# Package 'predhy'

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Author Yang Xu, Guangning Yu, Yanru Cui, Shizhong Xu, Chenwu Xu	
Maintainer Yang Xu <xuyang_89@126.com></xuyang_89@126.com>	
<b>Description</b> Performs genomic prediction of hybrid performance using eight GS methods including GBLUP, BayesB, RKHS, PLS, LASSO, Elastic net, LightGBM and XGBoost. It also provides fast cross-validation and mating design scheme for training population (Xu S et al (2016) <doi:10.1111 tpj.13242="">; Xu S (2017) <doi:10.1534 g3.116.038059="">).</doi:10.1534></doi:10.1111>	
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convertgen

Convert Genotype

#### **Description**

Convert genotypes in HapMap format or in numeric format for hypred package.

#### Usage

```
convertgen(
  input_geno,
  type = c("hmp1", "hmp2", "num"),
  missingrate = 0.2,
  maf = 0.05,
  impute = TRUE
)
```

#### **Arguments**

genotype in HapMap format or in numeric format. The names of individuals should be provided. Missing (NA) values are allowed.

type the type of genotype. There are three options: "hmp1" for genotypes in HapMap format with single bit, "hmp2" for genotypes in HapMap format with double bit, and "num" for genotypes in numeric format.

missingrate max missing percentage for each SNP, default is 0.2.

maf minor allele frequency for each SNP, default is 0.05.

impute logical. If TRUE, imputation. Default is TRUE.

#### Value

A matrix of genotypes in numeric format, coded as 1, 0, -1 for AA, Aa, aa. Each row represents an individual and each column represents a marker. The rownames of the matrix are the names of individuals.

```
## load genotype in HapMap format with double bit
data(input_geno)

## convert genotype for hypred package
inbred_gen <- convertgen(input_geno, type = "hmp2")</pre>
```

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```
## load genotype in numeric format
data(input_geno1)
head(input_geno1)

## convert genotype for hypred package
inbred_gen1 <- convertgen(input_geno1, type = "num")</pre>
```

crodesign

Generate Mating Design

#### **Description**

Generate a mating design for a subset of crosses based on a balanced random partial rectangle cross-design (BRPRCD) (Xu et al. 2016).

# Usage

```
crodesign(d, male_name, female_name, seed = 123)
```

#### **Arguments**

d an integer denoting 1/d percentage of crosses to be evaluated in the field.

male\_name a character string for the names of male parents.

female\_name a character string for the names of male parents.

seed the random number, default is 123.

#### Value

A data frame of mating design result with three columns. The first column is "crossID", the second column is the "male\_Name" and the third column is the "female\_Name".

#### References

Xu S, Xu Y, Gong L and Zhang Q. (2016) Metabolomic prediction of yield in hybrid rice. Plant J. 88, 219-227.

```
## generate a mating design with 100 male parents and 150 female parents
## for 1/d = 1/50 percentage of crosses to be evaluated in the field.
## the total number of potential crosses is 100*150 = 15000.
## The number of crosses to be field evaluated is 15000*(1/50) = 300.
male_name <- paste("m", 1:100, sep = "")</pre>
```

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```
female_name <- paste("f", 1:150, sep = "")
design <- crodesign(d = 50, male_name, female_name)</pre>
```

cv

Evaluate Trait Predictability via Cross Validation

# Description

The cv function evaluates trait predictability based on eight GS methods via k-fold cross validation. The trait predictability is defined as the squared Pearson correlation coefficient between the observed and the predicted trait values.

