# **ALStructure Manual**

February 20, 2018

alstructure

 ${\it Main function for execution of the ALStructure algorithm}$ 

# Description

Computes global ancestry estimates under the admixture model given a SNP data matrix  $\boldsymbol{X}$ . This function is based on the ALStrcture algorithm from (Cabreros and Storey 2017).

# Usage

```
alstructure(X, d_hat = NULL, svd_method = "base", tol = 1e-05,
    max_iters = 1000, order_method = "ave_admixture", P_init, Q_init)
```

# **Arguments**

Χ	The $m \times n$ SNP data matrix.
d_hat	Estimate of the latent space dimension $d$ . If left blank, this is estimated by the function $estimate_d()$
svd_method	One of "base" or "truncated_svd." If "base" is chosen, the base svd() function is used. If "truncated_svd" is chosen, the truncated svd algorithm propack.svd() from the svd package is used.
tol	The convergence criterion. If $RMSE(\hat{Q}_t - \hat{Q}_{t+1}) < tol$ , then the algorithm halts
max_iters	The maximum number of iterations (repetitions of steps (6) and (7) in Algorithm 1) to be executed
order_method	One of "ave_admixture" or "var_explained." If "ave_admixture," the $d$ populations are ordered by decreasing average admixture accross samples (i.e. $1/n\sum_j q_{ij}$ ). If "var_explained", the $d$ populations are ordered be decreasing variation explained. Specifically, we compute a modified version of the eigen- $R^2$ statistic from (L. S. Chen and Storey 2008). The statistic is modified in the following ways: 1) we treat rows of $Q$ as the response variables 2) we regress each row of $Q$ on the eigenvectors of $Q$ as the response variables 3) we take the weighted average only over the top $Q$ eigenvectors. and columns of $Q$ are ordered by amount of variation explained by each row of $Q$ by the function order_ $Q$
P_init	Optional initialization of $P$ . Only available for cALS method.
Q_init	Optional initialization of $Q$ . Only available for cALS method.

2 estimate\_F

#### Value

A list with the following elements:

P\_hat: The estimated P matrix. Each column of P may be interpreted as a vector of allele frequencies for a specific ancestral population.

 $\mathbf{Q}$ \_hat: The estimated  $\mathbf{Q}$  matrix. Each column of  $\mathbf{Q}$  may be interpreted as the admixture proportions of a specific individual.

**rowspace**: a list with the following elements:

**vectors**: The top d eigenvectors of the matrix G sorted by decreasing eigenvalue. These vectors approximate the subspace spanned by the rows of Q.

values: The top d eigenvalues of the matrix G sorted by decreasing eigenvalue.

#### References

Cabreros, I., and J. D. Storey. 2017. "A Nonparametric Estimator of Population Structure Unifying Admixture Models and Principal Components Analysis." BioRxiv. Cold Spring Harbor Laboratory. doi:10.1101/240812.

Hao, W., M. Song and J. D. Storey. 2015. "Ifa: Logistic Factor Analysis for Categorical Data." R package version 1.8.0, https://github.com/StoreyLab/Ifa.

estimate\_F

Estimates the individual-specific allele frequency matrix  $oldsymbol{F}$ 

#### **Description**

Estimates the  $m \times n$  individual-specific allele frequency matrix F using the method of Latent Subspace Estimation (X. Chen and Storey 2015) as described in (Cabreros and Storey 2017) in Section 2.3.

#### Usage

```
estimate_F(X, d, svd_method = "base")
```

#### **Arguments**

X The  $m \times n$  SNP data matrix

d The rank of F. This can be estimated using the function d\_estimate().

svd\_method One of "base" or "truncated\_svd." If "base" is chosen, the base svd() function is

used. If "truncated\_svd" is chosen, the truncated svd algorithm propack.svd()

from the svd package is used.

