# Package 'IBCF.MTME'

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**Title** Item Based Collaborative Filtering for Multi-Trait and Multi-Environment Data

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**Description** Implements the item based collaborative filtering (IBCF) method for continues phenotypes in the context of plant breeding where data are collected for various traits that were studied in various environments proposed by Montesinos-López et al. (2017) <doi:10.1534/g3.117.300309>.

**Depends** R (>= 3.0.0)

License LGPL-3

Encoding UTF-8

LazyData true

Type Package

RoxygenNote 6.1.1

URL https://github.com/frahik/IBCF.MTME

BugReports https://github.com/frahik/IBCF.MTME/issues/new

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**Imports** lsa, tidyr, dplyr

Suggests testthat, knitr, rmarkdown, covr

```
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# Contents

barplot.IBCFY 2
CV.RandomPart
getMatrixForm
getTidyForm
IBCF 5
IBCF.MTME
IBCF.Years
plot.IBCF
print.IBCF
print.IBCFY
summary.IBCF 9
summary.IBCFY 10
Wheat_IBCF         10
Year_IBCF 11

# Index

barplot.IBCFY barplot.IBCFY

# Description

Barplot of the results from IBCFY object

# Usage

```
## S3 method for class 'IBCFY'
barplot(height, select = "Pearson", ...)
```

# Arguments

height	IBCFY object IBCFY object, result of use the IBCF.Years() function
select	character By default ('Pearson'), plot the Pearson Correlations of the IBCF Object, else ('MAAPE'), plot the MAAPE of the IBCF Object.
	Further arguments passed to or from other methods.

12

CV.RandomPart

#### Description

This method consists of randomly dividing the training data set and the test data set. For each division, the approximation function is adjusted from the training data and calculates the output values for the test data set. The result corresponds to the arithmetic mean of the values obtained for the different divisions.

#### Usage

```
CV.RandomPart(DataSet, NPartitions = 10, PTesting = 0.35,
Traits.testing = NULL, Set_seed = NULL)
```

#### Arguments

DataSet	data.frame The data set object is a data.frame object that contains 4 column in the Tidy data format: \$Line is the Line or genotype identifier, and the nam of this column could change. \$Env is the name of the evaluated environmer (s). \$Trait is the name of the evaluated trait (s). \$Response Variable response obtained for the row corresponding to line and environment.	
NPartitions	integer Number of Partitions for the Cross-Validation. Is 10 by default.	
PTesting	Double Percentage of Testing for the Cross-Validation. Is 0.35 by default.	
Traits.testing	character By default is null and use all the traits to fit the model, else only part of the traits specified be used to fit the model.	
Set_seed	integer Number of seed for reproducible research. Is NULL by default.	

#### Value

List A list object with length of NPartitions, every index has a matrix  $n \times x$ , where n is the number of NLines and x is the number of NEnv  $\times$  NTraits. The values inside is 1 for training and 2 for testing.

# Examples

```
## Not run:
library(IBCF.MTME)
data('Wheat_IBCF')
CV.RandomPart(Wheat_IBCF, NPartitions = 10)
CV.RandomPart(Wheat_IBCF, Traits.testing = 'DH')
CV.RandomPart(Wheat_IBCF, NPartitions = 10, PTesting = .35)
CV.RandomPart(Wheat_IBCF, NPartitions = 10, Traits.testing = 'DH')
CV.RandomPart(Wheat_IBCF, NPartitions = 10, PTesting = .35, Set_seed = 5)
CV.RandomPart(Wheat_IBCF, NPartitions = 10, PTesting = .35, Set_seed = 5)
CV.RandomPart(Wheat_IBCF, NPartitions = 10, PTesting = .35, Traits.testing = 'DH')
```

CV.RandomPart(Wheat\_IBCF, NPartitions = 10, PTesting = .35, Traits.testing = 'DH', Set\_seed = 5 )
## End(Not run)

getMatrixForm Tidy data format to Matrix format

# Description

Tidy data format to Matrix format

# Usage

