# Package 'Pedixplorer'

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Title Pedigree Functions

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Description Routines to handle family data with a Pedigree object. The initial purpose was to create correlation structures that describe family relationships such as kinship and identity-by-descent, which can be used to model family data in mixed effects models, such as in the coxme function. Also includes a tool for Pedigree drawing which is focused on producing compact layouts without intervention. Recent additions include utilities to trim the Pedigree object with various criteria, and kinship for the X chromosome.

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BugReports https://github.com/LouisLeNezet/Pedixplorer/issues

URL https://louislenezet.github.io/Pedixplorer/

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Pedix	plorer-package The Pedixplorer package for pedigree data					

# Description

The Pedixplorer package for pedigree data an updated package of the kinship2 package. The kinship2 package was originally written by Terry Therneau and Jason Sinnwell. The Pedixplorer package is a fork of the kinship2 package with additional functionality and bug fixes.

#### **Details**

The package download, NEWS, and README are available on CRAN: Kinship2 for the previous version of the package.

#### **Functions**

Below are listed some of the most widely used functions available in arsenal:

Pedigree(): Contstructor of the Pedigree class, given identifiers, sex, affection status(es), and special relationships

kinship(): Calculates the kinship matrix, the probability having an allele sampled from two individuals be the same via IBD.

plot(): Method to transform a Pedigree object into a graphical plot. Allows extra information to be included in the id under the plot symbol. This method use the plot\_fromdf() function to transform the Pedigree object into a data frame of graphical elements, the same is done for the legend with the ped\_to\_legdf() function. When done, the data frames are plotted with the plot\_fromdf() function.

shrink(): Shrink a Pedigree to a specific bit size, removing non-informative members first.

bit\_size(): Approximate the output from SAS's PROC FREQ procedure when using the /list
option of the TABLE statement.

#### Data

- sampleped(): Pedigree example data sets with two pedigrees
- minnbreast(): Larger cohort of pedigrees from MN breast cancer study

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#### See Also

Useful links:

- https://louislenezet.github.io/Pedixplorer/
- Report bugs at https://github.com/LouisLeNezet/Pedixplorer/issues

# **Examples**

```
library(Pedixplorer)
```

align

Align a Pedigree object

### **Description**

Given a Pedigree, this function creates helper matrices that describe the layout of a plot of the Pedigree.

#### Usage

```
## S4 method for signature 'Pedigree'
align(
  obj,
  packed = TRUE,
  width = 10,
  align = TRUE,
  hints = NULL,
  missid = "NA_character_",
  align_parents = TRUE,
  force = FALSE,
  precision = 4
)
```

align 7

#### **Arguments**

obj	A Pedigree object
packed	Should the Pedigree be compressed. (i.e. allow diagonal lines connecting parents to children in order to have a smaller overall width for the plot.)
width	For a packed output, the minimum width of the plot, in inches.
align	For a packed Pedigree, align children under parents TRUE, to the extent possible given the page width, or align to to the left margin FALSE. This argument can be a two element vector, giving the alignment parameters, or a logical value. If TRUE, the default is c(1.5, 2), or if numeric the routine alignped4() will be called.
hints	A Hints object or a named list containing horder and spouse. If NULL then the Hints stored in <b>obj</b> will be used.
missid	A character vector with the missing values identifiers. All the id, dadid and momid corresponding to those values will be set to NA_character
align_parents	If align_parents = TRUE, go one step further and try to make both parents of each child have the same depth. (This is not always possible). It helps the drawing program by lining up pedigrees that 'join in the middle' via a marriage.
force	If force = TRUE, the function will return the depth minus min(depth) if depth reach a state with no founders is not possible.
precision	The number of significatif numbers to round the solution to.

#### **Details**

This is an internal routine, used almost exclusively by ped\_to\_plotdf().

The subservient functions auto\_hint(), alignped1(), alignped2(), alignped3(), and alignped4() contain the bulk of the computation.

If the **hints** are missing the auto\_hint() routine is called to supply an initial guess.

If multiple families are present in the **obj** Pedigree, this routine is called once for each family, and the results are combined in the list returned.

For more information you can read the associated vignette: vignette("pedigree\_alignment").

#### Value

A list with components

- n: A vector giving the number of subjects on each horizonal level of the plot
- nid: A matrix with one row for each level, giving the numeric id of each subject plotted. (A value of 17 means the 17th subject in the Pedigree).
- pos: A matrix giving the horizontal position of each plot point
- fam: A matrix giving the family id of each plot point. A value of 3 would mean that the two subjects in positions 3 and 4, in the row above, are this subject's parents.
- spouse: A matrix with values
  - 0 = not a spouse

- 1 = subject plotted to the immediate right is a spouse
- 2 = subject plotted to the immediate right is an inbred spouse
- twins: Optional matrix which will only be present if the Pedigree contains twins:
  - 0 = not a twin
  - -1 = sibling to the right is a monozygotic twin
  - -2 = sibling to the right is a dizygotic twin
  - 3 = sibling to the right is a twin of unknown zygosity

#### See Also

```
alignped1(), alignped2(), alignped3(), alignped4(), auto_hint()
```

### **Examples**

```
data(sampleped)
pedi <- Pedigree(sampleped)
align(pedi)</pre>
```

alignped1

Alignment first routine

### **Description**

First alignment routine which create the subtree founded on a single subject as though it were the only tree.

