Parallel G-quadruplex-mediated protein dimerization and activation

Truong, T. T. T., Cao, C., & Dang, D. T. (2020). Parallel G-quadruplex-mediated protein dimerization and activation. RSC Advances, 10, 29957-29960. https://doi.org/10.1039/d0ra06173e

Published in:
RSC Advances

Document Version:
Publisher's PDF, also known as Version of record

Queen's University Belfast - Research Portal:
Link to publication record in Queen's University Belfast Research Portal

Publisher rights
© 2020 The Royal Society of Chemistry.
This work is made available online in accordance with the publisher's policies. Please refer to any applicable terms of use of the publisher.

General rights
Copyright for the publications made accessible via the Queen's University Belfast Research Portal is retained by the author(s) and/or other copyright owners and it is a condition of accessing these publications that users recognise and abide by the legal requirements associated with these rights.

Take down policy
The Research Portal is Queen's institutional repository that provides access to Queen's research output. Every effort has been made to ensure that content in the Research Portal does not infringe any person's rights, or applicable UK laws. If you discover content in the Research Portal that you believe breaches copyright or violates any law, please contact openaccess@qub.ac.uk.
Parallel G-quadruplex-mediated protein dimerization and activation

Truong, T. T. T., Cao, C., & Dang, D. T. (2020). Parallel G-quadruplex-mediated protein dimerization and activation. RSC Advances, 10, 29957. https://doi.org/10.1039/d0ra06173e

Published in:
RSC Advances

Queen's University Belfast - Research Portal:
Link to publication record in Queen's University Belfast Research Portal

General rights
Copyright for the publications made accessible via the Queen's University Belfast Research Portal is retained by the author(s) and / or other copyright owners and it is a condition of accessing these publications that users recognise and abide by the legal requirements associated with these rights.

Take down policy
The Research Portal is Queen's institutional repository that provides access to Queen's research output. Every effort has been made to ensure that content in the Research Portal does not infringe any person's rights, or applicable UK laws. If you discover content in the Research Portal that you believe breaches copyright or violates any law, please contact openaccess@qub.ac.uk.
Parallel G-quadruplex-mediated protein dimerization and activation†

Tuom Tinh Thi Truong, a Cuong Cao b and Dung Thanh Dang a,c

We studied parallel G4-mediated protein dimerization and activation by incorporating a RHAU peptide with a fluorescent protein FRET pair CFP/YFP and an apoptotic casp9. Occurrence of energy transfer (from donor CFP to acceptor YFP) and enhancement of 60-fold cleavage efficiency of casp9 were observed in the presence of parallel G4, which indicated that parallel G4 can induce dimerization and activation of proteins. This novel approach holds a great promise for studying G4-targeting functional dimeric proteins in cellular biology.

G-quadruplex (G4) is a four-stranded structure formed by G-rich sequences stacking multiple G-tetrads.1–3 The G4 structure is highly polymorphic, and depends on orientation of the strand loops that can adopt parallel and nonparallel structure topologies.4,5 Computational analysis of the human genome identified more than 700 000 sequences that have potential to form G4 structure.6 G4 highly localizes at telomeres and promoter regions of genes.7,8 They have also been found in 5′-UTR of the encoded RNA and the long G-rich RNA transcripts of telomeric DNA.9,10 G4 location is not randomly distributed, correlating with functional genomic regions. The formation of G4 structure correlates with many cellular processes such as replication, transcription, translation and telomere maintenance that regulate cell proliferation.11,12 Therefore, G4 has recently emerged as a potential target for anti-cancer drug design.13–15

