

# Package ‘TSDFGS’

January 20, 2025

**Type** Package

**Title** Training Set Determination for Genomic Selection

**Version** 2.0

**Date** 2022-06-07

**Description** We propose an optimality criterion to determine the required training set, r-score, which is derived directly from Pearson’s correlation between the genomic estimated breeding values and phenotypic values of the test set <[doi:10.1007/s00122-019-03387-0](https://doi.org/10.1007/s00122-019-03387-0)>. This package provides two main functions to determine a good training set and its size.

**License** GPL (>= 3)

**Encoding** UTF-8

**Imports** dplyr, ggplot2, latex2exp, lifecycle, parallel, Rcpp (>= 1.0.8.3)

**LinkingTo** Rcpp, RcppEigen

**RoxygenNote** 7.2.0

**URL** <https://github.com/oumarkme/TSDFGS>

**BugReports** <https://github.com/oumarkme/TSDFGS/issues>

**Depends** R (>= 2.10)

**LazyData** true

**NeedsCompilation** yes

**Author** Jen-Hsiang Ou [aut, cre] (<<https://orcid.org/0000-0001-9305-2931>>),  
Po-Ya Wu [aut] (<<https://orcid.org/0000-0002-7342-2867>>),  
Chen-Tuo Liao [aut, ths] (<<https://orcid.org/0000-0001-9777-3701>>)

**Maintainer** Jen-Hsiang Ou <jen-hsiang.ou@imbim.uu.se>

**Repository** CRAN

**Date/Publication** 2022-06-07 14:00:11 UTC

## Contents

cd_score . . . . .	2
FGCM . . . . .	3
geno . . . . .	3
nt2r . . . . .	4
optTrain . . . . .	5
pev_score . . . . .	6
r_score . . . . .	7
SSDFGS . . . . .	7
subpop . . . . .	8
<b>Index</b>	<b>9</b>

---

cd_score	<i>CD-score</i>
----------	-----------------

---

### Description

This function calculate CD-score [doi:10.1186/1297-9686-28-4-359](https://doi.org/10.1186/1297-9686-28-4-359) by given training set and test set.

### Usage

```
cd_score(X, X0)
```

### Arguments

X	A numeric matrix. The training set genotypic information matrix can be given as genotype matrix (coded as -1, 0, 1) or principle component matrix (row: sample; column: marker).
X0	A numeric matrix. The test set genotypic information matrix can be given as genotype matrix (coded as -1, 0, 1) or principle component matrix (row: sample; column: marker).

### Value

A floating-point number, CD score.

### Author(s)

Jen-Hsiang Ou

### Examples

```
data(geno)
## Not run: cd_score(geno[1:50, ], geno[51:100])
```

---

 FGCM

*Fit logistic growth curve model*


---

**Description**

A function for fitting logisti growth model

**Usage**

```
FGCM(geno, nt = NULL, n_iter = NULL, multi.threads = TRUE)
```

**Arguments**

geno	Genotype information saved as a dataframe. Columns represent variants (SNPs or PCs).
nt	A numerical vector of training set sample size for estimating logistic growth curve parameters
n_iter	Number of simulation of each training set size. Automatically gave a suitable number by default.
multi.threads	Default: TRUE. Set as FALSE if you just want to run it by single thread.

**Value**

Estimation of parameters.

**Examples**

```
data(geno)
## Not run: FGCM(geno)
```

---

 geno

*Genotype information*


---

**Description**

A PCA matrix of rice genotype information. This data was published by Zhao et al. (2011) [doi: 10.1038/ncomms1467](https://doi.org/10.1038/ncomms1467)

**Usage**

```
geno
```

**Format**

A numeric matrix (PCA) with 404 rows (sample) and 404 columns (PCs).

**Source**

<http://www.ricediversity.org/data/>

**Examples**

```
data(geno)
```

---

nt2r

*Simulate r-scores of each training set size*

---

**Description**

Calculate r-scores (un-target) by in parallel.

**Usage**

```
nt2r(geno, nt, n_iter = 30, multi.threads = TRUE)
```

**Arguments**

geno	A numeric dataframe of genotype, column represent sites (genotype coding as 1, 0, -1)
nt	Numeric. Number of training set size
n_iter	Times of iteration. (default = 30)
multi.threads	Default: TRUE

**Value**

A vector of r-scores of each iteration

**Examples**

```
data(geno)
## Not run: nt2r(geno, 50)
```

---

optTrain	<i>Optimal training set determination</i>
----------	---

---

**Description**

This function is designed for determining optimal training set.