#### Usage

```
alignped1(idx, dadx, momx, level, horder, packed, spouselist)
```

#### **Arguments**

idx	Indexes of the subjects
dadx	Indexes of the fathers
momx	Indexes of the mothers

level Vector of the level of each subject

horder A named numeric vector with one element per subject in the Pedigree. It de-

termines the relative horizontal order of subjects within a sibship, as well as the relative order of processing for the founder couples. (For this latter, the female founders are ordered as though they were sisters). The names of the vector

should be the individual identifiers.

packed Should the Pedigree be compressed. (i.e. allow diagonal lines connecting par-

ents to children in order to have a smaller overall width for the plot.)

spouselist Matrix of spouses with 4 columns:

- 1: husband index
- · 2: wife index
- 3: husband anchor
- 4: wife anchor

#### Details

In this routine the **nid** array consists of the final nid array + 1/2 of the final spouse array. Note that the **spouselist** matrix will only contain spouse pairs that are not yet processed. The logic for anchoring is slightly tricky.

#### 1. Anchoring::

First, if col 4 of the spouselist matrix is 0, we anchor at the first opportunity. Also note that if spouselist[, 3] == spouselist[, 4] it is the husband who is the anchor (just write out the possibilities).

#### 2. Return values initialization::

Create the set of 3 return structures, which will be matrices with 1 + nspouse columns. If there are children then other routines will widen the result.

### 3. Create Ispouse and rspouse::

This two complimentary lists denote the spouses plotted on the left and on the right. For someone with lots of spouses we try to split them evenly. If the number of spouses is odd, then men should have more on the right than on the left, women more on the right. Any hints in the spouselist matrix override. We put the undecided marriages closest to **idx**, then add predetermined ones to the left and right. The majority of marriages will be undetermined singletons, for which **nleft** will be 1 for female (put my husband to the left) and 0 for male. In one bug found by plotting canine data, Ispouse could initially be empty but length(rspouse) > 1. This caused nleft > length(indx). A fix was to not let **indx** to be indexed beyond its length, fix by JPS 5/2013.

# 4. List the children::

For each spouse get the list of children. If there are any we call alignped2() to generate their tree and then mark the connection to their parent. If multiple marriages have children we need to join the trees.

#### 5. Splice the tree::

To finish up we need to splice together the tree made up from all the kids, which only has data from lev + 1 down, with the data here. There are 3 cases:

- 1. No children were found.
- 2. The tree below is wider than the tree here, in which case we add the data from this level onto theirs.
- 3. The tree below is narrower, for instance an only child.

#### Value

A list containing the elements to plot the Pedigree. It contains a set of matrices along with the spouselist matrix. The latter has marriages removed as they are processed.

• n : A vector giving the number of subjects on each horizonal level of the plot

• nid: A matrix with one row for each level, giving the numeric id of each subject plotted. (A value of 17 means the 17th subject in the Pedigree).

- pos: A matrix giving the horizontal position of each plot point
- fam: A matrix giving the family id of each plot point. A value of 3 would mean that the two subjects in positions 3 and 4, in the row above, are this subject's parents.
- spouselist: Spouse matrix with anchors informations

#### See Also

```
align()
```

#### **Examples**

```
data(sampleped)
pedi <- Pedigree(sampleped)
align(pedi)</pre>
```

alignped2

Alignment second routine

#### **Description**

Second of the four co-routines which takes a collection of siblings, grows the tree for each, and appends them side by side into a single tree.

#### Usage

```
alignped2(idx, dadx, momx, level, horder, packed, spouselist)
```

### **Arguments**

idx Indexes of the subjects
dadx Indexes of the fathers
momx Indexes of the mothers

level Vector of the level of each subject

horder A named numeric vector with one element per subject in the Pedigree. It de-

termines the relative horizontal order of subjects within a sibship, as well as the relative order of processing for the founder couples. (For this latter, the female founders are ordered as though they were sisters). The names of the vector

should be the individual identifiers.

packed Should the Pedigree be compressed. (i.e. allow diagonal lines connecting par-

ents to children in order to have a smaller overall width for the plot.)

spouselist Matrix of spouses with 4 columns:

· 1: husband index

- 2: wife index
- 3: husband anchor
- 4: wife anchor

#### **Details**

The input arguments are the same as those to alignped1() with the exception that **idx** will be a vector. This routine does nothing to the spouselist matrix, but needs to pass it down the tree and back since one of the routines called by alignped2() might change the matrix.

