SeleDiff v1.0 Manual

Xin Huang Friday, March 16, 2018

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SeleDiff is implemented with a probabilistic method for estimating and testing selection (coefficient) differences between populations¹.

If you use SeleDiff, please cite

Huang X, Jin L, He Y. 2018. SeleDiff: A fast and scalable tool for estimating and testing selection differences between populations. *In submission*.

1 The Model

Consider a bi-allelic locus in the population i, let $p_i(t)$ and $q_i(t)$ denote the derived and ancestral allele frequencies at time t, respectively. We can define the absolute fitness of the derived and ancestral alleles as w_D and w_A , respectively. We then define the relative fitness as

$$e^s = \frac{w_D}{w_A}.$$

Here, s is the (allele) selection coefficient. Based on our previous study¹, the selection (coefficient) difference between populations i and j is

$$\begin{aligned} d_{ij} &= s_i - s_j \\ &= \frac{1}{t} \left[\ln \frac{p_i(t)/q_i(t)}{p_j(t)/q_j(t)} + \Omega \right], \\ &= \frac{1}{t} \left[\ln OR + \Omega \right] \end{aligned}$$

where OR stands for odds ratio; Ω approximately follows a normal distribution with a mean of zero, and reflects the uncertainty of allele frequencies caused by factors other than selection; t is the divergence time from populations i and j to their most recent common ancestor. The expectation and variance of d_{ij} is

$$E(d_{ij}) = \frac{1}{t} \ln OR$$
$$\operatorname{var}(d_{ij}) = \frac{1}{t^2} [\operatorname{var}(\ln OR) + \operatorname{var}(\Omega)]$$

Because $\ln OR$ also approximately follows a normal distribution, d_{ij} is approximately normal. Its 95% confidence interval is $E(d_{ij}) \pm 1.96 \sqrt{\text{var}(d_{ij})}$. We proposed a statistic δ for testing selection difference in a locus:

$$\delta = \frac{\left[E\left(d_{ij}\right)\right]^2}{\operatorname{var}\left(d_{ij}\right)}.$$

Under the null hypothesis, $E(d_{ij}) = 0$, thus, δ follows a central χ^2 -distribution with one degree of freedom. Under the alternative hypothesis, δ follows a non-central χ^2 -distribution with non-centrality parameter $E(d_{ij})$ with one degree of freedom. Because

$$\delta = \frac{\left[E\left(d_{ij}\right)\right]^{2}}{\operatorname{var}\left(d_{ij}\right)}$$

$$= \frac{\ln^{2} OR}{\operatorname{var}\left(\ln OR\right) + \operatorname{var}\left(\Omega\right)} \sim \chi_{1}^{2}$$

and the median of χ_1^2 -distribution approximately equals to 0.455. Therefore, given a dataset with n loci, we assume most loci are neutral in both populations i and j, i.e. $s_i = 0 = s_j$, $E(d_{ij}) = 0$. Then we can estimate var (Ω) as

$$\operatorname{var}(\Omega) = \operatorname{median} \left\{ \frac{\ln^2 OR_k}{0.455} - \operatorname{var} \left(\ln OR_k \right), 1 \leq k \leq n \right\}.$$

where

$$\mathrm{var}(\ln OR) \approx \frac{1}{N_i \hat{p}_i(t)} + \frac{1}{N_i \hat{q}_i(t)} + \frac{1}{N_j \hat{p}_j(t)} + \frac{1}{N_j \hat{q}_j(t)}.$$

Here, N_i and N_j are the sample sizes of populations i and j. We add 0.5 to allele counts less than 5 for continuity correction.

2 Usages

2.1 Environment Setting

To use SeleDiff, you should install Java SE Development Kit 8 8 first.

After the installation, you can check Java version in the command line (command starts by ">" prompt).

```
> java -version
java version "1.8.0_25"
Java(TM) SE Runtime Environment (build 1.8.0_25-b17)
Java HotSpot(TM) 64-Bit Server VM (build 25.25-b02, mixed mode)
```

2.2 Installation

To install SeleDiff, you first clone the SeleDiff repository from GitHub.

2.2.1 Linux/Mac

In Linux, you can open the terminal and clone SeleDiff using git.

> git clone https://github.com/xin-huang/SeleDiff

Then you can enter the SeleDiff directory and use gradlew to install SeleDiff.

