Discovery of a Structural-Element Specific G-Quadruplex ‘‘Light-Up’’ Probe

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The development of a fluorescent probe capable of detecting and distinguishing the wide diversity of G-quadruplex structures is particularly challenging. Herein, we report a novel BODIPY-based fluorescent sensor (GQR) that shows unprecedented selectivity to parallel-stranded G-quadruplexes with exposed ends and four medium grooves. Mechanistic studies suggest that GQR associates with G-quadruplex grooves close to the end of the tetrad core, which may explain the dye’s specificity to only a subset of parallel structures. This specific recognition favours the disaggregation of GQR in aqueous solutions thereby recovering the inherent fluorescence of the dye. Due to its unique features, GQR represents a valuable tool for basic biological research and the rapid discovery of novel, specific ligands that target similar structural features of G-quadruplexes.

Nucleic acid sequences rich in guanines (G) have a propensity to arrange into G-quadruplexes (G4). These non-canonical structures have been implicated in transcription regulation and telomere maintenance; hence, they are targeted for cancer treatment and detection. Conversely, engineered G4 oligonucleotides can act as drug delivery vehicles, materials for nanodevices, support catalysts or even drugs for cancer, HIV and other diseases. For these reasons, there is strong impetus to develop probes to improve our understanding of G4 and their ligand binding characteristics.

Studies suggest that G4 structures are highly polymorphic and exhibit distinct differences between each other. These myriad of topologies arises from the combination of several well-defined structural elements such as nucleic acid type (DNA or RNA), molecularity (monomer, dimer or tetramer), strand orientation (parallel, antiparallel or hybrid), loops (orientation, sequence and length), grooves type (narrow, wide and/or medium), end-cap morphology and number of G-tetrads. Efficient probing of these motifs provides opportunities for discrimination and is also a prerequisite to the discovery and study of new G4 specific ligands. We were thus motivated to develop fluorescent probes that are specific for these fundamental elements.

Several fluorescent probes capable of discerning G4 from duplex DNA have been developed. These emissive probes are generally planar structures which achieve selectivity through end-stacking with G-tetrads. In contrast, structural specific G4 probes are considerably rarer. Moreover, to achieve higher specificity in distinguishing different G4s, simultaneous association with multiple structural elements is desired. Reported probes are conceived from ligands which have been found to interact with the grooves of G4 a priori. Therefore, such rational design strategies can have limited efficiency for discovering new G4 probes with novel selectivity. Conversely, employing the diversity-oriented fluorescent library approach (DOFLA) is a more effective strategy; screening of diversity oriented fluorescent libraries (DOFL) can accelerate the identification of fluorescent probes with the desired qualities without former insight on the structural guidelines needed.

Results
Sensor discovery. A collection of 5000 potential fluorescent sensors were thus gathered and an unbiased, high throughput screening was performed to uncover primary hits responding to G4 (Fig. S1). Additional modifications to improve the quantum yield of the hit after binding led to the discovery of a sensitive and highly selective fluorescent sensor for G4 (GQR, Fig. 1). In the presence of 93del – an interlocked, dimeric,
parallel-stranded G4 with 4 medium grooves26–29. GQR displayed up to 30-fold increase in fluorescence at 597 nm and a 12 nm bathochromic shift in emission maximum (Fig. 2). A Job plot analysis of the GQR-93del complex revealed a binding stoichiometry of 1:1 (Fig. S2); the dissociation constant \( K_d \) was thus determined as 25.18 \( \pm \) 0.02 \( \mu \)M (Fig. 2, inset and Table S1).

**Selectivity of GQR.** To investigate the uniqueness of GQR, we examined its selectivity towards various G4 oligonucleotides with different structural elements (Table 1). Like 93del, J19, T95, T95 and T95-2T form parallel-stranded G4 with exposed ends and four medium grooves (Fig. S3)20–29. When incubated with GQR, all four sequences elicited a fluorescence increase from the dye (Fig. 3). Specifically, the enhancement was most pronounced with 93del. In contrast, both c-kit1 and Pu24T are parallel-stranded but they possess additional snapback motifs that cap the G-tetrad core30,31. Interestingly, both sequences returned negligible fluorescence when mixed with GQR (Fig. 3 and Fig. S4). On the contrary, Oxy and HT adopt non-parallel-stranded conformations (Fig. S3)20,21. GQR similarly remained quenched when incubated with either G4 (Fig. 3 and Fig. S4). Likewise, GQR remained virtually non-fluorescent in the presence of other conventional nucleic acids (Fig. 3 and Fig. S4). Previous reports of parallel-stranded selective G4 sensors generally do not display selectivity for additional motifs within the parallel-stranded structures20,22,23. To the best of our knowledge, GQR is the first fluorescent dye that is able to discern different structural elements (Table 1). Like c-kit1 and Pu24T, GQR displayed up to 30-fold increase in fluorescence at 597 nm and a 12 nm bathochromic shift in emission maximum (Fig. 2). A Job plot analysis of the GQR-93del complex revealed a binding stoichiometry of 1:1 (Fig. S2); the dissociation constant \( K_d \) was thus determined as 25.18 \( \pm \) 0.02 \( \mu \)M (Fig. 2, inset and Table S1).