#### Usage

```
cv(
   fix = NULL,
   gena,
   gend = NULL,
   inbred_phe = NULL,
   y,
   method = "GBLUP",
   drawplot = TRUE,
   nfold = 5,
   nTimes = 1,
   seed = 1234,
   CPU = 1
)
```

# **Arguments**

fix a design matrix of the fixed effects.

gena a matrix (n x m) of additive genotypes for the training population.

gend a matrix (n x m) of domiance genotypes for the training population. Default is

NULL.

inbred\_phe a matrix (n x 2) of inbred\_phe phenotypic.Default is NULL.

y a vector(n x 1) of the phenotypic values.

method eight GS methods including "GBLUP", "BayesB", "RKHS", "PLS", "LASSO",

"EN", "XGBoost", "LightGBM". Users may select one of these methods or all

of them simultaneously with "ALL". Default is "GBLUP".

drawplot when method ="ALL", user may select TRUE for a barplot about eight GS meth-

ods. Default is TRUE.

nfold the number of folds. Default is 5.

nTimes the number of independent replicates for the cross-validation. Default is 1.

seed the random number. Default is 1234.

CPU the number of CPU.

cv\_fast 5

# Value

Trait predictability

# **Examples**

```
## load example data from hypred package
data(hybrid_phe)
data(input_geno)

## convert original genotype
inbred_gen <- convertgen(input_geno, type = "hmp2")

##additive model infer the additive and dominance genotypes of hybrids
gena <- infergen(inbred_gen, hybrid_phe)$add
gend <- infergen(inbred_gen, hybrid_phe)$dom

##additive model
R2<-cv(fix=NULL,gena,gend=NULL,y=hybrid_phe[,3],inbred_phe=NULL,method ="GBLUP")

##additive-dominance model
R2<-cv(fix=NULL,gena,gend,y=hybrid_phe[,3],inbred_phe=NULL,method ="GBLUP")</pre>
```

cv\_fast

Evaluate Trait Predictability via the HAT Method

# **Description**

The HAT method is a fast algorithm for the ordinary cross validation. It is highly recommended for large dataset (Xu et al. 2017).

# Usage

```
cv_fast(fix = NULL, y, kk, nfold = 5, seed = 123)
```

# **Arguments**

fix	a design matrix of the fixed effects. If not passed, a vector of ones is added for the intercept.
у	a vector of the phenotypic values.
kk	a list of one or multiple kinship matrices.
nfold	the number of folds, default is 5. For the HAT Method, nfold can be set as the sample size (leave-one-out CV) to avoid variation caused by random partitioning of the samples, but it is not recommended for cv.
seed	the random number, default is 123.

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#### Value

Trait predictability

#### References

Xu S. (2017) Predicted residual error sum of squares of mixed models: an application for genomic prediction. G3 (Bethesda) 7, 895-909.

# **Examples**

```
## load example data from hypred package
data(hybrid_phe)
data(input_geno)

## convert original genotype
inbred_gen <- convertgen(input_geno, type = "hmp2")

## infer the additive and dominance genotypes of hybrids
gena <- infergen(inbred_gen, hybrid_phe)$add
gend <- infergen(inbred_gen, hybrid_phe)$dom

## calculate the additive and dominance kinship matrix
ka <- kin(gena)
kd <- kin(gend)

##for the additive model
predictability <- cv_fast(y = hybrid_phe[,3], kk = list(ka))

##for the additive-dominance model
predictability <- cv_fast(y = hybrid_phe[,3], kk = list(ka,kd))</pre>
```

hybrid\_phe

Phenotypic data of hybrids

# **Description**

This dataset contains phenotypic data of 410 hybrids for grain yield in maize.

#### Usage

hybrid\_phe

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#### **Format**

A data frame with 410 rows and 3 variables:

M The names of male parents.

F The names of female parents.

GY The grain yield of hybrids.

infergen

Infer Genotype of Hybrids

# **Description**

Infer additive and dominance genotypes of hybrids based on their parental genotypes.

#### Usage

```
infergen(inbred_gen, hybrid_phe)
```

#### **Arguments**

inbred\_gen

a matrix for genotypes of parental lines in numeric format, coded as 1, 0 and

-1. The row.names of inbred\_gen must be provied. It can be obtained from the

original genotype using convertgen function.

hybrid\_phe

a data frame with three columns. The first column and the second column are the names of male and female parents of the corresponding hybrids, respectively; the third column is the phenotypic values of hybrids. The names of male and female parents must match the rownames of inbred\_gen. Missing (NA) values

are not allowed.

