Quadron: predictor of sequence-driven genomic DNA quadruplex formation propensity

The Quadron methodology, developed via gradient boosting machines trained on the experimental G4-seq data (Balasubramanian Laboratory), is described and validated in detail at http://dx.doi.org/ToBePublished.

Installation

- 1. Install the latest version of R programming language or skip to the next step. Step-wise instructions on R installation and upgrade for Linux and MacOSX can be found through the following links: 1, 2.
- **2.** Launch R from the command line and install the following R packages from within R, including *shiny* (required for the graphical user interface), *doMC*, *foreach* and *itertools* (required for a parallel execution of the program).

\$ R

```
> install.packages("plyr")
> install.packages("data.table")
> install.packages("caret")
> install.packages("shiny")
> install.packages("doMC")
> install.packages("foreach")
> install.packages("itertools")
```

3. Download the Quadron source code from the GitHub repository or its interfacing home page. You can also do that via a Linux/Unix/OSX command line, given that git is installed, by typing the following:

```
$ git clone https://github.com/aleksahak/Quadron
```

The downloaded folder has the following content:

- lib/ the subfolder containing all the source files,
- QuadronGUI/ the subfolder containing the graphical user interface,
- Quadron.R the interfacing R script used to execute Quadron from command line.
- **4.** Install the specific, 0.4-4, version of *xgboost* R library. This specific version is required since there has been a compatibility breach in mapping the model tree structures in subsequent versions of *xgboost*. The easiest way to install this version for R on Linux operating system, with all the necessary compilers available, is through the source code distributed through the native R CRAN archive. To do so, launch R and type the following command:

\$ R

```
# For Linux:
> install.packages(
    "http://cran.r-project.org/src/contrib/Archive/xgboost/xgboost_0.4-4.tar.gz",
    repos=NULL,
    type="source")
```

Theoretically, the above command should work with MacOSX as well, however, some versions of the clang compiler in MacOSX do not support the fopenmp compiler keyword, hence installation from binaries is a safer routh. The pre-compiled binaries of $xgboost_0.4-4$ are distributed with Quadron (also accessible from Revolution Analytics MRAN repository). They are located in the $lib/xgboost_0.4_4$ subdirectory. For the installation, execute the following commands for MacOSX and Windows operating systems respectively (from command prompt, while located in the $lib/xgboost_0.4_4$ subdirectory of the downloaded Quadron folder):

```
# For MacOSX:
$ cd lib/xgboost_0.4_4
$ R CMD INSTALL macosx_bin/xgboost_0.4-4.tgz

# For Windows, first launch R:
$ R

# Next, from within R (for Windows):
> install.packages(
    "windows_bin/xgboost_0.4-4.zip",
    repos=NULL,
    type="binary")
```

5. Finally, you need to bit compile the Quadron package by going into the **lib**/ subfolder and executing **bitcompile.R** code from within R.

```
$ cd lib/
$ R
> source("bitcompile.R")
```

This generates a single file, **Quadron.lib**, which encapsulates the main Quadron code and all its dependencies. At this stage, the subfolder **lib**/ can be safely removed. You might, however, want to copy the **test.fasta** file from inside **lib**/, in order to test the Quadron installation.

At this point, the Quadron installation folder should contain:

- Quadron.lib the bit-compiled Quadron program,
- QuadronGUI/ the subfolder containing the graphical user interface,
- Quadron.R the interfacing R script used to execute Quadron from command line,

and, if the test sequence file is preserved,

• test.fasta - the example DNA sequence fasta file.

Running Quadron from R

Quadron can be executed as an R program, either from within R, or from the Linux/Unix/OSX command line through R CMD BATCH or Rscript execution. The latter two options allow the usage of Quadron from the scripts written via programming languages other than R.

In R, as exemplified in the **Quadron.R** interfacing script, one should load the **Quadron.lib** bit-compiled file, then execute Quadron via the R function *Quadron()*. The latter accepts three arguments:

- FastaFile the relative or absolute path to the fasta file to analyse,
- OutFile the relative or absolute path to the output file to be saved,
- *nCPU* the number of CPUs to be used for the calculation, where the larger values can markedly speed up the PQS mapping stage of Quadron predictions for large genomes.

Alternatively, the **Quadron.R** file can be edited to set the desired arguments, and executed from the command line via R CMD BATCH or Rscript.

Running Quadron from GUI (Graphical User Interface)

Quadron features a browser-based graphical user interface (GUI), written with Shiny that can be executed locally on as many CPUs as desired. To launch the GUI, enter the **QuadronGUI**/ subfolder and double

click on **QuadronGUI** file. If the file fails to open a browser, make sure that the permissions are correctly set for the file (as executable):

```
$ cd QuadronGUI/
$ chmod 700 QuadronGUI # For setting the permissions to open. May require sudo rights.
```

If double clicking on **QuadronGUI** does not launch your browser after the above step, then, most probably, your Rscript utility of R is not installed on the default /usr/bin/Rscript path. To correct the QuadronGUI setup, first find out the installed path for Rscript by typing the following from the command line:

```
$ which Rscript
```

then open the **QuadronGUI** executable file via a usual plain text editor and correct the Rscript path stated at the first line.

Now double click on QuadronGUI. As soon as the browser opens, the rest is self explanatory.

License

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