**Mechanism studies.** **Disaggregation-induced emission of GQR.** Following experiments were aimed at understanding the sensing mechanism of GQR. First, we examined the photophysical properties of GQR. GQR exhibited strong fluorescence emission in organic solvents, but only negligible emission in buffer (Table 2, entries 13–16 vs. entry 1). Further measurements of the absorption spectra showed significant peak broadening and a bathochromic shifts in buffer (Fig. 4a). With higher concentrations of GQR, more pronounced red-shifts were observed suggesting that this phenomenon may be a result of dye aggregation (Fig. 5). Transmission electron microscope and dynamic light scattering analysis revealed the existence of GQR-aggregates with sizes related to the dye concentration (Fig. S6 and S7). Together, these results indicate that the low fluorescence of GQR in buffer is a result of aggregation-caused quenching (ACQ)35 thereby affirming the potential of GQR to behave as fluorescent turn-on sensors for G422,36,37. To confirm this potential, the absorption spectra of GQR with 93del were studied. Indeed, when GQR was mixed with increasing concentrations of the G4, the spectral shape gradually shifted towards that similar to organic solvents (disaggregated state) (Fig. 4b).

**Environmental sensitivity of GQR.** To investigate the contribution of environment polarity to the emission of GQR, we examined its emis-

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**Table 1 | Oligonucleotide sequences used in this work**

| Name      | Sequence (5’ → 3’) | Structural Elements                  | Origin                        |
|-----------|--------------------|--------------------------------------|-------------------------------|
| 93del     | GGGG TGGG AGGA GGGT | parallel, interlocked dimeric         | Aptamer, HIV-1 integrase      |
| J19       | GIGT GGGT GGGA GGGT | parallel, dimeric                     | Aptamer, HIV-1 integrase      |
| T95       | GGGT GGGT GGGA GGGA | parallel, dimeric                     | Aptamer, HIV-1 integrase      |
| T95-2T    | TT GGGT GGGT GGGA | parallel, monomeric                   | Aptamer, HIV-1 integrase      |
| c-kit1    | AGGG AAGG CGCT GGGA GGAG GG | snap-back parallel                  | c-kit oncogene promoter       |
| Pu24T     | TGAG GGGT GTGA GGGT GGGA AAGG | snap-back parallel                  | c-myc oncogene promoter       |
| Oxy       | GGGG TTTT GGGG | antiparallel                          | Oxytricha nova telomere       |
| HT        | TG GGTG AGGTA GGTTA GGGA | hybrid                              | Human telomere                |
| dsDNA     | SigmaD8515        | genomic dsDNA                        | Calf thymus                   |
| ssDNA     | SigmaD8899        | genomic ssDNA                        | Calf thymus                   |
| RNA       | SigmaR7250        | single-stranded                      | Calf liver                    |

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Figure 1 | The structures of GQR and its analogues.

Figure 2 | GQR reacts with 93del to give a turn-on fluorescence response. Fluorescence spectra of GQR (10 \( \mu \)M) upon incubation with serial dilutions of 93del (from 0–80 \( \mu \)M) in buffer (20 mM K2HPO4/KH2PO4, 100 mM KCl, pH 7.0). \( \lambda _{ex} \) 360 nm, \( \Phi _{0} \) (without 93del) = 0.014, \( \Phi _{1} \) (in 40 \( \mu \)M 93del) = 0.28. Inset: photographic image of GQR (10 \( \mu \)M) in the presence of 93del (from 0–40 \( \mu \)M) and data plot of fluorescent emission intensity upon addition of 93del (from 0–80 \( \mu \)M). \( F_{0} \) and \( F_{\text{max}} \) are the fluorescent maximum intensities of the GQR in the absence and presence of 93del respectively. \( K_d \) = 25.18 \( \pm \) 0.02 \( \mu \)M (one site specific binding model). Values are represented as means (\( n = 3 \)). Measurements were taken at room temperature (RT).
Figure 3 | Selectivity of GQR to oligonucleotides. (a) Fluorescence titration of GQR with various G-quadruplexes, DNA and RNA. Conditions: GQR (10 μM), nucleic acid (from 0 to 40 μM), buffer (20 mM K2HPO4/KH2PO4, 100 mM KCl, pH 7.0), RT. λex: 360 nm, λem: 600 nm. (b) Photographic image of GQR (10 μM) mixed with various nucleic acids (40 μM). Irradiation with a hand-held UV lamp at 365 nm.