The code below has one non-obvious special case. Suppose that two sibs marry. When the first sib is processed by alignped1 then both partners (and any children) will be added to the rval structure below. When the second sib is processed they will come back as a 1 element tree (the marriage will no longer be on the **spouselist**), which should be added onto rval. The rule thus is to not add any 1 element tree whose value (which must be idx[i] is already in the rval structure for this level.

#### Value

A list containing the elements to plot the Pedigree. It contains a set of matrices along with the spouselist matrix. The latter has marriages removed as they are processed.

- n : A vector giving the number of subjects on each horizonal level of the plot
- nid: A matrix with one row for each level, giving the numeric id of each subject plotted. (A value of 17 means the 17th subject in the Pedigree).
- pos: A matrix giving the horizontal position of each plot point
- fam: A matrix giving the family id of each plot point. A value of 3 would mean that the two subjects in positions 3 and 4, in the row above, are this subject's parents.
- spouselist: Spouse matrix with anchors informations

### See Also

```
align()
```

### **Examples**

```
data(sampleped)
pedi <- Pedigree(sampleped)
align(pedi)</pre>
```

alignped3	Alignment third routine	

#### **Description**

Third of the four co-routines to merges two pedigree trees which are side by side into a single object.

### Usage

```
alignped3(alt1, alt2, packed, space = 1)
```

### **Arguments**

alt1	Alignment of the first tree
alt2	Alignment of the second tree
packed	Should the Pedigree be compressed. (i.e. allow diagonal lines connecting parents to children in order to have a smaller overall width for the plot.)
space	Space between two subjects

#### **Details**

The primary special case is when the rightmost person in the left tree is the same as the leftmost person in the right tree; we need not plot two copies of the same person side by side. (When initializing the output structures do not worry about this, there is no harm if they are a column bigger than finally needed.) Beyond that the work is simple book keeping.

#### 1. Slide::

For the unpacked case, which is the traditional way to draw a Pedigree when we can assume the paper is infinitely wide, all parents are centered over their children. In this case we think if the two trees to be merged as solid blocks. On input they both have a left margin of 0. Compute how far over we have to slide the right tree.

#### 2. Merge::

Now merge the two trees. Start at the top level and work down.

#### Value

A list containing the elements to plot the Pedigree. It contains a set of matrices along with the spouselist matrix. The latter has marriages removed as they are processed.

- n : A vector giving the number of subjects on each horizonal level of the plot
- nid: A matrix with one row for each level, giving the numeric id of each subject plotted. (A value of 17 means the 17th subject in the Pedigree).
- pos: A matrix giving the horizontal position of each plot point
- fam: A matrix giving the family id of each plot point. A value of 3 would mean that the two subjects in positions 3 and 4, in the row above, are this subject's parents.
- spouselist: Spouse matrix with anchors informations

### See Also

```
align()
```

### **Examples**

```
data(sampleped)
pedi <- Pedigree(sampleped)
align(pedi)</pre>
```

alignped4

Alignment fourth routine

### **Description**

Last routines which attempts to line up children under parents and put spouses and siblings "close" to each other, to the extent possible within the constraints of page width.

### Usage

```
alignped4(rval, spouse, level, width, align, precision = 4)
```

#### **Arguments**

rval	A list with components n, nid, pos, and fam.
spouse	A boolean matrix with one row per level representing if the subject is a spouse or not.
level	Vector of the level of each subject
width	For a packed output, the minimum width of the plot, in inches.
align	For a packed Pedigree, align children under parents TRUE, to the extent possible given the page width, or align to to the left margin FALSE. This argument can be a two element vector, giving the alignment parameters, or a logical value. If TRUE, the default is c(1.5, 2), or if numeric the routine alignped4() will be called.
precision	The number of significatif numbers to round the solution to.

#### **Details**

The alignped4() routine is the final step of alignment. The current code does necessary setup and then calls the quadprog::solve.QP() function.

There are two important parameters for the function:

1. The maximum width specified. The smallest possible width is the maximum number of subjects on a line. If the user suggestion is too low it is increased to that amount plus one (to give just a little wiggle room).

2. The align vector of 2 alignment parameters a and b. For each set of siblings x with parents at p\_1 and p\_2 the alignment penalty is:

$$(1/k^a)\sum_{i=1}^k (x_i - (p_1 + p_2)/2)^2$$

where k is the number of siblings in the set.

Using the fact that when a = 1:

$$\sum (x_i - c)^2 = \sum (x_i - \mu)^2 + k(c - \mu)^2$$

then moving a sibship with k sibs one unit to the left or right of optimal will incur the same cost as moving one with only 1 or two sibs out of place.

If a = 0 then large sibships are harder to move than small ones. With the default value a = 1.5, they are slightly easier to move than small ones. The rationale for the default is as long as the parents are somewhere between the first and last siblings the result looks fairly good, so we are more flexible with the spacing of a large family. By tethering all the sibs to a single spot they tend to be kept close to each other.

The alignment penalty for spouses is  $b(x_1 - x_2)^2$ , which tends to keep them together. The size of b controls the relative importance of sib-parent and spouse-spouse closeness.

