# Package 'designGG'

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**Title** Computational tool for designing genetical genomics experiments.

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**Description** The package provides R scripts for designing genetical genomics experiments.

**Depends** R (>= 2.2.0)

License GPL

URL http://gbic.biol.rug.nl/designGG

# R topics documented:

acceptanceProbability	2
arrayUpdate	3
conditionAllocation	4
conditionCombination	5
conditionLevel	6
conditionUpdate	7
designGG	8
designScore	11
exampleArrayDesignTable	13
exampleConditionDesignTable	14
examplePlotObj	15
experimentDesignTable	16
genotype	18
initialDesign	18
interactionLevel	20
pairLevel	21
plotAllScores	22
temperatureStep	23
updateDesign	24
variableNames	25
variableNumber	26

Index 28

```
acceptanceProbability
```

Compute the acceptance probability for each updated design

# **Description**

Compute the acceptance probability for each updated design. It depends on the current temperature value of simulated annealing process. This is a subfunction needed for designGG, but is not directly used.

# Usage

# Arguments

```
designScore score of current design.

newDesignScore
score of updated design.

method either "SA" (simulated annealing) or "MH". (Metropolis Hastings)

temperature current temperature in simulated annealing process.
```

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#### References

E. Wit and J. McClure. Statistics for Microarrays: Design, Analysis and Inference. (2004) Chichester: Wiley.

Y. Li, R. Breitling and R.C. Jansen. Generalizing genetical genomics: the added value from environmental perturbation, Trends Genet (2008) 24:518-524.

Y. Li, M. Swertz, G. Vera, J. Fu, R. Breitling, and R.C. Jansen. designGG: An R-package and Web tool for the optimal design of genetical genomics experiments. (submitted) http://gbic.biol.rug.nl/designGG

### See Also

designGG

arrayUpdate 3

|--|

### **Description**

Update the allocation of samples on the arrays. This is a subfunction needed for updateDesign, but is not directly used.

### Usage

```
arrayUpdate(array.allocation, condition.allocation, nRILs, nSlides)
```

### **Arguments**

```
array.allocation
```

matrix with nArray rows and nRIL columns. Elements of 1/0 indicate this RIL

(or strain) is/not selected for this array.

condition.allocation

matrix with nCondition rows and nRIL columns. Elements of 1/0 indicate this

RIL (or strain) is/not selected for this condition.

nRILs number of RILs or strains available for the experiment.

nSlides total number of slides available for experiment.

#### **Details**

This function is used only for designing a dual-channel experiment where samples need to be paired.

# Value

```
A list with the following two elements:
```

```
new.array.allocation: an updated array allocation table new.condition.allocation: an updated condition allocation table
```

### Author(s)

```
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```

# References

Y. Li, R. Breitling and R.C. Jansen. Generalizing genetical genomics: the added value from environmental perturbation, Trends Genet (2008) 24:518-524.

Y. Li, M. Swertz, G. Vera, J. Fu, R. Breitling, and R.C. Jansen. designGG: An R-package and Web tool for the optimal design of genetical genomics experiments. (submitted) http://gbic.biol.rug.nl/designGG

#### See Also

```
updateDesign
```

4 conditionAllocation

conditionAllocation

Allocate the selected RILs into different conditions

### **Description**

This is a subfunction used by initialDesign but is not directly used. In the experiment where samples are profiled in pairs, the samples are firstly selected and paired on each array and then the selected samples are randomly allocated into different conditions.

#### Usage

conditionAllocation( selectedRILs, genotype, nConditions, nSlides, nTuple )

#### **Arguments**

selectedRILs the index of the selected RILs or strains among all that are available for the

experiment.

genotype genotype data: a nMarker-by-nRILs matrix with two allels being 0 and 1 (or A

and B) or three allels being 0, 0.5 and 1 (or, A, H, and B), where 0.5 (or H)

represents heterozygous allele.

nConditions number of all possible combination of all environmental factors. It should be

larger than 1.

nSlides total number of slides available for the experiment. It should be a non-zero

integer.

nTuple average number of RILs to be assigned onto each condition.

 $\verb"nTuple" should be a real number which is larger than 1.$ 

if nTuple < 1, the algorithm will stop and show a message as below,

warning: "The number of slides is too small to perform

the experiment."

### **Details**

This function is only called by initialDesign function when btwoColorArray is TRUE.

#### Value

A matrix with nCondition rows and nRIL columns. Elements of 1/0 indicate that this RIL (or strain) is/not selected for this condition.

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# References

Y. Li, R. Breitling and R.C. Jansen. Generalizing genetical genomics: the added value from environmental perturbation, Trends Genet (2008) 24:518-524.

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#### See Also

initialDesign

conditionCombination

Generate a matrix indicating all possible levels for environmental factors

# Description

Generate a matrix indicating all possible levels for environmental factors with dimension nConditions × nEnvFactors. This is a subfunction needed for designScore, but is not directly used.

# Usage

