# Package 'polymapR'

July 23, 2025

Type Package **Title** Linkage Analysis in Outcrossing Polyploids Version 1.1.6 Date 2024-05-30 Description Creation of linkage maps in polyploid species from marker dosage scores of an F1 cross from two heterozygous parents. Currently works for outcrossing diploid, autotriploid, autotetraploid and autohexaploid species, as well as segmental allotetraploids. Methods are described in a manuscript of Bourke et al. (2018) <doi:10.1093/bioinformatics/bty371>. Since version 1.1.0, both discrete and probabilistic genotypes are acceptable input; for more details on the latter see Liao et al. (2021) <doi:10.1007/s00122-021-03834-x>. **Depends** R (>= 3.5.0) License GPL **Imports** doParallel, foreach, graphics, grDevices, igraph, knitr, MDSMap, stats, utils RoxygenNote 7.3.1 Suggests ggplot2, Hmisc, RColorBrewer, reshape2, rmarkdown, polyRAD, updog, mappoly VignetteBuilder knitr **Encoding UTF-8** LazyData TRUE NeedsCompilation no Author Peter Bourke [aut, cre], Geert van Geest [aut], Roeland Voorrips [ctb], Yanlin Liao [ctb] Maintainer Peter Bourke <pbourkey@gmail.com>

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add_d	dup_markers Add back duplicate markers after mapping	

# Description

Often there will be duplicate markers that can be put aside to speed up mapping. These may be added back to the maps afterwards.

# Usage

```
add_dup_markers(maplist, bin_list, marker_assignments = NULL)
```

# Arguments

maplist A list of maps. Output of MDSMap\_from\_list.

bin\_list A list of marker bins containing marker duplicates. One of the list outputs of screen\_for\_duplicate\_markers

marker\_assignments

Optional argument to include the marker\_assignments (output of check\_marker\_assignment). If included, marker assignment information will also be copied.

4 ALL\_dosages

### Value

A list with the following items:

maplist List of maps, now with duplicate markers addedmarker\_assignments If required, marker assignment list with duplicate markers added

ALL\_dosages

A dosage matrix for a random pairing tetraploid with five linkage groups.

# Description

A dosage matrix for a random pairing tetraploid with five linkage groups.

# Usage

```
ALL_dosages
segregating_data
screened_data
screened_data2
screened_data3
TRI_dosages
```

#### **Format**

A matrix

An object of class matrix (inherits from array) with 2873 rows and 209 columns.

An object of class matrix (inherits from array) with 1417 rows and 209 columns.

An object of class matrix (inherits from array) with 1417 rows and 207 columns.

An object of class matrix (inherits from array) with 1417 rows and 200 columns.

An object of class matrix (inherits from array) with 250 rows and 202 columns.

all\_linkages\_list\_P1 5

```
all_linkages_list_P1 A (nested) list of linkage data frames classified per linkage group and homologue
```

# **Description**

A (nested) list of linkage data frames classified per linkage group and homologue

# Usage

```
all_linkages_list_P1
all_linkages_list_P1_split
all_linkages_list_P1_subset
```

#### **Format**

```
An object of class list of length 5.
An object of class list of length 5.
An object of class list of length 5.
```

# **Description**

assign\_linkage\_group quantifies per marker number of linkages to a linkage group and evaluates to which linkage group (and homologue(s)) the marker belongs.

```
assign_linkage_group(
  linkage_df,
  LG_hom_stack,
  SN_colname = "marker_a",
  unassigned_marker_name = "marker_b",
  phase_considered = "coupling",
  LG_number,
  LOD_threshold = 3,
  ploidy,
  assign_homologue = T,
  log = NULL
)
```

6 assign\_SN\_SN

### **Arguments**

linkage\_df A linkage data.frame as output of linkage.

LG\_hom\_stack A data.frame with markernames ("SxN\_Marker"), linkage group ("LG") and

homologue ("homologue")

SN\_colname The name of the column in linkage\_df harbouring the 1.0 markers

unassigned\_marker\_name

The name of the column in linkage\_df harbouring the marker that are to be

assigned.

phase\_considered

The phase that is used to assign the markers (deprecated)

LG\_number The number of chromosomes (linkage groups) in the species.

LOD\_threshold The LOD score at which a linkage to a linkage group is significant.

ploidy The ploidy of the plant species.

assign\_homologue

Logical. Should markers be assigned to homologues? If FALSE markers will be

assigned to all homologues

log Character string specifying the log filename to which standard output should be

written. If NULL log is send to stdout.

#### Value

Output is a data.frame with at least the following columns:

Assigned\_LG The assigned linkage group

Assigned\_hom1 The homologue with most linkages

The columns LG1 - LGn and Hom1 - Homn give the number of hits per marker for that linkage group/homologue. Assigned\_hom2 .. gives the nth homologue with most linkages.

### **Examples**

assign\_SN\_SN

Assign (leftover) 1.0 markers

# **Description**

Some 1.0 markers might have had ambiguous linkages, or linkages with low LOD scores leaving them unlinked to a linkage group. assign\_SN\_SN finds 1.0 markers unlinked to a linkage group and tries to assign them.

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# Usage

```
assign_SN_SN(
    linkage_df,
    LG_hom_stack,
    LOD_threshold,
    ploidy,
    LG_number,
    log = NULL
)
```

# Arguments

linkage\_df A data.frame as output of linkage with arguments markertype1=c(1,0) and

markertype2=NULL.

LG\_hom\_stack A data.frame with markernames ("SxN\_Marker"), linkage group ("LG") and

homologue ("homologue")

LOD\_threshold A LOD score at which linkages between markers are significant.

ploidy Integer. The ploidy level of the plant species.

LG\_number Integer. Number of chromosomes (linkage groups)

log Character string specifying the log filename to which standard output should be

written. If NULL log is send to stdout.

### Value

Returns a data. frame with the following columns:

SxN\_Marker The markername

Assigned\_hom1 The assigned homologue

Assigned\_LG The assigned linkage group

### **Examples**

8 bridgeHomologues

bridgeHomologues

Use bridge markers to cluster homologues into linkage groups

### **Description**

Clustering at high LOD scores results in marker clusters representing homologues. bridgeHomologues clusters these (pseudo)homologues to linkage groups using linkage information between 1.0 and bridge markers within a parent (e.g. 2.0 for a tetraploid). If parent-specific bridge markers (e.g. 2.0) cannot be used, biparental markers can also be used (e.g. 1.1, 1.2, 2.1, 2.2 and 1.3 markers). The linkage information between 1.0 and biparental markers can be combined.

#### Usage

```
bridgeHomologues(
  cluster_stack,
  cluster_stack2 = NULL,
  linkage_df,
  linkage_df2 = NULL,
  LOD_threshold = 5,
  automatic_clustering = TRUE,
  LG_number,
  parentname = "",
  min_links = 1,
  min_bridges = 1,
  only_coupling = FALSE,
  log = NULL
)
```

### **Arguments**

cluster\_stack A data.frame with a column "marker" specifying markernames, and a column

"cluster" specifying marker cluster

cluster\_stack2 Optional. A cluster\_stack for the other parent. Use this argument if cross-

parent markers are used (e.g. when using 1.1 markers).

linkage\_df A linkage data.frame as output of linkage between bridge (e.g. 1.0 and 2.0)

markers.

linkage\_df2 Optional. A linkage\_df specifying linkages between 1.0 and cross-parent

markers in the other parent. Use this argument if cross-parent markers are used (e.g. when using 1.1, 2.1, 1.2 and/or 2.2 markers). The use of multiple types of

cross-parent markers is allowed.

LOD\_threshold Integer. The LOD threshold specifying at which LOD score a link between 1.0

and bridging-type marker (e.g. 2.0) is used for clustering homologues.

automatic\_clustering

Logical. Should clustering be executed without user input?

LG\_number Integer. Expected number of chromosomes (linkage groups)

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parentname Name of the parent. Used in the main title of the plot. The minimum number of links between a bridge marker and a cluster for that min\_links bridge to be considered. In the case of a 2x0 marker for example, this argument means that the 2x0 marker must have at least min\_links linkages of at least a LOD of LOD\_threshold with markers from each of the clusters involved, to be considered a single bridging link. Make this number higher if there are a lot of spurious links. The minimum number of bridge markers needed to assign two homologues tomin\_bridges gether as coming from the same chromosomal linkage group. See argument min\_links for further details. only\_coupling Logical, should only coupling linkages be used in the process? By default FALSE Character string specifying the log filename to which standard output should be log written. If NULL log is send to stdout.

#### Value

A data frame with markers classified by homologue and linkage group.

#### **Examples**

calcSegtypeInfo

Build a list of segregation types

### **Description**

For each possible segregation type in an F1 progeny with given parental ploidy (and ploidy2, if parent2 has a different ploidy than parent1) information is given on the segregation ratios, parental dosages and whether the segregation is expected under polysomic, disomic and/or mixed inheritance.

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### Usage

calcSegtypeInfo(ploidy, ploidy2=NULL)

### Arguments

ploidy The ploidy of parent 1 (must be even, 2 (diploid) or larger).

ploidy2 The ploidy of parent 2. If omitted (default=NULL) it is assumed to be equal to

ploidy.

#### **Details**

The names of the segregation types consist of a short sequence of digits (and sometimes letters), an underscore and a final number. This is interpreted as follows, for example segtype 121\_0: 121 means that there are three consecutive dosages in the F1 population with frequency ratios 1:2:1, and the 0 after the underscore means that the lowest of these dosages is nulliplex. So 121\_0 means a segregation of 1 nulliplex: 2 simplex: 1 duplex. A monomorphic F1 (one single dosage) is indicated as e.g. 1\_4 (only one dosage, the 4 after the underscore means that this is monomorphic quadruplex). If UPPERCASE letters occur in the first part of the name these are interpreted as additional digits with values of A=10 to Z=35, e.g. 18181\_0 means a segregation of 1:8:18:8:1 (using the I as 18), with the lowest dosage being nulliplex.

With higher ploidy levels higher numbers (above 35) may be required. In that case each unique ratio number above 35 is assigned a lowercase letter. E.g. one segregation type in octaploids is 9bcb9\_2: a 9:48:82:48:9 segregation where the lowest dosage is duplex.

Segregation types with more than 5 dosage classes are considered "complex" and get codes like c7e\_1 (again in octoploids): this means a complex type (the first c) with 7 dosage classes; the e means that this is the fifth type with 7 classes. Again the \_1 means that the lowest dosage is simplex. It is always possible (and for all segtype names with lowercase letters it is necessary) to look up the actual segregation ratios in the intratio item of the segtype. For octoploid segtype c7e\_1 this shows 0:1:18:69:104:69:18:1:0 (the two 0's mean that nulli- and octoplexes do not occur).

#### Value

A list with for each different segregation type (segtype) one item. The names of the items are the names of the segtypes. Each item is itself a list with components:

**freq** A vector of the ploidy+1 fractions of the dosages in the F1

intratios An integer vector with the ratios as the simplest integers

**expgeno** A vector with the dosages present in this segtype

allfrq The allele frequency of the dosage allele in the F1

polysomic Boolean: does this segtype occur with polysomic inheritance?

**disomic** Boolean: does this segtype occur with disomic inheritance?

**mixed** Boolean: does this segtype occur with mixed inheritance (i.e. with polysomic inheritance in one parent and disomic inheritance in the other)?