- > cd ./SeleDiff
- > ./gradlew build
- > ./gradlew install

The runnable SeleDiff is in ./build/install/SeleDiff/bin/. You can add this directory into your system environment variable PATH by

> export PATH="/path/to/SeleDiff/build/install/SeleDiff/bin/": \$PATH

You can get help information by typing

> SeleDiff

and you will get the following:

Usage: SeleDiff [command] [command options]

Commands:

compute-var Sub-command for estimating variances of population

demography parameters

Usage: compute-var [options]

Options:

--geno

The EIGENSTRAT GENO file stores allele counts: 0, zero copy of the reference allele; 1, one copy of the reference allele and one copy of the alternative allele; 2, two copies of the reference allele; 9, missing values.

* --ind

The EIGENSTRAT IND file stores information of individuals and populations.

* --output

The output file.

--snp

The EIGENSTRAT SNP file stores information of variants.

--vcf

The VCF file stores SNP information and genotype data.

compute-diff Sub-command for estimating selection differences of loci Usage: compute-diff [options]

Options:

--geno

The EIGENSTRAT GENO file stores allele counts: 0, zero copy of the reference allele; 1, one copy of the reference allele and one copy of the alternative allele; 2, two copies of the reference allele; 9, missing values.

* --ind

The EIGENSTRAT IND file stores information of individuals and populations.

* --output

The output file.

--snp

The EIGENSTRAT SNP file stores information of variants.

* --time

The file stores divergence times between populations. A divergence time file is space delimited without header, where the first column is the population ID of the first population, the second column is the population ID of the second population, the third column is the divergence time of this population pair. This file is needed when estimating selection differences.

* --var

The file stores variances of Omegas,

which is space delimited without header the first column is the first population ID the second column is the second population ID the third column is the variance of drift of this population pair. This file is needed when estimating selection differences.

--vcf

The VCF file stores SNP information and genotype data.

You can use gradlew to remove SeleDiff.

^{*} indicates required options

> ./gradlew clean

2.2.2 Windows

In Windows, you can download the SeleDiff repository directly from GitHub using the green button Clone or download at the upright corner. Please make sure your environment variable JAVA_HOME correctly point to you JDK directory. After download and uncompression, you can open cmd and enter the directory of SeleDiff in cmd. Please use gradlew.bat to build and install SeleDiff.

```
> cd /path/to/SeleDiff
> gradlew.bat build
> gradlew.bat install
```

And run SeleDiff.bat in ./build/install/SeleDiff/bin/

- > cd /build/install/SeleDiff/bin/
- > SeleDiff.bat

You can use gradlew.bat to remove SeleDiff.

- > cd /path/to/SeleDiff
- > gradlew.bat clean

There are two sub-commands in SeleDiff. The first sub-command compute-var is used for estimating variances of Ω^1 , which are required for the second sub-command compute-diff.

2.3 Input Files

SeleDiff assumes bi-allelic genetic data and will not perform any checks on this assumption. All input files can be compressed by gzip.

2.3.1 EIGENSTRAT

SeleDiff accepts EIGENSTRAT format of genetic data as inputs. EIGENSOFT provides several functions to convert other formats to EIGENSTRAT format.

For EIGENSTRAT format, there are 3 files: **SNP** file, **IND** file and **GENO** file. Consider an example dataset containing 3 unrelated individuals (Ind1, Ind2 & Ind3) from 3 populations (Pop1, Pop2 & Pop3) that were typed on 3 SNPs (SNP1, SNP2 & SNP3):

```
        SNP1
        SNP2
        SNP3

        Ind1
        T/T
        A/T
        T/T

        Ind2
        C/G
        C/G
        C/G

        Ind3
        C/C
        A/A
        ?/?
```

The SNP file describes the information of each SNP. The SNP file corresponding to the example dataset is:

```
SNP1 1 0.1 100 A T
SNP2 1 0.2 101 C G
SNP3 1 0.3 103 C A
```

Each row corresponds to a SNP. The 6 columns are:

- 1. SNP ID
- 2. Chromosome number
- 3. SNP genetic position
- 4. SNP physical position
- 5. Reference allele

6. Alternative allele

The IND file describes the information of each individual. The IND file corresponding to the example dataset is:

```
Ind1 M pop1
Ind2 F pop2
Ind3 U pop3
```

Each row corresponds to an individual. The 3 columns are:

- 1. Individual ID
- 2. Sex: M for male, F for female and U for unknown
- 3. Population ID

The **GENO** file contains genetic data. The GENO file corresponding to the example dataset is:

010

111

209

Each row corresponds to a SNP, and each column corresponds to an individual. The characters, 0, 1, 2, 9, correspond to an individual's genotype:

- 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.