```
conditionCombination( nEnvFactors, nLevels, Level, envFactorNames )
```

# **Arguments**

nEnvFactors

number of environmental factors, an integer bewteen 1 and 3. When nEnvFactors is 1 and the number of levels for the environmental factor (nLevels) is 1, there is one condition in the experiment (i.e. no environmental perturbation) and thus only genetic factor will be considered in the algorithm. When nEnvFactors is 1 and nLevels is larger than 1 or nEnvFactors is larger than 1, all main factor(s) and interacting facotr(s) will be included. Examples: If there is a temperature perturbation, then nEnvFactors is 1; If there is both temperature and drug treatment perturbation, then nEnvFactors is 2.

nLevels

number of levels for each factor, a vector with each component being integer. The length should be equal to nEnvFactors.

Level

a list which specifies the levels for each factor in the experiment. There are in total nEnvFactors elements in the list and each element corresponds to certain environmental factor. The element is a vector describing all levels of the environmental factor. Default setting for the level of each factor is 1, 2, ..., nLevels[i]. (Here nLevels[i] is the ith element of nLevels, which tells the total number of levels for i environmental factor).

envFactorNames

a vector with names for all environmental factor(s). For example, for an experiment with two environmental factors of temperature and drug treatment: envFactorNames <- c( "Temperature", "Dosage") Default = NULL, then the output will use "F1" and "F2" to indicate the environmental factors.

# Details

Currently this function works only when nEnvFactors is between 1 and 3.

### Value

A matrix with dimension of nConditions  $\times$  nEnvFactors. Each element in the matrix indicates the levels of corresponding environmental factor.

6 conditionLevel

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#### See Also

designScore

conditionLevel

Levels of all environmental factors

# Description

Describe the levels of all environmental factors for each RIL/strain in the experiment. This is a subfunction needed for designScore, but is not directly used.

# Usage

# **Arguments**

array.allocation

a matrix with nArray rows and nRIL columns. Elements of 1/0 indicates this RIL (or strain) is/not selected for this array.

condition.allocation

a matrix with nCondition rows and nRIL columns. Elements of 1/0 indicates this RIL (or strain) is/not selected for this condition.

condition.combination

a matrix indicating all possible levels for environmental factors, with dimension of nConditions by nEnvFactors.

nEnvFactors

number of environmental factors, an integer bewteen 1 and 3. When nEnvFactors is 1 and nLevels is 1, there is one condition in the experiment (i.e. no environmental perturbation) and thus only genetic factor will be considered in the algorithm. When nEnvFactors is 1 and nLevels is larger than 1 or nEnvFactors is larger than 1, all main factor(s) and interacting facotr(s) will be included.

#### **Details**

For single-channel experiment, array.allocation is NULL. Then the conditionLevel is decided by condition.allocation. For dual-channel experiment, array.allocation decides which RILs are selected and then the condition.allocation indicates which condition this RIL will be put in for the experiment.

conditionUpdate 7

#### Value

A matrix with dimension of nRILs by nEnvFactors, each element indicates the level of a certain environmental factor to which the RIL (or strain) is exposed in the experiment.

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#### See Also

designScore, conditionCombination

conditionUpdate

Update condition allocation

# **Description**

Update the allocation of samples onto different conditions. This is a subfunction needed for updateDesign, but is not directly used.

### Usage

```
conditionUpdate( condition.allocation, nTuple )
```

### **Arguments**

condition.allocation

a matrix with nCondition rows and nRIL columns. elements of 1/0 indicate this

RIL (or strain) is/not selected for this condition.

nTuple average number of RILs (or strains) to be assigned onto each condition

nTuple should be a real number which is larger than 1.

if nTuple < 1, the algorithm will stop and show the message,

warning: "The number slides is too less to perform the

experiment."

#### **Details**

This function will be used both in single and dual channel experiment design.

#### Value

An updated condition.allocation table.

8 designGG

#### Author(s)

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### See Also

arrayUpdate, designGG

designGG

Optimal design for genetical genomics experiments

# Description

Main function to search and display A- and D- optimal designs for single- or two-channel genetical genomics experiments. Simulated annealing or Metropolis Hastings used to find the best design.

#### Usage

# **Arguments**

genotype	genotype data: a nMarker-by-nRILs matrix with two allels being 0 and 1 (or A and B) or three allels being 0, 0.5 and 1 (or, A, H, and B), where 0.5 (or H) represents heterozygous allele.
nSlides	total number of slides available for the experiment.
nTuple	average number of RILs (or strains) to be assigned onto each condition.  nTuple should be a real number which is larger than 1.  If nTuple < 1, the algorithm will stop and show the message,  warning: "The number of slides is too small to perform the experiment."
nEnvFactors	number of environmental factors, an integer bewteen 1 and 3. When $nEnvFactors$ is 1 and the number of levels for the environmental factor ( $nLevels$ ) is 1, there is one condition in the experiment (i.e. no environmental perturbation) and thus

is 1 and the number of levels for the environmental factor (nLevels)is 1, there is one condition in the experiment (i.e. no environmental perturbation) and thus only genetic factor will be considered in the algorithm. When nEnvFactors is 1 and nLevels is larger than 1 or nEnvFactors is larger than 1, all main factor(s) and interacting facotr(s) will be included. Examples: If there is a temperature perturbation, then nEnvFactors is 1; If there is both temperature and drug treatment perturbation, then nEnvFactors is 2.

designGG 9

nLevels number of levels for each factor, a vector with each component being an integer.

The length of it should equal nEnvFactors.

Level a list which specifies the levels for each factor in the experiment. There are

in total nEnvFactors elements in the list and each element corresponds to a certain environmental factor. The element is a vector describing all levels of the environmental factor. default setting for the level of each factor is 1, 2, ...., nlevels[i]. (Here nLevels[i] is the *i*th element of nLevels, which gives the total

number of levels for i environmental factor).

bTwoColorArray

binary variable indicating experiment type:

bTwoColorArray <- T #for dual channel experiment bTwoColorArray <- F #for single channel experiment

initial the starting design matrix for the algorithm. If specified, this should be a list

with 2 matrices:

 $\verb|condition.allocation: allocate RILs (or strains) into different condition.|\\$ 

tional (nrow = nCondition, ncol= nRILs)

array.allocation: pair RILs (or strains) into sldies (nrow = nSlide, ncol

= nRILs)

However, the algorithm does not require that a starting matrix is specified. De-

fault = NULL.

weight a vector with length of variable Number which is calculated from function

variableNumber. Default = 1 (which means the parameters to be estimated

are all equally important during optimization). See details below.

region genome region of biological interest. Default = NULL (which means the entire

genome will considered).

optimality type of optimality, i.e. "A" (A-optimality) or "D" (D-optimality). A-optimality

minimizes  $Trace((X'X)^{-1})$ , which corresponds to minimum average variance of the parameter estimates. D-optimality minimizes  $det(X'X)^{-1}$ , which corre-

sponds to minimum generalized variance of the parameter estimates.

method method for searching for an optimal design. "SA" uses simulated annealing.

"MH" uses Metropolis Hasting. Default = "SA".

nIterations number of iterations of the simulated annealing method. Default = 3000.

n.search number of times for simulated annealing optimaization with different initial de-

sign, default = 2. Here it is suggested to be between 1 and 5. It should not to be

too large because of the reaching computational burden.

endTemp ending temperature of simulated annealing process. An important optimization

parameter. Default =  $1e^{-10}$ .

 $\verb|startTemp| starting temperature of simulated annealing process. Default = 1.$ 

maxTempStep maximum temperature decreasing step for simulated annealing process. The

parameter ensures that the multiplicative cooling factor is not smaller than that. If nlterations is too small, the preferred final temperature (endTemp) may

not be reached. See Wit and McClure (2004) for details. Default = 0.9.

plot Scores If TRUE (default) it produces a plot of the optimazation by SA using the function

plotAllScores.

directory It tells where the resulting optimal design tables are to be stored. If NULL (de-

fault), it will take currect working directory.

fileName the final optimal design table(s) in csv format and a plot (in png format) of all

scores during SA process (if plotScores = T) will be produced. The users can specify the table and plot name by setting fileName. If NULL (default) it

produces files starting with "myDesignGG".

10 designGG

envFactorNames

a vector with names for all environmental factor(s). For example, for the experiment with two environmental factors of temperature and drug treatment:  $\verb|envFactorNames| <- c( "Temperature", "Dosage") |$ 

Default = NULL, then the output will use "F1" and "F2" to indicate the environmental factors.

writingProcess

If TRUE, it prints how much computation work has been finished in a file called "processing.txt". Default = TRUE.

#### **Details**

Given the genetic information of samples available for the experiment (genotype) and the information about experimental settings (nEnvFactors, nSlides,nLevels etc.), the algorithm searches for an A-optimal or D-optimal (see optimality) using simulated annealing (see method). A plot of the scores at each iterations can also be given using the plotAllScores function. It also contains a number of the arguments:

region is used to specify the genome region that are of major interest to experimenters. weight is used to define the weight of genetic and environmental factors, and interaction terms. Prior knowledge about expected effect sizes of interesting factors can also be incorporated as weight parameters for the algorithm. The weight is inversely proportional to the expected effect size of the corresponding parameter. Example parameter settings: Suppose to design an experiment with two environmental factors (F1, F2) and there are two diffferent levels for each environment. The levels are 16 and 24 for F1, and 5 and 10 for F2. Thus the following command can be used:

```
nEnvFactors <- 2 nLevels <- c ( 2, 2 ) levels <- list ( c(16, 24), c(5, 10) ) The length of parameter weight is dependent on the number of environmental factors: When nEnvFactor = 0, weight is 1 as there is only one parameter of interest (genotype). When nEnvFactor = 1, weight = c(w_Q, w_{F1}, w_{QF1}) When nEnvFactor = 2, weight = c(w_Q, w_{F1}, w_{F2}, w_{QF1}, w_{QF2}, w_{F1F2}, w_{QF1F2}) When nEnvFactor = 3, weight = c(w_Q, w_{F1}, w_{F2}, w_{F2}, w_{F2}, w_{F2}, w_{F1F2}, w_{F1F2}, w_{F1F3}, w_{F2F3}, w_{F2F3}, w_{F1F2}, w_{F1F3}, w_{F2F3}, w_{F1F2}, w_{F1F2}, w_{F1F3}, w_{F2F3}, w_{F1F2}, w_{F1F2}, w_{F1F3}, w_{F2F3}, w_{F1F2}, w_{F1F2}, w_{F1F3}, w_{F1F3}, w_{F1F2}, w_{F1F3}, w_{F1F3}, w_{F1F3}, w_{F1F2}, w_{F1F3}, w_{
```

Here  $w_Q$  represents the weight for genotype effect,  $w_{F1}$  represent the weight for F1 effect and  $w_{QF1}$  represent the weight for interaction between genotype and F1 effect, etc.

# Value

An array design table (arrayDesign.csv) and a condition design table (conditionDesign.csv) are generated.

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### References

Y. Li, M. Swertz, G. Vera, J. Fu, R. Breitling, and R.C. Jansen. designGG: An R-package and Web tool for the optimal design of genetical genomics experiments. (submitted)

designScore 11

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E. Wit and J. McClure. Statistics for Microarrays: Design, Analysis and Inference. (2004) Chichester: Wiley.

#### See Also

```
initialDesign, designScore, updateDesign, acceptanceProbability,
experimentDesignTable, plotAllScores,
exampleArrayDesignTable,exampleConditionDesignTable,
```

### **Examples**

