# Package 'ImDiallel'

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Imports multcomp, plyr, sommer, tidyr	
<b>Description</b> Several service functions to be used to analyse datasets obtained from diallel experiments within the frame of linear models in R, as described in Onofri et al (2020) <doi:10.1007 s00122-020-03716-8="">.</doi:10.1007>	
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block	latrixDiagonal Creates block diagonal matrix. It is used internally.	

# Description

This function takes a list of matrices and creates a block diagonal matrix. It is used to fit multi-environment diallel models

# Usage

blockMatrixDiagonal(matList)

# Arguments

matList It is a list of matrices to be combined

#### Value

Returns a matrix object

## Author(s)

Andrea Onofri, Niccolo' Terzaroli, Luigi Russi

## References

DD 3

#### **Examples**

```
a <- matrix(1:16, 8, 2)
b <- matrix(1:9, 3, 3)
c <- list(a, b)
blockMatrixDiagonal(c)</pre>
```

DD

Dominant Deviation effect

## **Description**

DD effect to fit Hayman2 model with 1m function

## Usage

```
DD(P1, P2, type = "fix", data)
```

## **Arguments**

## **Details**

a design matrix of n-1 parentals

#### Value

A design matrix for the DD effect

# Author(s)

Andrea Onofri, Niccolo' Terzaroli, Luigi Russi

#### References

4 diallelMET

## **Examples**

diallelMET

Factitious dataset for Diallel analysis

# Description

Multi-environment half-diallel dataset with six parentals, in five blocks and ten environments; the dataset is factitious and was obtained by Monte Carlo simulation.

## Usage

```
data("diallelMET")
```

#### **Format**

A data frame with 1050 observations on the following 5 variables.

Env environment, a factor with 10 levels Block block, a factor with 5 levels Par1 male parent, a factor with 6 levels Par2 female parent, a factor with 6 levels Yield yield, a numeric vector

## Author(s)

Andrea Onofri, Niccolo' Terzaroli, Luigi Russi

#### **Source**

Onofri, A., Terzaroli, N. & Russi, L. Linear models for diallel crosses: a review with R functions. Theor Appl Genet (2020). https://doi.org/10.1007/s00122-020-03716-8

#### References

Onofri, A., Terzaroli, N. & Russi, L. Linear models for diallel crosses: a review with R functions. Theor Appl Genet (2020). https://doi.org/10.1007/s00122-020-03716-8

```
data("diallelMET")
```

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diallelMod-methods

Functions to retreive full list of genetical effects

## **Description**

Diallel model parameters are estimated under a set of restrictions and, therefore, the methods 'coef' and 'summary' do not return the full list of genetical parameters. Therefore, the 'glht.diallelMod' method can be used, which works by way of a series of helper functions, providing the necessary contrast matrices.

## Usage

```
## S3 method for class 'diallelMod'
glht(model, linfct, ...)
```

#### **Arguments**

model a model object (OPTIONAL)
linfct a diellel.eff() function
... Other optional arguments

#### **Details**

...

## Value

summary Returns the full list of genetical parameters

#### References

Onofri, A., Terzaroli, N. & Russi, L. Linear models for diallel crosses: a review with R functions. Theor Appl Genet (2020). https://doi.org/10.1007/s00122-020-03716-8

6 expand.diallel

expand.diallel

Create a Data Frame from All Combinations of Parentals

# Description

This is a modification of the 'expand.grid()' function working specifically with diallel experiments. It creates a data frame from all combinations of the supplied vector of parents, depending on the mating scheme.

## Usage

```
expand.diallel(pars, mating = 1)
```

#### **Arguments**

pars a vector of parentals

mating The type of mating scheme. 1: full diallel experiment; 2: no reciprocals; 3: no

selfs; 4: no reciprocals and no selfs

## Value

returns a data.frame object

## Author(s)

Andrea Onofri, Niccolo' Terzaroli, Luigi Russi

#### References

Onofri, A., Terzaroli, N. & Russi, L. Linear models for diallel crosses: a review with R functions. Theor Appl Genet (2020). https://doi.org/10.1007/s00122-020-03716-8

```
pars <- LETTERS[1:4]
expand.diallel(pars, mating = 3)</pre>
```