- 1. We start by adding in these penalties. The total number of parameters in the alignment problem (what we hand to quadprog) is the set of sum(n) positions. A work array myid keeps track of the parameter number for each position so that it is easy to find. There is one extra penalty added at the end. Because the penalty amount would be the same if all the final positions were shifted by a constant, the penalty matrix will not be positive definite; solve.QP() does not like this. We add a tiny amount of leftward pull to the widest line.
- 2. If there are k subjects on a line there will be k+1 constraints for that line. The first point must be  $\geq 0$ , each subsequent one must be at least 1 unit to the right, and the final point must be  $\leq$  the max width.

#### Value

The updated position matrix

#### See Also

align()

# Examples

```
data(sampleped)
pedi <- Pedigree(sampleped)
align(pedi)</pre>
```

ancestors 15

ancestors

Ancestors indexes of a subject

# Description

Given the index of one or multiple individual(s), this function iterate through the mom and dad indexes to list out all the ancestors of the said individual(s). This function is use in the align() function to identify which spouse pairs has a common ancestor and therefore if they need to be connected with a double line (i.e. inbred).

### Usage

```
ancestors(idx, momx, dadx)
```

# Arguments

idx Indexes of the subjects
momx Indexes of the mothers
dadx Indexes of the fathers

#### Value

A vector of ancestor indexes

### See Also

```
align()
```

# **Examples**

```
ancestors(c(1), c(3, 4, 5, 6), c(7, 8, 9, 10)) ancestors(c(1, 2), c(3, 4, 5, 6), c(7, 8, 9, 10))
```

anchor\_to\_factor

Anchor variable to ordered factor

# Description

Anchor variable to ordered factor

### Usage

```
anchor_to_factor(anchor)
```

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### **Arguments**

anchor

A character, factor or numeric vector corresponding to the anchor of the individuals. The following values are recognized:

```
• character() or factor(): "0", "1", "2", "left", "right", "either"
```

```
• numeric(): 1 = "left", 2 = "right", 0 = "either"
```

### Value

An ordered factor vector containing the transformed variable "either" < "left" < "right"

# **Examples**

```
Pedixplorer:::anchor_to_factor(c(1, 2, 0, "left", "right", "either"))
```

auto\_hint

Initial hint for a Pedigree alignment

### **Description**

Compute an initial guess for the alignment of a Pedigree

# Usage

```
## S4 method for signature 'Pedigree'
auto_hint(obj, hints = NULL, packed = TRUE, align = FALSE, reset = FALSE)
```

### Arguments

obj	A Pedigree object
hints	A Hints object or a named list containing horder and spouse. If NULL then the Hints stored in <b>obj</b> will be used.
packed	Should the Pedigree be compressed. (i.e. allow diagonal lines connecting parents to children in order to have a smaller overall width for the plot.)
align	For a packed Pedigree, align children under parents TRUE, to the extent possible given the page width, or align to to the left margin FALSE. This argument can be a two element vector, giving the alignment parameters, or a logical value. If TRUE, the default is c(1.5, 2), or if numeric the routine alignped4() will be called.
reset	If TRUE, then even if the Ped object has Hints, reset them to the initial values.

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#### **Details**

A Pedigree structure can contain a Hints object which helps to reorder the Pedigree (e.g. left-to-right order of children within family) so as to plot with minimal distortion. This routine is used to create an initial version of the hints. They can then be modified if desired.

This routine would not normally be called by a user. It moves children within families, so that marriages are on the "edge" of a set children, closest to the spouse. For pedigrees that have only a single connection between two families this simple-minded approach works surprisingly well. For more complex structures hand-tuning of the hints may be required.

When auto\_hint() is called with a a vector of numbers as the **hints** argument, the values for the founder females are used to order the founder families left to right across the plot. The values within a sibship are used as the preliminary order of siblings within a family; this may be changed to move one of them to the edge so as to match up with a spouse. The actual values in the vector are not important, only their order.

#### Value

The initial Hints object.

#### See Also

```
align(), best_hint()
Hints
```

#### **Examples**

```
data(sampleped)
pedi <- Pedigree(sampleped[sampleped$famid == 1, ])
Pedixplorer:::auto_hint(pedi)</pre>
```

best\_hint

Best hint for a Pedigree alignment

# Description

When computer time is cheap, use this routine to get a *best* Pedigree alignment. This routine will try all possible founder orders, and return the one with the least **stress**.

# Usage

```
## S4 method for signature 'Pedigree'
best_hint(obj, wt = c(1000, 10, 1), tolerance = 0)
```

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### **Arguments**

obj A Pedigree object
wt A vector of three weights for the three error measures. Default is c(1000, 10,

- 1. The number of duplicate individuals in the plot
- 2. The sum of the absolute values of the differences in the positions of duplicate individuals
- 3. The sum of the absolute values of the differences between the center of the children and the parents.

tolerance The maximum stress level to accept. Default is 0

#### **Details**

The auto\_hint() routine will rearrange sibling order, but not founder order. This calls auto\_hint() with every possible founder order, and finds that plot with the least "stress". The stress is computed as a weighted sum of three error measures:

- nbArcs The number of duplicate individuals in the plot
- lgArcs The sum of the absolute values of the differences in the positions of duplicate individuals
- lgParentsChilds The sum of the absolute values of the differences between the center of the children and the parents

```
stress = wt[1]*nbArcs + wt[2]*lgArcs + wt[3]*lgParentsChilds
```

If during the search, a plot is found with a stress level less than tolerance, the search is terminated.