```
library(designGG)
#load genotype data
data (genotype)
#Example: single-channel experiment with 2 environmental factors,
#each with 2 level, and there will be four samples per condition(nTuple=4).
optimalDesign <- designGG ( genotype, nSlides=NULL, nTuple=4, nEnvFactors=2,
                      nLevels=c(2,2), Level=list(c(16,24),c(5,10)), bTwoColorArray=F,
                      initial=NULL, weight=1, region=seq(1,20), optimality="A",
                      method="SA", nIterations=100, n.search=2, endTemp=1e-10,
                      startTemp=1, maxTempStep=0.9, plotScores=T,
                      directory=NULL, fileName=NULL, envFactorNames=NULL,
                      writingProcess=F )
#Example 2: dual-channel experiment with 2 environmental factors,
#each with 2 level. There are 50 slides available.
optimalDesign <- designGG ( genotype, nSlides=50, nTuple=NULL, nEnvFactors=2,
                      nLevels=c(2,2), Level=list(c(16,24),c(5,10)), bTwoColorArray=T,
                      initial=NULL, weight=1, region=seq(1,20), optimality="A",
                      method="SA", nIterations=100, n.search=2, endTemp=1e-10,
                      startTemp=1, maxTempStep=0.9, plotScores=T,
                      directory=NULL, fileName=NULL, envFactorNames=NULL,
                      writingProcess=F )
#result
optimalDesign$arrayDesign
optimalDesign$conditionDesign
plotAllScores(optimalDesign$plot.obj)
#Use the following commands to see example output tables:
data(exampleArrayDesignTable)
exampleArrayDesignTable
data(exampleConditionDesignTable)
exampleConditionDesignTable
```

designScore

Calculate the A- or D- optimality score based on current experimental design

# **Description**

According to the current experimental design, the Fisher information matrix is obtained and then either the A- or D- optimality score is computed.

12 designScore

#### **Usage**

#### **Arguments**

genotype

genotype data: a nMarker-by-nRILs matrix with two allels being 0 and 1 (or A and B) or three allels being 0, 0.5 and 1 (or, A, H, and B), where 0.5 (or H) represents heterozygous allele.

array.allocation

matrix with nArray rows and nRIL columns. Elements of 1/0 indicate this RIL (or strains) is/not selected for this array.

condition.allocation

matrix with nCondition rows and nRIL columns. Elements of 1/0 indicate this RIL (or strains) is/not selected for this condition.

nEnvFactors

number of environmental factors, an integer bewteen 1 and 3. When nEnvFactors is 1 and the number of levels for the environmental factor (nLevels)is 1, there is one condition in the experiment (i.e. no environmental perturbation) and thus only genetic factor will be considered in the algorithm. When nEnvFactors is 1 and nLevels is larger than 1 or nEnvFactors is larger than 1, all main factor(s) and interacting facotr(s) will be included. Examples: If there is a temperature perturbation, then nEnvFactors is 1; If there is both temperature and drug treatment perturbation, then nEnvFactors is 2.

nLevels

number of levels for each factor, a vector with each component being an integer. The length of it should equal nEnvFactors.

Level

a list which specifies the levels for each factor in the experiment. There are in total nEnvFactors elements in the list and each element correpsond to certain environmental factor. The emlemet is a vector describing all levels of the environmental factor. default setting for the level of each factor is 1, 2, ... nLevels[i]. (Here nLevels[i] is the *i*th element of nLevels, which gives the total number of levels for *i* environmental factor).

nConditions

number of all possible combination of all environmental factors.

weight

a vector with length of variableNumber which is calculated from function variableNumber. Default = 1 (which means the parameters to be estimated are equally important during optimization.)

optimality

type of optimality, i.e. "A" (A-optimality) or "D" (D-optimality). A-optimality minimizes  $Trace((X'X)^{-1})$ , which corresponds to minimum average variance of the parameter estimates. D-optimality minimizes  $det(X'X)^{-1}$ , which corresponds to minimum generalized variance of the parameter estimates.

bTwoColorArray

binary variable indicating experiment type:

bTwoColorArray <- T #for dual channel experiment bTwoColorArray <- F #for single channel experiment

envFactorNames

a vector with names for all environmental factor(s). For example, for the experiment with two environmental factors of temperature and drug treatment: envFactorNames <- c( "Temperature", "Dosage")

Default = NULL, then the output will use "F1" and "F2" to indicate the environmental factors.

#### **Details**

Example parameter settings:

 $w_{QF1F3}, w_{QF2F3}, w_{QF1F2F3}$ )

Suppose to design an experiment with two environmental factors (F1, F2) and there are two diffferent levels for each environment. The levels are 16 and 24 for F1, and 5 and 10 for F2. Thus the following command can be used:

```
nEnvFactors <- 2 nLevels <- c ( 2, 2 ) levels <- list ( c(16, 24), c(5, 10) ) The length of parameter weight is dependent on the number of environmental factors: When nEnvFactor = 0, weight is 1 as there is only one parameter of interest (genotype). When nEnvFactor = 1, weight = c(w_Q, w_{F1}, w_{QF1}) When nEnvFactor = 2, weight = c(w_Q, w_{F1}, w_{F2}, w_{QF1}, w_{QF2}, w_{F1F2}, w_{QF1F2}) When nEnvFactor = 3, weight = c(w_Q, w_{F1}, w_{F2}, w_{F2}, w_{F2}, w_{QF1}, w_{QF2}, w_{QF3}, w_{F1F2}, w_{F1F3}, w_{F2F3}, w_{QF1F2}, weight = c(w_Q, w_{F1}, w_{F2}, w_{F2}, w_{F2}, w_{F1}, w_{F2}, w_{F1F2}, w_{F1F3}, w_{F2F3}, w_{F1F2}, w_{F1F2}, w_{F1F3}, w_{F2F3}, w_{F1F2}, w_{F1F2}, w_{F1F3}, w_{F2F3}, w_{F1F2}, w_{F1F2}, w_{F1F3}, w_{F2F3}, w_{F1F2}, w_{F1F3}, w_{F1F2}, w_{F1F3}, w_{F2F3}, w_{F1F2}, w_{F1F3}, w_{F2F3}, w_{F1F2}, w_{F1F3}, w_{F1F3}, w_{F1F2}, w_{F1F3}, w_{F1F2}, w_{F1F3}, w_{F1F3}, w_{F1F2}, w_{F1F3}, w_{F1F3}, w_{F1F2}, w_{F1F3}, w_{F1F2}, w_{F1F3}, w_{F1F3}
```

Here  $w_Q$  represents the weight for genotype effect,  $w_{F1}$  represent the weight for F1 effect and  $w_{QF1}$  represent the weight for interaction between genotype and F1 effect, etc.

#### Value

The score is defined as the "double" sum of the variances, summed over all parameters and over all markers.

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E. Wit and J. McClure. Statistics for Microarrays: Design, Analysis and Inference. (2004) Chichester: Wiley.

### See Also

designGG

exampleArrayDesignTable

Example output of ArrayDesignTable data

#### **Description**

exampleArrayDesignTable: Example data of exampleArrayDesignTable for a hypothetical dual-channel microarray experiment in which there are 100 strains (e.g. recombinant inbred lines) and 27 arrays available. Two environmental factors (temperature and cell type) are considered in this experiment. There are three levels for temperature (15, 24 and 29) and four levels for cell types (A,B,C,D). This table tells how to pair samples into arrays.

