A short manual for sNMF (command-line version)

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Please, print this reference manual only if it is necessary.

This short manual aims to help users to run sNMF command-line engine on Mac and Linux.

1 Description

Inference of individual admixture coefficients, which is important for population genetic and association studies, is commonly performed using compute-intensive likelihood algorithms. With the availability of large population genomic data sets, fast versions of likelihood algorithms have attracted considerable attention. Reducing the computational burden of estimation algorithms remains, however, a major challenge. Here, we present a fast and efficient method for estimating individual admixture coefficients based on sparse non-negative matrix factorization algorithms. We implemented our method in the computer program sNMF, and applied it to human and plant genomic data sets. The performances of sNMF were then compared to the likelihood algorithm implemented in the computer program ADMIXTURE. Without loss of accuracy, sNMF computed estimates of admixture coefficients within run-times approximately 10 to 30 times faster than those of ADMIXTURE

Eric Frichot, François Mathieu, Théo Trouillon, Guillaume Bouchard, Olivier François. Fast Inference of Admixture Coefficients Using Sparse Non-negative Matrix Factorization Algorithms, submitted.

2 Installation

To install sNMF CL version, you just have to execute the install script (install.command) in sNMF main directory. To execute it in a terminal shell, go to sNMF main directory and write "./install.command". If the script is not executable, type "chmod +x install.command" and then "./install.command". A set of binaries should be created in sNMF main directory.

3 Data format

Input file is composed a genotype file.

eigenstratgeno (example.eigenstratgeno)

The **genotype file** contains 1 line per SNP. Each line contains 1 character per individual: 0 means zero copies of reference allele. 1 means one copy of reference allele. 2 means two copies of reference allele. 9 means missing data.

Below, an example of genotype file for n=3 individuals and L=5 loci.

140		
112		
040		
010		
201		
091		
112 010 091 121		
121		

There are 2 output files.

• The first file (with extension $.\mathbf{Q}$) contains the individual admixtures coefficients. It is a matrix a matrix of n lines (the number of individuals) and K columns (the number of ancestral populations).

• The second file (with extension .F) contains the ancestral allele frequencies. It is a matrix a matrix of $nc \times L$ lines (the number of alleles times the number of SNPs) and K columns (the number of ancestral populations). For each SNP, the first line contains the ancestral frequencies for allele 0, the second line for allele 1,

4 Run the programs

The program is executed by a command line. The format is:

```
./sNMF -g genotype_file.geno -K number_of_ancestral_populations
```

All the previous options are mandatory. There is no order for the options in the command line. Here is a more precise description of the options:

- -g genotype_file.geno is the path for the genotype file (in .geno format).
- ullet -K number_of_ancestral_populations is the number of K of ancestral populations.

Other options are not mandatory:

- -p p is the number of processes that you choose to use if you run the algorithm in parrallel. Be careful, the number of process has to be lower or equal than the number of physical processes available on your computer (default: 1).
- -i iteration_number is the max number of iterations in algorithm (default: 1000). The algorithm should not go until the max number of iterations. The stopping criteria should depend on the tolerance error only.
- -a alpha is the value of the regularization parameter (by default: 100). Results can depend on the value of this parameter, especially for small data sets (see the associated paper).
- -e tolerance is the tolerance error (by default: 0.0001).
- -s seed is the initialization for the random parameter (by default: random).
- -m ploidy 1 if haploid, 2 if diploid (default: 2).

If you need a summary of the options, you can use the -h option by typing the command line

```
./sNMF -h
```

A full example is available at the end of this note.

5 Cross-Entropy criteria

We also provide two other programs:

• the first program creates a data set with a given percentage of missing data from your original data set. The command line format is:

```
./createDataSet -g genotype_file.geno
```

The mandatory option is:

- -g genotype_file.geno is the path for the genotype file (in .geno format).

It will create by default, a file with around 5 % of missing data with the name genotype_file_**I**.geno with a **I** extension to differentiate this file from the original file.

Other options are not mandatory:

- -r percentage is the percentage of missing data in your data set (default: 0.05).
- -e tolerance is the tolerance error (by default: 0.0001).
- -s seed is the initialization for the random parameter (by default: random).
- -m ploidy is 1 if haploid, 2 if diploid (default: 2).

• the second program calculates the cross-entropy criterion for all data and for missing data from the output of sNMF. The cross-entropy criteria is useful to choose the best run for different number of ancestral populations (K) and different values of the regularization parameter (α) . A smaller value of the cross-entropy with missing data means a better prediction of the data. The command line format is:

```
./crossEntropy -g genotype_file.geno -K number_of_ancestral_populations
```

The mandatory option is:

- -g genotype_file.geno is the path for the genotype file (in .geno format).
- -K number_of_ancestral_populations is the number of K of ancestral populations.

In this case, the output from sNMF, the files with missing data, the original files and the results files have to b in the same directory.

Other options are not mandatory:

- -m ploidy 1 if haploid, 2 if diploid (default: 2).

6 Tutorial

6.1 Data set

The data set that we analyze in this tutorial is an Asian human data set of SNPs data. This data is a worldwide sample of genomic DNA (10757 SNPs) from 934 individuals, taken from the Harvard Human Genome Diversity Project - Centre Etude Polymorphism Humain (Harvard HGDP-CEPH)2. In those data, each marker has been ascertained in samples of Mongolian ancestry (referenced population HGDP01224) [1].

6.2 Create a data set with missing data

In the main directory, type:

```
./createDataSet -g examples/panel11.geno
```

A file with 5 % of missing data with path examples/panel11_I.geno has been created.

6.3 Run sNMF

Then, run sNMF for the data set with 5 % of missing data (with K=5 for example):

```
./sNMF -g examples/panel11_I.geno -K 5
```

The results files examples/panel11_I.Q examples/panel11_I.F have been created.

6.4 Calculate the Cross-Entropy

Finally, calculate the cross-entropy criteria:

```
./crossEntropy -g examples/panel_11.geno -K 5
```

With this procedure, you can calculate a criteria for each of your analyses. It is a way to choose the best run for different number of ancestral populations (K) and different values of the regularization parameter (α) .

7 Contact

If you need assistance, do not hesitate to send me an email (efrichot@gmail.com or eric.frichot@imag.fr). A FAQ (Frequently Asked Questions) section is available on our webpage (ttp://membres-timc.imag.fr/Olivier.Francois/snmf.html). sNMF software is still under development. All your comments and feedbacks are more than welcome.

References

[1] Nick J. Patterson, Priya Moorjani, Yontao Luo, Swapan Mallick, Nadin Rohland, Yiping Zhan, Teri Genschoreck, Teresa Webster, and David Reich. Ancient admixture in human history. *Genetics*, doi:10.1534/genetics.112.145037, 2012.