**pardosage** Integer matrix with 2 columns and as many rows as there are parental dosage combinations for this segtype; each row has one possible combination of dosages for parent 1 (1st column) and parent 2 (2nd column)

**parmode** Logical matrix with 3 columns and the same number of rows as pardosage. The 3 columns are named polysomic, disomic and mixed and tell if this parental dosage combination will generate this segtype under polysomic, disomic and mixed inheritance

### **Examples**

```
si4 <- calcSegtypeInfo(ploidy=4) # two 4x parents: a 4x F1 progeny
print(si4[["11_0"]])
si3 <- calcSegtypeInfo(ploidy=4, ploidy2=2) # a 4x and a diplo parent: a 3x progeny
print(si3[["11_0"]])</pre>
```

checkF1

*Identify the best-fitting F1 segregation types* 

# Description

For a given set of F1 and parental samples, this function finds the best-fitting segregation type using either discrete or probabilistic input data. It can also perform a dosage shift prior to selecting the segregation type.

```
checkF1(
  input_type = "discrete",
  dosage_matrix,
  probgeno_df,
 parent1,
 parent2,
 F1,
  ancestors = character(0),
 polysomic,
 disomic,
 mixed,
 ploidy,
 ploidy2,
 outfile = "",
  critweight = c(1, 0.4, 0.4),
 Pvalue_threshold = 1e-04,
  fracInvalid_threshold = 0.05,
  fracNA_threshold = 0.25,
  shiftmarkers,
  parentsScoredWithF1 = TRUE,
  shiftParents = parentsScoredWithF1,
  showAll = FALSE,
  append_shf = FALSE
)
```

#### **Arguments**

input\_type Can be either one of 'discrete' or 'probabilistic'. For the former (default), a

dosage\_matrix must be supplied, while for the latter a probgeno\_df must be

supplied.

dosage\_matrix An integer matrix with markers in rows and individuals in columns.

probgeno\_df A data frame as read from the scores file produced by function saveMarkerModels

of R package fitPoly, or alternatively, a data frame containing the following

columns:

**SampleName** Name of the sample (individual)

**MarkerName** Name of the marker **P0** Probabilities of dosage score '0'

P1... Probabilities of dosage score '1' etc. (up to max dosage, e.g. P4 for

tetraploid population)

maxP Maximum genotype probability identified for a particular individual and

marker combination

maxgeno Most probable dosage for a particular individual and marker combi-

nation

geno Most probable dosage for a particular individual and marker combination,

if maxP exceeds a user-defined threshold (e.g. 0.9), otherwise NA

parent1 character vector with the sample names of parent 1
parent2 character vector with the sample names of parent 2

F1 character vector with the sample names of the F1 individuals

ancestors character vector with the sample names of any other ancestors or other samples

of interest. The dosages of these samples will be shown in the output (shifted if shiftParents TRUE) but they are not used in the selection of the segregation type.

polysomic if TRUE at least all polysomic segtypes are considered; if FALSE these are not

specifically selected (but if e.g. disomic is TRUE, any polysomic segtypes that

are also disomic will still be considered)

disomic if TRUE at least all disomic segtypes are considered (see polysomic)

mixed if TRUE at least all mixed segtypes are considered (see polysomic). A mixed

segtype occurs when inheritance in one parent is polysomic (random chromosome pairing) and in the other parent disomic (fully preferential chromosome

pairing)

ploidy The ploidy of parent 1 (must be even, 2 (diploid) or larger).

ploidy2 The ploidy of parent 2. If omitted it is assumed to be equal to ploidy.

outfile the tab-separated text file to write the output to; if NA a temporary file checkF1.tmp

is created in the current working directory and deleted at end

critweight NA or a numeric vector containing the weights of three quality criteria; do not

need to sum to 1. If NA, the output will not contain a column qall\_weights. Else the weights specify how qall\_weights will be calculated from quality parameters

q1, q2 and q3.

Pvalue\_threshold

a minimum threshold value for the Pvalue of the bestParentfit segtype (with a

smaller Pvalue the q1 quality parameter will be set to 0)

fracInvalid\_threshold

a maximum threshold for the fracInvalid of the bestParentfit segtype (with a larger fraction of invalid dosages in the F1 the q1 quality parameter will be set to 0)

fracNA\_threshold

a maximum threshold for the fraction of unscored F1 samples (with a larger fraction of unscored samples in the F1 the q3 quality parameter will be set to 0)

shiftmarkers

if specified, shiftmarkers must be a data frame with columns MarkerName and shift; for the markernames that match exactly (upper/lowercase etc) those in the input (either dosage\_matrix or probgeno\_df), the dosages are increased by the amount specified in column shift, e.g. if shift is -1, dosages 2..ploidy are converted to 1..(ploidy-1) and dosage 0 is a combination of old dosages 0 and 1, for all samples. The segregation check is then performed with the shifted dosages. A shift=NA is allowed, these markers will not be shifted. The sets of markers in the input (either dosage\_matrix or probgeno\_df) and shiftmarkers may be different, but markers may occur only once in shiftmarkers. A column shift is added at the end of the returned data frame.

If parameter shiftParents is TRUE, the parental and ancestor scores are shifted as the F1 scores, if FALSE they are not shifted.

parentsScoredWithF1

TRUE if parents are scored in the same experiment and the same fitPoly run as the F1, else FALSE. If TRUE, their fraction missing scores and conflicts tell something about the quality of the scoring. If FALSE (e.g. when the F1 is triploid and the parents are diploid and tetraploid) the quality of the F1 scores can be independent of that of the parents.

If not specified, TRUE is assumed if ploidy2 == ploidy and FALSE if ploidy2 != ploidy

shiftParents

only used if parameter shiftmarkers is specified. If TRUE, apply the shifts also to the parental and ancestor scores. By default TRUE if parentsScoredWithF1 is

(default FALSE) if TRUE, for each segtype 3 columns are added to the returned data frame with the frqInvalid, Pvalue and matchParents values for these segtype (see the description of the return value)

append\_shf

showA11

if TRUE and parameter shiftmarkers is specified, \_shf is appended to all marker names where shift is not 0. This is not required for any of the functions in this package but may prevent duplicated marker names when using other software.

# Details

For each marker is tested how well the different segregation types fit with the observed parental and F1 dosages. The results are summarized by columns bestParentfit (which is the best fitting segregation type, taking into account the F1 and parental dosages) and columns qall\_mult and/or qall\_weights (how good is the fit of the bestParentfit segtype: 0=bad, 1=good).

Column bestfit in the results gives the segtype best fitting the F1 segregation without taking account of the parents. This bestfit segtype is used by function correctDosages, which tests for possible "shifts" in the marker models.

In case the parents are not scored together with the F1 (e.g. if the F1 is triploid and the parents are diploid and tetraploid) dosage\_matrix should be edited to contain the parental as well

as the F1 scores. In case the diploid and tetraploid parent are scored in the same run of function saveMarkerModels (from package fitPoly) the diploid is initially scored as nulliplex-duplex-quadruplex (dosage 0, 2 or 4); that must be converted to the true diploid dosage scores (0, 1 or 2). Similar corrections are needed with other combinations, such as a diploid parent scored together with a hexaploid population etc.

#### Value

A list containing two elements, checked\_F1 and meta. meta is itself a list that stores the parameter settings used in running checkF1 which can be useful for later reference. The first element (checked\_F1) contains the actual results: a data frame with one row per marker, with the following columns:

- m: the sequential number of the marker (as assigned by fitPoly)
- MarkerName: the name of the marker, with \_shf appended if the marker is shifted and append\_shf is TRUE
- parent1: consensus dosage score of the samples of parent 1
- parent2: consensus dosage score of the samples of parent 2
- F1\_0 ... F1\_<ploidy>: the number of F1 samples with dosage scores 0 ... <ploidy>
- F1\_NA: the number of F1 samples with a missing dosage score
- sample names of parents and ancestors: the dosage scores for those samples
- bestfit: the best fitting segtype, considering only the F1 samples
- frqInvalid\_bestfit: for the bestfit segtype, the frequency of F1 samples with a dosage score that is invalid (that should not occur). The frequency is calculated as the number of invalid samples divided by the number of non-NA samples
- Pvalue\_bestfit: the chisquare test P-value for the observed distribution of dosage scores vs the expected fractions. For segtypes where only one dosage is expected (1\_0, 1\_1 etc) the binomial probability of the number of invalid scores is given, assuming an error rate of seg invalidrate (hard-coded as 0.03)
- matchParent\_bestfit: indication how the bestfit segtype matches the consensus dosages of parent 1 and 2: "Unknown"=both parental dosages unknown; "No"=one or both parental dosages known and conflicting with the segtype; "OneOK"= only one parental dosage known, not conflicting with the segtype; "Yes"=both parental dosages known and combination matching with the segtype. This score is initially assigned based on only high-confidence parental consensus scores; if low-confidence dosages are confirmed by the F1, the matchParent for (only) the selected segtype is updated, as are the parental consensus scores.
- bestParentfit: the best fitting segtype that does not conflict with the parental consensus scores
- frqInvalid\_bestParentfit, Pvalue\_bestParentfit, matchParent\_bestParentfit: same as the corresponding columns for bestfit. Note that matchParent\_bestParentfit cannot be "No".
- q1\_segtypefit: a value from 0 (bad) to 1 (good), a measure of the fit of the bestParentfit segtype based on Pvalue, invalidP and whether bestfit is equal to bestParentfit
- q2\_parents: a value from 0 (bad) to 1 (good), based either on the quality of the parental scores (the number of missing scores and of conflicting scores, if parentsScoredWithF1 is TRUE) or on matchParents (No=0, Unknown=0.65, OneOK=0.9, Yes=1, if parentsScoredWithF1 is FALSE)

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• q3\_fracscored: a value from 0 (bad) to 1 (good), based on the fraction of F1 samples that have a non-missing dosage score

- qall\_mult: a value from 0 (bad) to 1 (good), a summary quality score equal to the product q1\*q2\*q3. Equal to 0 if any of these is 0, hence sensitive to thresholds; a natural selection criterion would be to accept all markers with qall\_mult > 0
- qall\_weights: a value from 0 (bad) to 1 (good), a weighted average of q1, q2 and q3, with weights as specified in parameter critweight. This column is present only if critweight is specified. In this case there is no "natural" threshold; a threshold for selection of markers must be obtained by inspecting XY-plots of markers over a range of qall\_weights values
- shift: if shiftmarkers is specified a column shift is added with for all markers the applied shift (for the unshifted markers the shift value is 0)

qall\_mult and/or qall\_weights can be used to compare the quality of the SNPs within one analysis and one F1 population but not between analyses or between different F1 populations.

If parameter showAll is TRUE there are 3 additional columns for each segtype with names frqInvalid\_<segtype>, Pvalue\_<segtype> and matchParent\_<segtype>; see the corresponding columns for bestfit for an explanation. These extra columns are inserted directly before the bestfit column.

### **Examples**

```
## Not run:
data("ALL_dosages")
chk1<-checkF1(input_type="discrete",dosage_matrix=ALL_dosages,parent1="P1",parent2="P2",
F1=setdiff(colnames(ALL_dosages),c("P1","P2")),polysomic=T,disomic=F,mixed=F,
ploidy=4)
data("gp_df")
chk1<-checkF1(input_type="probabilistic",probgeno_df=gp_df,parent1="P1",parent2="P2",
F1=setdiff(levels(gp_df$SampleName),c("P1","P2")),polysomic=T,disomic=F,mixed=F,
ploidy=4)
## End(Not run)</pre>
```

check\_map

Check the quality of a linkage map

# Description

Perform a series of checks on a linkage map and visualise the results using heatplots. The difference between the pairwise and multi-point r estimates are also plotted against the LOD of the pairwise estimate. The weighted root mean square error of these differences (weighted by the LOD scores) is printed on the console.

```
check_map(
  linkage_list,
  maplist,
```

```
mapfn = "haldane",
lod.thresh = 5,
detail = 1,
plottype = c("", "pdf", "png")[1],
prefix = ""
)
```

# **Arguments**

A named list with r and LOD of markers within linkage groups. linkage\_list A list of maps. In the first column marker names and in the second their position. maplist The map function used in generating the maps, either one of "haldane" or "kosambi". mapfn By default "haldane" is assumed. lod.thresh Numeric. Threshold for the LOD values to be displayed in heatmap, by default 5 (set at 0 to display all values) detail Level of detail for heatmaps, by default 1 cM. Values less than 0.5 cM can have serious performance implications. Option to specify graphical device for plotting, (either png or pdf), or by default plottype "", in which case plots are directly plotted within R Optional prefix appended to plot names if outputting plots. prefix

# **Examples**

```
## Not run:
data("maplist_P1","all_linkages_list_P1")
check_map(linkage_list = all_linkages_list_P1, maplist = maplist_P1)
## End(Not run)
```

check\_marker\_assignment

Check for consistent marker assignment between both parents

### **Description**

Function to ensure there is consistent marker assignment to chromosomal linkage groups for biparental markers

```
check_marker_assignment(
  marker_assignment.P1,
  marker_assignment.P2,
  log = NULL,
  verbose = TRUE
)
```

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### **Arguments**

marker\_assignment.P1

A marker assignment matrix for parent 1 with markernames as rownames and at least containing the column "Assigned\_LG"; the output of homologue\_lg\_assignment.

marker\_assignment.P2

A marker assignment matrix for parent 2 with markernames as rownames and at

least containing the column "Assigned\_LG"; the output of homologue\_lg\_assignment.

log Character string specifying the log filename to which standard output should be

written. If NULL (by default) log is send to stdout.

verbose Should messages be sent to stdout or log?

#### Value

Returns a list of matrices with corrected marker assignments.