```
data(exampleArrayDesignTable)
exampleArrayDesignTable[1:5,]
```

	Channel 1	Channel 2
array1	Strain28	Strain92
array2	Strain70	Strain47
array3	Strain22	Strain89
array4	Strain45	Strain15
array5	Strain52	Strain41

### Usage

```
data(exampleArrayDesignTable)
```

#### **Format**

exampleArrayDesignTable: 27 arrays by two channels.

#### Author(s)

Yang Li <yang.li@rug.nl>, Gonzalo Vera <gonzalo.vera.rodriguez@gmail.com> Rainer Breitling <r.breitling@rug.nl>, Ritsert Jansen <r.c.jansen@rug.nl>

### **Examples**

```
##load the data
data(exampleArrayDesignTable)
##view part of the the data
exampleArrayDesignTable[1:5,]
```

```
exampleConditionDesignTable
```

Example ConditionDesignTable data

# **Description**

exampleConditionDesignTable: Example data of exampleConditionDesignTable for a hypothetical dual-channel microarray experiment in which there are 100 strains (e.g. recombinant inbred lines ) and 27 arrays available. Two environmental factors (temperature and cell type) are considered in this experiment. There are three levels for temperature (15, 24 and 29) and four levels for cell types (A, B, C, D). This table tells how to allocate samples into  $12 = 3 \times 4$  different conditions. On average there are  $4.5 = 27 \times 2/12$  samples per condition.

examplePlotObj 15

- > data(exampleConditionDesignTable)
- > exampleConditionDesignTable[1:5,]

	Temperature	Cell Type	Selected Strains				
condition1	15	A	Strain28	Strain81	Strain18	Strain61	
condition2	24	A	Strain72	Strain40	Strain83	Strain44	Strain10
condition3	29	A	Strain22	Strain89	Strain3	Strain30	Strain58
condition4	15	В	Strain70	Strain47	Strain4	Strain59	
condition5	24	В	Strain93	Strain97	Strain49	Strain14	

### Usage

data(exampleConditionDesignTable)

#### **Format**

exampleConditionDesignTable: 12 combination of conditions from three temepratures and four cell types.

# Author(s)

Yang Li <yang.li@rug.nl>, Gonzalo Vera <gonzalo.vera.rodriguez@gmail.com> Rainer Breitling <r.breitling@rug.nl>, Ritsert Jansen <r.c.jansen@rug.nl>

examplePlotObj Example PlotObj data

# **Description**

examplePlotObj: Example data of examplePlotObj for plot all scores and cooling at each iteration during simulated annealing process.

```
data(examplePlotObj)
plotAllScores(examplePlotObj)
```

# Usage

```
data(examplePlotObj)
```

#### **Format**

examplePlotObj: a list which contains the following elements: (1) scores (2) cooling (3) start-Temp (4) temperature (5) temperature.step (6) nIterations (7) optimality.

# Author(s)

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experimentDesignTable

Make experiment table based design matrix

#### **Description**

This function generates two .csv files which descibe how samples are allocated samples into different conditions and paired on arrays.

### Usage

# **Arguments**

array.allocation

matrix with nArray rows and nRIL columns. Elements of 1/0 indicate this RIL is/not selected for this array.

condition.allocation

matrix with nCondition rows and nRIL columns. Elements of 1/0 indicate this RIL is/not selected for this condition.

nEnvFactors

number of environmental factors, an integer bewteen 1 and 3. When nEnvFactors is 1 and the number of levels for the environmental factor (nLevels)is 1, there is one condition in the experiment (i.e. no environmental perturbation) and thus only genetic factor will be considered in the algorithm. When nEnvFactors is 1 and nLevels is larger than 1 or nEnvFactors is larger than 1, all main factor(s) and interacting facotr(s) will be included. Examples: If there is a temperature perturbation, then nEnvFactors is 1; If there is both temperature and drug treatment perturbation, then nEnvFactors is 2.

nLevels

number of levels for each factor, a vector with each component being integer. The length of it should equal nEnvFactors.

Level

a list which specifies the levels for each factor in the experiment. There are in total nEnvFactors elements in the list and each element correpsond to certain environmental factor. The emlemet is a vector describing all levels of the environmental factor. default setting for the level of each factor is 1, 2, ...nLevels[i]. (Here nLevels[i] is the *i*th element of nLevels, which gives the total number of levels for *i* environmental factor).

fileName

the final optimal design table(s) in <code>csv</code> format and a plot (in <code>png</code> format) of the all scores during SA process (if <code>plotScores = T</code>) will be produced. The users can specify the table and plot name by setting <code>fileName</code>. If <code>NULL</code> (default) it produces files starting with "myDesignGG".

envFactorNames

a vector with names for all environmental factor(s). For example, for the experiment with two environmental factors of temperature and drug treatment: envFactorNames <- c("Temperature", "Dosage")

Default =  $\mathtt{NULL}$ , then the output will use "F1" and "F2" to indicate the environmental factors.

directory

It tells where the resulting optimal design tables are to be stored. If NULL (default), it will use the currect working directory.

#### **Details**

Based on nEnvFactors and nLevels, nConditions is calculated.

#### Value

Two tables report the results: table "pair design" which is only used for two-channel experiments and describes how samples are paired together on the slide (e.g. microarray chip), and table "environment design" which is used when there are more environments evolved in the experiment. With these two tables, the experimenters can set up the environmental treatment and follow-up profiling measurement.

### Examples:

1. conditionDesign.csv

	Temperature	Cell Type			Selected Samples		
condition1	15	A	RIL28	RIL81	RIL18	RIL61	
condition2	24	A	RIL72	RIL40	RIL83	RIL44	RIL10
condition3	29	Α	RIL22	RIL89	RIL3	RIL30	RIL58
condition4	15	В	RIL70	RIL47	RIL4	RIL59	
condition5	24	В	RIL93	RIL97	RIL49	RIL14	

# 2. arrayDesign.csv

	Channel 1	Channel 2
array1	RIL28	RIL92
array2	RIL70	RIL47
array3	RIL22	RIL89
array4	RIL45	RIL15
array5	RIL52	RIL41

### Note

The optimal design results are described in two tables. One is called "array design" which is only used for two-channel experiments. It describes how samples are paired together on the slide (e.g. microarray chip). The other table is called "condition design" which is used when there is more than one environmental factor involved in the experiment. Each cell in condition design table represents a combination of different levels of environmental factors and the selected sample names (e.g. RIL names) for this condition are shown. Based on these two tables, the experimenters can set up the environmental treatment and follow-up profiling measurement.

### Author(s)

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# References

Y. Li, R. Breitling and R.C. Jansen. Generalizing genetical genomics: the added value from environmental perturbation, Trends Genet (2008) 24:518-524.

Y. Li, M. Swertz, G. Vera, J. Fu, R. Breitling, and R.C. Jansen. designGG: An R-package and Web tool for the optimal design of genetical genomics experiments. (submitted) http://gbic.biol.rug.nl/designGG 18 initialDesign

#### See Also

 ${\tt designGG, exampleArrayDesignTable, exampleConditionDesignTable}$ 

genotype Example genotype data

# **Description**

genotype: example data of genotypes for each marker (rownames) and 100 strains such as recombinant inbred lines (RIL) (columnames), with numeric values 1 and 0 (or A and B).

