T\_G and 5'-TTAG\_T\_T\_T\_G\_T\_G, respectively (Fig. 4F). The straightforward relationship between the V\_0 values and the percentage of dimeric GQs is unambiguously demonstrated in Fig. 4G, without being a simple linear correlation (Fig. 4H). Of note is that we demonstrated the specificity of the K\(^+\) cations by replacing them with Li\(^+\), Na\(^+\), NH\(_4\)^+ or Mg\(^{2+}\), which resulted in a decrease of both GQ dimerization and GQ-DNAzyme activities (Fig. S5\_7). We also performed these experiments at pH 7.5 and 25 °C since these conditions were found to be optimal for GQ-DNAzyme catalysis (Fig. S6\_7).

We finally assessed whether the 5'-TTAG\_T\_T\_G\_T\_G dimer withstood dilution, in order to operate in the low micromolar concentration range routinely used for DNAzyme-type experiments. SEC-
HPLC and PAGE analyses showed that the percentage of the dimeric GQ did not change with dilution (Fig. S7A-C and Table S2†). The optimal concentration for GQ–DNAzyme experiments was thus determined (Fig. S7D†); the catalytic activity was found to increase rapidly with the GQ concentration in the 0–2.5 μM range and to reach a plateau after 5.0 μM. This concentration was thus chosen for subsequent studies.

**Kinetics and scope of non-covalent dimeric GQ–DNAzymes**

The kinetics of GQ–DNAzymes with 3′-homodimers were studied via the measurement of $V_0$ values for reactions performed with increasing $H_2O_2$ concentration (0 to 6 mM). The kinetic curves of 5′-TTAG₃T-3′, 5′-TTAG₃T-3′, 5′-TTAG₃p-3′ and 5′-TTAG₅p-3′ are seen in Fig. 5A and B (those of 5′-TTAG₃T-3′, 5′-TTAG₃T-3′ and 5′-TTAG₃T-3′ as well as 5′-TTAG₅p-3′, 5′-TTAG₅p-3′ and 5′-TTAG₅p-3′ in Fig. S8A†). Michaelis–Menten fitting gave the kinetic parameters $k_{cat}$, $K_m$ and $k_{cat}/K_m$ (summarized in Fig. S8B and Table S3†); the blunt-end dimeric GQs 5′-TTAG₃T-3′ (n = 3, 4 or 5) were found to be far more efficient ($k_{cat}/K_m$ between 1.2 and 1.6 s⁻¹ mM⁻¹, Fig. 5C) than any other sequences, with $k_{cat}/K_m$ between 4.3- and 5.4-fold higher than those of the monomeric GQs 5′-TTAG₃T-3′ (n = 3, 4 or 5). The best results were obtained with 5 μM 5′-TTAG₃T-3′ (denoted by * in Fig. 5C), with $k_{cat} = 8.2$ s⁻¹, $K_m = 3.8$ mM and $k_{cat}/K_m = 2.2$ s⁻¹ mM⁻¹, which represent a 82-fold ($k_{cat}$) and 7.4-fold ($k_{cat}/K_m$) enhancement as compared to those of monomeric 5′-TTAG₃T-3′. These results positioned the 3′-homodimeric system in the upper range of the reported $k_{cat}$ examples [1.1, 4.2, 4.3 and 2.3 s⁻¹ for PS2.2,† c-Myc,† EAD2 (ref. 60) and c-Myc3C-2A,† respectively], but still lower than that of horseradish peroxidase (HRP, 330 s⁻¹), and in the high range of $k_{cat}/K_m$ values (e.g., 2.1 s⁻¹ mM⁻¹ for c-Myc3C-2A), but still far lower than that of HRP (10⁴ s⁻¹ mM⁻¹).†

To further investigate the scope of applications of this new system, we studied the catalytic oxidations of eight different substrates (Fig. 6), *i.e.*, luminol, TMB, D-tyrosine, L-tyrosine, D-serine, L-serine, dopamine and NADH are naturally occurring substrates whose oxidation products are implicated in oxidative stress; second, L-, D-tyrosine, L-dopa, dopamine and NADH are naturally occurring substrates whose oxidation products are implicated in oxidative stress; third, most of these substrates are an illustrative panel of organic contaminants from industrial activities. Oxidative enzymes such as HRP are indeed used for remediation purposes, allowing for the treatment of wastes and contaminated soils, with this enzyme being effective with a wide range of substrates, from organic dyes to phenolic contaminants. This series of dyes (luminol, ABTS and TMB), phenolic (tyrosine) and biphenolic compounds (dopamine) and a thiol derivative (thioanisole) was used to assess the efficiency of both 3′-homodimers (5′-TTAG₃T-3′, 5′-TTAG₃T-3′ and 5′-TTAG₃T-3′) and 5′-homodimers (5′-G₃T₄₃-3′, 5′-G₃T₄₃-3′ and 5′-G₃T₄₃-3′) as tools for degrading environmental pollutants. We found that the 5′-TTAG₃T-3′ dimer strongly accelerated all these oxidations, while the dT-containing GQ and the monomeric GQs only moderately (Fig. S9†). We also checked whether some of these catalysts were asymmetric but no inductions of chirality were observed for L- and D-tyrosine (Fig. S9.3 and S9.4†), and thioanisole (see chiral HPLC traces in the ESI†), which is not unexpected given the lack of a chiral environment provided by the homodimers.