Specific recognition and stabilization of G4 by peptide or protein is a promising approach for regulation of various biological processes. In cells, several helicases such as BLM, FANCJ, PIF1 and RHAU were showed to selectively bind and resolve G4 structure.16–19 A G4-specific binding motif was also identified in N-terminus of RHAU (RNA helicase associated with AU rich element) protein. RHAU peptide selectively binds and stabilizes only parallel G4 structure including DNA and RNA G4s.20,21 NMR structure solution of a complex between an 18-residue peptide fragment (RHAU18) consisting a G4-specific binding motif and a parallel DNA G4 has showed that the RHAU18 peptide forms an α-helix that specifically recognizes the G-tetrad platform of G4. Interestingly, the parallel DNA G4 can selectively bind two RHAU peptides at the 3′ end G-tetrads.20 The peptide covers the G-tetrad and clamps the G4 with three-anchor-point electrostatic interactions between negatively charged phosphate groups of the G4 and three positively charged amino acids of the peptide.20,21 Recently, specific recognition of G4 by RHAU peptide has been applied for chemical biology applications, i.e. incorporating the RHAU peptide with a fluorescent protein provided a useful protein probe for distinguishing different G4 topologies.21 Generation of new ribonuclease by incorporating the RHAU peptide with the catalytic domain of RNase H1 that can target G4 and efficiently cleave the single-stranded RNA in a site-specific manner.22 However, G4 has not been applied as a target molecule for dimerization and activation of protein. Herein, we studied on characterization of G4-mediated dimerization by incorporating a RHAU peptide with a fluorescent protein pair: cyan fluorescent protein/yellow fluorescent protein (CFP/YFP) that was physically detected by the fluorescence resonance energy transfer (FRET) technique (Fig. 1). In addition, we also introduced a RHAU peptide in an apoptotic casp9 to study G4-enhanced enzymatic activity. Such approach holds a great promise for inactive monomeric proteins to specifically target G4 and play a function at dimeric form in cellular biology.

Fig. 1 Schematic representation of parallel G4-mediated protein dimerization that can be physically detected by hetero-FRET with excitation at 400 nm and emission at 527 nm.
Protein dimerization is a ubiquitous mechanism to regulate activity of protein in a broad range of cellular processes including receptor clustering, signal transduction, and apoptosis. Therefore, control over protein dimerization is highly desirable. Casp9 is an apoptotic cysteine protease which is primarily present in its inactive monomeric form under normal physiological conditions. It becomes active upon induced dimerization by auxiliary factors and plays a key role in the apoptosis pathway, cleaving proteins at specific aspartate residues. Engineering of the casp9 dimerization interface by using specific mutations could enhance enzymatic activity in cell. Principle limitations of this engineered approach are the unknown effects of the point mutations on the conformation of the active site and lack of controllable reversibility. In addition, introduction of an N-terminal phenylalanine–glycine–glycine (FGG) motif in casp9 allowed cucurbit[8]uril to induce dimerization of casp9, resulting in an enhancement of cleavage enzyme activity. However, poor water solubility and permeability of cucurbit[8]uril limited applications of molecules-induced dimerization and activation of casp9 in cell. Therefore, G4 with high solubility and permeability would be a potential target molecule for inducing dimerization and activation of protein in cellular processes.

It has been evident that a short 16-aa RHAU peptide (aa 53–68) is sufficient for specific recognition of a parallel G4, the length of the RHAU peptides significantly influences binding affinity. Herein, we designed a fluorescent protein FRET pair RHAU–CFP/RHAU–YFP for characterization of G4-mediated protein dimerization by incorporating an engineered RHAU peptide 30-aa (consisting of RHAU specific binding motif) with the CFP and YFP, respectively. In addition, G4-mediated activation of protein was also proofed by incorporating this RHAU peptide with inactive monomeric casp9. DNA sequences coding RHAU–CFP, RHAU–YFP and RHAU–casp9 were confirmed by DNA sequencing. All proteins were expressed in E. coli BL21 (DE3) under regulation of IPTG. Proteins consisting His-tag at N-terminus were purified by the His-tag chromatography column. Subsequently, the purified proteins were evaluated by SDS-PAGE (Fig. S2, ESI†).

