Documentation for **Hetero2**

*A program to simulate the evolution of DNA under*

*the mixture models of heterogeneity across lineage*

*and heterogeneity across sites*

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JACOBI\_EIGENVALUE.C (http://people.sc.fsu.edu/~jburkardt/c\_src/jacobi\_eigenvalue/jacobi\_eigenvalue.c)

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Credits

This software was developed by the bioinformatics and phylogenomics team in Ecosystem Sciences, The Commonwealth Scientific and Industrial Research Organisation (CSIRO), Canberra, Australia.

**Introduction**

We present a software to simulate the evolution of nucleotide sequences under mixture models of heterogeneity across lineages and heterogeneity across sites. Hetero2 allows users to assign lineage-specific (and also site-specific) differences in the rate matrices used to describe the evolutionary process.

**Installation of the software**

The software was written in C++, and it has been tested under linux and MacOS platform. You need to have C++ compiler installed in the machine in order to compile the source codes. The compilation steps are shown as follows:

$ tar -zxvf Hetero-2.2.tar.gz

$ cd Hetero-2.2

$ make

Then the executable file named *Hetero2* will appear.

**Usage of Hetero2**

Syntax:

./Hetero2 <tree file> <site info file> <param file list> <other options>

./Hetero2 -h

<tree file> : "Tree file" lists the tree of each site category, the

edge lengths and the labels of terminal/internal nodes

<site info file> : "Site info file" lists the detailed information of

each site category, including the site proportion and

the nucleotide distribution at the root

<param file list> : "Param file list" shows the name of the parameter file

of each variant site category

Other options:

-l <sequence length> : The length of sequences to be simulated

(default: 50,000)

-f <output format> : The format of simulated multiple sequence alignment

1 - FASTA format

2 - Sequential PHYLIP format (default)

-o <output prefix> : Prefix for output files

(default: <tree file> w/o .ext)

-h : The help page

The output file will be:

<output prefix>.out, which stores the simulated multiple sequence alignment file. By default, it is in sequential PHYLIP format, and the user can select FASTA as the output format.

**Example files**

The following example files are available for reference:

|  |  |
| --- | --- |
| trees.txt | An example of "tree file" which lists the tree of each site category, the edge lengths and the labels of terminal/internal nodes |
| site\_info\_file.txt | An example of "Site info file" lists the detailed information of each site category, including the site proportion and the nucleotide distribution at the root |
| param\_file\_list.txt | An example of "parameter list file" showing the name of the parameter file of each variant site category |
| parameter\_1.txt  parameter\_2.txt | An example of parameter file showing the detailed parameters of the rate matrix of each edge leading to the corresponding node |

1. **Example tree file – tree.txt**

The “Tree file” should list the tree of each site category, the edge lengths and the labels of the terminal and the internal nodes. Also, the edge lengths represent the average number of substitutions per site.

*Format:*

[name of variant site category] [newick tree format with internal node labels]

Category\_1 (A:0.30941,(B:0.33809,(C:0.29115,(D:0.04607,(E:0.11096,(F:0.06955,(G:0.03861,

H:0.04429)1:0.01701)2:0.02719)3:0.04063)4:0.27458)5:0.02226)6:0.20349);

Category\_2 (A:0.47928,(B:0.06142,(C:0.06739,(D:0.00915,(E:0.00227,(F:0.00033,(G:0.00114,

H:0.00059)1:0.00076)2:0.00018)3:0.00069)4:0.05522)5:0.06595)6:0.09457);

*Note:*

1. “A,B,C,D,E,F,G,H” are terminal nodes and “1,2,3,4,5,6” are internal nodes.
2. All the trees have to be in the same topology, and with the same set of terminal/internal nodes.
3. **Example site info file – site\_info\_file.txt**

The “site info file” should show the detailed information of each site category, including the site proportion and the nucleotide distribution at the root.

*Format:*

[name of site category] [variant/invariant] [proportion] [freq(A)] [freq(C)] [freq(G)] [freq(T)]

Constant\_site invariant 0.48100 0.30606 0.16447 0.14259 0.38688

Category\_1 variant 0.13214 0.51059 0.18069 0.12901 0.17971

Category\_2 variant 0.38686 0.32819 0.26728 0.10713 0.29740

*Note:*

1. The names of the variant categories are same as the names of variant site categories in the tree file.
2. There is at most one invariant category.
3. **Example parameter list file – param\_file\_list.txt**

The "parameter list file" shows the name of the corresponding parameter file for each variant site category.

*Format:*

[name of variant site category] [parameter file name]

Category\_1 parameter\_1.txt

Category\_2 parameter\_2.txt

*Note:* The names of the variant categories are same as the names of variant site categories in the tree file.

1. **Example parameter file – parameter\_1.txt**

The “parameter file” shows the detailed parameters of the rate matrix of each edge leading to the corresponding node.

*Format:*

Node S1 S2 S3 S4 S5 S6 Pi1 Pi2 Pi3 Pi4

A 1.91928 2.13915 1.59805 2.58998 1.50701 1.00000 0.75553 0.12277 0.04531 0.07639

B 1.54547 3.27634 1.64800 4.70045 3.30819 1.00000 0.57448 0.15236 0.16912 0.10404

C 2.67365 4.14835 1.73040 4.36875 2.94216 1.00000 0.63957 0.16518 0.10018 0.09507

D 1.54547 3.27634 1.64800 4.70045 3.30819 1.00000 0.57448 0.15236 0.16912 0.10404

E 2.95455 13.8448 2.56968 7.04199 16.4082 1.00000 0.49983 0.14356 0.17577 0.18084

F 2.95455 13.8448 2.56968 7.04199 16.4082 1.00000 0.49983 0.14356 0.17577 0.18084

G 2.95455 13.8448 2.56968 7.04199 16.4082 1.00000 0.49983 0.14356 0.17577 0.18084

H 2.95455 13.8448 2.56968 7.04199 16.4082 1.00000 0.49983 0.14356 0.17577 0.18084

1 2.95455 13.8448 2.56968 7.04199 16.4082 1.00000 0.49983 0.14356 0.17577 0.18084

2 2.95455 13.8448 2.56968 7.04199 16.4082 1.00000 0.49983 0.14356 0.17577 0.18084

3 2.95455 13.8448 2.56968 7.04199 16.4082 1.00000 0.49983 0.14356 0.17577 0.18084

4 2.67365 4.14835 1.73040 4.36875 2.94216 1.00000 0.63957 0.16518 0.10018 0.09507

5 2.67365 4.14835 1.73040 4.36875 2.94216 1.00000 0.63957 0.16518 0.10018 0.09507

6 1.54547 3.27634 1.64800 4.70045 3.30819 1.00000 0.57448 0.15236 0.16912 0.10404

*Note:*

1. Pi1, Pi2, Pi3, Pi4 are the equilibrium distribution of A, C, G, T respectively. Their sum should equal to 1.
2. The labels of the terminal and the internal nodes have to be same as those in the tree file.
3. The value of S6 does not need to be 1.0, although it is in this example.

**To run *Hetero2* for these example files**

$./Hetero2 trees.txt site\_info\_file.txt param\_file\_list.txt

The simulated multiple sequence alignment would be in the file: "trees.out".

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