2.3.2 VCF

SeleDiff also accepts VCF format of genetic data as inputs, and assumes genotypes of each individual are encoded with 0 and 1. Because VCF format contains no population information of each individual, users should provide an additional file following EIGENSTRAT IND format.

2.3.3 Var File

The Var file is the output file from the first sub-command compute-var, which stores variances of pairwise Ω . When using sub-command compute-diff to estimate selection differences, SeleDiff uses --var option to accept a a SPACE delimited file without header that specifies variances of Ω between two populations.

```
YRI CEU 1.547660
YRI CHS 1.639591
CEU CHS 0.989241
```

The first two columns are the population IDs, and the third column is the variances of Ω of the two populations.

2.3.4 Divergence Time File

When using sub-command compute-diff to estimate selection differences, SeleDiff uses --time option to accept a SPACE delimited file without header that specifies divergence times between two populations.

```
YRI CEU 5000
YRI CHS 5000
CEU CHS 3000
```

The first two columns are the population IDs, and the third column is the divergence times of the two populations.

2.4 Output File

The output file from SeleDiff is TAB delimited. The first row is a header that describes the meaning of each column.

Column	Column Name	Description
1	SNP ID	The name of a SNP
2	Ref	The reference allele
3	Alt	The alternative allele
4	Population1	The first population ID
5	Population2	The second population ID
6	Selection difference	The selection difference between the first and
		second populations
7	Std	The standard deviation of the selection difference
8	Lower bound of 95% CI	Lower bound of 95% confidence interval of the selection difference
9	Upper bound of 95% CI	Upper bound of 95% confidence interval of the selection difference
10	Delta	The δ statistic for selection difference
11	p-value	The p -value of the δ statistic

3 An Example

Here is an example to show how SeleDiff estimates and tests selection differences between populations. 4 populations (YRI, CEU, CHB, CHD) from HapMap3 (release3) were extracted. CHB and CHD were merged into one population called CHS. PLINK 1.7 were used to remove correlated individuals and SNPs with minor allele frequencies less than 0.05 and strong linkage disequilibrium. These genome-wide data are stored in ./examples/data/example.geno and used for estimating variances of Ω .

Two alternative alleles (rs1800407 and rs12913832) associated with blue eyes were identified in genes *HERC2* and *OCA2*.² These candidate data are stored in ./examples/data/example.candidates.geno and used for estimating selection differences of these SNPs between populations.

The counts of alleles in our example data were summarized in below.

SNP ID	Population	Reference Allele Count	Alternative Allele Count
rs1800407	YRI	290	0
rs1800407	CEU	207	17
rs1800407	CHS	486	4
rs12913832	YRI	294	0
rs12913832	CEU	47	177
rs12913832	CHS	491	1

We assume the divergence time of YRI-CEU and YRI-CHS are both 5000 generations, while the divergence time of CEU-CHS is 3000 generations. This information is stored in ./examples/data/example.time.

First, we estimate variances of Ω using sub-command compute-var.

To estimate selection differences, we use the sub-command compute-diff.

The result is stored in ./examples/results/example.candidates.results. The main result is in below.

SNP ID	Population1	Population2	Selection difference	Std	delta	p-value
rs1800407	YRI	CEU	-0.000773	0.000380	4.129	0.042154
rs1800407	YRI	CHS	-0.000336	0.000393	0.731	0.392559
rs1800407	CEU	CHS	0.000728	0.000377	3.730	0.053443
rs12913832	YRI	CEU	-0.001541	0.000378	16.583	0.000047
rs12913832	YRI	CHS	-0.000117	0.000415	0.080	0.777297
$\mathrm{rs}12913832$	CEU	CHS	0.002372	0.000433	30.062	0.000000

From the result, we can see the selection coefficient of rs12913832 in CEU is significantly larger than that in YRI or CHS, which indicates rs12913832 is under directional selection in CEU. While the selection coefficient of rs1800407 in CEU is marginal significantly larger than that in YRI or CHS.

4 Dependencies

- Java 1.8
- Apache Commons Math 3.6
- JCommander 1.72
- t-digest 3.1

5 References

- 1. He Y, Wang M, Huang X, Li R, Xu H, Xu S, Jin L. 2015. A probabilistic method for testing and estimating selection differences between populations. *Genome Res*, **25**: 1903-1909.
- 2. Sturm RA, Duffy DL, Zhao ZZ, Leite FP, Stark MS, Hayward NK, Martin NG, Montgomery GW. A single SNP in an evolutionary conserved region within intron 86 of the *HERC2* gene determines human blue-brown eye color. *Am J Hum Genet*, **82**: 424-431.