```
getMatrixForm(Tidy_DataSet, onlyTrait = FALSE)
```

# Arguments

Tidy_DataSet	data.frame object that contains 4 columns: \$Line: Line or genotype identi- fier, and the name of this column could change. \$Env: Name of the evaluated
	environment (s). \$Trait: Name of the evaluated trait (s). \$Response: Variable response obtained for the row corresponding to line and environment.
onlyTrait	logical by default is FALSE, if is TRUE only the column \$Trait is transformed.

# Value

A data.frame object with the \$Response divided by \$Traits columns.

# Examples

```
## Not run:
    data('Wheat_IBCF')
    M <- getMatrixForm(Wheat_IBCF)
## End(Not run)
## Not run:
    data('Year_IBCF')
    M.Y <- getMatrixForm(Year_IBCF, onlyTrait = T)
## End(Not run)
```

4

getTidyForm

#### Description

Matrix format to Tidy data format

#### Usage

getTidyForm(Matrix\_DataSet, onlyTrait = FALSE)

# Arguments

Matrix_DataSet	A data.frame object with the response values divided in n environments or traits	
	columns	
onlyTrait	logical by default is FALSE, if is TRUE only is considered the \$Trait column.	

#### Value

A data.frame object with the \$Response divided by \$Traits columns.

# Examples

```
## Not run:
    data('Wheat_IBCF')
    M <- getMatrixForm(Wheat_IBCF)
    Tidy <- getTidyForm(M)
## End(Not run)
## Not run:
    data('Year_IBCF')
    M.Y <- getMatrixForm(Year_IBCF, onlyTrait = T)
    Tidy.Y <- getTidyForm(M.Y, onlyTrait = T)
## End(Not run)
```

IBCF

IBCF

# Description

Item Based Collaborative Filtering for multi-trait and multi-environment data.

# Usage

IBCF(object, dec = 4)

#### Arguments

object	list CrossValidation object, is obtained from CV.RandomPartition function.
dec	integer Number of decimals to print in the results.

#### Value

A list with the next components

NPartitions integer Number of partitions used for testing dat		
predictions_Summary		
	${\tt data.frame}\ A$ data.frame with the results of the test	
Predictions	list A list with the predicted results for each partition	

# Examples

```
## Not run:
library(IBCF.MTME)
data('Wheat_IBCF')
```

```
CV <- CV.RandomPart(Wheat_IBCF)
IBCF(CV)
```

## End(Not run)

IBCF.MTME

*IBCF.MTME: Item Based Collaborative Filtering for Multi-Trait and Multi-Environment Data.* 

# Description

The Item Based Collaborative Filtering for Multi-Trait and Multi-Environment Data (IBCF.MTME) package was developed to implement the item based collaborative filtering (IBCF) method for continues phenotypes in the context of plant breeding where data are collected for various traits that were studied in various environments. It is important to point out that the main difference of this package with the available packages that can implement IBCF is that this package was developed for continuous phenotypes which cannot be implemented in the current packages that can implement IBCF that only work for binary and ordinary phenotypes. IBCF.Years

#### Description

Item Based Collaborative Filtering for Years data

# Usage

```
IBCF.Years(DataSet, colYears = 1, colID = 2, Years.testing = "",
Traits.testing = "", dec = 4)
```

# Arguments

DataSet	data.frame A data set in Matrix Form.
colYears	string or integer A name or the position of the 'Years' column just in case that is not the first column.
colID	string or integer A name or the position of the 'ID' column, just in case that is not the second column.
Years.testing	vector A vector with the names of the years to use in test.
Traits.testing	vector A vector with the names of the traits to use in test.
dec	integer Number of decimals to print in the results.

#### Value

A list with the next components

Year.testing	vector a vector with the Years used for the testing data	
Traits.testing	vector a vector with the Traits used for the testing data	
Data_Obs_Pred	data.frame Contains the values observed and predicted (the predicted values has '.1' after the name)	
predictions_Summary		
	data.frame Contains the summary of the correlation of the predictions and the MAAPE	

# Examples

```
## Not run:
library(IBCF.MTME)
data('Year_IBCF')
DataSet <- getMatrixForm(Year_IBCF, onlyTrait = TRUE)
IBCF.Years(DataSet , Years.testing = c('2015', '2016'), Traits.testing = c('T5', 'T6'))
```

## End(Not run)

plot.IBCF

# Description

Plot from IBCF object

# Usage