# **Examples**

```
data("marker_assignments_P1"); data("marker_assignments_P2")
check_marker_assignment(marker_assignments_P1,marker_assignments_P2)
```

check\_maxP

check your dataset's maxP distribution

#### **Description**

Function to assess the distribution of maximum genotype probabilities (maxP), if these are available. The function plots a violin graph showing the distribution of the samples' maxP.

### Usage

```
check_maxP(probgeno_df)
```

## **Arguments**

probgeno\_df

A data frame as read from the scores file produced by function saveMarkerModels of R package fitPoly, or alternatively, a data frame containing the following columns:

**SampleName** Name of the sample (individual)

MarkerName Name of the marker

**P0** Probabilities of dosage score '0'

**P1...** Probabilities of dosage score '1' etc. (up to max dosage, e.g. P4 for tetraploid population)

**maxP** Maximum genotype probability identified for a particular individual and marker combination

**maxgeno** Most probable dosage for a particular individual and marker combination

**geno** Most probable dosage for a particular individual and marker combination, if maxP exceeds a user-defined threshold (e.g. 0.9), otherwise NA

18 cluster\_per\_LG

# Value

This function does not return any value, is simply a visualisation tool to help assess data quality.

### **Examples**

```
data("gp_df")
check_maxP(gp_df)
```

chk1

Example output of the checkF1 function

# Description

Example output of the checkF1 function

### Usage

chk1

### **Format**

An object of class list of length 2.

cluster\_per\_LG

Cluster 1.0 markers into correct homologues per linkage group

# **Description**

Clustering at one LOD score for all markers does usually not result in correct classification of homologues. Usually there are more clusters of (pseudo)homologues than expected. This function lets you inspect every linkage group separately and allows for clustering at a different LOD threshold per LG.

```
cluster_per_LG(
   LG,
   linkage_df,
   LG_hom_stack,
  LOD_sequence,
   modify_LG_hom_stack = FALSE,
   nclust_out = NULL,
   network.layout = c("circular", "stacked", "n"),
   device = NULL,
   label.offset = 1,
```

cluster\_per\_LG

```
cex.lab = 0.7,
log = NULL,
...
)
```

#### **Arguments**

LG Integer. Linkage group to investigate. linkage\_df A data frame as output of linkage with arguments marker type 1 = c(1, 0) and markertype2=NULL. LG\_hom\_stack A data.frame with columns "SxN\_Marker" providing 1.0 markernames and "LG" and "homologue" providing linkage group and homologue respectively. LOD\_sequence A numeric or vector of numerics giving LOD threshold(s) at which clustering should be performed. modify\_LG\_hom\_stack Logical. Should LG\_hom\_stack be modified and returned? Number of clusters in the output. If there are more clusters than this number nclust\_out only the nclust\_out largest clusters are returned. network.layout Network layout: "circular" or "stacked". If "n" no network is plotted. Function of the graphics device to plot to (e.g. pdf, png, jpeg). The active device device is used when NULL label.offset Offset of labels. Only used if network.layout="circular". cex.lab label character expansion. Only for network.layout="circular". Character string specifying the log filename to which standard output should be log written. If NULL log is send to stdout.

#### Value

A modified  $LG_{hom\_stack}$  data. frame if  $modify_LG_{hom\_stack} = TRUE$ 

Arguments passed to device.

### **Examples**

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cluster\_SN\_markers

Cluster 1.0 markers

# **Description**

cluster\_SN\_markers clusters simplex nulliplex at different LOD scores.

# Usage

```
cluster_SN_markers(
  linkage_df,
  LOD_sequence = 7,
  independence_LOD = FALSE,
  LG_number,
  ploidy,
  parentname = "",
  plot_network = FALSE,
  min_clust_size = 1,
  plot_clust_size = TRUE,
  max_vertex_size = 5,
  min_vertex_size = 2,
  phase_considered = "All",
  log = NULL
)
```

# Arguments

linkage\_df A linkage data.frame as output of linkage calculating linkage between 1.0

markers.

LOD\_sequence A numeric vector. Specifying a sequence of LOD thresholds at which clustering

is performed.

independence\_LOD

Logical. Should the LOD of independence be used for clustering? (by default,

FALSE.)

LG\_number Expected number of chromosomes (linkage groups)

ploidy Ploidy level of the parent for which clustering is to be performed

parentname Name of parent

plot\_network Logical. Should a network be plotted. Recommended FALSE with large number

of marker combinations.

size of 1 is used, meaning all markers are returned. Setting this to a higher number can be useful for cleaning out mini-clusters that don't show strong linkage

to the rest of the marker set.

plot\_clust\_size

Logical. Should exact cluster size be plotted as vertex labels?

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```
max_vertex_size

Integer. The maximum vertex size. Only used if plot_clust_size=FALSE.

min_vertex_size

Integer. The minimum vertex size. Only used if plot_clust_size=FALSE.

phase_considered

Character string. By default all phases are used, but "coupling" or "repulsion" are also allowed.

log

Character string specifying the log filename to which standard output should be written. If NULL log is send to stdout (console).
```

#### Value

A (named) list of cluster stacks, each of which is a data.frame with columns "marker" and "cluster"

### **Examples**

```
data("SN_SN_P1")
cluster_list<-cluster_SN_markers(SN_SN_P1,LOD_sequence=c(4:10),parentname="P1",ploidy=4,LG_number=5)

compare_maps

Compare linkage maps, showing links between connecting markers
common to neighbouring maps
```

# **Description**

This function allows the visualisation of connections between different maps, showing them side by side.

#### Usage

```
compare_maps(
  maplist,
  chm.wd = 0.2,
  bg.col = "white",
  links.col = "grey42",
  thin.links = NULL,
  type = "karyotype",
  ...
)
```

# **Arguments**

maplist

A list of maps. This is probably most conveniently built on-the-fly in the function call itself. If names are assigned to different maps (list items) these will appear above the maps. In cases of multiple comparisons, for example comparing 1 map of interest to 3 others, the map of interest can be supplied multiple times in the list, interspersed between the other maps. See the example below for details.

chm.wd	The width in inches that linkage groups should be drawn. By default 0.2 inches is used.
bg.col	The background colour of the maps, by default white. It can be useful to use a different background colour for the maps. In this case, supply bg.col as a vector of colour identifiers, with the same length as maplist and corresponding to its elements in the same order. See the example below for details.
links.col	The colour with which links between maps are drawn, by default grey.
thin.links	Option to thin the plotting of links between maps, which might be useful if there are very many shared markers in a small genetic region. By default NULL, otherwise supply a value (in cM) for the minimum genetic distance between linking-lines (e.g. 0.5).
type	Plot type, by default "karyotype". If "scatter" is requested a scatter plot is drawn, but only if the comparison is between 2 maps.
	option to supply arguments to the plot function (e.g. main = to add a title to the plot)

### Value

NULL

# **Examples**

```
\label{lem:data("map1", "map2", "map3")} $$ compare_maps(maplist=list("1a"=map1,"c08"=map2,"1b"=map3),bg.col=c("thistle","white","skyblue")) $$ $$ colered ("thistle","white","skyblue") $$ $$ $$ ("thistle","white","skyblue") $$ $$ ("thistle","white","skyblue") $$ $$ ("thistle","white","skyblue") $$ ("thistle","white","white","white","white","white","white "thistle","white "t
```

 ${\tt consensus\_LG\_assignment}$ 

Consensus LG assignment

# Description

Assign markers to an LG based on consensus between two parents.

```
consensus_LG_assignment(
  P1_assigned,
  P2_assigned,
  LG_number,
  ploidy,
  consensus_file = NULL,
  log = NULL
)
```

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# **Arguments**

P1\_assigned A marker assignment file of the first parent. Should contain the number of link-

ages per LG per marker.

P2\_assigned A marker assignment file of the second parent. Should be the same markertype

as first parent and contain the number of linkages per LG per marker.

LG\_number Number of linkage groups (chromosomes).

ploidy Ploidy level of plant species.

consensus\_file Filename of consensus output. No output is written if NULL.

log Character string specifying the log filename to which standard output should be

written. If NULL log is send to stdout.

### Value

Returns a list containing the following components:

P1\_assigned A (modified) marker assignment matrix of the first parent.

P2\_assigned A (modified) marker assignment matrix of the second parent.

## **Examples**

```
data("P1_SxS_Assigned", "P2_SxS_Assigned_2")
SxS_Assigned_list <- consensus_LG_assignment(P1_SxS_Assigned,P2_SxS_Assigned_2,5,4)</pre>
```

consensus\_LG\_names

Find consensus linkage group names

### **Description**

Chromosomes that should have same number, might have gotten different numbers between parents during clustering. consensus\_LG\_names uses markers present in both parents (usually 1.1 markers) to modify the linkage group numbers in one parent with the other as template

```
consensus_LG_names(
  modify_LG,
  template_SxS,
  modify_SxS,
  merge_LGs = TRUE,
  log = NULL
)
```

### **Arguments**

modify_LG	A data.frame with markernames, linkage group ("LG") and homologue ("homologue"), in which the linkage group numbers will be modified
template_SxS	A file with assigned markers of which (at least) part is present in both parents of the template parent.
modify_SxS	A file with assigned markers of which (at least) part is present in both parents of the parent of which linkage group number are modified.
merge_LGs	Logical, by default TRUE. If FALSE, any discrepency in the number of linkage groups will not be merged, but removed instead. This can be needed if the number of chromosomes identified is not equal between parents, and the user wishes to proceed with a core set.
log	Character string specifying the log filename to which standard output should be written. If NULL log is send to stdout.

### Value

A modified modified\_LG according to the template\_SxS linkage group numbering

# **Examples**

```
data("LGHomDf_P2_2", "P1_SxS_Assigned", "P2_SxS_Assigned")
consensus_LGHomDf<-consensus_LG_names(LGHomDf_P2_2, P1_SxS_Assigned, P2_SxS_Assigned)</pre>
```

```
convert_marker_dosages
```

Convert marker dosages to the basic types.

# Description

Convert marker dosages to the basic types which hold the same information and for which linkage calculations can be performed.

```
convert_marker_dosages(
  dosage_matrix,
  ploidy,
  ploidy2 = NULL,
  parent1 = "P1",
  parent2 = "P2",
  marker_conversion_info = FALSE,
  log = NULL
)
```

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### **Arguments**

dosage\_matrix An integer matrix with markers in rows and individuals in columns.

ploidy ploidy level of the plant species. If parents have different ploidy level, ploidy of

parent1.

ploidy2 ploidy level of the second parent. NULL if both parents have the same ploidy

level.

parent1 Character string specifying the first (usually maternal) parentname.

parent2 Character string specifying the second (usually paternal) parentname.

marker\_conversion\_info

Logical, by default FALSE. Should marker conversion information be returned? This output can be useful for later map phasing step, if original marker coding

is desired (which is most likely the case).

log Character string specifying the log filename to which standard output should be

written. If NULL log is send to stdout.

#### Value

A modified dosage matrix. If marker\_conversion\_info = TRUE, this function returns a list, with both the converted dosage\_matrix, and information on the marker conversions performed per marker.

# **Examples**

```
data("ALL_dosages")
conv<-convert_marker_dosages(dosage_matrix=ALL_dosages, ploidy = 4)</pre>
```

convert\_polyRAD Convert (probabilistic) genotype calling results from polyRAD to input compatible with polymapR

# **Description**

Convert (probabilistic) genotype calling results from polyRAD to input compatible with polymapR

### Usage

```
convert_polyRAD(RADdata)
```

### **Arguments**

RADdata An RADdata (S3 class) object; output of the function PipelineMapping2Parents

having followed the prior steps needed in the polyRAD pipeline. See the polyRAD

vignette for details.

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#### Value

A data frame which include columns: MarkerName, SampleName,P0 ~ Pploidy (e.g. P0 ~ P4 for tetraploid, which represents the probability assigning to this dosage), maxgeno (the most likely dosage), and maxP (the maximum probability)

# **Examples**

```
data("exampleRAD_mapping")
convert_polyRAD(RADdata = exampleRAD_mapping)
```

convert\_updog

Convert (probabilistic) genotype calling results from updog to input compatible with polymapR.

### **Description**

Convert (probabilistic) genotype calling results from updog to input compatible with polymapR.

### Usage

```
convert_updog(mout, output_type = "discrete", min_prob = 0.7)
```

### **Arguments**

mout An object of class multidog; output of the function multidog.

output\_type Output genotypes can be either "discrete" or "probabilistic", defaults to discrete.

min\_prob If genotypes are being discretised, sets the minimum posterior probability in

order to call a genotype with confidence. If maxpostprob < min\_prob, that genotype is made missing. A default of 0.7 is suggested with no particular

motivation.

#### Value

If output\_type is discrete, the function returns a dosage matrix with rownames given by marker names. Columns are organised as parent 1 genotype, parent 2 genotype and then F1 individuals. If output\_type is probabilistic, then the output is a data frame which include columns: MarkerName, SampleName,P0 ~ Pploidy (e.g. P0 ~ P4 for tetraploid, which represents the probability assigning to this dosage), maxgeno (the most likely dosage), and maxP (the maximum probability)

# **Examples**

