Sequence analysis

G4Killer web application: a tool to design G-quadruplex mutations

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Abstract

Motivation: G-quadruplexes (G4) are important regulatory non-B DNA structures with therapeutic potential. A tool for rational design of mutations leading to decreased propensity for G4 formation should be useful in studying G4 functions. Although tools exist for G4 prediction, no easily accessible tool for the rational design of G4 mutations has been available.

Results: We developed a web-based tool termed G4Killer that is based on the G4Hunter algorithm. This new tool is a platform-independent and user-friendly application to design mutations crippling G4 propensity in a parsimonious way (i.e., keeping the primary sequence as close as possible to the original one). The tool is integrated into our DNA analyzer server and allows for generating mutated DNA sequences having the desired lowered G4Hunter score with minimal mutation steps.

Availability and implementation: The G4Killer web tool can be accessed at: http://bioinformatics.ibp.cz.

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Supplementary information: Supplementary data are available at Bioinformatics online.

1 Introduction

Recently, an increasing number of articles have pointed to important biological roles of local non-B DNA structures. Several algorithms are accessible to analyze such various local DNA structures as cruciforms, triplexes, and G-quadruplexes (G4). Increasing efforts are being devoted to the study of G4 due to their possible involvement in serious pathologies and great stability under physiological conditions. G4 play important roles in transcription, translation and RNA, genomic stability, telomere biology, protein recognition and replication origin definition.

G4-forming sequences are common in many regulatory regions of the human genome (Huppert and Balasubramanian, 2007). To evaluate the role of a particular G4-forming sequence in the genome, targeted mutations that do not change length and minimally perturb the primary sequence are required. We therefore designed the ‘G4Killer’ web application, a new and easily accessible tool for the evaluation and mutation of G4-forming sequences. Our algorithm allows defining conditions for G4Hunter score requirements and determining the minimal number of point mutations abolishing G4 propensity. This platform-independent tool has an easy-to-use, web-based graphical interface, freely available at http://bioinformatics.ibp.cz.

2 Features

G4Killer is an application with a web interface. A user inserts either a single sequence or multiple sequences (up to 10, with an upper limit for sequence length of 200 bp) separated by a newline character. The next step is to specify a maximum G4Hunter score (G4HS; default value = 1) so that the proposed mutant sequence has a G4HS equal to or lower than this threshold. The result is a sequence or sequences with the minimal number of mutations and lowest G4Hunter score achievable below the threshold. The number of sequences depends on the input. If the input sequence contains multiple G-tracks of identical length and/or longer repeats of Gs, then multiple equivalent results with mutations in different bases but having the same score are possible and displayed. This effect results from the way G4HS is calculated. For example, a GGGG run can be mutated to a single G and a double G by changing either the second or third position to A or T (the resulting GAGG, GTGG, GGAG or GGTT motifs will have the same G4HS). G=C mutations that would further decrease G4HS is not considered here in order to avoid GC-rich strands that may form stable competing hairpins. The user can see all possible solutions in single sequence mode, while only one possible solution is displayed in multiple sequence mode. The sequence with the lowest score is shown; if there are multiple
equivalent sequences, one is selected randomly. For C-rich sequences with negative G4Hunter scores, there is an option to perform calculations with the complementary sequence where the input is transformed into complementary sequence and then standard calculations are performed. The mutations required for reducing the G4Hunter score are highlighted and the calculated G4Hunter score is presented in the Result section. The user also can apply custom rules for mutated bases replacement in the details view (Fig. 1). More examples are shown in Supplementary Material S1.

3 Validation
The algorithm itself produces only candidate sequences with mutated bases that are then evaluated using our rewrite of G4Hunter published and validated earlier (Brázdá et al., 2019). The algorithm is based on a breadth-first search graph algorithm. It takes the original sequence, changes each individual G-base into an A or T and calculates the score of mutated sequences. The result of this step is N copies of the original sequence with one mutated base, where N is the number of G bases. These mutated sequences are stored, and the same approach is repeated for each of them. This is repeated until the score is lower than or equal to the specified target (e.g. 1.0). The set of sequences from the last step is presented as a result. Sequences are ordered by their scores. Although it has greater memory requirements, using a breadth-first search algorithm ensures finding the best solution to the problem with the least number of steps.

4 Conclusions
We developed a web tool, the G4Killer application, for finding the minimal mutation of G4-forming sequences. Our webserver allows quick and platform-independent usage for single and multiple G4-forming sequences, including their visualization. This tool is also integrated into the G4Hunter server application. That means sequences found by the G4Hunter server can be analyzed directly by G4Killer with just one click and no requirement to copy and edit.

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