GCA 7

GCA

General Combining Ability effect

# Description

GCA effect to fit Hayman1 & 2 and Griffing 1 & 2 models with 1m function

## Usage

```
GCA(P1,P2,type = "fix", data)
```

# Arguments

P1 a variable for the first parent P2 a variable for the second parent

type a variable for model selection. May be "fix" (fixed model) or "random"

(random model).

data a 'data.frame' where to look for explanatory variables

#### **Details**

a design matrix of n-1 parentals

#### Value

A design matrix for the GCA effect

## Author(s)

Andrea Onofri, Niccolo' Terzaroli, Luigi Russi

#### References

Onofri, A., Terzaroli, N. & Russi, L. Linear models for diallel crosses: a review with R functions. Theor Appl Genet (2020). https://doi.org/10.1007/s00122-020-03716-8

8 GCAC

**GCAC** 

General Combining Ability without considering the selfed parents

## **Description**

Design matrix for GCAC, useful to fit Gardner & Eberhart model 3 (GE3) with 1m function

## Usage

```
GCAC(P1,P2,type = "fix", data)
```

## **Arguments**

P1 a variable for the first parent
P2 a variable for the second parent

type a variable for model selection. May be "fix" (fixed model) or "random"

(random model).

data a 'data.frame' where to look for explanatory variables

## **Details**

a design matrix of n-1 parentals

#### Value

A design matrix for the GCAC effect

#### Author(s)

Andrea Onofri, Niccolo' Terzaroli, Luigi Russi

## References

Onofri, A., Terzaroli, N. & Russi, L. Linear models for diallel crosses: a review with R functions. Theor Appl Genet (2020). https://doi.org/10.1007/s00122-020-03716-8

```
data("hayman54")
GCAC(Par1,Par2, data=hayman54)
```

griffing56

griffing56

Griffing's dataset for diallel analysis

## Description

Data for a diallel in maize, with no selfs and no selfed parents. Data are the means of several replicates.

## Usage

```
data("griffing56")
```

#### **Format**

A data frame with 36 observations on the following 5 variables

Par1 male parent, a factor with 8 levels

Par2 female parent, a factor with 8 levels

Yield Maize Yield

Cob Cob weight

Shelled Shelled corn weight

## Author(s)

Andrea Onofri, Niccolo' Terzaroli, Luigi Russi

#### Source

Griffing, B., 1956. Concept of general and specific combining ability in relation to diallel crossing systems. Australian Journal of Biological Science 9, 463–493.

#### References

Onofri, A., Terzaroli, N. & Russi, L. Linear models for diallel crosses: a review with R functions. Theor Appl Genet (2020). https://doi.org/10.1007/s00122-020-03716-8

10 H.BAR

H.BAR	Average heterosis effect

# Description

H.BAR effect to fit GE2 and GE3 models with 1m function

## Usage

```
H.BAR(P1, P2, type = "fix", data)
```

## **Arguments**

P1	a variable for the first parent
P2	a variable for the second parent
type	a variable for model selection. May be "fix" (fixed model) or "random" (random model).
data	a 'data.frame' where to look for explanatory variables

#### Value

A design matrix for the H.BAR effect

## Author(s)

Andrea Onofri, Niccolo' Terzaroli, Luigi Russi

## References

Onofri, A., Terzaroli, N. & Russi, L. Linear models for diallel crosses: a review with R functions. Theor Appl Genet (2020). https://doi.org/10.1007/s00122-020-03716-8

hayman54

hayman54

Hayman dataset for diallel analysis

## **Description**

Data for a diallel in tobacco with 2 reps

## Usage

```
data(hayman54)
```

#### **Format**

A data frame with 128 observations on the following 4 variables

Block block, a factor with 2 levels

Par1 male parent, a factor with 8 levels

Par2 female parent, a factor with 8 levels

Ftime mean flowering time (days), a numeric vector

## Author(s)

Andrea Onofri, Niccolo' Terzaroli, Luigi Russi

#### **Source**

B. I. Hayman (1954a). The Analysis of Variance of Diallel Tables. Biometrics, 10, 235-244. Table 5, page 241. http://doi.org/10.2307/3001877