```
data(genotypes)
genotypes[1:5,1:5]
```

	Strain1	Strain2	Strain3	Strain4	Strain5
C1M1	1	0	0	0	1
C1M2	1	0	0	0	1
C1M3	1	0	0	0	1
C1M4	1	0	0	1	1
C1M5	1	0	0	1	1

# Usage

data(genotypes)

### **Format**

genotypes: 120 markers by 100 samples (Strains).

# Author(s)

Yang Li <yang.li@rug.nl>, Gonzalo Vera <gonzalo.vera.rodriguez@gmail.com> Rainer Breitling <r.breitling@rug.nl>, Ritsert Jansen <r.c.jansen@rug.nl>

initialDesign

Initialize an experiment design matrix

# **Description**

Allocate RILs (or strains) into different conditional and pair RILs (or strains) into slides.

# Usage

initialDesign 19

#### **Arguments**

genotype genotype data: a nMarker-by-nRILs matrix with two allels being 0 and 1 (or A

and B) or three allels being 0, 0.5 and 1 (or, A, H, and B), where 0.5 (or H)

represents heterozygous allele.

nRILs total number of RILs ((or strains) available for the experiment.

nSlides total number of slides available for the experiment.

nConditions number of all possible combination of all environmental factors.

nTuple average number of RILs (or strains) to be assigned onto each condition

nTuple should be a real number which is larger than 1.

if nTuple < 1, the algorithm will stop and shw the message below,

warning: "The number of slides is too small to perform

the experiment."

bTwoColorArray

binary variable indicating experiment type:

bTwoColorArray <- T #for dual channel experiment bTwoColorArray <- F #for single channel experiment

#### **Details**

For two-color array experiments, randomly choose a RIL (or strain) and pair it with the genetically most different RIL (or strain) on one array.

For one-color array experiments, array.allocation is NULL as there is no need to pair samples.

#### Value

a list with 2 matrices:

condition.allocation: allocate RILs (or strains) into different conditional (nCondition  $\times$  nRILs)

array.allocation: pair RILs (or strains) into sldies ( $nSlides \times nRILs$ )

#### Note

This function calls conditionAllocation function to allocate selected RILs (or strains) into different conditions.

# Author(s)

Yang Li <yang.li@rug.nl>, Gonzalo Vera <gonzalo.vera.rodriguez@gmail.com> Rainer Breitling <r.breitling@rug.nl>, Ritsert Jansen <r.c.jansen@rug.nl>

### References

Y. Li, R. Breitling and R.C. Jansen. Generalizing genetical genomics: the added value from environmental perturbation, Trends Genet (2008) 24:518-524.

Y. Li, M. Swertz, G. Vera, J. Fu, R. Breitling, and R.C. Jansen. designGG: An R-package and Web tool for the optimal design of genetical genomics experiments. (submitted) http://gbic.biol.rug.nl/designGG

#### See Also

designGG

20 interactionLevel

#### **Examples**

```
genotype <- read.table("genotype.txt")</pre>
nEnvFactors <- 2
nLevels <- c(2, 2)
levels <- list ( c(16, 24), c(5, 10) )
nSlides <- 100
nTuple <- 25
bTwoColorArray <- TRUE
initialDesign( genotype, nRILs, nSlides, nConditions, nTuple, bTwoColorArray )
```

Generate levels for all interacting factors interactionLevel

# **Description**

Generate levels for all interacting factors for all RILs (or strains). This is a subfunction needed for designScore, but is not directly used.

# Usage

```
interactionLevel( genotype.level, condition.level, markerIndex,
                  nEnvFactors )
```

### **Arguments**

genotype.level

levels of genetic factor for each RIL (or strain) in the experiment.

condition.level

levels of all environmental factors for each RIL (or strain)in the experiment.

markerIndex indicate which genome position that level of genetic factor corresponds to.

nEnvFactors

number of environmental factors, an integer bewteen 1 and 3. When nEnvFactors is 1 and the number of levels for the environmental factor (nLevels)is 1, there is one condition in the experiment (i.e. no environmental perturbation) and thus only genetic factor will be considered in the algorithm. When nEnvFactors is 1 and nLevels is larger than 1 or nEnvFactors is larger than 1, all main factor(s) and interacting facotr(s) will be included. Examples: If there is a temperature perturbation, then nEnvFactors is 1; If there is both temperature and drug treatment perturbation, then nEnvFactors is 2.

#### **Details**

markerIndex indicates the genome position that genotype.level corresponds to. An experiment design is defined to be optimal over all markers if the sum of scores, e.g. Aoptimality criterion over all markers is minimized.

#### Value

a matrix with nRILs rows. The number columns depends on nEnvFactors. For example: If nEnvFactors = 1, there is only one interaction term.

If nEnvFactors = 2, there are three pair-wise two-way interaction terms and one three-way interaction term.

pairLevel 21

#### Author(s)

Yang Li <yang.li@rug.nl>, Gonzalo Vera <gonzalo.vera.rodriguez@gmail.com> Rainer Breitling <r.breitling@rug.nl>, Ritsert Jansen <r.c.jansen@rug.nl>

#### References

Y. Li, R. Breitling and R.C. Jansen. Generalizing genetical genomics: the added value from environmental perturbation, Trends Genet (2008) 24:518-524.

Y. Li, M. Swertz, G. Vera, J. Fu, R. Breitling, and R.C. Jansen. designGG: An R-package and Web tool for the optimal design of genetical genomics experiments. (submitted) http://gbic.biol.rug.nl/designGG

#### See Also

designScore, conditionLevel

pairLevel

Pair levels for paired RILs (or strains)

#### **Description**

Pair levels for two RILs (or strains) allocated into one slide (bTwoColorArray=T). It is a subfunction needed for designScore function, but is not directly used.

#### Usage