FRET technology has emerged as a powerful tool for determining protein–protein interaction and molecules-induced protein dimerization. Physically, FRET involves the excitation of an acceptor molecule by the emission of a donor molecule within a distance range of 1–10 nm. Herein, we used a hetero-FRET system (energy transfer from donor CFP to acceptor YFP) to elucidate G4-mediated dimerization of RHAU–CFP and RHAU–YFP. Addition of parallel G4s (1 μM) to an equimolar mixture of RHAU–CFP and RHAU–YFP (both at 1 μM) resulted in a strong hetero-FRET signal (Fig. 2A and S3, ESI†). The strong FRET is more notable, considering that G4-induced protein heterodimerization in this case is probably accompanied by 50% homodimerization of protein, which does not contribute to the hetero-FRET. The addition of TERRA (parallel RNA G4) and T95-2T (parallel DNA G4) (Table 1) to a solution of RHAU–CFP/RHAU–YFP mixture resulted in an increase of the peak ratio 527 nm/475 nm from 0.55 to 1.68 and 0.55 to 0.8, respectively (Fig. 2B). The different increase of the peak ratio showed energy transfer of CFP/YFP in the system of TERRA-induced protein dimerization was greater than that of T95-2T-induced protein dimerization. That may explain that RHAU binds RNA G4 with a somewhat greater affinity than DNA G4 (ref. 29) or TERRA-induced protein dimerization may be more optimal for position and orientation of protein fluorescence which may impact the FRET signal. In contrast, addition of Htelo2 (nonparallel DNA G4, Table 1) (1 μM) to a mixture of RHAU–CFP and RHAU–YFP (both at 1 μM) did not result in an increase of the peak ratio 527 nm/475 nm (from 0.55 to 0.54) (Fig. 2B and S3, ESI†). These results showed parallel G4s (TERRA and T95-2T) are capable of selectively binding and inducing protein dimerization.

Parallel G4-mediated dimerization of protein approach was applied for dimer-driven activation of casp9 (Fig. 3A). Casp9 exits primarily inactive monomeric form and becomes active dimeric form under biophysical condition, resulting in apoptosis. In the pathway of apoptosis, casp9 catalyzes, amongst others, the activation of caspase3 (casp3) through

![Fig. 2. FRET studies with RHAU–CFP/RHAU–YFP protein pair under the G4s. (A) Representative spectra of a mixture of RHAU–CFP and RHAU–YFP (both at 1 μM) in the absence (blue) and presence (yellow) of TERRA (1 μM). (B) Comparison of the 527 nm/475 nm FRET ratios observed without (blue bars) and with G4s (1 μM) (yellow bars) in mixture of RHAU–CFP and RHAU–YFP (both at 1 μM). The excitation of wavelength was at 400 nm.](image)

Table 1. DNA and RNA sequences used in this study

| Name   | Sequences (5’–3’)     | Structure† |
|--------|-----------------------|------------|
| TERRA  | r(UAGGGUUAGGGUUAGGGUU) | Parallel G4|
| T95-2T | d(TTGGTTGGTTGGTTGGT)  | Parallel G4|
| Htelo2 | d(TAGGGTTAGGGTTAGGGTT) | Non-parallel G4|
Parallel G4-mediated protein dimerization system is applied for activation of inactive monomeric casp-9, an apoptotic enzyme. Results show that enzymatic activity of casp-9 significantly increase in the presence of TERRA (parallel RNA G4). It is notable that high activity of casp-9 depends on (i) dimerization of protein and (ii) rearrangement of active site. Therefore, TERRA can play as a target molecule for inducing both dimerization and rearrangement of the active site of RHau–casp9. Such approach would be used to promote apoptosis of cancer cells by G4-mediated dimerization and activation of casp9. Induction of protein dimerization by G4 is crucial for studies in function of protein and interplay between protein oligomerization state and activation, not only casp-9, but also many other protein homodimerization events.

In conclusion, the results show that parallel G4 can act as a target-inducer of protein dimerization and activation, thereby leading to energy transfer from donor RHau–CFP to acceptor RHau–YFP and optimal protein reorganization for enzymatic activity of RHau–casp9. Specific recognition of parallel G4 by two RHau peptides allows inactive monomeric proteins fusing with RHau peptide to specifically target parallel G4 and play a dimer-driven activation of proteins. We believe that parallel G4-mediated protein dimerization and activation hold great promises for studying not only caspases, but also many other protein homodimerization events such as dimerizing enzymes and membrane receptor proteins in cellular processes.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

This research is funded by Vietnam National Foundation for Science and Technology Development (NAFOSTED) under grant number 108.02-2017.305. We would like to thank Prof. Anh Tuấn Phan (Nanyang Technological University, Singapore) for scientific discussion and kindly giving plasmids.