```
data("mout")
convert_updog(mout)
```

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# Description

fitPoly sometimes uses a "shifted" model to assign dosage scores (e.g. all samples are assigned a dosage one higher than the true dosage). This happens mostly when there are only few dosages present among the samples. This function checks if a shift of +/-1 is possible.

# Usage

```
correctDosages(chk, dosage_matrix, parent1, parent2, ploidy,
polysomic=TRUE, disomic=FALSE, mixed=FALSE,
absent.threshold=0.04)
```

### **Arguments**

•	3	
	chk	data frame returned by function checkF1 when called without shiftmarkers
	dosage_matrix	An integer matrix with markers in rows and individuals in columns.
	parent1	character vector with names of the samples of parent 1
	parent2	character vector with names of the samples of parent 2
	ploidy	ploidy of parents and F1 (correctDosages must not be used for F1 populations where the parents have a different ploidy, or where the parental genotypes are not scored together with the F1); same as used in the call to checkF1 that generated data.frame chk
	polysomic	if TRUE at least all polysomic segtypes are considered; if FALSE these are not specifically selected (but if e.g. disomic is TRUE, any polysomic segtypes that are also disomic will still be considered); same as used in the call to checkF1 that generated data.frame chk
	disomic	if TRUE at least all disomic segtypes are considered (see param polysomic); same as used in the call to checkF1 that generated data.frame chk
	mixed	if TRUE at least all mixed segtypes are considered (see param polysomic). A

absent.threshold

chk

the threshold for the fraction of ALL samples that has the dosage that is assumed to be absent due to mis-fitting of fitPoly; should be at least the assumed error rate of the fitPoly scoring assuming the fitted model is correct

mixed segtype occurs when inheritance in one parent is polysomic (random chromosome pairing) and in the other parent disomic (fully preferential chromosome pairing); same as used in the call to checkF1 that generated data.frame

#### **Details**

A shift of -1 (or +1) is proposed when (1) the fraction of all samples with dosage 0 (or ploidy) is below absent.threshold, (2) the bestfit (not bestParentfit!) segtype in chk has one empty dosage on the low (or high) side and more than one empty dosage at the high (or low) side, and (3) the shifted consensus parental dosages do not conflict with the shifted segregation type.

The returned data.frame (or a subset, e.g. based on the values in the fracNotOk and parNA columns) can serve as parameter shiftmarkers in a new call to checkF1.

Based on the quality scores assigned by checkF1 to the original and shifted versions of each marker the user can decide if either or both should be kept. A data frame combining selected rows of the original and shifted versions of the checkF1 output (which may contain both a shifted and an unshifted version of some markers) can then be used as input to compareProbes or writeDosagefile.

#### Value

a data frame with columns

- markername
- segtype: the bestfit (not bestParentfit!) segtype from chk
- parent1, parent2: the consensus parental dosages; possibly low-confidence, so may be different from those reported in chk
- shift: -1, 0 or 1: the amount by which this marker should be shifted

The next fields are only calculated if shift is not 0:

- fracNotOk: the fraction of ALL samples that are in the dosage (0 or ploidy) that should be empty if the marker is indeed shifted.
- parNA: the number of parental dosages that is missing (0, 1 or 2)

createTetraOriginInput

Create input files for TetraOrigin using an integrated linkage map list and marker dosage matrix

### **Description**

createTetraOriginInput is a function for creating an input file for TetraOrigin, combining map positions with marker dosages.

```
createTetraOriginInput(
  maplist,
  dosage_matrix,
  bin_size = NULL,
  bounds = NULL,
  remove_markers = NULL,
```

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```
outdir = "TetraOrigin",
  output_stem = "TetraOrigin_input",
  plot_maps = TRUE,
  log = NULL
)
```

#### **Arguments**

maplist A list of maps. In the first column marker names and in the second their position.

dosage\_matrix An integer matrix with markers in rows and individuals in columns. Either pro-

vide the unconverted dosages (i.e. before using the <code>convert\_marker\_dosages</code> function), or converted dosages (i.e. screened data), in matrix form. The analysis and results are unaffected by this choice, but it may be simpler to understand the results if converted dosages are used. Conversely, it may be advantageous to use the original unconverted dosages if particular marker alleles are being

tracked for (e.g.) the development of selectable markers afterwards.

bin\_size Numeric. Size (in cM) of the bins to include. If NULL (by default) then all

markers are used (no binning).

bounds Numeric vector. If NULL (by default) then all positions are included, however

if specified then output is limited to a specific region, which is useful for later

fine-mapping work.

remove\_markers Optional vector of marker names to remove from the maps. Default is NULL.

outdir Output directory to which input files for TetraOrigin are written.

plot\_maps Logical. Plot the marker positions of the selected markers using plot\_map.

log Character string specifying the log filename to which standard output should be

written. If NULL log is send to stdout.

# **Examples**

```
## Not run:
data("integrated.maplist","ALL_dosages")
createTetraOriginInput(maplist=integrated.maplist,dosage_matrix=ALL_dosages,bin_size=10)
## End(Not run)
```

create\_phased\_maplist Create a phased homologue map list using the original dosages

# **Description**

create\_phased\_maplist is a function for creating a phased maplist, using integrated map positions and original marker dosages.

### Usage

```
create_phased_maplist(
  input_type = "discrete",
  maplist,
  dosage_matrix.conv,
  dosage_matrix.orig = NULL,
  probgeno_df,
  chk,
  remove_markers = NULL,
  original_coding = FALSE,
 N_{linkages} = 2,
  lower_bound = 0.05,
  ploidy,
  ploidy2 = NULL,
 marker_assignment.1,
 marker_assignment.2,
  parent1 = "P1",
  parent2 = "P2",
  marker_conversion_info = NULL,
  log = NULL,
  verbose = TRUE
)
```

#### **Arguments**

input\_type

Can be either one of 'discrete' or 'probabilistic'. For the former (default), at least dosage\_matrix.conv must be supplied, while for the latter chk must be supplied.

maplist

A list of maps. In the first column marker names and in the second their position. dosage\_matrix.conv

Matrix of marker dosage scores with markers in rows and individuals in columns.

Note that dosages must be in converted form, i.e. after having run the convert\_marker\_dosages function. Errors may result otherwise.

dosage\_matrix.orig

Optional, by default NULL. The unconverted dosages (i.e. raw dosage data before using the convert\_marker\_dosages function). Required if original\_coding is TRUE.

probgeno\_df

Probabilistic genotypes, for description see e.g. gp\_overview. Required if probabilistic genotypes are used.

chk

Output list as returned by function checkF1. Required if probabilistic genotypes are used.

original\_coding

remove\_markers Optional vector of marker names to remove from the maps. Default is NULL.

Logical. Should the phased map use the original marker coding or not? By default FALSE.

N\_linkages

Number of significant linkages (as defined in homologue\_lg\_assignment) required for high-confidence linkage group assignment.

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lower\_bound Numeric. Lower bound for the rate at which homologue linkages (fraction of

total for that marker) are recognised.

ploidy Integer. Ploidy of the organism.

ploidy2 Optional integer, by default NULL. Ploidy of parent 2, if different from parent 1.

marker\_assignment.1

A marker assignment matrix for parent 1 with markernames as rownames and at

least containing the column "Assigned\_LG".

marker\_assignment.2

A marker assignment matrix for parent 2 with markernames as rownames and at

least containing the column "Assigned\_LG".

parent1 character vector with names of the samples of parent 1
parent2 character vector with names of the samples of parent 2

marker\_conversion\_info

One of the list elements (named 'marker\_conversion\_info') generated by the function convert\_marker\_dosages when the argument marker\_conversion\_info was set to TRUE (not the default, so a user will typically have to re-run this step

first). Required if original\_coding is TRUE.

log Character string specifying the log filename to which standard output should be

written. If NULL log is send to stdout.

verbose Logical, by default TRUE. Should details of the phasing process be given?

# **Examples**

define\_LG\_structure

Generate linkage group and homologue structure of SxN markers

# Description

Function which organises the output of cluster\_SN\_markers into a data frame of numbered linkage groups and homologues. Only use this function if it is clear from the graphical output of cluster\_SN\_markers that there are LOD scores present which define both chromosomes (lower LOD) and homologues (higher LOD).

```
define_LG_structure(cluster_list, LOD_chm, LOD_hom, LG_number, log = NULL)
```

# Arguments

cluster_list	A list of cluster_stacks, the output of cluster_SN_markers.
LOD_chm	Integer. The LOD threshold specifying at which LOD score the markers divide into chromosomal groups
LOD_hom	Integer. The LOD threshold specifying at which LOD score the markers divide into homologue groups
LG_number	Integer. Expected number of chromosomes (linkage groups). Note that if this number of clusters are not present at LOD_chm, the function will abort.
log	Character string specifying the log filename to which standard output should be written. If NULL log is send to stdout.

# Value

A data frame with markers classified by homologue and linkage group.

# Examples

```
\label{local-continuity} $$  data("P1\_homologues") $$  ChHomDf<-define\_LG\_structure(cluster\_list=P1\_homologues,LOD\_chm=3.5,LOD\_hom=5,LG\_number=5) $$  $$  (a) $$  (a) $$  (b) $$  (b) $$  (c) $$  (c
```

exampleRAD_mapping	Example output dataset of polyRAD::PipelineMapping2Parents function

# Description

Example output dataset of polyRAD::PipelineMapping2Parents function

# Usage

```
exampleRAD_mapping
```

### **Format**

An object of class RADdata of length 23.

```
finish_linkage_analysis
```

Linkage analysis between all markertypes within a linkage group.

## **Description**

finish\_linkage\_analysis is a wrapper for linkage, or in the case of probabilistic genotypes, linkage.gp. The function performs linkage calculations between all markertypes within a linkage group.

# Usage

```
finish_linkage_analysis(
  input_type = "discrete",
 marker_assignment,
 dosage_matrix,
  probgeno_df,
  chk,
 marker_combinations = NULL,
 parent1 = "P1",
 parent2 = "P2",
 which_parent = 1,
  ploidy,
 ploidy2 = NULL,
  convert_palindrome_markers = TRUE,
  pairing = "random",
  prefPars = c(0, 0),
  LG_number,
  verbose = TRUE,
  log = NULL,
)
```

### **Arguments**

input\_type

Can be either one of 'discrete' or 'probabilistic'. For the former (default), dosage\_matrix must be supplied, while for the latter probgeno\_df and chk must be supplied.

marker\_assignment

A marker assignment matrix with markernames as rownames and at least containing the column "Assigned\_LG".

dosage\_matrix

A named integer matrix with markers in rows and individuals in columns.

probgeno\_df

A data frame as read from the scores file produced by function saveMarkerModels of R package fitPoly, or alternatively, a data frame containing the following columns:

**SampleName** Name of the sample (individual)

MarkerName Name of the marker

P0 Probabilities of dosage score '0'

**P1...** Probabilities of dosage score '1' etc. (up to max dosage, e.g. P4 for tetraploid population)

**maxP** Maximum genotype probability identified for a particular individual and marker combination

maxgeno Most probable dosage for a particular individual and marker combination

**geno** Most probable dosage for a particular individual and marker combination, if maxP exceeds a user-defined threshold (e.g. 0.9), otherwise NA

chk Output list as returned by function checkF1. This argument is only needed if probabilistic genotypes are used.

marker\_combinations

A matrix with four columns specifying marker combinations to calculate linkage. If NULL all combinations are used for which there are rf functions. Dosages of markers should be in the same order as specified in the names of rf functions. E.g. if using  $1.0\_2.0$  and  $1.0\_3.0$  types use: matrix(c(1,0,2,0,1,0,3,0), byrow = TRUE, ncol = 4)

parent1 Character string specifying the identifier of parent 1, by default "P1"

parent2 Character string specifying the identifier of parent 2, by default "P2"

which\_parent Integer, either 1 or 2, with default 1, where 1 or 2 refers to parent1 or parent2

respectively.

ploidy Integer ploidy level of parent1, and also by default parent2. Argument ploidy2

can be used if parental ploidies differ.

ploidy2 Integer, by default NULL. If parental ploidies differ, use this to specify the ploidy

of parent2.

convert\_palindrome\_markers

Logical. Should markers that behave the same for both parents be converted to a workable format for that parent? E.g.: should 3.1 markers be converted to 1.3?

pairing Type of pairing at meiosis, with options "random" or "preferential". By

default, random pairing is assumned.

prefPars The estimates for preferential pairing parameters for parent 1 and 2, in range

 $0 \le p \le 2/3$ . By default this is c(0,0) (so, no preferential pairing). See the

function test\_prefpairing and the vignette for more details.