```
pairLevel( xxx, rilNames )
```

### **Arguments**

can be genotype.level, condition.level or interaction.level rilNames names for all RILs (or strains) that have been selected for the experiment

#### **Details**

This function is used only for two-color array.

# Author(s)

Yang Li <yang.li@rug.nl>, Gonzalo Vera <gonzalo.vera.rodriguez@gmail.com> Rainer Breitling <r.breitling@rug.nl>, Ritsert Jansen <r.c.jansen@rug.nl>

# References

Y. Li, R. Breitling and R.C. Jansen. Generalizing genetical genomics: the added value from environmental perturbation, Trends Genet (2008) 24:518-524.

Y. Li, M. Swertz, G. Vera, J. Fu, R. Breitling, and R.C. Jansen. designGG: An R-package and Web tool for the optimal design of genetical genomics experiments. (submitted) http://gbic.biol.rug.nl/designGG

# See Also

See Also designScore

22 plotAllScores

AllScores Plot scores profiles	
es Plot scores profiles	

# **Description**

Plot all scores and the temperature at each iteration during the simulated annealing process.

# Usage

```
plotAllScores(plot.obj,fileName=NULL)
```

# Arguments

plot.obj	a list containing: scores, cooling, startTemp, temperature, temperature.step, nIterations and optimality. Details can be found below.
scores	A- or D- optimality score of all accepted designs during optimization process.
cooling	describes the cooling step in the Simulated Annealing, defined as (new.score $-$ now.score)/ now.score.
startTemp	starting temperature of the simulated annealing process.
temperature	final temperature that the simulated annealing reaches.
temperatureS	tep
	temperature decreasing step in the simulated annealing (SA) process.
nIterations	number of iterations in the simulated annealing method.
optimality	type of optimality, i.e. "A" (A-optimality) or "D" (D-optimality). A-optimality minimizes $Trace((X'X)^{-1})$ , which corresponds to minimum average variance of the parameter estimates. D-optimality minimizes $det(X'X)^{-1}$ , which corresponds to minimum generalized variance of the parameter estimates.
fileName	the final optimal design table(s) in csv format and a plot (in png format) of the all scores during SA process (if plotScores = T) will be produced. The users can specify the table and plot name by setting fileName. If NULL (default) it produces files starting with "myDesignGG".

#### Value

Draw a plot that visualizeds the scores (y-axis) at each iteration during the simulated annealing process (x-axis is time of moving)

# Note

The calculation of score is dependent on the choice of optimality. Cooling is defined as (newScore - nowScore)/nowScore.

# Author(s)

Yang Li <yang.li@rug.nl>, Gonzalo Vera <gonzalo.vera.rodriguez@gmail.com> Rainer Breitling <r.breitling@rug.nl>, Ritsert Jansen <r.c.jansen@rug.nl>

temperatureStep 23

#### References

Y. Li, R. Breitling and R.C. Jansen. Generalizing genetical genomics: the added value from environmental perturbation, Trends Genet (2008) 24:518-524.

Y. Li, M. Swertz, G. Vera, J. Fu, R. Breitling, and R.C. Jansen. designGG: An R-package and Web tool for the optimal design of genetical genomics experiments. (submitted) http://gbic.biol.rug.nl/designGG

temperatureStep Calculate the temperature decreasing step for simulated annealing process

### **Description**

Calculate the temperature decreasing step for simulated annealing process. This is a subfunction needed for designGG, but is not directly used.

### Usage

temperatureStep(startTemp, maxTempStep, endTemp, nIterations)

# **Arguments**

startTemp starting temperature of simulated annealing process.

maxTempStep maximum temperature decreasing step for simulated annealing process. The parameter ensures that the multiplicative cooling factor is not smaller than this value. If nIterations is too small, the preferred final temperature (endTemp) may not be reached. See Wit and McClure (2004) for details.

endTemp ending temperature of simulated annealing process. An important optimization parameter. Setting this parameter closer to zero. See Wit and McClure (2004) for details

nIterations number of iterations in the simulated annealing method.

#### Value

A temperature decreasing step in the simulated annealing process.

#### Author(s)

Yang Li <yang.li@rug.nl>, Gonzalo Vera <gonzalo.vera.rodriguez@gmail.com> Rainer Breitling <r.breitling@rug.nl>, Ritsert Jansen <r.c.jansen@rug.nl>

# References

Y. Li, M. Swertz, G. Vera, J. Fu, R. Breitling, and R.C. Jansen. designGG: An R-package and Web tool for the optimal design of genetical genomics experiments. (submitted) http://gbic.biol.rug.nl/designGG

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E. Wit and J. McClure. Statistics for Microarrays: Design, Analysis and Inference. (2004) Chichester: Wiley.

24 updateDesign

#### See Also

designGG

updateDesign

Updates current design

#### **Description**

Updates current experimental design (including array.allocation and condition.allocation).

#### Usage

### **Arguments**

array.allocation

matrix with nArray rows and nRIL columns. Elements of 1/0 indicate this RIL

(or strain) is/not selected for this array.

condition.allocation

matrix with nCondition rows and nRIL columns. Elements of 1/0 indicate this

RIL (or strain) is/not selected for this condition.

nRILs number of RILs (or strains) available for the experiment.

nSlides total number of slides available for experiment.

nEnvFactors number of environmental factors, an integer bewteen 1 and 3. When nEnvFactors

is 1 and the number of levels for the environmental factor (nLevels) is 1, there is one condition in the experiment (i.e. no environmental perturbation) and thus only genetic factor will be considered in the algorithm. When nEnvFactors is 1 and nLevels is larger than 1 or nEnvFactors is larger than 1, all main factor(s) and interacting facotr(s) will be included. Examples: If there is a temperature perturbation, then nEnvFactors is 1; If there is both temperature and

drug treatment perturbation, then nEnvFactors is 2.

nTuple average number of RILs (or strains) to be assigned onto each condition.

nTuple should be a real number which is larger than 1. If nTuple < 1, the algorithm will stop and show the message,

warning: "The number of slides is too small to perform

the experiment."

bTwoColorArray

binary variable indicating experiment type:

bTwoColorArray <- T #for dual channel experiment bTwoColorArray <- F #for single channel experiment

# Details

This function calls two subfunctions: conditionUpdate and arrayUpdate.

### Value

a list with two elements, array.allocation and condition.allocation.

variableNames 25

#### Author(s)

Yang Li <yang.li@rug.nl>, Gonzalo Vera <gonzalo.vera.rodriguez@gmail.com> Rainer Breitling <r.breitling@rug.nl>, Ritsert Jansen <r.c.jansen@rug.nl>

#### References

Y. Li, R. Breitling and R.C. Jansen. Generalizing genetical genomics: the added value from environmental perturbation, Trends Genet (2008) 24:518-524.

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variableNames

Generate variable names for all factors

### **Description**

Generate variable names for genetic, environmental factors and interacting terms.

#### Usage

variableNames (nEnvFactors)

#### **Arguments**

nEnvFactors number of environmental factors, an integer bewteen 1 and 3. When nEnvFactors is 1 and the number of levels for the environmental factor (nLevels)is 1, there is one condition in the experiment (i.e. no environmental perturbation) and thus only genetic factor will be considered in the algorithm. When nEnvFactors is 1 and nLevels is larger than 1 or nEnvFactors is larger than 1, all main factor(s) and interacting facotr(s) will be included. Examples: If there is a temperature perturbation, then nEnvFactors is 1; If there is both temperature and drug treatment perturbation, then nEnvFactors is 2.

# **Details**

generates names for variables, a vector with the length of (variableNumber+1).

#### Value

When nEnvFactors = 1 and nLevels = 1, there is no environmetal pertubation in the experimental. Then we re-define nEnvFactors to be 0 within the algorithm. Accordingly, variable Number = 1, and variableNames is one genetic factor "Q".

When nEnvFactors = 1, variableNumber = 3, and variableNames are one genetic factor "Q", one environmental factor "F", and one interacting factor "QxF".

When nEnvFactors = 2, variableNumber = 7, and variableNames are one genetic factor "Q", two environmental factors "F1" and "F2", three two-way interacting factors "QF1", "QF2", "F1F2", and one three way interacting factors "QxF1xF2".

When nEnvFactors = 3, variableNumber = 15, and variableNames are one genetic factor "Q", three environmental factors "F1", "F2" and "F3", six two-way interacting factors "QF1", "QF2", "QF3", "F1F2", "F2F3" and "F1F3", four three-way interacting factors "QxF1xF2", "QxF1xF3", "QxF2xF3", "F1xF2xF3" and one four-way interacting factors "QxF1xF2xF3".

26 variableNumber

#### Author(s)

Yang Li <yang.li@rug.nl>, Gonzalo Vera <gonzalo.vera.rodriguez@gmail.com> Rainer Breitling <r.breitling@rug.nl>, Ritsert Jansen <r.c.jansen@rug.nl>

#### References

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Y. Li, M. Swertz, G. Vera, J. Fu, R. Breitling, and R.C. Jansen. designGG: An R-package and Web tool for the optimal design of genetical genomics experiments. (submitted) http://gbic.biol.rug.nl/designGG

#### See Also

variableNumber

variableNumber

Compute the number of variables in the experiment

#### **Description**

When nEnvFactors = 1 and nLevels = 1, there is no environmetal pertubation in the experimental. Then we re-define nEnvFactors to be 0 within the algorithm. nEnvFactors = 0, only genetic factor is considered.

nEnvFactors > 1, genetic and environmental facotrs, and all possible interacting factors are considered.

### Usage