Table 2 | Fluorescence quantum yields (ΦF) of respective probes in various environments

| Entry | Probe | Solvent | Analyte | ΦFb |
|-------|-------|---------|---------|------|
| 1     | GQR   | Buffer  | -       | 0.014|
| 2     | GQR   | Buffer  | 93del   | 0.28 |
| 3     | GQR   | Buffer  | J19     | 0.22 |
| 4     | GQR   | Buffer  | T95     | 0.19 |
| 5     | GQR   | Buffer  | T95-2T  | 0.14 |
| 6     | GQR   | Buffer  | c-kit1  | 0.009|
| 7     | GQR   | Buffer  | Pu24T   | 0.005|
| 8     | GQR   | Buffer  | Oxy     | 0.027|
| 9     | GQR   | Buffer  | HT      | 0.007|
| 10    | GQR   | Buffer  | dsDNA   | 0.011|
| 11    | GQR   | Buffer  | ssDNA   | 0.008|
| 12    | GQR   | Buffer  | RNA     | 0.016|
| 13    | GQR   | DMSO    | -       | 0.55 |
| 14    | GQR   | MeOH    | -       | 0.57 |
| 15    | GQR   | PEG-400 | -       | 0.46 |
| 16    | GQR   | Toluene | -       | 0.30 |
| 17    | GQR   | DMSO    | -       | 0.30 |
| 18    | GQR   | DMSO    | -       | 0.36 |
| 19    | GQR   | DMSO    | -       | 0.39 |
| 20    | GQR   | DMSO    | -       | 0.21 |
| 21    | GQR   | Buffer  | -       | 0.013|
| 22    | GQR   | Buffer  | -       | 0.030|
| 23    | GQR   | Buffer  | -       | 0.33 |
| 24    | GQR   | Buffer  | -       | 0.24 |
| 25    | GQR   | Buffer  | 93del   | 0.20 |
| 26    | GQR   | Buffer  | 93del   | 0.21 |
| 27    | GQR   | Buffer  | 93del   | 0.34 |
| 28    | GQR   | Buffer  | 93del   | 0.23 |

αAnalyte concentration: 40 μM.
βValues are represented as means (n = 3). Measurements were taken at room temperature (RT). Buffer: 20 mM KH2PO4/K2HPO4, 100 mM KCl, pH 7.0.

Discussion

Common BODIPY dyes have a tendency to aggregate which results in self-quenching. This ACQ effect often limits the label-to-analyte ratio and narrows the practical applications. However, BODIPY can recover its high fluorescence in response to a target molecule through a recognition-induced disassembly of the aggregates. In this study, GQR-aggregates exhibit remarkable fluorescence enhancement in the presence of 93del. However, water-soluble analogues show no such response under the same conditions. The discovery of GQR as an effective fluorescent sensor demonstrates that the conventional ACQ problem can, in reality, be a general mechanism for probe development.

The structural polymorphism of G-quadruplexes has been supported by NMR and X-ray crystallography. Nonetheless, a simple method to distinguish different G-quadruplex structures conveniently and sensitively is highly desirable. The use of specific fluorescent probes, such as GQR, is one such strategy. Fluorescent probes can be potentially employed for mapping and tracking G-quadruplexes-based structure motifs and can also be used for rapid discovery of novel, specific anticancer drugs that recognize similar G-quadruplex structural elements to the probe.

In summary, we have described the systematic discovery of a novel fluorescent dye – GQR – which specifically “light-up” when bound to parallel G4 with end exposed medium grooves. Specific recognition and favourable binding to G4 induces the disaggregation of GQR which results in recovery of the inherent fluorescence of the dye. Preliminary mechanistic studies suggest a groove-binding mode close to the end of the quadruplex core, thereby accounting for the
high specificity. Therefore, GQR represents a promising tool for the future discovery of similar G4 groove binding ligands that can be important for disease studies, G4-based drug deliveries and nanotechnology.

Methods

Probe synthesis. Detailed description of the synthesis of each probe can be found in the Supplementary Information. Each step was characterized by high-resolution mass spectra, 1H and 13C NMR.

Nucleic acids. DNA oligonucleotides were synthesized and structurally verified by NMR spectroscopy as previously reported. dsDNA, ssDNA and RNA were purchased from Sigma Aldrich (product number D8515, D8899 and D7250 respectively) and used as such. The nucleic acids were dissolved in buffer (20 mM K2HPO4/KH2PO4, 100 mM KCl, pH 7.0). DNA concentration was expressed in strand molarity using a nearest-neighbor approximation for the absorption coefficients of the unfolded species.