#### Value

The best Hints object out of all the permutations

#### See Also

```
auto_hint(), align()
```

#### **Examples**

```
data(sampleped)
pedi <- Pedigree(sampleped[sampleped$famid == 1,])
best_hint(pedi)</pre>
```

bit\_size 19

bit_size	Bit size of a Pedigree	

# Description

Utility function used in the shrink() function to calculate the bit size of a Pedigree.

### Usage

```
## S4 method for signature 'character_OR_integer'
bit_size(obj, momid, missid = NA_character_)

## S4 method for signature 'Pedigree'
bit_size(obj)

## S4 method for signature 'Ped'
bit_size(obj)
```

### Arguments

obj	A Ped or Pedigree object or a vector of fathers identifiers
momid	A vector containing for each subject, the identifiers of the biologicals mothers.
missid	A character vector with the missing values identifiers. All the id, dadid and
	momid corresponding to those values will be set to NA_character

# **Details**

The bit size of a Pedigree is defined as:

```
2 \times NbNonFounders - NbFounders
```

Where NbNonFounders is the number of non founders in the Pedigree (i.e. individuals with identified parents) and NbFounders is the number of founders in the Pedigree (i.e. individuals without identified parents).

### Value

A list with the following components:

- bit\_size The bit size of the Pedigree
- nFounder The number of founders in the Pedigree
- nNonFounder The number of non founders in the Pedigree

### See Also

```
shrink()
```

20 check\_columns

# **Examples**

```
data(sampleped)
pedi <- Pedigree(sampleped)
bit_size(pedi)</pre>
```

char\_to\_date

Convert a character to a date

# Description

Convert a character to a date

### Usage

```
char_to_date(date, date_pattern = "%Y-%m-%d")
```

# Arguments

date A character vector of dates

date\_pattern The pattern of the date

#### Value

A date vector

# **Examples**

```
Pedixplorer:::char_to_date("2020-01-01", "%Y-%m-%d")
Pedixplorer:::char_to_date("01/01/20", "%d/%m/%y")
```

check\_columns

Check columns presence in a dataframe

# Description

Check for presence / absence of columns names depending on their need

check\_columns 21

#### Usage

```
check_columns(
   df,
   cols_needed = NULL,
   cols_used = NULL,
   cols_to_use = NULL,
   others_cols = FALSE,
   cols_used_init = FALSE,
   cols_to_use_init = FALSE,
   cols_used_del = FALSE,
   verbose = FALSE
)
```

### **Arguments**

df	The dataframe to use	
cols_needed	A vector of columns needed	
cols_used	A vector of columns that are used by the script and that will be overwritten.	
cols_to_use	A vector of optional columns that are authorized.	
others_cols	Boolean defining if non defined columns should be allowed.	
cols_used_init	Boolean defining if the columns that will be used should be initialised to NA.	
cols_to_use_init		
	Boolean defining if the optional columns should be initialised to NA.	
cols_used_del	Boolean defining if the columns that will be used should be deleted.	
verbose	Should message be prompted to the user	

### **Details**

3 types of columns are here checked:

- cols\_needed: those columns need to be present if any is missing an error will be prompted and the script will stop
- cols\_used: those columns will be used in the script and will be overwritten to NA.
- cols\_to\_use: those columns are optional and will be recognise if present. The last two types of columns can be initialised to NA if needed.

#### Value

Dataframe with only the column allowed and all the column correctly initialised.

# **Examples**

```
data.frame
df <- data.frame(
    ColN1 = c(1, 2), ColN2 = 4,
    ColU1 = 'B', ColU2 = '1',
    ColTU1 = 'A', ColTU2 = 3,</pre>
```

check\_col\_config

```
ColNR1 = 4, ColNR2 = 5
)
tryCatch(
   Pedixplorer:::check_columns(
          df,
          c('ColN1', 'ColN2'), c('ColU1', 'ColU2'),
          c('ColTU1', 'ColTU2')
), error = function(e) print(e))
```

check\_col\_config

Check column configuration

### **Description**

This function checks the validity of the column configuration provided to the data\_col\_sel\_server function.

### Usage

```
check_col_config(col_config)
```

### **Arguments**

col\_config

A list of column definitions. It must contain a list for each column, with the following keys: 'alternate' and 'mandatory'.

#### **Details**

The list names must correspond to the column names of the dataframe to be selected. Each list must contain two keys: 'alternate' and 'mandatory'. The 'alternate' key must contain a character vector of column names that can be selected as an alternative to the main column. The 'mandatory' key must contain a logical value (TRUE/FALSE) to indicate whether the column is required to be selected.