```
variableNumber( nEnvFactors )
```

### **Arguments**

nEnvFactors number of environmental factors, an integer.

When nEnvFactors is between 0 and 3, all main factors and interacting factors will be included.

# Value

```
nEnvFactors = 1, variableNumber = 3 (one genetic factor Q, one environmental factor F,
and one interacting factor QxF)
```

```
nEnvFactors = 2, variableNumber = 7
nEnvFactors = 3, variableNumber = 15
```

# Author(s)

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variableNumber 27

#### References

Y. Li, R. Breitling and R.C. Jansen. Generalizing genetical genomics: the added value from environmental perturbation, Trends Genet (2008) 24:518-524.

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# See Also

variableNames

# **Index**

genotype, 18

```
*Topic datasets
                                          initialDesign, 4, 11, 18
    exampleArrayDesignTable, 13
                                          interactionLevel, 20
   exampleConditionDesignTable,
                                          pairLevel, 21
       14
                                          plotAllScores, 11, 22
   examplePlotObj, 15
   genotype, 18
                                          temperatureStep, 23
*Topic method
   acceptanceProbability, 1
                                          updateDesign, 3, 11, 24
   arrayUpdate, 2
   conditionAllocation, 3
                                          variableNames, 25, 27
   conditionCombination, 4
                                          variableNumber, 26, 26
   conditionLevel, 6
   conditionUpdate, 7
   designGG, 8
   designScore, 11
   experimentDesignTable, 16
   initialDesign, 18
   interactionLevel, 20
   pairLevel, 21
   plotAllScores, 22
   updateDesign, 24
   variableNames, 25
   variableNumber, 26
acceptanceProbability, 1, 11
arrayUpdate, 2, 8
conditionAllocation, 3
conditionCombination, 4, 7
conditionLevel, 6, 21
conditionUpdate, 7
design genetical genomics
       experiment (designGG), 8
designGG, 2, 8, 8, 13, 18, 19, 24
designScore, 5, 7, 11, 11, 21
exampleArrayDesignTable, 11, 13, 18
exampleConditionDesignTable, 11,
       14, 18
examplePlotObj, 15
experimentDesignTable, 11, 16
```