#### Value

A list with following information is returned: \$add additive genotypes of hybrids \$dom dominance genotypes of hybrids

```
## load example data from hypred package
data(hybrid_phe)
head(hybrid_phe)
data(input_geno)

## convert original genotype
inbred_gen <- convertgen(input_geno, type = "hmp2")
gena <- infergen(inbred_gen, hybrid_phe)$add
gend <- infergen(inbred_gen, hybrid_phe)$dom</pre>
```

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input\_geno

Genotype in Hapmap Format

# Description

Genotypic data of 348 maize inbred lines in Hapmap format with double bit.

# Usage

input\_geno

# **Format**

A data frame with 4979 rows and 359 columns.

input\_geno1

Genotype in Numeric Format

# Description

Genotypic data of 50 rice inbred lines with 1000 SNPs.

# Usage

input\_geno1

#### **Format**

A data frame with 1000 rows and 50 variables.

kin

Calculate Kinship Matrix

# **Description**

Calculate the additive and dominance kinship matrix.

# Usage

kin(gen)

# **Arguments**

gen

a matrix for genotypes, coded as 1, 0, -1 for AA, Aa, aa. Each row represents an individual and each column represents a marker.

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#### Value

```
a kinship matrix
```

#### **Examples**

```
## random population with 100 lines and 1000 markers gen <- matrix(rep(0,100*1000),100,1000) gen <- apply(gen,2,function(x)\{x <- sample(c(-1,0,1), 100, replace = TRUE)\}) ## generate 100*100 kinship matrix k <- kin(gen)
```

mixed

Solve Mixed Model

#### **Description**

Solve linear mixed model using restricted maximum likelihood (REML). Multiple variance components can be estimated.

#### Usage

```
mixed(fix = NULL, y, kk)
```

# **Arguments**

fix	a design matrix of the fixed effects. If not passed, a vector of ones is added for the intercept.
У	a vector of the phenotypic values.
kk	a list of one or multiple kinship matrices.

# Value

A list with following information is returned:

\$v\_i the inverse of the phenotypic variance-covariance matrix

\$var estimated variance components of genetic effects

\$ve estimated residual variance

\$beta estimated fixed effects

#### References

Xu S, Zhu D and Zhang Q. (2014) Predicting hybrid performance in rice using genomic best linear unbiased prediction. Proc. Natl. Acad. Sci. USA 111, 12456-12461.

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# **Examples**

```
## load example data from hypred package
data(hybrid_phe)
data(input_geno)

## convert original genotype
inbred_gen <- convertgen(input_geno, type = "hmp2")

## infer the additive and dominance genotypes of hybrids
gena <- infergen(inbred_gen, hybrid_phe)$add
gend <- infergen(inbred_gen, hybrid_phe)$dom

## calculate the additive and dominance kinship matrix
ka <- kin(gena)
kd <- kin(gend)

## for the additive model
parm <- mixed(y = hybrid_phe[,3], kk = list(ka))

## for the additive-dominance model
parm <- mixed(y = hybrid_phe[,3], kk = list(ka, kd))</pre>
```

predhy.predict

Predict the Performance of Hybrids

#### **Description**

Predict all potential crosses of a given set of parents using a subset of crosses as the training sample.

# Usage

```
predhy.predict(
  inbred_gen,
  hybrid_phe,
  inbred_phe = NULL,
  method = "GBLUP",
  model = "A",
  select = "top",
  number = "100"
)
```

#### **Arguments**

inbred\_gen

a matrix for genotypes of parental lines in numeric format, coded as 1, 0 and -1. The row.names of inbred\_gen must be provied. It can be obtained from the original genotype using convertgen function.