**Usage**

```
optTrain(  
  geno,  
  cand,  
  n.train,  
  subpop = NULL,  
  test = NULL,  
  method = "rScore",  
  min.iter = NULL  
)
```

**Arguments**

geno	A numeric matrix of principal components (rows: individuals; columns: PCs).
cand	An integer vector of which rows of individuals are candidates of the training set in the geno matrix.
n.train	The size of the target training set. This could be determined with the help of the ssdfgp function provided in this package.
subpop	A character vector of sub-population's group name. The algorithm will ignore the population structure if it remains NULL.
test	An integer vector of which rows of individuals are in the test set in the geno matrix. The algorithm will use an un-target method if it remains NULL.
method	Choices are rScore, PEV and CD. rScore will be used by default.
min.iter	Minimum iteration of all methods can be appointed. One should always check if the algorithm is converged or not. A minimum iteration will set by considering the candidate and test set size if it remains NULL.

**Value**

This function will return 3 information including OPTtrain (a vector of chosen optimal training set), TOPscore (highest scores of before iteration), and ITERscore (criteria scores of each iteration).

**Author(s)**

Jen-Hsiang Ou

## Examples

```
data(geno)
## Not run: optTrain(geno, cand = 1:404, n.train = 100)
```

---

pev_score	<i>PEV score</i>
-----------	------------------

---

## Description

This function calculate prediction error variance (PEV) score [doi:10.1186/s12711-015-0116-6](https://doi.org/10.1186/s12711-015-0116-6) by given training set and test set.

## Usage

```
pev_score(X, X0)
```

## Arguments

X	A numeric matrix. The training set genotypic information matrix can be given as genotype matrix (coded as -1, 0, 1) or principle component matrix (row: sample; column: marker).
X0	A numeric mareix. The test set genotypic information matrix can be given as genotype matrix (coded as -1, 0, 1) or principle component matrix (row: sample; column: marker).

## Value

A floating-point number, PEV score.

## Author(s)

Jen-Hsiang Ou

## Examples

```
data(geno)
## Not run: pev_score(geno[1:50, ], geno[51:100])
```

---

r_score	<i>r-score</i>
---------	----------------

---

**Description**

This function calculate r-score [doi:10.1007/s00122-019-03387-0](https://doi.org/10.1007/s00122-019-03387-0) by given training set and test set.

**Usage**

```
r_score(X, X0)
```

**Arguments**

X	A numeric matrix. The training set genotypic information matrix can be given as genotype matrix (coded as -1, 0, 1) or principle component matrix (row: sample; column: marker).
X0	A numeric matrix. The test set genotypic information matrix can be given as genotype matrix (coded as -1, 0, 1) or principle component matrix (row: sample; column: marker).

**Value**

A floating-point number, r-score.

**Author(s)**

Jen-Hsiang Ou

**Examples**

```
data(geno)
## Not run: r_score(geno[1:50, ], geno[51:100])
```

---

SSDFGS	<i>Sample size determination for genomic selection</i>
--------	--

---

**Description**

This function is designed to generate an operating curve for sample size determination

**Usage**

```
SSDFGS(geno, nt = NULL, n_iter = NULL, multi.threads = TRUE)
```

**Arguments**

<code>geno</code>	A numeric data frame carried genotype information (column: PCs, row: sample)
<code>nt</code>	A numeric vector carried training set sizes for r-score simulation.
<code>n_iter</code>	Number of iterations for estimating parameters.
<code>multi.threads</code>	Default ( <code>multi.threads = TRUE</code> ) use 75% of threads if the computer has more than 4 threads.

**Value**

An operating curve and its information.

**Author(s)**

Jen-Hsiang Ou & Po-Ya Wu

**Examples**

```
data(geno)
## Not run: SSDFGS(geno)
```

---

subpop	<i>Sub-population information</i>
--------	-----------------------------------

---

**Description**

Sub-population information of samples. This data was published by Zhao et al. (2011) [doi:10.1038/ncomms1467](https://doi.org/10.1038/ncomms1467)

**Usage**

```
subpop
```

**Format**

A character vector.

**Source**

<http://www.ricediversity.org/data/>

**Examples**

```
data(subpop)
```



# Index

- \* **datasets**
  - geno, [3](#)
  - subpop, [8](#)
- cd\_score, [2](#)
- FGCM, [3](#)
- geno, [3](#)
- nt2r, [4](#)
- optTrain, [5](#)
- pev\_score, [6](#)
- r\_score, [7](#)
- SSDFGS, [7](#)
- subpop, [8](#)