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URL: https://github.com/bsjodin/mADMIXTURE

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DESCRIPTION

mADMIXTURE is a bash program for running ADMIXTURE across a range of k and/or over multiple iterations in a single command. It creates a set of organized directories based upon the chosen options (ie, using multiple k or iterations). This still requires ADMIXTURE to be installed, as ADMIXTURE is still doing all the heavy lifting. All arguments that can be passed to the standalone ADMIXTURE program can be supplied through this program as well. Please reference the ADMIXTURE manual for detailed descriptions of these arguments.

If you have any questions/concerns/recommendations for improvement, please contact me at bryson.sjodin@ubc.ca.

BEFORE USE

Before you begin using **mADMIXTURE**, you must first install **ADMIXTURE**, which can be downloaded from the following website:

http://www.genetics.ucla.edu/software/admixture/

Refer to the ADMIXTURE manual for installation instructions.

Please ensure you cite ADMIXTURE in all publications:

Alexander, D. H., Novembre, J., & Lange, K. (2009). Fast model-based estimation of ancestry in unrelated individuals. *Genome Research*, 19(9), 1655–1664.

DOWNLOAD & INSTALLATION

DOWNLOAD:

You can download mADMIXTURE from the GitHub directly at:

https://github.com/bsjodin/mADMIXTURE

Alternatively, you can clone the git repository by entering the following into a new terminal window:

\$ git clone https://github.com/bsjodin/mADMIXTURE

INSTALLATION:

While **mADMIXTURE** does not technically need to be installed, you must make a few modifications to use the program.

First, change the file permissions:

\$ cd mADMIXTURE

\$ chmod 777 mADMIXTURE

Next, choose one of the following options for "installation."

Option 1:

Copy this script into a PATH directory (eg. /usr/local/bin). This an be accomplished in terminal with:

\$ sudo cp mADMIXTURE /path/to/directory/

Option 2:

Alternatively, you can add the directory where this script is saved to your PATH variable in terminal with:

\$ cd ~

Then one of:

\$ nano .bash profile

OR

\$ nano .bashrc

depending on your operating system.

Scroll to the very end of the file and add the line:

export PATH=/path/to/directory:\$PATH

Save and exit using Ctrl+c. Next type into terminal:

\$ source .bash profile

OR

\$ source .bashrc

depending on which file you edited earlier.

If neither of the above options work, you can simply run the program from the command line with:

\$ bash mADMIXTURE ...

USAGE

Run mADMIXTURE from the command line using the following:

\$ madmixture -f input_file.ped -k integer [optional arguments]

mADMIXTURE requires that you specify both an input file (-f flag) as well as a minimum k value (-k flag). The input file should be a PLINK file with extension .ped, and alleles should be coded in the 1-2 format (see **ADMIXTURE** and PLINK manuals for reference).

In addition to the required arguments, **mADMIXTURE** can pass along arguments to any **ADMIXTURE** setting. See below for a complete list of options. For more details on each option, please reference the **ADMIXTURE** manual.

REQUIRED ARGUMENTS:

-f|--file <filename> : Input PLINK file with extension (must be in current working directory).

-k-minK <int> : Minimum k. If -K is not used, this will be the only k tested.

OPTIONAL ARGUMENTS:

-K|--maxK <int> : Maximum k. Optional. This must be larger than -k

-i|--iterations <int> :Number of iterations. If not set, default is 1. -i|--threads <int> :Enables multi-threaded mode use n threads.

-cv <float/int> : Enables cross-validation and sets the threshold. If no value is given with

the -cv flag, threshold defaults to 10.

-B|--bootstraps <int>: Enables bootstrapping and specifies number of replicates.

-S|--supervised : Enables supervised mode. Must include a ".pop" file in current working

directory.

-P|--projection : Enables projection analysis. Must include a reference ".P.in" file in

current working directory.

-s|--seed <int/str> : Sets the random seed. If no value is given with the -s flag, defaults to

shell variable \$RANDOM.

-EM|--algorithm : Use EM algorithm instead of block-relaxation.

-C|--majC <float/int> : Sets the major termination criterion. -c|--minC <float/int> : Sets the minor termination criterion. -a|--accel <str> : Changes the acceleration method used

-e|--epsilon <float> : Enables penalized estimation and sets epsilon (must be less than 1).

-ll--lambda <float> : Enables penalized estimation and sets lambda.

--haploid : Use haploid data (default is diploid).

--help : Displays help message.

--version : Displays version information.

OUTPUTS

If multiple iterations are specified, the results will be split into separate directories labeled "run_1, run_2, etc". If a range of K is specified, the **ADMIXTURE** outputs will be sorted based on their extension to separate directories (ie, .Q files will be moved to the directory Q_outs, .P files to P_outs, and log files to logs). This makes it easier to sort through files when many k or iterations are performed.