LG\_number Number of linkage groups (chromosomes).

verbose Should messages be sent to stdout or log?

log Character string specifying the log filename to which standard output should be

written. If NULL log is send to stdout.

... (Other) arguments passed to linkage

#### Value

Returns a matrix with marker assignments. Number of linkages of 1.0 markers are artificial.

### **Examples**

get\_markertype\_combinations

Visualize and get all markertype combinations for which there are functions in polymapR

# **Description**

Visualize and get all markertype combinations for which there are functions in polymapR

#### **Usage**

```
get_markertype_combinations(ploidy, pairing, nonavailable_combinations = TRUE)
```

# **Arguments**

```
ploidy Ploidy level
pairing Type of pairing. Either "random" or "preferential".
nonavailable_combinations
```

Logical. Should nonavailable combinations be plotted with grey lines?

### Value

A matrix with two columns. Each row represents a function with the first and second markertype.

# **Examples**

```
get_markertype_combinations(ploidy = 4, pairing = "random")
```

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gp\_df

An example of a genotype probability data frame

# **Description**

An example of a genotype probability data frame

### Usage

gp\_df

#### **Format**

Data frame

gp\_overview

 $gp\_overview$ 

### **Description**

Function to generate an overview of genotype probabilities across a population

### Usage

```
gp_overview(probgeno_df, cutoff = 0.7, alpha = 0.1)
```

# **Arguments**

probgeno\_df

A data frame as read from the scores file produced by function saveMarkerModels of R package fitPoly, or equivalently, a data frame containing the following columns:

SampleName Name of the sample (individual)

MarkerName Name of the marker

**P0** Probabilities of dosage score '0'

**P1...** Probabilities of dosage score '1' etc. (up to max dosage, e.g. P4 for tetraploid population)

**maxP** Maximum genotype probability identified for a particular individual and marker combination

**maxgeno** Most probable dosage for a particular individual and marker combination

**geno** Most probable dosage for a particular individual and marker combination, if maxP exceeds a user-defined threshold (e.g. 0.9), otherwise NA

gp\_vignette\_data 37

cutoff a filtering threshold, by default 0.7, to identify individuals with more than alpha

non-missing (maximum) genotype probabilities falling below this cut-off. In other words, by using this default settings (cutoff = 0.7 and alpha = 0.1), you require that 90 in one of the possible genotype dosage classes. This can help identify problematic individuals with many examples of diffuse genotype calls.

Lowering the threshold allows more diffuse calls to be accepted.

alpha Option to specify the quantile of an individuals' scores that will be used to test

against cutoff, by default 0.1.

#### Value

a list with the following elements:

probgeno\_df Input data, filtered based on chosen cutoff

**population\_overview** data.frame containing summary statistics of each individual's genotyping scores

# Examples

```
## Not run:
data("gp_df")
gp_overview(gp_df)
## End(Not run)
```

gp\_vignette\_data

A list of objects needed to build the probabilistic genotype vignette

# Description

A list of objects needed to build the probabilistic genotype vignette

# Usage

```
gp_vignette_data
```

# Format

An object of class list of length 15.

homologue\_lg\_assignment

Assign markers to linkage groups and homologues.

## Description

This is a wrapper combining linkage (or linkage.gp) and assign\_linkage\_group. It is used to assign all marker types to linkage groups by using linkage information with 1.0 markers. It allows for input of marker assignments for which this analysis has already been performed.

# Usage

```
homologue_lg_assignment(
  input_type = "discrete",
  dosage_matrix,
  probgeno_df,
  chk,
  assigned_list,
  assigned_markertypes,
  SN_functions = NULL,
  LG_hom_stack,
  parent1 = "P1"
 parent2 = "P2",
 which_parent = 1,
  ploidy,
  ploidy2 = NULL,
  convert_palindrome_markers = TRUE,
  pairing = "random",
  LG_number,
  LOD_{threshold} = 3,
 write_intermediate_files = TRUE,
  log = NULL,
)
```

# Arguments

input\_type

Can be either one of 'discrete' or 'probabilistic'. For the former (default), dosage\_matrix must be supplied, while for the latter probgeno\_df and chk must be supplied.

dosage\_matrix An integer i

An integer matrix with markers in rows and individuals in columns.

probgeno\_df

A data frame as read from the scores file produced by function saveMarkerModels of R package fitPoly, or alternatively, a data frame containing the following columns:

**SampleName** Name of the sample (individual)

MarkerName Name of the marker

**P0** Probabilities of dosage score '0'

**P1...** Probabilities of dosage score '1' etc. (up to max dosage, e.g. P4 for tetraploid population)

**maxP** Maximum genotype probability identified for a particular individual and marker combination

maxgeno Most probable dosage for a particular individual and marker combination

**geno** Most probable dosage for a particular individual and marker combination, if maxP exceeds a user-defined threshold (e.g. 0.9), otherwise NA

chk Output list as returned by function checkF1. This argument is only needed if probabilistic genotypes are used.

assigned\_list List of data.frames with marker assignments for which the assignment analysis is already performed.

assigned\_markertypes

List of integer vectors of length 2. Specifying the markertypes in the same order as assigned\_list.

SN\_functions A vector of function names to be used. If NULL all remaining linkage functions with SN markers are used.

LG\_hom\_stack A data.frame with markernames ("SxN\_Marker"), linkage group ("LG") and homologue ("homologue")

parent1 A character string specifying name of parent1.

parent2 A character string specifying the name of parent2.

which\_parent Integer, either 1 or 2, with default 1, where 1 or 2 refers to parent1 or parent2 respectively.

ploidy Ploidy level of parent 1. If parent 2 has the same ploidy level, then also the ploidy level of parent 2.

Integer, by default NULL. If parental ploidies differ, use this to specify the ploidy of parent 2. Note that in cross-ploidy situations, ploidy2 must be smaller than ploidy.

convert\_palindrome\_markers

ploidy2

Logical. Should markers that behave the same for both parents be converted to a workable format for that parent? E.g.: should 3.1 markers be converted to 1.3?

pairing Type of pairing. Either "random" or "preferential". By default random pairing is assumed.

LOD\_threshold LOD threshold at which a linkage is considered significant.

write\_intermediate\_files

Logical. Write intermediate linkage files to working directory?

Character string specifying the log filename to which standard output should be written. If NULL log is send to stdout.

. . Arguments passed to linkage

40 LGHomDf\_P1\_1

#### Value

A data. frame specifying marker assignments to linkage group and homologue.

# **Examples**

# **Description**

A nested list with integrated maps

# Usage

```
integrated.maplist
```

#### **Format**

An object of class list of length 5.

LGHomDf\_P1\_1

A data. frame specifying the assigned homologue and linkage group number per SxN marker

#### **Description**

A data.frame specifying the assigned homologue and linkage group number per SxN marker

```
LGHomDf_P1_1
LGHomDf_P2_1
LGHomDf_P2_2
```

linkage 41

#### **Format**

- SxN\_Marker. Markername of simplex nulliplex marker
- homologue. Assigned homologue number
- LG Assigned. linkage group number

An object of class data.frame with 195 rows and 3 columns.

An object of class data. frame with 195 rows and 3 columns.

linkage

Calculate recombination frequency, LOD and phase

# **Description**

linkage is used to calculate recombination frequency, LOD and phase within one type of marker or between two types of markers.

## Usage

```
linkage(
  dosage_matrix,
 markertype1 = c(1, 0),
 markertype2 = NULL,
 parent1 = "P1",
  parent2 = "P2",
 which_parent = 1,
  ploidy,
 ploidy2 = NULL,
 G2_test = FALSE,
  convert_palindrome_markers = TRUE,
 LOD_threshold = 0,
  pairing = "random",
 prefPars = c(0, 0),
  combinations_per_iter = NULL,
  iter_RAM = 500,
  ncores = 1,
  verbose = TRUE,
  full_output = FALSE,
  log = NULL
)
```

#### **Arguments**

dosage\_matrix An integer matrix with markers in rows and individuals in columns.

markertype1

A vector of length 2 specifying the first markertype to compare. The first element specifies the dosage in which\_parent (see below), the second in the other parent.

42 linkage

markertype2 A vector of length 2 specifying the first markertype to compare. This argument

is optional. If not specified, the function will calculate linkage within the markertype as specified by markertype1. The first element specifies the dosage in

which\_parent (see below), the second in the other parent.

parent1 Character string specifying the name of parent1 as provided in the column-

names of dosage\_matrix. By default, "P1".

parent2 Character string specifying the other parent as provided in the column-names of

dosage\_matrix. By default, "P2".

which\_parent Integer, either 1 or 2, with default 1, where 1 or 2 refers to parent1 or parent2

respectively. For example, if you wish to estimate linkage between markers with alleles that are polymorphic (i.e. segregating) and originates from parent1, then which\_parent = 1. A bi-parental marker is a marker such as a 1x1 marker, so having a segregating allele in both parents. For linkage estimation between pairs of bi-parental markers, the result does not depend on this argument. For linkage estimation between e.g. a 1x0 and 1x1 marker, then which\_parent should be 1. Similarly, to calculate linkage between 0x1 and 1x1 markers, which\_parent

should be 2.

ploidy Integer. The ploidy of the parent 1. If parent2 has the same ploidy level, then

also the ploidy level of parent 2.

ploidy2 Integer, by default NULL. If parental ploidies differ, use this to specify the ploidy

of parent2.

G2\_test Apply a G2 test (LOD of independence) in addition to the LOD of linkage.

convert\_palindrome\_markers

Logical. Should markers that behave the same for both parents be converted to a workable format for that parent? E.g.: should 3.1 markers be converted to 1.3?

If unsure, set to TRUE.

LOD\_threshold Minimum LOD score of linkages to report. Recommended to use for large num-

ber (> millions) of marker comparisons in order to reduce memory usage.

pairing Type of chromosomal pairing behaviour during meiosis, either "random" or

"preferential". By default, random pairing is assumed (i.e. polysomic inheritance) is assumed. Note that this default does not affect linkage estimation

in a diploid, where pairing is arguably not random.

prefPars The estimates for preferential pairing parameters for the target and other parent,

respectively, in range  $0 \le p \le 2/3$ . By default this is c(0,0) (so, no preferential pairing). See the function test\_prefpairing and the vignette for more details.

combinations\_per\_iter

Optional integer. Number of marker combinations per iteration.

iter\_RAM A (very) conservative estimate of working memory in megabytes used per core.

It only takes the size frequency matrices into account. Actual usage is more, especially with large number of linkages that are reported. Reduce memory

usage by using a higher LOD\_threshold.

ncores Number of cores to use. Works both for Windows and UNIX (using doParallel).

Use parallel::detectCores() to find out how many cores you have available.

verbose Should messages be sent to stdout?

linkage.gp 43

full_output	Logical, by default FALSE. If TRUE, the complete output over all phases and showing marker combination counts is returned.
log	Character string specifying the log filename to which standard output should be written. If NULL log is send to stdout.

# Value

Returns a data.frame with columns:

marker\_a first marker of comparison. If markertype2 is specified, it has the type of markertype1.marker\_b second marker of comparison. It has the type of markertype2 if specified.

 ${f r}$  (estimated) recombinations frequency

```
LOD (estimated) LOD score phase phase between markers
```

# **Examples**

linkage.gp

Calculate recombination frequency, LOD and phase using genotype probabilities

# Description

linkage.gp is used to calculate recombination frequency, LOD and phase within one type of marker or between two types of markers.

```
linkage.gp(
  probgeno_df,
  chk,
  pardose = NULL,
  markertype1 = c(1, 0),
  markertype2 = NULL,
  target_parent = match.arg(c("P1", "P2")),
  G2_test = FALSE,
  LOD_threshold = 0,
  prefPars = c(0, 0),
```

44 linkage.gp

```
combinations_per_iter = NULL,
  iter_RAM = 500,
 ncores = 2,
  verbose = TRUE,
  check_gall_mult = FALSE,
 method = "approx",
 log = NULL
)
```

#### **Arguments**

probgeno\_df

A data frame as read from the scores file produced by function saveMarkerModels of R package fitPoly, or alternatively, a data frame containing the following columns:

**SampleName** Name of the sample (individual)

MarkerName Name of the marker **P0** Probabilities of dosage score '0'

P1... Probabilities of dosage score '1' etc. (up to max dosage, e.g. P4 for tetraploid population)

maxP Maximum genotype probability identified for a particular individual and marker combination

maxgeno Most probable dosage for a particular individual and marker combination

geno Most probable dosage for a particular individual and marker combination, if maxP exceeds a user-defined threshold (e.g. 0.9), otherwise NA

Output list as returned by function checkF1 chk

Option to include the most likely (discrete) parental dosage scores, used mainly pardose

for internal calls of this function. By default NULL

A vector of length 2 specifying the first markertype to compare. The first elemarkertype1

ment specifies the dosage in target\_parent (and the second in the other par-

A vector of length 2 specifying the first markertype to compare. This argument

is optional. If not specified, the function will calculate linkage within the markertype as specified by markertype1. The first element specifies the dosage in

target\_parent (and the second in the other parent).

target\_parent Which parent is being targeted (only acceptable options are "P1" or "P2"), ie.