#### Value

A list with the following elements:

**F\_hat**: The  $m \times n$  matrix  $\hat{F}$ .

**rowspace**: a list with the following elements:

**vectors**: The top d eigenvectors of the matrix G sorted by decreasing eigenvalue. These vectors approximate the subspace spanned by the rows of Q.

values: The top d eigenvalues of the matrix G sorted by decreasing eigenvalue.

estimate\_d 3

#### References

Cabreros, I., and J. D. Storey. 2017. "A Nonparametric Estimator of Population Structure Unifying Admixture Models and Principal Components Analysis." BioRxiv. Cold Spring Harbor Laboratory. doi:10.1101/240812.

Chen, X., and J. D. Storey. 2015. "Consistent Estimation of Low-Dimensional Latent Structure in High-Dimensional Data." ArXiv E-Prints, October.

estimate\_d

Estimate the latent space dimension

#### **Description**

Estimates the dimension of the rowspace of Q (equivalently, the rank of the matrix F). This estimate  $\hat{d}$  is based on the estimator from (Leek 2011), page 6.

# Usage

```
estimate_d(X)
```

#### **Arguments**

Χ

the  $m \times n$  SNP data matrix

#### Value

an estimate  $\hat{d}$  of the dimension of the latent space dimension.

# References

Leek, J. T. 2011. "Asymptotic conditional singular value decomposition for high-dimensional genomic data." Biometrics 67 (4): 344–52.

factor\_F

A fast algorithm for factoring  $\hat{F}$ .

# **Description**

An algorithm for finding approximate factors  $\hat{P}$  and  $\hat{Q}$  of the matrix  $\hat{F}$  that obey constraints of the admixture model:

```
1. p_{ij} \in [0, 1] \ \forall (i, j)
2. q_{ij} \ge 0 \ \forall (i, j) \ \text{and} \ \sum_i q_{ij} = 1 \ \forall j
```

This algorithm is described in Algorithm 2 in (Cabreros and Storey 2017). While it lacks the theoretical guarantees of the cALS algorithm, it is much faster.

#### Usage

```
factor_F(F_hat, d, tol = 1e-05, max_iters = 1000)
```

4 Ise

#### **Arguments**

F_hat	The estimate of the matrix $F$ to be factored
d	The dimension of the latent space. This can be estimated by the function $d_{\tt estimate}$ .
tol	The convergence criterion. If $RMSE(\hat{Q}_t - \hat{Q}_{t+1}) < tol$ , then the algorithm halts
max_iters	The maximum number of iterations (repetitions of steps (6) and (7) in Algorithm 1) to be executed

#### Value

A list with the following elements:

P\_hat: The estimated P matrix. Each column of P may be interpreted as a vector of allele frequencies for a specific ancestral population.

 $\mathbf{Q}$ —hat: The estimated  $\mathbf{Q}$  matrix. Each column of  $\mathbf{Q}$  may be interpreted as the admixture proportions of a specific individual.

@references Cabreros, I., and J. D. Storey. 2017. "A Nonparametric Estimator of Population Structure Unifying Admixture Models and Principal Components Analysis." BioRxiv. Cold Spring Harbor Laboratory. doi:10.1101/240812.

lse

Estimates the latent subspace

# **Description**

Estimates the rowspace of Q using the method of latent subspace estimation. The function returns the top d eigenvalues and vectors of the matrix

$$\boldsymbol{G} = \frac{1}{m} \boldsymbol{X}^T \boldsymbol{X} - \boldsymbol{D}$$

where the matrix D is a diagonal matrix with each diagonal entry  $d_{ii}$  an estimate of the average of the variances of the random variables in the i column of X. As is proven in (X. Chen and Storey 2015), the span of the top d eigenvectors of G span the same space as the rows of G. The eigenvectors are returned in order of decreasing eigenvalue.

# Usage

lse(X, d, svd\_method = "base")

# **Arguments**

Χ	The $m \times n$ SNP data matrix
d	The rank of ${\pmb F}$ . This can be estimated using the function d_estimate(). When $d=n$ , all eigenvectors of ${\pmb G}$ are returned.
svd_method	One of "base" or "truncated_svd." If "base" is chosen, the base svd() function is used. If "truncated_svd" is chosen, the truncated svd algorithm propack.svd() from the svd package is used.

order\_pops 5

#### Value

A list with the following elements:

**vectors**: The top d eigenvectors of the matrix G sorted by decreasing eigenvalue. These vectors approximate the subspace spanned by the rows of Q.

values: The top d eigenvalues of the matrix G sorted by decreasing eigenvalue.