#### References

Onofri, A., Terzaroli, N. & Russi, L. Linear models for diallel crosses: a review with R functions. Theor Appl Genet (2020). https://doi.org/10.1007/s00122-020-03716-8

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Ηi

Average heterosis effect

# **Description**

H.i effect to fit GE2 Model with 1m function

## Usage

```
Hi(P1, P2, type = "fix", data)
```

## **Arguments**

```
P1 a variable for the first parent
P2 a variable for the second parent
type a variable for model selection. May be "fix" (fixed model) or "random"
(random model).
```

data a 'data.frame' where to look for explanatory variables

#### **Details**

a design matrix of n-1 parentals

#### Value

A design matrix for the Hi effect

## Author(s)

Andrea Onofri, Niccolo' Terzaroli, Luigi Russi

#### References

Onofri, A., Terzaroli, N. & Russi, L. Linear models for diallel crosses: a review with R functions. Theor Appl Genet (2020). https://doi.org/10.1007/s00122-020-03716-8

int.matrix 13

int.matrix	Utilities for fitting diallel models.

## **Description**

These functions are used internally by the package, but they can also called from the outside, to fit specific needs

## Usage

```
int.matrix(Xa, Xb)
checkScheme(P1, P2)
emm.diallel(obj)
```

#### **Arguments**

Ха	Incidence matrix of genetic effects
Xb	Incidence matrix for an external factor
P1	A vector with parentals
P2	A vector with parentals
obj	A glht object

#### **Details**

The function 'int.matrix()' produces the incidence matrix for the interaction between two main effects; 'Xa' and 'Xb' are two incidence matrices for two main effects. The function 'checkScheme()' takes two vectors containing the codings for parentals (P1 and P2), retrieves the mating scheme and detects whether there are missing crosses. The function 'emm.diallel()' is used with multi-environment diallel experiments to obtain the expected marginal means for genetic effects across environments.

#### Value

The function 'int.matrix()' returns an incidence matrix. The function 'checkScheme()' returns a list, containing the main traits of the mating scheme. The function 'emm.diallel()' retrns a data.frame with the marginal means, standard errors and t-test statistics.

## Note

No further notes

## Author(s)

Andrea Onofri

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#### References

Onofri, A., Terzaroli, N. & Russi, L. Linear models for diallel crosses: a review with R functions. Theor Appl Genet (2020). https://doi.org/10.1007/s00122-020-03716-8

#### **Examples**

```
data(griffing56)
head(griffing56)
checkScheme(griffing56$Par1, griffing56$Par2)
```

lm.diallel

Fitting diallel linear models

## **Description**

Wrapper function for lm.fit and diallel models. It can be used to carry out several powerful methods for linear models, such as 'summary()', anova() or 'glht()' in the 'multcomp' package.

## **Usage**

```
lm.diallel(formula, Block, Env, fct = "GRIFFING2", data)
```

## **Arguments**

fct

formula an object of class "formula" (or one that can be coerced to that class): a symbolic description of the model to be fitted.' formula' uses the regular R syntax to specify the response variable and the two variables for parentals (e.g., Yield ~ Par1 + Par2)

Block used to specify an optional variable coding for blocks Env used to specify an optional variable coding for environments

a string variable coding for the selected model. 8 main diallel models: Hayman's model 1 (="HAYMAN1"), Hayman's model 2 (="HAYMAN2"), Griffing's model 1 (="GRIFFING1"), Griffing's model 2 (="GRIFFING2"), Griffing's model 3 (="GRIFFING3"), Griffing's model 4 (="GRIFFING4"), Gardner-Eberhart model 2 (="GE2") and Gardner-Eberhart model 3 (="GE3"). The strings "GE2r" and "GE3r" can be used to specify the 'enhanced' GE2 and GE3

models, including the effect of reciprocals (REC).

data a 'data.frame' where to look for explanatory variables

## **Details**

Notations for the 8 models

Model name in 'lm.diallel()' Model notation in 'lm()'

HAYMAN1 Y ~ GCA(Par1, Par2) + tSCA(Par1, Par2) + RGCA(Par1, Par2) + RSCA(Par1, Par2)