### Value

TRUE if the configuration is valid.

#### **Examples**

```
Pedixplorer:::check_col_config(list(
   ColA = list(alternate = c("A"), mandatory = TRUE),
   ColB = list(alternate = c("B"), mandatory = FALSE)
))
```

check\_num\_na 23

check\_num\_na

Is numeric or NA

### **Description**

Check if a variable given is numeric or NA

# Usage

```
check_num_na(var, na_as_num = TRUE)
```

### **Arguments**

var Vector of value to test

na\_as\_num Boolean defining if the NA string should be considered as numerical values

#### **Details**

Check if the values in var are numeric or if they are NA in the case that na\_as\_num is set to TRUE.

#### Value

A vector of boolean of the same size as var

check\_slot\_fd

Check if the fields are present in an object slot

### **Description**

Check if the fields are present in an object slot

#### Usage

```
check_slot_fd(obj, slot = NULL, fields = character())
```

### Arguments

obj An object. slot A slot of object.

fields A character vector with the fields to check.

#### Value

A character vector with the errors if any.

24 circfun

	-	
check	val	ues

Check values in a slot

# Description

Check if the all the values in a slot are in a vector of values.

# Usage

```
check_values(val, ref, name = NULL, present = TRUE)
```

### **Arguments**

val A vector of values to check.
ref A vector of reference values.

name A character vector with the name of the values to check.

present A logical value indicating if the values should be present or not

#### Value

A character vector with the errors if any.

circfun

Circular element

### **Description**

Create a list of x and y coordinates for a circle with a given number of slices.

# Usage

```
circfun(nslice, n = 50, start = 0)
```

### **Arguments**

nslice Number of slices in the circle

n Total number of points in the circle

#### Value

A list of x and y coordinates per slice.

# Examples

```
Pedixplorer:::circfun(1)
Pedixplorer:::circfun(1, 10)
Pedixplorer:::circfun(4, 50)
```

color\_picker\_ui 25

picker	

Shiny modules to select colours

### **Description**

This function allows to select different colours for an array of variables.

### Usage

```
color_picker_ui(id)
color_picker_server(id, colors = NULL)
color_picker_demo()
```

# Arguments

id A string to identify the module.

colors A list of variables and their default colours.

#### Value

A reactive list with the selected colours.

### **Examples**

```
if (interactive()) {
    color_picker_demo()
}
```

create\_text\_column

Create a text column

### Description

Aggregate multiple columns into a single text column separated by a newline character.

#### Usage

```
create_text_column(df, title = NULL, cols = NULL, na_strings = c("", "NA"))
```

### **Arguments**

df A dataframe

title The title of the text column

cols A vector of columns to concatenate

na\_strings A vector of strings that should be considered as NA

26 data\_col\_sel\_ui

#### Value

The concatenated text column

#### **Examples**

```
df <- data.frame(a = seq_len(3), b = c("4", "NA", 6), c = c("", "A", 2))
Pedixplorer:::create_text_column(df, "a", c("b", "c"))
```

data\_col\_sel\_ui

Shiny modules to select columns from a dataframe

### **Description**

This function allows to select columns from a dataframe and rename them to a set of names present in a configuration list. This generate a Shiny module that can be used in a Shiny app. The function is composed of two parts: the UI and the server. The UI is called with the function data\_col\_sel\_ui() and the server with the function data\_col\_sel\_server().

### Usage

```
data_col_sel_ui(id, ui_col_nb = 1)

data_col_sel_server(
   id,
   df,
   col_config,
   title,
   na_omit = TRUE,
   others_cols = TRUE,
   ui_col_nb = 1,
   by_row = FALSE
)

data_col_sel_demo(ui_col_nb = 2, by_row = FALSE)
```

### **Arguments**

id A string to ic	dentify the module.
-------------------	---------------------

df A reactive dataframe.

col\_config A named list of column definitions. It must contain a list for each column,

with the following keys: 'alternate' and 'mandatory'. The 'alternate' key must contain a character vector of column names that can be selected as an alternative to the main column. The 'mandatory' key must contain a logical value (TRUE/FALSE) to indicate whether the column is required to be selected.

title A string to display in the selectInput.

na\_omit A boolean to allow or not the selection of NA.

others\_cols A boolean to authorize other columns to be present in the output datatable.

data\_download\_ui 27

### Value

A reactive dataframe with the selected columns renamed to the names present in the configuration list.

### **Examples**

```
if (interactive()) {
    data_col_sel_demo()
}
```

data\_download\_ui

Shiny modules to download a dataframe

### **Description**

This function allows to download a dataframe as a csv file. This generate a Shiny module that can be used in a Shiny app. The function is composed of two parts: the UI and the server. The UI is called with the function data\_download\_ui() and the server with the function data\_download\_server().

### Usage

```
data_download_ui(id)

data_download_server(
   id,
   df,
   filename,
   label = NULL,
   helper = TRUE,
   title = "Data download"
)

data_download_demo()
```

### **Arguments**

id A string to identify the module.

df A reactive dataframe.
filename A string to name the file.

label A string to display in the download button.
helper A boolean to display a helper message.