**Model of the non-covalent dimeric GQ–DNAzyme**

The actual structure of the DNAzyme system formed from the dimerization of the GQ formed using 5′-TTAG₃T-3′, and
The dimeric GQ is of high spin and the six-coordination (HS/6C)-type, similar to what was described with PS2-M and MetMb.74,68 The negligible changes of these UV-Vis bands after addition of the monomeric control 5′-TAG3T-3′ and 5′-TAG3-3′ at low potassium concentration demonstrated again the privileged binding of hemin to dimeric GQs (Fig. S11†). The apparent dissociation constant $K_d$ decreased with the steric hindrance on the 3′-end ($K_d = 5.96, 3.91$ and $2.84 \mu M$ for 5′-TTAG3T-3′, 5′-TTAG3-3′ and 5′-TAG3-3′, respectively, Table S4 and Fig. S12†), and with increasing $K^+$ concentrations ($K_d = 5.68, 4.16, 2.84$ and $0.91 \mu M$ for 5′-TTAG3-3′ at $25, 50, 100$ and $300 \text{mM } K^+$, respectively, Table S4 and Fig. S12†), thus further crediting the hypothesis whereby dimeric GQs provide a better binding pocket to hemin than monomeric GQs.

We next measured the formation rate of reactive intermediates (high-valent iron species) via the H$_2$O$_2$-mediated oxidative degradation of GQ/hemin complexes monitored via the time-dependent evolution of the Soret band in the absence of a reducing substrate (Fig. S14†).66 GQ/hemin complexes formed from 10 selected sequences (Fig. 8A) were tested: the initial degradation velocities ($V_0$) determined indicated that complexes from the 5′-TTAG3-3′ and 5′-TTAG5-3′ series were degraded more quickly than the corresponding monomers, with the notable exception of the 5′-TTAG3-3′ series (vide infra). To further illustrate this, we plotted the $V_0$ values versus the $k_{cat}/K_m$ determined above: the representation seen in Fig. 8B shows that the most catalytically competent sequences experience faster degradation. The case of 5′-TTAG3-3′ is interesting since it exhibits a high catalytic activity and a low degradation rate, thereby opening new avenues towards the design of ever more competent and robust GQ–DNAzyme systems.

A mechanism of why and how GQ dimerization enhances the DNAzyme activity of the resulting complexes can be built in agreement and on the basis of previously published models26,19,62,65,66,72,73 (Fig. 9): at the 3′-3′ interface, one of the guanines can flip out from the proximal G-quartet (a ‘wobbling’ guanine)66,19,66,74 to coordinate axially to the iron atom and activate hemin while the distal G-quartet interacts with iron-bound H$_2$O and H$_2$O$_2$. This H-bonding contributes to the heterolysis of O–H via an electron-withdrawing activation. The orchestrated action of both proximal and distal G-quartets...
uniquely provides a finely tuned hemin binding site, close to what is found in hemoproteins, which is more defined than in monomeric GQs in which one activator is the G-quartet and the other one a structurally labile adjacent nucleobase or externally added nucleotides. The presence of two G-quartets uniquely allows for a concerted mechanism (Fig. 9, right red box) that enhances the catalytic efficiency of the resulting complex via a concomitant activation of the cofactor and the oxidant.

Conclusion

We report here a new prototypic GQ–DNAzyme system comprising two GQ units that self-assemble in a non-covalent manner. We demonstrate that dimeric GQs provide a more defined binding pocket to hemin than monomeric GQs, in which all reaction partners (cofactor and oxidant) are concomitantly activated. The sequence used to validate this proof-of-concept derives from the human telomeric sequence 5′-TTAGGG-3′, in which the 5′-end was left naked on purpose, to make the resulting tetramolecular GQ self-assemble to form a 3′-homodimer, in light of the capacity of blunt-ended GQs to interact with each other, drawn in like magnets through hydrophobic, electrostatic and π-stacking interactions. This new system referred to as the non-covalent dimeric GQ–DNAzyme exhibits high catalytic efficiency, with $k_{\text{cat}}$ values up to 8 s$^{-1}$ for the ABTS oxidation. This system contributes to a new strategy to design efficient GQ–DNAzymes, which can be used to catalyse various types of biotransformations: the scope and versatility of this system were demonstrated via a selection of substrates that are representative of organic pollutants of anthropogenic origin. This provides interesting perspectives for the use of DNA catalysts as biosourced tools for green chemistry purposes and potential sustainable solutions to critical environmental concerns.

Conflicts of interest

There are no conflicts to declare.

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