Notes and references

1 M. Gellert, M. N. Lipsett and D. R. Davies, Proc. Natl. Acad. Sci. U.S.A., 1962, 48, 2013–2018.
2 D. Sen and W. Gilbert, Nature, 1988, 334, 364–366.
3 C. Weldon, I. Behm-Ans Mant, L. H. Hurley, G. A. Burley, C. Branlant, I. C. Eperon and C. Dominguez, Nat. Chem. Biol., 2017, 13, 18–20.
4 S. Burge, G. N. Parkinson, P. Hazel, A. K. Todd and S. Neidle, Nucleic Acids Res., 2006, 34, 5402–5415.
5 D. J. Patel, A. T. Phan and V. Kuryayi, Nucleic Acids Res., 2007, 35, 7429–7455.
6 V. S. Chambers, G. Marsico, J. M. Boutell, M. Di Antonio, G. P. Smith and S. Balasubramanian, Nat. Biotechnol., 2015, 33, 877–881.
7 S. Balasubramanian, L. H. Hurley and S. Neidle, Nat. Rev. Drug Discov., 2011, 10, 261–275.
8. G. Biffi, D. Tannahill, J. McCafferty and S. Balasubramanian, *Nat. Chem.*, 2013, 5, 182–186.
9. J. D. Beaudoin and J. P. Perreault, *Nucleic Acids Res.*, 2010, 38, 7022–7036.
10. A. T. Phan, *FEBS J.*, 2010, 277, 1107–1117.
11. N. Maizels and L. T. Gray, *PLoS Genet.*, 2013, 9(4), e1003468.
12. D. Rhodes and H. J. Lipps, *Nucleic Acids Res.*, 2015, 43, 8627–8637.
13. S. M. Kerwin, B. Mamiya, C. Brian, T. Fletcher, J. T. Kern and P. W. Thomas, *Abstracts of Papers of the American Chemical Society*, 2000, vol. 219, p. U6.
14. H. Y. Han and L. H. Hurley, *Trends Pharmacol. Sci.*, 2000, 21, 136–142.
15. J. L. Mergny and C. Helene, *Nature Medicine*, 1998, 4, 1366–1367.
16. P. Mohaghegh, J. K. Karow, R. M. Brosh Jr, V. A. Bohr and I. D. Hickson, *Nucleic Acids Res.*, 2001, 29, 2843–2849.
17. J. P. Vaughn, S. D. Creacy, E. D. Routh, C. Joyner-Butt, G. S. Jenkins, S. Pauli, Y. Nagamine and S. A. Akman, *J. Biol. Chem.*, 2005, 280, 38117–38120.
18. C. Ribeyre, J. Lopes, J. B. Boule, A. Piazza, A. Guedin, V. A. Zakian, J. L. Mergny and A. Nicolas, *PLoS Genet.*, 2009, 5, e1000475.
19. O. Mendoza, A. Bourdoncele, J. B. Boule, R. M. Brosh Jr and J. L. Mergny, *Nucleic Acids Res.*, 2016, 44, 1989–2006.
20. B. Heddí, V. V. Cheong, H. Martadinata and A. T. Phan, *Proc. Natl. Acad. Sci. U.S.A.*, 2015, 112, 9608–9613.
21. D. T. Dang and A. T. Phan, *ChemBioChem*, 2016, 17, 42–45.
22. D. T. Dang and A. T. Phan, *Sci. Rep.*, 2019, 9, 7432.
23. N. J. Marianayagam, M. Sunde and J. M. Matthews, *Trends Biochem. Sci.*, 2004, 29, 618–625.
24. C. Pop, J. Timmer, S. Sperandio and G. S. Salvesen, *Mol. Cell.*, 2006, 22, 269–275.
25. S. B. Bratton and G. S. Salvesen, *J. Cell Sci.*, 2010, 123, 3209–3214.
26. Y. Chao, E. N. Shiozaki, S. M. Srinivasula, D. J. Rigotti, R. Fairman and Y. Shi, *PLoS Biol.*, 2005, 3, e183.
27. D. T. Dang, H. D. Nguyen, M. Merkx and L. Brunsveld, *Angew. Chem. Int. Ed.*, 2013, 52, 2915–2919.
28. H. D. Nguyen, D. T. Dang, J. L. van Dongen and L. Brunsveld, *Angew. Chem. Int. Ed.*, 2010, 49, 895–898.
29. S. D. Creacy, E. D. Routh, F. Iwamoto, Y. Nagamine, S. A. Akman and J. P. Vaughn, *J. Biol. Chem.*, 2008, 283, 34626–34634.