#### Value

A shiny module to export a dataframe.

28 data\_import\_ui

### **Examples**

```
if (interactive()) {
    data_download_demo()
}
```

data\_import\_ui

Shiny modules to import data files

### **Description**

This module allow to import multiple type of data. The file type currently supported are csv, txt, xls, xslx, rda and tab. The server dynamically create a selection input if multiple dataframe are present in the file selected. This module is composed of two parts: the UI and the server. The UI is called with the function data\_import\_ui() and the server with the function data\_import\_server(). Different options are available to the user to import the data.

# Usage

```
data_import_ui(id)

data_import_server(
   id,
   label = "Select data file",
   dftest = datasets::mtcars,
   max_request_size = 30
)

data_import_demo(options = list())
```

#### **Arguments**

 $\begin{array}{ll} \text{id} & A \text{ string.} \\ \\ \text{label} & A \text{ string use to prompt the user} \\ \\ \text{dftest} & A \text{ dataframe to test the function} \\ \\ \text{max\_request\_size} \end{array}$ 

A number to define the maximum size of the file that can be uploaded.

### Value

A reactive dataframe selected by the user.

#### **Examples**

```
if (interactive()) {
    data_import_demo()
}
```

descendants 29

descendants	Descendants of individuals
-------------	----------------------------

### Description

Find all the descendants of a particular list of individuals given a Pedigree object.

# Usage

```
## S4 method for signature 'character_OR_integer, character_OR_integer'
descendants(idlist, obj, dadid, momid)

## S4 method for signature 'character_OR_integer, Pedigree'
descendants(idlist, obj)

## S4 method for signature 'character_OR_integer, Ped'
descendants(idlist, obj)
```

### **Arguments**

idlist	List of individuals identifiers to be considered
obj	A Ped or Pedigree object or a vector of the individuals identifiers.
dadid	A vector containing for each subject, the identifiers of the biologicals fathers.
momid	A vector containing for each subject, the identifiers of the biologicals mothers.

#### Value

Vector of all descendants of the individuals in idlist. The list is not ordered.

# **Examples**

```
data("sampleped")
pedi <- Pedigree(sampleped)
descendants(c("1_101", "2_208"), pedi)</pre>
```

distribute\_by

Distribute elements by group

# Description

This function distributes elements by group for a given number of elements. The distribution can be done by row or by column.

#### Usage

```
distribute_by(nb_group, nb_elem, by_row = FALSE)
```

30 draw\_arc

### **Arguments**

nb\_group The number of group.

nb\_elem The number of elements to distribute.

by\_row A boolean to distribute by row or by column.

#### Value

A vector of group indices.

### **Examples**

```
Pedixplorer:::distribute_by(3, 10)
Pedixplorer:::distribute_by(3, 10, by_row = TRUE)
```

draw\_arc

Draw arcs

### **Description**

Draw arcs

### Usage

```
draw_arc(
    x0,
    y0,
    x1,
    y1,
    p = NULL,
    ggplot_gen = FALSE,
    lwd = par("lwd"),
    lty = 2,
    col = "black"
)
```

# Arguments

```
x0
                  x coordinate of the first point
y0
                  y coordinate of the first point
                  x coordinate of the second point
х1
                  y coordinate of the second point
у1
                  ggplot object
р
                  If TRUE add the segments to the ggplot object
ggplot_gen
                  Line width
lwd
                  Line type
1ty
                  Line color
col
```

draw\_arrow 31

# Value

Plot the arcs to the current device or add it to a ggplot object

# Description

Draw arrows

# Usage

```
draw_arrow(
    x0,
    y0,
    x1,
    y1,
    p = NULL,
    ggplot_gen = FALSE,
    lwd = par("lwd"),
    lty = 1,
    col = "black"
)
```

# Arguments

```
x0
                  x coordinate of the first point
y0
                  y coordinate of the first point
                  x coordinate of the second point
х1
                  y coordinate of the second point
у1
                  ggplot object
                  If TRUE add the segments to the ggplot object
ggplot_gen
lwd
                  Line width
                  Line type
lty
                  Line color
col
```

# Value

Plot the arrows to the current device or add it to a ggplot object

32 draw\_polygon

draw\_point

Draw arrows

# Description

Draw arrows

# Usage

```
draw_point(
    x,
    y,
    p = NULL,
    ggplot_gen = FALSE,
    cex = par("lwd"),
    pch = 1,
    col = "black"
)
```

# Arguments

```
p ggplot objectggplot_gen If TRUE add the segments to the ggplot objectcol Line color
```

# Value

Plot the arrows to the current device or add it to a ggplot object

draw\_polygon

Draw a polygon

# Description

Draw a polygon

# Usage

```
draw_polygon(
    x,
    y,
    p = NULL,
    ggplot_gen = FALSE,
    fill = "grey",
    border = "black",
```

draw\_segment 33

```
density = NULL,
angle = 45,
lwd = par("lwd"),
tips = NULL
)
```