#### References

Chen, X., and J. D. Storey. 2015. "Consistent Estimation of Low-Dimensional Latent Structure in High-Dimensional Data." ArXiv E-Prints, October.

order\_pops

Orders the d populations

# **Description**

Orders the d populations according to one of two methods: "ave\_admixture" or "var\_explained." Function returns matrices P and Q with permuted columns and rows according to the determined ordering of populations.

#### Usage

```
order_pops(P, Q, method = "ave_admixture", Q_space = NULL)
```

# **Arguments**

P The  $m \times d$  loadings matrix with columns to be ordered

Q The  $d \times n$  admixture matrix with rows to be ordered

method One of "ave\_admixture" or "var\_explained." If "ave\_admixture," the d popula-

tions are ordered by decreasing average admixture accross samples (i.e.  $1/n\sum_j q_{ij}$ ). If "var\_explained", the d populations are ordered be decreasing variation explained. Specifically, we compute a modified version of the eigen- $R^2$  statistic from (L. S. Chen and Storey 2008). The statistic is modified in the following ways: 1) we treat rows of Q as the response variables 2) we regress each row of Q on the eigenvectors of G rather than the eigenvectors of the data matrix itself

3) we take the weighted average only over the top d eigenvectors.

Q\_space Only required for "var\_explained" methodThe list containing the top d eigen-

vectors and their corresponding eigenvalues of the G. These may be obtained

through the function 1se.

#### Value

A list with the following elements:

 $\mathbf{P}$ \_ordered: The permuted P  $\mathbf{Q}$ \_ordered: The permuted Q

 $\mathbf{perm\_mat}$ : The permutation matrix A such that  $PA^T = P_{\mathrm{ord}}$  and  $AQ = Q_{\mathrm{ord}}$ .

6 simulate\_admixture

#### References

Chen, L. S., and J. D. Storey. 2008. "Eigen-R2 for dissecting variation in high-dimensional studies." Bioinformatics 24 (19): 2260–2.

simulate\_admixture

Simulates data from the PSD model

#### **Description**

Creates a data frame that contains the parameters of the admixture model (F, P, Q) as well as a single draw X such that  $x_{ij} \sim \text{Bernoulli}(f_{ij})$ . The Q matrix is drawn from the Dirichlet distribution with parameter  $\alpha$ , and the P matrix is simulated from the Balding-Nichols model. The parameter  $\alpha$  is supplied by the user. The  $m \times 2$  matrix of Balding-Nichols parameters is optional. If BN\_params is not supplied, the parameters are derived from a random sample of estimated Balding-Nichols parameters from the Human Genomes Diversity Project (HGDP) dataset. The Balding-Nichols parameter estimates are provided by (Gopalan et al. 2016), and included in this package in the object hgdpBN.

# Usage

```
simulate_admixture(m, n, d, alpha, BN_params = NA, seed = NA)
```

#### **Arguments**

m number of SNPs
n number of individuals
d number of groups

alpha dirichlet parameter; length(alpha) = d

BN\_params a  $m \times 2$  matrix of parameters. The first column contains  $F_{ST}$  for each SNP,

while the second column contains the allele frequency.

#### Value

a list with the following elements:

 $\mathbf{P}$ : the  $m \times d$  matrix of loadings

 $\mathbf{Q}$ : the  $d \times n$  matrix of latent admixture components

 $\mathbf{F}$ : the  $m \times n$  matrix PQ

**X**: a random draw such that  $x_{ij} \sim \text{Bernoulli}(f_{ij}, 2)$ .

## References

Gopalan, P., W. Hao, D. M. Blei, and J. D. Storey. 2016. "Scaling probabilistic models of genetic variation to millions of humans." Nature Genetics 48 (12): 1587–90.

# Index

```
alstructure, 1
estimate_d, 3
estimate_F, 2
factor_F, 3
lse, 4
order_pops, 5
simulate_admixture, 6
```