**GRIFFING1**  $Y \sim GCA(Par1, Par2) + tSCA(Par1, Par2) + REC(Par1, Par2)$  Im.diallel 15

```
\begin{array}{lll} GRIFFING2 & Y \sim GCA(Par1, Par2) + tSCA(Par1, Par2) \\ HAYMAN2 & Y \sim GCA(Par1, Par2) + MDD(Par1, Par2) + DD(Par1, Par2) + SCA(Par1, Par2) + RGCA(Par1, Par2) + WE.i(Par1, Par2) + WE.i(Par1, Par2) + WE.i(Par1, Par2) + SCA(Par1, Par2) + SCA(Par1, Par2) \\ GE3 & Y \sim H.BAR(Par1, Par2) + SP(Par1, Par2) + GCAC(Par1, Par2) + SCA(Par1, Par2) \\ GE2r & Y \sim H.BAR(Par1, Par2) + VE.i(Par1, Par2) + H.i(Par1, Par2) + SCA(Par1, Par2) + RGCA(Par1, Par2) \\ GE3r & Y \sim H.BAR(Par1, Par2) + SP(Par1, Par2) + GCAC(Par1, Par2) + SCA(Par1, Par2) + RGCA(Par1, Par2) \\ \end{array}
```

#### Value

lm.diallel returns an object of class c("diallel", "lm"), that is a list containing at least the following components:

coefficients a named vector of coefficients

residuals the residuals, that is response minus fitted values

fitted.values the fitted mean values

rank the numeric rank of the fitted linear models

weights (only for weighted fits) the specified weights

df.residual the residual degrees of freedom

call the matched call

terms the terms object used

contrasts (only where relevant) the contrasts used

xlevels (only where relevant) a record of the levels of the factors used in fitting

call the matched call

offset the offset used (missing if none were used)

y if requested, the response used

x if requested, the model matrix used

model if requested (the default), the model frame used

na.action (where relevant) information returned by model.frame on the special handling of NAs

## Author(s)

Andrea Onofri, Niccolo' Terzaroli, Luigi Russi

#### References

Onofri, A., Terzaroli, N. & Russi, L. Linear models for diallel crosses: a review with R functions. Theor Appl Genet (2020). https://doi.org/10.1007/s00122-020-03716-8

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lmDiallel-methods

Methods for diallel model fitting

## Description

The object returned by the 'lm.diallel()' function is of classes 'lm' and 'diallel'. Specific methods were devised to explore the 'diallel' object.

#### Usage

```
## S3 method for class 'diallel'
summary(object, MSE, dfr, ...)
## S3 method for class 'diallel'
vcov(object, MSE, ...)
## S3 method for class 'diallel'
anova(object, MSE, dfr, type = 1, ...)
## S3 method for class 'diallel'
model.matrix(object, ...)
```

## Arguments

object an object of class diallel.

MSE Mean Square Error, when it cannot be derived from model fit

dfr Residual degrees of freedom, when they cannot be derived from model fit

type It is used to select between Type I (sequential) or Type III (marginal) F tests in

**ANOVA** 

... Other optional arguments

## **Details**

To be defined

lonnquist61

#### Value

vcov.diallel: a variance-covariance matrix summary.diallel: a data.frame of estimated parameters with standard errors anova.diallel: an ANOVA table predict.diallel: a vector of predictions from a diallel model model.matrix.diallel: a design matrix for the fitted diallel model

#### References

Onofri, A., Terzaroli, N. & Russi, L. Linear models for diallel crosses: a review with R functions. Theor Appl Genet (2020). https://doi.org/10.1007/s00122-020-03716-8

## **Examples**

lonnquist61

Half diallel of maize dataset

## **Description**

Diallel experiment with six maize varieties and no reciprocals. The data here are means adjusted for block effects.

## Usage

```
data("lonnquist61")
```

## **Format**

A data.frame with 21 observations on the following 3 variables.