```
## S3 method for class 'IBCF'
plot(x, select = "Pearson", ...)
```

# Arguments

х	IBCF object IBCF object, result of use the IBCF() function
select	character By default ('Pearson'), plot the Pearson Correlations of the IBCF Object, else ('MAAPE'), plot the MAAPE of the IBCF Object.
	Further arguments passed to or from other methods.

ſ	orint.IBCF	Print IBCF information object

# Description

Print IBCF information object

# Usage

```
## S3 method for class 'IBCF'
print(x, ...)
```

# Arguments

х	IBCF object
	Further arguments passed to or from other methods.

# Value

printeable object

print.IBCFY

# Description

Print IBCFY information object

#### Usage

## S3 method for class 'IBCFY'
print(x, ...)

# Arguments

Х	IBCFY object
	Further arguments passed to or from other methods.

#### Value

printeable object

summary.IBCF Summary

# Description

Summary of IBCF object

# Usage

```
## S3 method for class 'IBCF'
summary(object, information = "compact", digits = 4,
    ...)
```

# Arguments

object	IBCF object IBCF object, result of use the IBCF() function
information	string
digits	numeric
	Further arguments passed to or from other methods.

summary.IBCFY Summary

# Description

Summary of IBCFY object

#### Usage

## S3 method for class 'IBCFY'
summary(object, digits = 4, ...)

#### Arguments

object	IBCFY object IBCFY object, result of use the IBCF.Years() function
digits	numeric Number of digits of the output.
	Further arguments passed to or from other methods.

#### Description

The package includes a data set based on a portion of the data used in the study of Montesinos-Lopez, O. A.; Montesinos-Lopez, A.; Crossa, J.; Toledo, F. H.; Montesinos-Lopez, J. C.; Singh, P. & Salinas-Ruiz, J. (2017). A Bayesian Poisson-lognormal Model for Count Data for Multiple-Trait Multiple-Environment Genomic-Enabled Prediction. G3: GeneslGenomeslGenetics 7(5):1595–1606. http://doi.org/10.1534/g3.117.039974. The data set consists of 250 wheat lines evaluated in 3 environments and 4 distinct traits, i.e. 3000 observations.

#### Usage

data(Wheat\_IBCF)

# Format

a TidyData format, 3000 row per 4 columns.

#### Author(s)

Montesinos-Lopez, O. A.

Year\_IBCF

#### Description

Dataset based on simulated data with the next code:

```
set.seed(2)
A <- matrix(0.65,ncol=12,nrow=12)
diag(A) <- 1
Sdv <- diag(c(0.9^0.5,0.8^0.5,0.9^0.5,0.8^0.5,0.86^0.5,0.7^0.5,0.9^0.5,0.8^0.5,0.9^0.5,0.7^0.5,0.7^0
Sigma <- Sdv
No.Lines <- 60
Z <- rmvnorm(No.Lines,mean=c(5,5.5,6,5.5,7,6.5,6.0,7,6.6,8,6.3,8),sigma=Sigma)
Years <- c(rep(2014,20),rep(2015,20),rep(2016,20))
Gids <- c(1:No.Lines)
Data.Final <- data.frame(cbind(Years,Gids,Z))
colnames(Data.Final) <- c("Years", "Gids", "T1", "T2", "T3", "T4", "T5", "T6", "T7", "T8", "T9", "T10", "T11", "T
head(Data.Final)
Year_IBCF <- getTidyForm(Data.Final, onlyTrait = T)</pre>
```

#### Usage

data(Year\_IBCF)

#### Format

a TidyData format, 750 row per 4 columns.

#### Author(s)

Montesinos-Lopez, O. A.

# Index

barplot.IBCFY, 2 CV.RandomPart, 3 getMatrixForm, 4 getTidyForm, 5 IBCF, 5 IBCF.MTME, 6 IBCF.MTME-package (IBCF.MTME), 6 IBCF.Years, 7 plot.IBCF, 8 print.IBCF, 8 print.IBCF, 9 summary.IBCF, 9 summary.IBCF, 10

Wheat\_IBCF, 10

Year\_IBCF, 11