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hybrid\_phe a data frame with three columns. The first column and the second column are the

names of male and female parents of the corresponding hybrids, respectively; the third column is the phenotypic values of hybrids. The names of male and female parents must match the rownames of inbred\_gen. Missing (NA) values

are not allowed.

inbred\_phe a matrix (n x 2) of inbred\_phe phenotypic.Default is NULL.

method eight GS methods including "GBLUP", "BayesB", "RKHS", "PLS", "LASSO",

"EN", "XGBoost", "LightGBM". Users may select P of these methods. Default

is "GBLUP".

model the prediction model. There are four options: model = "A" for the additive

model, model = "AD" for the additive-dominance model,model = "A-P" for the additive-phenotypic model,model = "AD-P" for the additive-dominance-

phenotypic model. Default is model = "A".

select the selection of hybrids based on the prediction results. There are three options:

select = "all", which selects all potential crosses. select = "top", which selects the top n crosses. select = "bottom", which selects the bottom n crosses. The n

is determined by the param number.

number the number of selected top or bottom hybrids, only when select = "top" or select

= "bottom".

#### Value

a data frame of prediction results with two columns. The first column denotes the names of male and female parents of the predicted hybrids, and the second column denotes the phenotypic values of the predicted hybrids.

#### **Examples**

```
## load example data from hypred package
data(hybrid_phe)
data(input_geno)
inbred_gen <- convertgen(input_geno, type = "hmp2")

## infer the additive and dominance genotypes of hybrids
gena <- infergen(inbred_gen, hybrid_phe)$add
gend <- infergen(inbred_gen, hybrid_phe)$dom

pred<-predhy.predict(inbred_gen,hybrid_phe,method="LASSO",model="A",select="top",number="100")
pred<-predhy.predict(inbred_gen,hybrid_phe,method="LASSO",model="AD",select="all")</pre>
```

# Description

Predict all potential crosses of a given set of parents using a subset of crosses as the training sample.

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#### Usage

```
predhy.predict_NCII(
   inbred_gen,
   hybrid_phe,
   inbred_phe = NULL,
   male_name = hybrid_phe[, 1],
   female_name = hybrid_phe[, 2],
   method = "GBLUP",
   model = "A",
   select = "top",
   number = "100"
)
```

#### **Arguments**

inbred\_gen a matrix for genotypes of parental lines in numeric format, coded as 1, 0 and

-1. The row.names of inbred\_gen must be provied. It can be obtained from the

original genotype using convertgen function.

hybrid\_phe a data frame with three columns. The first column and the second column are the

names of male and female parents of the corresponding hybrids, respectively; the third column is the phenotypic values of hybrids. The names of male and female parents must match the rownames of inbred\_gen. Missing (NA) values

are not allowed.

inbred\_phe a matrix (n x 2) of inbred\_phe phenotypic.Default is NULL.

male\_name a vector of the names of male parents.

female\_name a vector of the names of female parents.

method eight GS methods including "GBLUP", "BayesB", "RKHS", "PLS", "LASSO",

"EN", "XGBoost", "LightGBM". Users may select P of these methods. Default

is "GBLUP".

model the prediction model. There are two options: model = "A" for the additive

model, model = "AD" for the additive-dominance model, model = "A-P" for the additive-phenotypic model, model = "AD-P" for the additive-dominance-

phenotypic model. Default is model = "A".

select the selection of hybrids based on the prediction results. There are three options:

select = "all", which selects all potential crosses. select = "top", which selects the top n crosses. select = "bottom", which selects the bottom n crosses. The n

is determined by the param number.

number the number of selected top or bottom hybrids, only when select = "top" or select

= "bottom".

#### Value

a data frame of prediction results with two columns. The first column denotes the names of male and female parents of the predicted hybrids, and the second column denotes the phenotypic values of the predicted hybrids.

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```
## load example data from hypred package
data(hybrid_phe)
data(input_geno)
inbred_gen <- convertgen(input_geno, type = "hmp2")

## infer the additive and dominance genotypes of hybrids
gena <- infergen(inbred_gen, hybrid_phe)$add
gend <- infergen(inbred_gen, hybrid_phe)$dom

pred<-predhy.predict_NCII(inbred_gen,hybrid_phe,method="LASSO",model="A")
pred<-predhy.predict_NCII(inbred_gen,hybrid_phe,method="LASSO",model = "AD",select="all")</pre>
```

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