> which parent is of specific interest? If this is the maternal parent, please specify as "P1". If the paternal parent, please use "P2". The actual identifiers of the two parents are entered using the arguments parent1\_replicates and

parent2\_replicates.

G2\_test Apply a G2 test (LOD of independence) in addition to the LOD of linkage.

Minimum LOD score of linkages to report. Recommended to use for large num-

ber (> millions) of marker comparisons in order to reduce memory usage.

prefPars The estimates for preferential pairing parameters for parent 1 and 2, in range

 $0 \le p \le 2/3$ . By default this is c(0,0) (so, no preferential pairing). See the

function test\_prefpairing and the vignette for more details.

markertype2

LOD\_threshold

linkage.gp 45

combinations\_per\_iter

Optional integer. Number of marker combinations per iteration.

iter\_RAM A (very) conservative estimate of working memory in megabytes used per core.

It only takes the size frequency matrices into account. Actual usage is more, especially with large number of linkages that are reported. Reduce memory

usage by using a higher LOD\_threshold.

ncores Number of cores to use. Works both for Windows and UNIX (using doParallel).

Use parallel::detectCores() to find out how many cores you have available.

verbose Should messages be sent to stdout?

check\_qall\_mult

Check the qall\_mult column of chk, and filter out markers with qall\_mult =

0. By default FALSE.

method Either "approx" or "mappoly". If "approx" (the default method), then an ap-

proximated estimator is used which introduces a small amount of bias in the estimator of recombination frequency. If method "mappoly" is specified, the full likelihood is used in the estimation, leading to an unbiased estimator (this has been implemented in the mappoly package of Marcelo Mollinari). The mappoly method has higher computational demands which may introduce problems for

larger datasets, but will lead to higher accuracy overall.

log Character string specifying the log filename to which standard output should be

written. If NULL log is send to stdout.

## Value

Returns a data.frame with columns:

marker\_a: first marker of comparison. If markertype2 is specified, it has the type of markertype1.

marker\_b: second marker of comparison. It has the type of markertype2 if specified.

**r:** recombination frequency

LOD: LOD score associated with r

**phase:** phase between markers

46 map3

map1

A sample map

# Description

A sample map

# Usage

map1

# **Format**

An object of class data. frame with 100 rows and 2 columns.

map2

A sample map

# Description

A sample map

# Usage

map2

# **Format**

An object of class data. frame with 100 rows and 2 columns.

map3

A sample map

# Description

A sample map

# Usage

map3

#### **Format**

An object of class data. frame with 60 rows and 2 columns.

maplist\_P1 47

maplist\_P1

A list of maps of one parent

# Description

A list of maps of one parent

# Usage

```
maplist_P1
maplist_P1_subset
maplist_P2_subset
```

#### **Format**

An object of class list of length 5. An object of class list of length 5. An object of class list of length 5.

marker\_binning

Perform binning of markers.

# Description

marker\_binning allows for binning of very closely linked markers and choses one representative.

```
marker_binning(
  dosage_matrix,
  linkage_df,
  r_thresh = NA,
  lod_thresh = NA,
  target_parent = "P1",
  other_parent = "P2",
  max_marker_nr = NULL,
  max_iter = 10,
  log = NULL
)
```

## **Arguments**

dosage\_matrix A dosage matrix.
linkage\_df A linkage data.frame.

r\_thresh Numeric. Threshold at which markers are binned. Is calculated if NA. lod\_thresh Numeric. Threshold at which markers are binned. Is calculated if NA.

target\_parent A character string specifying the name of the target parent.

other\_parent A character string specifying the name of the other parent.

max\_marker\_nr The maximum number of markers per homologue. If specified, LOD threshold

is optimized based on this number.

max\_iter Maximum number of iterations to find optimum LOD threshold. Only used if

max\_marker\_nr is specified.

log Character string specifying the log filename to which standard output should be

written. If NULL log is send to stdout.

#### Value

A list with the following components:

binned\_df A linkage data.frame with binned markers removed.

removed A data frame containing binned markers and their representatives.

left Integer. Number markers left.

# **Examples**

```
data("screened_data3", "all_linkages_list_P1_split")
binned_markers<-marker_binning(screened_data3, all_linkages_list_P1_split[["LG2"]][["homologue3"]])</pre>
```

marker\_data\_summary Summarize marker data

# **Description**

Gives a frequency table of different markertypes, relative frequency per markertype of incompatible offspring and the names of incompatible progeny.

```
marker_data_summary(
  dosage_matrix,
  ploidy,
  ploidy2 = NULL,
  pairing = c("random", "preferential"),
  parent1 = "P1",
  parent2 = "P2",
```

marker\_data\_summary 49

```
progeny_incompat_cutoff = 0.1,
  verbose = TRUE,
  shortform = FALSE,
  log = NULL
)
```

#### **Arguments**

dosage\_matrix An integer matrix with markers in rows and individuals in columns.

ploidy Integer. Ploidy of parent 1, and .

ploidy2 Ploidy of parent 2, by default NULL, as it is assumed ploidy2 equals ploidy.

pairing Type of pairing. "random" or "preferential".

parent1 Column name of first parent. Usually maternal parent.

Column name of second parent. Usually paternal parent.

Column name of second parent. Usually paternal parent.

progeny\_incompat\_cutoff

The relative number of incompatible dosages per genotype that results in reporting this genotype as incompatible. Incompatible dosages are greater than maximum number of alleles than can be inherited or smaller than the minimum

number of alleles that can be inherited.

verbose Logical, by default TRUE - should intermediate messages be written to stout?

shortform Logical, by default FALSE. Returns only a shortened output with parental dosage

summary, used internally by some functions.

log Character string specifying the log filename to which standard output should be

written. If NULL log is send to stdout.

#### Value

Returns a list containing the following components:

parental\_info frequency table of different markertypes. Names start with parentnames, and

behind that the dosage score.

offspring\_incompatible

Rate of incompatible ("impossible") marker scores (given as percentages of the

total number of observed marker scores per marker class)

progeny\_incompatible

progeny names having incompatible dosage scores higher than threshold at progeny\_incompat\_cutoff.

```
data("ALL_dosages")
summary_list<-marker_data_summary(dosage_matrix = ALL_dosages, ploidy = 4)</pre>
```

50 MDSMap\_from\_list

MDSMap\_from\_list Wrapper fu

Wrapper function for MDSMap to generate linkage maps from list of pairwise linkage estimates

# Description

Create multidimensional scaling maps from a list of linkages

#### Usage

```
MDSMap_from_list(
  linkage_list,
  write_to_file = FALSE,
  mapdir = "mapping_files_MDSMap",
  plot_prefix = "",
  log = NULL,
  ...
)
```

#### **Arguments**

linkage\_list A named list with r and LOD of markers within linkage groups.

write\_to\_file Should output be written to a file? By default FALSE, if TRUE then output, includ-

ing plots from MDSMap are saved in the same directory as the one used for input files. These plots are currently saved as pdf images. If a different plot format is required (e.g. for publications), then run the MDSMap function <code>estimate.map</code> (or similar) directly and save the output with a different plotting function as wrapper

around the map function call.

mapdir Directory to which map input files are initially written. Also used for output if

write\_to\_file=TRUE

plot\_prefix prefix for the filenames of output plots.

log Character string specifying the log filename to which standard output should be

written. If NULL log is send to stdout.

... Arguments passed to estimate.map.

```
## Not run:
data("all_linkages_list_P1")
maplist_P1 <- MDSMap_from_list(all_linkages_list_P1[1])
## End(Not run)</pre>
```

merge\_homologues 51

## **Description**

Based on additional information, homologue fragments, separated during clustered should be merged again. merge\_homologues allows to merge homologues per linkage group based on user input.

#### Usage

```
merge_homologues(LG_hom_stack, ploidy, LG, mergeList = NULL, log = NULL)
```

#### **Arguments**

LG\_hom\_stack A data.frame with markernames, linkage group ("LG") and homologue ("homologue")

ploidy The ploidy level of the plant species.

LG The linkage group where the to be merged homologue fragments are in.

mergeList A list of vectors of length 2, specifying the numbers of the homologue fragments

to be merged. User input is asked if NULL.

log Character string specifying the log filename to which standard output should be

written. If NULL log is send to stdout.

## Value

A modified LG\_hom\_stack

#### **Examples**

```
data("LGHomDf_P2_1")
merged<-merge_homologues(LGHomDf_P2_1,ploidy=4,LG=2,mergeList=list(c(1,5)))</pre>
```

mout

Example output dataset of updog::multidog function

#### Description

Example output dataset of updog::multidog function

#### Usage

mout

#### **Format**

An object of class multidog of length 2.

52 overviewSNlinks

overviewSNlinks

Plotting 1.0 links between homologues

# **Description**

overviewSNlinks is written to enable merging of homologue fractions. Fractions of homologues will have more markers in coupling than in repulsion, whereas separate homologues will only have markers in repulsion.

# Usage

```
overviewSNlinks(
   linkage_df,
   LG_hom_stack,
   LG,
   LOD_threshold,
   ymax = NULL,
   log = NULL
)
```

# **Arguments**

linkage\_df A data.frame as output of linkage with arguments markertype1=c(1,0) and

markertype2=NULL.

LG\_hom\_stack A data.frame with a column "SxN\_Marker" specifying markernames, a column

"homologue" specifying homologue cluster and "LG" specifying linkage group.

LG Integer. Linkage group number of interest.

LOD\_threshold Numeric. LOD threshold of linkages which are plotted.

ymax Maximum y-limit of the plots.

log Character string specifying the log filename to which standard output should be

written. If NULL log is send to stdout.

P1\_homologues 53

P1\_homologues

A list of cluster stacks at different LOD scores

# **Description**

A list of cluster stacks at different LOD scores

# Usage

```
P1_homologues
P2_homologues
```

P2\_homologues\_triploid

#### **Format**

A list with with LOD thresholds as names. The list contains dataframes with the following format:

- marker. markername
- pseudohomologue. name of (pseudo)homologue

An object of class list of length 10.

An object of class list of length 15.

P1\_SxS\_Assigned

A data.frame with marker assignments

# Description

A data. frame with marker assignments

```
P1_SxS_Assigned
P2_SxS_Assigned
P2_SxS_Assigned_2
P1_DxN_Assigned
P2_DxN_Assigned
marker_assignments_P1
marker_assignments_P2
```

54 p4\_functions

#### **Format**

A data.frame with at least the following columns:

- Assigned\_LG. The assigned linkage group
- Assigend\_hom1. The homologue with most linkages

The columns LG1 - LGn and Hom1 - Homn give the number of hits per marker for that linkage group/homologue. Assigned\_hom2 .. gives the nth homologue with most linkages.

An object of class matrix (inherits from array) with 301 rows and 14 columns.

An object of class matrix (inherits from array) with 301 rows and 14 columns.

An object of class matrix (inherits from array) with 111 rows and 14 columns.

An object of class matrix (inherits from array) with 101 rows and 14 columns.

An object of class matrix (inherits from array) with 1094 rows and 16 columns.

An object of class matrix (inherits from array) with 1127 rows and 16 columns.

p4_functions	Calculate recombination frequency, LOD and log-likelihood from fre-
	quency tables in a preferential pairing tetraploid

# Description

This group of functions is called by linkage.

# **Arguments**

X	A frequency table of the different classes of dosages in the progeny. The column names start with "n_". Followed by the dosage of the first marker and then of the second.
p1	Preferential pairing parameter for parent 1, numeric value in range $0 \le p1 \le 2/3$
p2	Preferential pairing parameter for parent 2, numeric value in range $0 \le p2 \le 2/3$
ncores	Number of cores to use for parallel processing (deprecated).