Par1 male parent, a factor with 6 levels
Par2 female parent, a factor with 6 levels
Yield mean across blocks, a numeric vector

# Author(s)

Andrea Onofri, Niccolo' Terzaroli, Luigi Russi

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#### Source

J. H. Lonnquist, C. O. Gardner. (1961) Heterosis in Intervarietal Crosses in Maize and Its Implication in Breeding Procedures. Crop Science, 1, 179-183. Table 1.

#### References

Onofri, A., Terzaroli, N. & Russi, L. Linear models for diallel crosses: a review with R functions. Theor Appl Genet (2020). https://doi.org/10.1007/s00122-020-03716-8

Mohring, Melchinger, Piepho. (2011). REML-Based Diallel Analysis. Crop Science, 51, 470-478. http://doi.org/10.2135/cropsci2010.05.0272

C. O. Gardner and S. A. Eberhart. 1966. Analysis and Interpretation of the Variety Cross Diallel and Related Populations. Biometrics, 22, 439-452. http://doi.org/10.2307/2528181

## **Examples**

matBlock

Design matrix for blocks

## **Description**

It creates a disign matrix for block effects (with sum-to-zero constraint). It is used internally

## Usage

```
matBlock(formula)
```

## **Arguments**

formula

a formula containing the block variables

## Value

A design matrix for the block effect

## Author(s)

Andrea Onofri, Niccolo' Terzaroli, Luigi Russi

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#### References

Onofri, A., Terzaroli, N. & Russi, L. Linear models for diallel crosses: a review with R functions. Theor Appl Genet (2020). https://doi.org/10.1007/s00122-020-03716-8

## **Examples**

```
data("hayman54")
matBlock(~hayman54$Block)
```

MDD

Mean Dominance Deviation effect

## **Description**

It relates to the difference between the average yield of selfed parents and the average yield of crosses. DD effect to fit Hayman2 model with 1m function

# Usage

```
MDD(P1, P2, type = "fix", data)
```

## **Arguments**

P1 a variable for the first parent P2 a variable for the second parent

type a variable for the model

data a 'data.frame' where to look for explanatory variables

#### Value

A design matrix for the MDD effect

# Author(s)

Andrea Onofri, Niccolo' Terzaroli, Luigi Russi

#### References

Onofri, A., Terzaroli, N. & Russi, L. Linear models for diallel crosses: a review with R functions. Theor Appl Genet (2020). https://doi.org/10.1007/s00122-020-03716-8

```
data("hayman54")
MDD(Par1, Par2, data = hayman54)
```

20 mmer.diallel

mmer.diallel	Fitting random diallel linear models	
mmer . drdirer	T titing random diditer tinear models	

# Description

Wrapper function for the function 'mmer()' in the 'sommer' package. It can be used to fit random diallel models and retreive variance components for main effects.

# Usage

```
mmer.diallel(formula, Block, Env, fct, data, type = "all")
```

# Arguments

an object of class "formula" (or one that can be coerced to that class): a symbolic description of the model to be fitted. formula' uses the regular R syntax to specify the response variable and the two variables for parentals (e.g., Yield ~ Par1 + Par2)
used to specify an optional variable coding for blocks
used to specify an optional variable coding for environments
a 'data.frame' where to look for explanatory variables
a string variable coding for the selected model. 8 main diallel models: Hayman's model 1 (="HAYMAN1"), Hayman's model 2 (="HAYMAN2"), Griffing's model 1 (="GRIFFING1"), Griffing's model 2 (="GRIFFING2"), Griffing's model 3 (="GRIFFING3"), Griffing's model 4 (="GRIFFING4"), Gardner-Eberhart model 2 (="GE2") and Gardner-Eberhart model 3 (="GE3"). The strings "GE2r" and "GE3r" can be used to specify the 'enhanced' GE2 and GE3 models, including the effect of reciprocals (REC).
a string variable coding for the selected model. It is only used for multi-environment experiments and it is equal to "all" when both the environment and genetical effects are random or "environment" when the environment is random and genetical effects are fixed.