Absorbance. UV/Vis absorption spectra of dyes and nucleic acids in buffer (20 mM K2HPO4/KH2PO4, 100 mM KCl, pH 7.0) were recorded from 200 to 700 nm using a SpectraMax M2 spectrophotometer.

Fluorescence. Fluorescence measurements were carried out on a SpectraMax M2 spectrophotometer in 96-well plates by scanning the emission spectra between 540 and 700 nm ($\lambda_{ex} = 360$ nm). All experiments were repeated three times. Data analysis was performed using Origin 8.0 (OriginLab Corporation, MA).

Quantum yield measurements. Quantum yields were calculated by measuring the integrated emission area of the fluorescent spectra in its respective solvents and comparing to the area measured for Coumarin 1 (reference) (10 $\mu$M), $\Phi_0 = 0.73$ in ethanol ($\eta = 1.36$), $\lambda_{ex} = 360$ nm. Quantum yields were calculated using the equation:

$$\Phi_F^{\text{sample}} = \Phi_F^{\text{reference}} \left( \frac{\text{Abs}^{\text{sample}}}{\text{Abs}^{\text{reference}}} \right)^2 \frac{\Phi_0^{\text{reference}}}{\Phi_0^{\text{sample}}}$$

where $F$ represents the area of fluorescent emission, $\eta$ is the refractive index of the solvent, and Abs is absorbance at the excitation wavelength. Emission was integrated between 540 and 700 nm.

Dissociation constant measurements. The $K_d$ of GQR to the respective G-quadruplexes were analyzed by Origin 8.0 (OriginLab Corporation, MA) using the following equation for a one site specific binding model:

$$y = \frac{y_{max} \times x}{K_d + x}$$

where $y$ represents the fluorescence fold change of GQR, $y_{max}$ the fluorescence fold change of GQR when saturated with G-quadruplex and $x$ the concentration of the G-quadruplex in $\mu$M.

Transmission electron microscope. GQR (10 $\mu$M) was first prepared in buffer and deposited on a thin copper-support film, followed by drying in vacuo. Images of the samples were obtained with JEOL JEM 3010 HRTEM microscope and operated at 100 kV without any contrast agent.

Dynamic light scattering. The dynamic light scattering of different concentration GQR and other compounds were measured at 25 C in buffer using quartz cell. All measurements were performed in triplicate in Zetasizer Nano ZS.
Nuclear magnetic resonance spectroscopy. Titration with GQR was performed on a 600 MHz NMR Bruker spectrometer equipped with a cryoprobe at 25 °C. Oligonucleotides (200 μM) were dissolved in buffer (20 mM K2HPO4/KH2PO4, 70 mM KCl, D2O/H2O (1:9), pH 7.0). The spectra were recorded immediately after each addition of QWR. Water suppression was achieved using excitation sculpting.

Molecular modeling. The coordinates of 93dQ structures were retrieved from the Protein Data Bank (ID code 1Y8D). DNA structures were prepared for docking. The GQR structure was optimized using the Gaussian03 program (B3LYP/6-31G* level). By using Autodock 4.0, docking studies were carried out with the Lamarckian genetic algorithm following the procedure developed for G-quadruplex DNA and ligand docking[1]. Two rounds of simulation were performed. In the first round, simulated annealing was used to find a rough binding mode of GQR with 50 runs while keeping all other parameters at the default values. The search space was subsequently reduced and another 200 runs were conducted to get a more precise result. Following the docking studies of GQR with 93del quadruplex DNA, all the figures were rendered using PyMOL v0.99.

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Acknowledgments
This study was supported by intramural funding and grant 10/1/21/19/656 from the A*STAR (Agency for Science, Technology and Research, Singapore) Biomedical Research Council. L.Z. acknowledges the financial support from Natural Science Foundation of China (No. 11101550) and the Special Financial Grant from China Postdoctoral Science Foundation (No. 2013T60061). J.C.E. and J.Y. designed, synthesized and characterized all dye compounds used in this study. K.K.G. designed and synthesized the diversity-oriented fluorescent libraries. W.J.C. performed the DLS analysis. W.Z. performed the molecular docking studies. All authors contributed to data analysis and manuscript writing.

Additional information
Supplementary information accompanies this article at http://www.nature.com/scientificreports
How to cite this article: Zhang, L.Y. et al. Discovery of a Structural-Element Specific G-Quadruplex "Light-Up" Probe. Sci. Rep. 4, 3776; DOI:10.1038/srep03776 (2014).