## Value

A list with the following items:

r\_mat A matrix with recombination frequencies for the different phases

LOD\_mat A matrix with LOD scores for the different phases

logL\_mat A matrix with log likelihood ratios for the different phases

phasing\_strategy

A character string specifying the phasing strategy. "MLL" for maximum likelihood en "MINR" for minimum recombination frequency.

parental\_quantities 55

```
possible_phases
```

The phases between markers that are possible. Same order and length as column names of output matrices.

parental\_quantities Calculate frequency of each markertype.

# **Description**

Plots and returns frequency information for each markertype.

# Usage

```
parental_quantities(
  dosage_matrix,
  parent1 = "P1",
  parent2 = "P2",
  log = NULL,
  ...
)
```

# Arguments

dosage\_matrix An integer matrix with markers in rows and individuals in columns.

parent1 Character string specifying the first (usually maternal) parentname.

parent2 Character string specifying the second (usually paternal) parentname.

Character string specifying the log filename to which standard output should be written. If NULL log is send to stdout.

Arguments passed to barplot

#### Value

A named vector containing the frequency of each markertype in the dataset.

```
data("ALL_dosages", "screened_data")
parental_quantities(dosage_matrix=ALL_dosages)
parental_quantities(dosage_matrix=screened_data)
```

56 phased.maplist

PCA_progeny	Perform a PCA on progeny

# **Description**

Principal component analysis in order to identify individuals that deviate from the population.

# Usage

```
PCA_progeny(dosage_matrix, highlight = NULL, colors = NULL, log = NULL)
```

#### **Arguments**

dosage\_matrix An integer matrix with markers in rows and individuals in columns.

highlight A list of character vectors specifying individual names that should be high-

lighted

colors Highlight colors. Vector of the same length as highlight.

log Character string specifying the log filename to which standard output should be

written. If NULL log is send to stdout.

#### **Details**

Missing values are imputed by taking the mean of marker dosages per marker.

# **Examples**

```
data("ALL_dosages")
PCA_progeny(dosage_matrix=ALL_dosages, highlight=list(c("P1", "P2")), colors="red")
```

phased.maplist A list of phased maps

# Description

A list of phased maps

# Usage

```
phased.maplist
```

#### **Format**

An object of class list of length 5.

phase\_SN\_diploid 57

phase\_SN\_diploid

Phase 1.0 markers at the diploid level

# **Description**

phase\_SN\_diploid phases simplex x nulliplex markers for a diploid parent.

#### Usage

```
phase_SN_diploid(
   linkage_df,
   cluster_list,
   LOD_chm = 3.5,
   LG_number,
   independence_LOD = FALSE,
   log = NULL
)
```

# Arguments

written. If NULL log is send to stdout (console).

#### Value

A data.frame with markers classified by homologue and linkage group.

```
\label{lem:continuous} data("SN_SN_P2\_triploid", "P2\_homologues\_triploid") \\ cluster_list2 <-phase_SN_diploid(SN_SN_P2\_triploid, P2\_homologues\_triploid, LOD\_chm=5, LG\_number = 3) \\
```

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plot\_hom\_vs\_LG

Plot homologue position versus integrated positions

## **Description**

Plot homologue position versus integrated positions

# Usage

```
plot_hom_vs_LG(map_df, maplist_homologue)
```

# **Arguments**

```
map_df A dataframe of a map that defines a linkage group.
maplist_homologue
```

A list of maps were each item represents a homoloogue.

#### **Examples**

plot\_map

Plot linkage maps

#### **Description**

Makes a simple plot of a list of generated linkage maps

```
plot_map(
    maplist,
    highlight = NULL,
    bg_col = "grey",
    highlight_col = "yellow",
    colname_in_mark = NULL,
    colname_beside_mark = NULL,
    palette_in_mark = colorRampPalette(c("white", "purple")),
    palette_beside_mark = colorRampPalette(c("white", "green")),
    color_by_type = FALSE,
    dosage_matrix = NULL,
    parent1 = "P1",
    parent2 = "P2",
```

plot\_map 59

```
legend = FALSE,
...,
legend.args = list(x = 1, y = 120)
)
```

#### **Arguments**

maplist A list of maps. In the first column marker names and in the second their position.

highlight A list of the same length of maplist with vectors of length 2 that specifies the

limits in cM from and to which the plotted chromosomes should be highlighted.

bg\_col The background colour of the map.

highlight\_col The color of the highlight. Only used if highlight is specified.

colname\_in\_mark

Optional. The column name of the value to be plotted as marker color.

colname\_beside\_mark

Optional. The column name of the value to be plotted beside the markers.

palette\_in\_mark, palette\_beside\_mark

Color palette used to plot values. Only used if colnames of the values are speci-

fied

color\_by\_type Logical. Should the markers be coloured by type? If TRUE, dosage\_matrix

should be specified.

dosage\_matrix Optional (by default NULL). Dosage matrix of marker genotypes, input of linkage

parent1 Character string specifying the first (usually maternal) parentname.

parent2 Character string specifying the second (usually paternal) parentname.

legend Logical. Should a legend be drawn?

... Arguments passed to plot

legend.args Optional extra arguments to pass to legend, by default a list with x = 1 and y

= 120 (position of the legend). Additional arguments should be passed using name = value, i.e. as a named list. Note that arguments 1ty (= 1) and 1wd (= 2) have already been used internally (as well as legend and col), so cannot be

re-specified without causing an error.

r2\_functions

plot_phased_maplist	Visualise the phased homologue maplist
---------------------	--

## **Description**

plot\_phased\_maplist is a function for visualising a phased maplist, the output of create\_phased\_maplist

#### Usage

```
plot_phased_maplist(
   phased.maplist,
   ploidy,
   ploidy2 = NULL,
   cols = c("black", "darkred", "navyblue"),
   width = 0.2,
   mapTitles = NULL
)
```

# Arguments

```
phased.maplist A list of phased linkage maps, the output of create_phased_maplist

ploidy Integer. Ploidy of the organism.

ploidy2 Optional integer, by default NULL. Ploidy of parent 2, if different from parent 1.

cols Vector of colours for the integrated, parent1 and parent2 maps, respectively.

width Width of the linkage maps, by default 0.2

mapTitles Optional vector of titles for maps, by default names of maplist, or titles LG1, LG2 etc. are used.
```

# **Examples**

```
data("phased.maplist")
plot_phased_maplist(phased.maplist, ploidy = 4)
```

r2_functions	Calculate recombination frequency, LOD and log-likelihood from fre-
	quency tables in a random pairing diploid cross.

#### **Description**

This group of functions is called by linkage.

r3\_functions 61

#### Usage

```
r2_1.0_1.0(x, ncores = 1)
r2_1.0_1.1(x, ncores = 1)
r2_1.1_1.1(x, ncores = 1)
```

#### **Arguments**

x A frequency table of the different classes of dosages in the progeny. The column

names start with "n\_". Followed by the dosage of the first marker and then of

the second.

ncores Number of cores to use for parallel processing (deprecated).

#### Value

A list with the following items:

r\_mat A matrix with recombination frequencies for the different phases

LOD\_mat A matrix with LOD scores for the different phases

logL\_mat A matrix with log likelihood ratios for the different phases

phasing\_strategy

A character string specifying the phasing strategy. "MLL" for maximum likeli-

hood en "MINR" for minimum recombination frequency.

possible\_phases

The phases between markers that are possible. Same order and length as column

names of output matrices.

r3\_functions Calculate recombination frequency, LOD and log-likelihood from frequency tables in a random pairing triploid from a 4x2 or 2x4 cross.

# Description

This group of functions is called by linkage.

```
r3_2_1.0_1.0(x, ncores = 1)
r3_2_1.0_1.1(x, ncores = 1)
r3_2_1.0_1.2(x, ncores = 1)
r3_2_1.2_1.2(x, ncores = 1)
```

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#### **Arguments**

x A frequency table of the different classes of dosages in the progeny. The column

names start with "n\_". Followed by the dosage of the first marker and then of

the second.

ncores Number of cores to use for parallel processing (deprecated).

#### Value

A list with the following items:

r\_mat A matrix with recombination frequencies for the different phases

LOD\_mat A matrix with LOD scores for the different phases

logL\_mat A matrix with log likelihood ratios for the different phases

phasing\_strategy

A character string specifying the phasing strategy. "MLL" for maximum likeli-

hood en "MINR" for minimum recombination frequency.

possible\_phases

The phases between markers that are possible. Same order and length as column

names of output matrices.

r4\_functions Calculate recombination frequency, LOD and log-likelihood from fre-

quency tables in a random pairing tetraploid

# Description

This group of functions is called by linkage.

#### **Arguments**

x A frequency table of the different classes of dosages in the progeny. The column

names start with "n\_". Followed by the dosage of the first marker and then of

the second.

ncores Number of cores to use for parallel processing (deprecated).

# Value

A list with the following items:

r\_mat A matrix with recombination frequencies for the different phases

LOD\_mat A matrix with LOD scores for the different phases

logL\_mat A matrix with log likelihood ratios for the different phases

phasing\_strategy

A character string specifying the phasing strategy. "MLL" for maximum likeli-

hood en "MINR" for minimum recombination frequency.

r6\_functions 63

```
possible_phases
```

The phases between markers that are possible. Same order and length as column names of output matrices.

r6\_functions

Calculate recombination frequency, LOD and log-likelihood from frequency tables in a random pairing hexaploid

#### **Description**

This group of functions is called by linkage.

#### **Arguments**

Х

A frequency table of the different classes of dosages in the progeny. The column names start with "n\_". Followed by the dosage of the first marker and then of the second.

#### Value

A list with the following items:

r\_mat A matrix with recombination frequencies for the different phases

LOD\_mat A matrix with LOD scores for the different phases

logL\_mat A matrix with log likelihood ratios for the different phases

phasing\_strategy

A character string specifying the phasing strategy. "MLL" for maximum likeli-

hood en "MINR" for minimum recombination frequency.

possible\_phases

The phases between markers that are possible. Same order and length as column

names of output matrices.

r\_LOD\_plot

Plot r versus LOD

## Description

r\_LOD\_plot plots r versus LOD, colour separated for different phases.

```
r_LOD_plot(
  linkage_df,
  plot_main = "",
  chm = NA,
  r_max = 0.5,
  tidyplot = TRUE,
  nbins = 200
)
```

#### **Arguments**

linkage\_df A linkage data.frame as output of linkage.

Plot\_main A character string specifying the main title

chm Integer specifying chromosome

r\_max Maximum r value to plot

tidyplot If TRUE (by default), an attempt is made to reduce the plot density using hexago-

nal binning from the ggplot2 package. This is recommended for large datasets,

where the number of pairwise estimates becomes high.

nbins The number of bins in each direction, passed to ggplot2::geom\_hex. Only used

if tidyplot = TRUE. Increasing this number can lead to slower but more accurate

plotting.

# Examples

```
data("SN_SN_P1")
r_LOD_plot(SN_SN_P1)
```

```
screen_for_duplicate_individuals
```

Screen for duplicate individuals

## Description

screen\_for\_duplicate\_individuals identifies and merges duplicate individuals.

# Usage

```
screen_for_duplicate_individuals(
  dosage_matrix,
  cutoff = NULL,
  plot_cor = TRUE,
  log = NULL
)
```

#### **Arguments**

dosage\_matrix An integer matrix with markers in rows and individuals in columns.

cutoff Correlation coefficient cut off. At this correlation coefficient, individuals are

merged. If NULL user input will be asked after plotting.

plot\_cor Logical. Should correlation coefficients be plotted? Can be memory/CPU in-

tensive with high number of individuals.

log Character string specifying the log filename to which standard output should be

written. If NULL log is send to stdout.

#### Value

A matrix similar to dosage\_matrix, with merged duplicate individuals.

#### **Examples**

```
## Not run:
 #user input:
 data("segregating_data")
 screen_for_duplicate_individuals(dosage_matrix=segregating_data,cutoff=0.9,plot_cor=TRUE)
 ## End(Not run)
screen_for_duplicate_individuals.gp
```

Screen for duplicate individuals using weighted genotype probabilities

## **Description**

screen\_for\_duplicate\_individuals.gp identifies and merges duplicate individuals based on probabilistic genotypes. See screen\_for\_duplicate\_individuals for the original function.