#### Value

Returns a data frame of variance components with standard errors

# Author(s)

Andrea Onofri, Niccolo' Terzaroli, Luigi Russi

# References

Covarrubias-Pazaran, G., 2016. Genome-Assisted Prediction of Quantitative Traits Using the R Package sommer. PLOS ONE 11, e0156744. https://doi.org/10.1371/journal.pone.0156744

model.matrixDiallel 21

## **Examples**

model.matrixDiallel

Incidence matrices for Diallel model parametrisation

## **Description**

model.matrixDiallel is useful to build design matrices, according to the user-defined (or default) parameterisation for lm function. It shares the same syntax of the lm.diallel function.

# **Arguments**

formula	an object of class "formula" (or one that can be coerced to that class): a symbolic description of the model to be fitted.'formula' uses the regular $R$ syntax to specify the response variable and the two variables for parentals (e.g., Yield $\sim$ Par1 + Par2)	
Block	used to specify an optional variable coding for blocks	
Env	used to specify an optional variable coding for environments	
data	a 'data.frame' where to look for explanatory variables	
fct	a string variable coding for the selected model. 6 main diallel models: H man's model 1 (="HAYMAN1"), Hayman's model 2 (="HAYMAN2"), Gring's model 1 (="GRIFFING1"), Griffing's model 2 (="GRIFFING2"), Gardr Eberhart model 2 (="GE2") and Gardner-Eberhart model 3 (="GE3"). Strings "GE2r" and "GE3r" can be used to specify the 'enhanced' GE2 and G	

## **Details**

model.matrixDiallel creates a design matrix for a diallel model, as specified in the 'fct' argument.

models, including the effect of reciprocals (REC).

## Value

The design matrix for a diallel model as specified in the 'fct' argument.

#### Author(s)

Andrea Onofri, Niccolo' Terzaroli, Luigi Russi

## References

22 REC

## **Examples**

REC

Reciprocal Effect not parted into RGCA and RSCA

## **Description**

Build incidence matrix to fit reciprocal effects in Griffing's model 1, 2, 4 (REC) and 3 (REC.G3) with 1m function

## Usage

```
REC(P1, P2, type = "fix", data)
```

# Arguments

P1 a variable for the first parent P2 a variable for the second parent

type a variable for model selection. May be "fix" (fixed model) or "random"

(random model).

data a 'data.frame' where to look for explanatory variables

#### **Details**

a design matrix of n-1 parentals

## Value

A design matrix for the reciprocal (REC) effect

## Author(s)

Andrea Onofri, Niccolo' Terzaroli, Luigi Russi

#### References

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## **Examples**

**RGCA** 

Reciprocal General Combining Ability

# Description

RGCA effect to fit Hayman1 & 2 models with 1m function

## Usage

```
RGCA(P1,P2,type = "fix", data)
```

## Arguments

P1 a variable for the first parent P2 a variable for the second parent

type a variable for model selection. May be "fix" (fixed model) or "random"

(random model).

data a 'data.frame' where to look for explanatory variables

#### **Details**

a design matrix of n-1 parentals

## Value

A design matrix for the RGCA effect

#### Author(s)

Andrea Onofri, Niccolo' Terzaroli, Luigi Russi

#### References

24 RSCA

## **Examples**

**RSCA** 

Reciprocal Specific Combining Ability

# Description

RSCA effect to fit Hayman 1 & 2 models with 1m function

## Usage

```
RSCA(P1,P2,type = "fix",data)
```

## Arguments

P1 a variable for the first parent P2 a variable for the second parent

type a variable for model selection. May be "fix" (fixed model) or "random"

(random model).

data a 'data.frame' where to look for explanatory variables

#### **Details**

a design matrix of all possible combinations between parentals with no selfs and no reciprocals

## Value

A design matrix for the RSCA effect

#### Author(s)

Andrea Onofri, Niccolo' Terzaroli, Luigi Russi

#### References

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## **Examples**

SCA

Specific Combining Ability

## **Description**

SCA effect to fit Hayman2, Griffing3 (SCA.G3), GE2 and GE3 (SCA.GE) models with 1m function

## Usage

```
SCA(P1, P2, type = "fix", data)
SCA.G3(P1, P2, type = "fix", data)
```

## **Arguments**

P1	a variable for the first parent
P2	a variable for the second parent
type	a variable for model selection. May be "fix" (fixed model) or "random" (random model).
data	a 'data.frame' where to look for explanatory variables