# Usage

```
screen_for_duplicate_individuals.gp(
  probgeno_df,
  ploidy,
  parent1 = "P1"
  parent2 = "P2",
 F1,
  cutoff = 0.95,
 plot_cor = TRUE,
  log = NULL
)
```

# **Arguments**

probgeno\_df

A data frame as read from the scores file produced by function saveMarkerModels of R package fitPoly, or alternatively, a data frame containing the following columns:

**SampleName** Name of the sample (individual)

MarkerName Name of the marker

**P0** Probabilities of dosage score '0'

P1... Probabilities of dosage score '1' etc. (up to max dosage, e.g. P4 for tetraploid population)

maxP Maximum genotype probability identified for a particular individual and marker combination

maxgeno Most probable dosage for a particular individual and marker combination

**geno** Most probable dosage for a particular individual and marker combination, if maxP exceeds a user-defined threshold (e.g. 0.9), otherwise NA

ploidy The ploidy of parent 1

parent1 character vector with the sample names of parent 1

parent2 character vector with the sample names of parent 2

F1 character vector with the sample names of the F1 individuals

cutoff Correlation coefficient cut off to declare duplicates. At this correlation coefficient, individuals are merged. If NULL user input will be asked after plotting.

plot\_cor Logical. Should correlation coefficients be plotted? Can be memory/CPU intensive with high number of individuals.

Character string specifying the log filename to which standard output should be

written. If NULL log is send to stdout.

#### Value

log

A data frame similar to input probgeno\_df, but with duplicate individuals merged.

```
screen\_for\_duplicate\_markers \\ Screen for and remove duplicated markers
```

# Description

screen\_for\_duplicate\_markers identifies and merges duplicate markers.

```
screen_for_duplicate_markers(
  dosage_matrix,
  merge_NA = TRUE,
  plot_cluster_size = TRUE,
  ploidy,
  ploidy2 = NULL,
  LG_number,
  estimate_bin_size = FALSE,
  log = NULL
)
```

screen\_for\_NA\_values 67

#### **Arguments**

dosage\_matrix An integer matrix with markers in rows and individuals in columns.

merge\_NA Logical. Should missing values be imputed if non-NA in duplicated marker? By

default, TRUE. If FALSE the dosage scores of representing marker are represented

in the filtered\_dosage\_matrix.

plot\_cluster\_size

Logical. Should an informative plot about duplicate cluster size be given? By

default, TRUE.

ploidy Ploidy level of parent 1. Only needed if estimate\_bin\_size is TRUE

ploidy2 Integer, by default NULL. If parental ploidies differ, use this to specify the ploidy

of parent 2. Only needed if estimate\_bin\_size is TRUE

LG\_number Expected number of chromosomes (linkage groups). Only needed if estimate\_bin\_size

is TRUE

estimate\_bin\_size

Logical, by default FALSE. If TRUE, a very rudimentary calculation is made to estimate the average size of a marker bin, assuming a uniform distribution of

cross-over events and on average one cross-over per bivalent.

log Character string specifying the log filename to which standard output should be

written. If NULL log is send to stdout.

#### Value

A list containing:

**bin\_list** list of binned markers. The list names are the representing markers. This information can later be used to enrich the map with binned markers.

**filtered\_dosage\_matrix** dosage\_matrix with merged duplicated markers. The markers will be given the name of the marker with least missing values.

#### **Examples**

```
data("screened_data3")
dupmscreened <- screen_for_duplicate_markers(screened_data3)</pre>
```

## Description

screen\_for\_NA\_values identifies and can remove rows or columns of a marker dataset based on the relative frequency of missing values.

#### Usage

```
screen_for_NA_values(
  dosage_matrix,
  margin = 1,
  cutoff = NULL,
  parentnames = c("P1", "P2"),
  plot_breakdown = FALSE,
  log = NULL,
  print.removed = TRUE
)
```

#### **Arguments**

dosage\_matrix An integer matrix with markers in rows and individuals in columns.

margin An integer at which margin the missing value frequency will be calculated. A

value of 1 means rows (markers), 2 means columns (individuals)

cutoff Missing value frequency cut off. At this frequency, rows or columns are re-

moved from the dataset. If NULL user input will be asked after plotting the

missing value frequency histogram.

parentnames A character vector of length 2, specifying the parent names.

plot\_breakdown Logical. Should the percentage of markers removed as breakdown per marker-

type be plotted? Can only be used if margin = 1.

log Character string specifying the log filename to which standard output should be

written. If NULL log is send to stdout.

print.removed Logical. Should removed instances be printed?

#### Value

A matrix similar to dosage\_matrix, with rows or columns removed that had a higher missing value frequency than specified.

# **Examples**

```
data("segregating_data","screened_data")
screened_markers<-screen_for_NA_values(dosage_matrix=segregating_data, margin=1, cutoff=0.1)
screened_indiv<-screen_for_NA_values(dosage_matrix=screened_data, margin=2, cutoff=0.1)</pre>
```

SNSN\_LOD\_deviations Identify deviations in LOD scores between pairs of simplex x nulliplex markers

#### **Description**

SNSN\_LOD\_deviations checks whether the LOD scores obtained in the case of pairs of simplex x nulliple markers are compatible with expectation. This can help identify problematic linkage estimates which can adversely affect marker clustering.

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#### Usage

```
SNSN_LOD_deviations(
  linkage_df,
  ploidy,
  N,
  plot_expected = TRUE,
  alpha = c(0.05, 0.2),
  phase = c("coupling", "repulsion")
)
```

#### **Arguments**

linkage\_df A linkage data.frame as output of linkage.

ploidy Integer. The ploidy level of the species.

N Numeric. The number of F1 individuals in the mapping population.

plot\_expected Logical. Plot the observed and expected relationship between r and LOD.

alpha Numeric. Vector of upper and lower tolerances around expected line.

phase Character string. Specify which phase to examine for deviations (usually this is

"coupling" phase).

#### Value

A vector of deviations in LOD scores outside the range defined by tolerances input alpha

#### **Examples**

```
data("SN_SN_P1")
SNSN_LOD_deviations(SN_SN_P1,ploidy = 4, N = 198)
```

SN\_SN\_P1

A linkage data.frame.

# **Description**

A linkage data. frame.

# Usage

```
SN_SN_P1
```

SN\_SN\_P2

SN\_SS\_P1

SN\_SS\_P2

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```
SN_DN_P1
SN_DN_P2
SN_SN_P2_triploid
```

#### **Format**

- marker\_a. First marker in comparison
- marker\_b. Second marker in comparison
- r. recombination frequency
- · LOD. LOD score
- phase. The phase between markers

An object of class linkage\_df (inherits from data.frame) with 19306 rows and 5 columns.

An object of class linkage\_df (inherits from data.frame) with 53152 rows and 5 columns.

An object of class linkage\_df (inherits from data.frame) with 59494 rows and 5 columns.

An object of class linkage\_df (inherits from data.frame) with 19536 rows and 5 columns.

An object of class linkage\_df (inherits from data.frame) with 19897 rows and 5 columns.

An object of class data. frame with 6655 rows and 5 columns.

test\_prefpairing

Check for and estimate preferential pairing

#### **Description**

Identify closely-mapped repulsion-phase simplex x nulliplex markers and test these for preferential pairing, including estimating a preferential pairing parameter.

```
test_prefpairing(
  dosage_matrix,
  maplist,
  LG_hom_stack,
  target_parent = "P1",
  other_parent = "P2",
  ploidy,
  min_cM = 0.5,
  adj.method = "fdr",
  verbose = TRUE
)
```

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## Arguments

An integer matrix with markers in rows and individuals in columns. dosage\_matrix A list of integrated chromosomal maps, as generated by e.g. MDSMap\_from\_list. maplist In the first column marker names and in the second their position. LG\_hom\_stack A data.frame with markernames ("SxN\_Marker"), linkage group ("LG") and homologue ("homologue"), the output of define\_LG\_structure or bridgeHomologues usually. Character string specifying the parent to be tested for preferential pairing as target\_parent provided in the columnnames of dosage\_matrix, by default "P1". other\_parent The other parent, by default "P2" ploidy The ploidy level of the species, by default 4 (tetraploid) is assumed. The smallest distance to be considered a true distance on the linkage map, by min\_cM default distances less than 0.5 cM are considered essentially zero. adj.method Method to correct p values of Binomial test for multiple testing, by default the FDR correction is used, other options are available, inherited from p. adjust verbose Should messages be sent to stdout? If NULL log is send to stdout.

# **Examples**

```
data("ALL_dosages","integrated.maplist","LGHomDf_P1_1")
P1pp <- test_prefpairing(ALL_dosages,integrated.maplist,LGHomDf_P1_1,ploidy=4)</pre>
```

write.mct

Write MapChart file

#### **Description**

Write a .mct file of a maplist for external plotting with MapChart software (Voorrips ).

```
write.mct(
  maplist,
  mapdir = "mapping_files_MDSMap",
  file_info = paste("; MapChart file created on", Sys.Date()),
  filename = "MapFile",
  precision = 2,
  showMarkerNames = FALSE
)
```

72 write.pwd

## **Arguments**

maplist A list of maps. In the first column marker names and in the second their position. All map data are compiled into a single MapChart file. Directory to which .mct files are written, by default the same directory as for mapdir MDSMap\_from\_list file\_info A character string added to the first lines of the .mct file, by default a datestamp is recorded. filename Character string of filename to write the .mct file to, by default "MapFile" To how many decimal places should marker positions be specified (default = 2)? precision showMarkerNames

Logical, by default FALSE, if TRUE, the marker names will be diplayed in the MapChart output as well.

# **Examples**

```
## Not run:
data("integrated.maplist")
write.mct(integrated.maplist)
## End(Not run)
```

write.pwd

Write a JoinMap compatible .pwd file from linkage data.frame.

# Description

Output of this function allows to use JoinMap to perform the marker ordering step.

## Usage

```
write.pwd(linkage_df, pwd_file, file_info, log = NULL)
```

# **Arguments**

linkage\_df A linkage data.frame. pwd\_file A character string specifying a file open for writing.

file\_info A character string added to the first lines of the .pwd file. log

Character string specifying the log filename to which standard output should be

written. If NULL log is send to stdout.

```
## Not run:
data("all_linkages_list_P1_split")
write.pwd(all_linkages_list_P1_split[["LG3"]][["homologue1"]],
           "LG3_homologue1_P1.pwd",
           "Please feed me to JoinMap")
## End(Not run)
```

write.TSNPM 73

write.TSNPM	Write TetraploidSNPMap input file	

# Description

Output the phased linkage map files into format readable by TetraploidSNPMap (Hackett et al. 2017) to perform QTL analysis.

# Usage

```
write.TSNPM(
  phased.maplist,
  outputdir = "TetraploidSNPMap_QTLfiles",
  filename = "TSNPM",
  ploidy,
  verbose = FALSE
)
```

# Arguments

Phased maps in list format, the output of create_phased_maplist
Directory to which TetraploidSNPMap files are written, by default written to "TetraploidSNPMap_QTLfiles" folder
Character string of filename stem to write the output files to, by default "TSNPM" with linkage groups names appended
The ploidy of the species, currently only 4 is supported by TetraploidSNPMap
Should messages be sent to stdout?

# Value

NULL

```
## Not run:
data("phased.maplist")
write.TSNPM(phased.maplist,ploidy=4)
## End(Not run)
```

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write\_nested\_list

Write out a nested list

# **Description**

Write a nested list into a directory structure

# Usage

```
write_nested_list(
  nested_list,
  directory,
  save_as_object = FALSE,
  object_prefix = directory,
  extension = if (save_as_object) ".Rdata" else ".txt",
  ...
)
```

## Arguments

```
nested_list A nested list.

directory Character string. Directory name to which to write the structure.

save_as_object Logical. Save as R object?

object_prefix Character. Prefix of R object. Only used if save_as_object = TRUE.

extension Character. File extension. Default is ".txt".

Arguments passed to write.table
```

# **Examples**

write\_pwd\_list

Write pwd files from a nested list

## **Description**

A wrapper for write.pwd, which allows to write multiple pwd files with a directory structure according to the nested linkage list.

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# Usage

```
write_pwd_list(
   linkages_list,
   target_parent,
   binned = FALSE,
   dir = getwd(),
   log = NULL
)
```

# Arguments

linkages\_list A nested list with linkage group on the first level and homologue on the sec-

ond.

target\_parent A character string specifying the name of the target parent.

binned Logical. Are the markers binned? This information is used in the pwd header.

dir A character string specifying the directory in which the files are written. De-

faults to working directory.

log Character string specifying the log filename to which standard output should be

written. If NULL log is send to stdout.

```
## Not run:
data("all_linkages_list_P1_split")
write_pwd_list(all_linkages_list_P1_split, target_parent="P1", binned=FALSE)
## End(Not run)
```

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