#### **Details**

a design matrix of all possible combinations between parentals with no selfs and no reciprocals

#### Value

A design matrix for the SCA effect

# Author(s)

Andrea Onofri, Niccolo' Terzaroli, Luigi Russi

#### References

26 SP

#### **Examples**

SP

Selfed Parents effect

## **Description**

SP effect to fit GE3 model with 1m function

## Usage

```
SP(P1, P2, type = "fix", data)
```

# Arguments

P1 a variable for the first parent
P2 a variable for the second parent
type a variable for model selection. May be "fix" (fixed model) or "random"
(random model).

a 'data.frame' where to look for explanatory variables

## Details

data

a design matrix of n-1 parentals

#### Value

A design matrix for the SP effect

## Author(s)

Andrea Onofri, Niccolo' Terzaroli, Luigi Russi

#### References

Onofri, A., Terzaroli, N. & Russi, L. Linear models for diallel crosses: a review with R functions. Theor Appl Genet (2020). https://doi.org/10.1007/s00122-020-03716-8

```
data("hayman54")
SP(Par1,Par2, data=hayman54)
```

tSCA 27

tSCA

Total Specific Combining Ability

## **Description**

Total SCA to fit Hayman1, Griffing1 and Griffing2 models with 1m function

# Usage

```
tSCA(P1,P2,type = "fix", data)
```

## Arguments

P1 a variable for the first parent
P2 a variable for the second parent
type a variable for model selection. May be "fix" (fixed model) or "random"
(random model).

data a 'data.frame' where to look for explanatory variables

#### **Details**

a design matrix of all possible combinations between parentals with selfs but no reciprocals

## Value

A design matrix for the tSCA effect

#### Author(s)

Andrea Onofri, Niccolo' Terzaroli, Luigi Russi

## References

Onofri, A., Terzaroli, N. & Russi, L. Linear models for diallel crosses: a review with R functions. Theor Appl Genet (2020). https://doi.org/10.1007/s00122-020-03716-8

28 VEi

VEi Variety Effect

# **Description**

VE.i effect to fit GE2 model with 1m function

## Usage

```
VEi(P1, P2, type = "fix", data)
```

## **Arguments**

P1 a variable for the first parent
P2 a variable for the second parent

type a variable for model selection. May be "fix" (fixed model) or "random"

(random model).

data a 'data.frame' where to look for explanatory variables

#### **Details**

a design matrix of n-1 parentals

#### Value

A design matrix for the VEi effect

## Author(s)

Andrea Onofri, Niccolo' Terzaroli, Luigi Russi

#### References

Onofri, A., Terzaroli, N. & Russi, L. Linear models for diallel crosses: a review with R functions. Theor Appl Genet (2020). https://doi.org/10.1007/s00122-020-03716-8

zhang05

zhang05

Data for diallel analysis from Zhang (2005)

## **Description**

Data collected in XXX with 5 parents, 2 reps and 2 environments

# Usage

```
data("zhang05")
```

#### **Format**

A data frame with 60 observations on the following 6 variables.

Par1 male parent, a factor with 5 levels

Par2 female parent, a factor with 5 levels

Block block, a factor with 2 levels

Combination combination between environment and block, an integer vector

Env environment, a factor with 2 levels

Yield yield, a numeric vector

## Author(s)

Andrea Onofri, Niccolo' Terzaroli, Luigi Russi

## **Source**

Zhang, Y., Kang, M.S. and Lamkey, K.R. (2005), DIALLEL-SAS05: A Comprehensive Program for Griffing's and Gardner&Eberhart Analyses. Agron. J., 97: 1097-1106. https://doi.org/10.2134/agronj2004.0260

## References

Onofri, A., Terzaroli, N. & Russi, L. Linear models for diallel crosses: a review with R functions. Theor Appl Genet (2020). https://doi.org/10.1007/s00122-020-03716-8

30 zhang05

```
 \label{eq:dMod2} $$ dMod2 <- lm(Yield ~ Env/Block + H.BAR(Par1, Par2) + VEi(Par1, Par2) + Hi(Par1, Par2) + SCA(Par1, Par2), data = zhang05) $$ summary(dMod2)$$ coefficients
```

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