# Arguments

```
x coordinates
                  y coordinates
у
                  ggplot object
р
ggplot_gen
                  If TRUE add the segments to the ggplot object
fill
                  Fill color
border
                  Border color
density
                  Density of shading
angle
                  Angle of shading
lwd
                  Line width
tips
                  Text to be displayed when hovering over the polygon
```

### Value

Plot the polygon to the current device or add it to a ggplot object

draw\_segment

Draw segments

### **Description**

Draw segments

### Usage

```
draw_segment(
    x0,
    y0,
    x1,
    y1,
    p = NULL,
    ggplot_gen = FALSE,
    col = par("fg"),
    lwd = par("lwd"),
    lty = par("lty")
)
```

34 draw\_text

# Arguments

x0	x coordinate of the first point
y0	y coordinate of the first point
x1	x coordinate of the second point
y1	y coordinate of the second point
р	ggplot object
ggplot_gen	If TRUE add the segments to the ggplot object
col	Line color
lwd	Line width
lty	Line type

### Value

Plot the segments to the current device or add it to a ggplot object

# Description

Draw texts

# Usage

```
draw_text(
    x,
    y,
    label,
    p = NULL,
    ggplot_gen = FALSE,
    cex = 1,
    col = NULL,
    adjx = 0.5,
    adjy = 0.5,
    tips = NULL
)
```

# Arguments

```
    x x coordinates
    y y coordinates
    label Text to be displayed
    p ggplot object
```

duporder 35

ggplot_gen	If TRUE add the segments to the ggplot object
cex	Character expansion of the text
col	Text color
adjx	x adjustment
adjy	y adjustment
tips	Text to be displayed when hovering over the text

# Value

Plot the text to the current device or add it to a ggplot object

C	luporder	Find the duplicate pairs of a subject

# Description

Find the duplicate pairs of a subject

# Usage

```
duporder(idlist, plist, lev, obj)
```

# Arguments

idlist	List of individuals identifiers to be considered
plist	The alignment structure representing the Pedigree layout. See align() for details.
lev	The generation level of the subject
obj	A Pedigree object

# **Details**

This routine is used by auto\_hint(). It finds the duplicate pairs of a subject and returns them in the order they should be plotted.

#### Value

A matrix of duplicate pairs

# See Also

```
auto_hint()
```

exclude\_stray\_marryin Exclude stray marry-ins

# Description

Exclude any founders who are not parents.

### Usage

```
exclude_stray_marryin(id, dadid, momid)
```

# Arguments

id	A character vector with the identifiers of each individuals
dadid	A vector containing for each subject, the identifiers of the biologicals fathers.
momid	A vector containing for each subject, the identifiers of the biologicals mothers.

### Value

Returns a data frame of subject identifiers and their parents. The data frame is trimmed of any founders who are not parents.

#### See Also

```
shrink()
```

```
exclude_unavail_founders
```

Exclude unavailable founders

# Description

Exclude any unavailable founders.

# Usage

```
exclude_unavail_founders(id, dadid, momid, avail, missid = NA_character_)
```

# Arguments

id	A character vector with the identifiers of each individuals
dadid	A vector containing for each subject, the identifiers of the biologicals fathers.
momid	A vector containing for each subject, the identifiers of the biologicals mothers.
avail	A logical vector with the availability status of the individuals (i.e. FALSE = not available, TRUE = available, NA = unknown).
missid	A character vector with the missing values identifiers. All the id, dadid and momid corresponding to those values will be set to NA_character

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## Value

Returns a list with the following components:

- n\_trimmed Number of trimmed individuals
- id\_trimmed Vector of IDs of trimmed individuals
- id Vector of subject identifiers
- dadid Vector of father identifiers
- momid Vector of mother identifiers

#### See Also

```
shrink()
```

family\_check

Check family

# Description

Error check for a family classification

# Usage

```
## S4 method for signature 'character_OR_integer'
family_check(obj, dadid, momid, famid, newfam)

## S4 method for signature 'Pedigree'
family_check(obj)

## S4 method for signature 'Ped'
family_check(obj)
```

# Arguments

obj	A character vector with the id of the individuals or a data. frame with all the informations in corresponding columns.
dadid	A vector containing for each subject, the identifiers of the biologicals fathers.
momid	A vector containing for each subject, the identifiers of the biologicals mothers.
famid	A character vector with the family identifiers of the individuals. If provide, will be aggregated to the individuals identifiers separated by an underscore.
newfam	The result of a call to make_famid(). If this has already been computed by the user, adding it as an argument shortens the running time somewhat.

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#### **Details**

Given a family id vector, also compute the familial grouping from first principles using the parenting data, and compare the results.

The make\_famid() function is used to create a de novo family id from the parentage data, and this is compared to the family id given in the data.

If there are any joins, then an attribute 'join' is attached. It will be a matrix with family as row labels, new-family-id as the columns, and the number of subjects as entries.

#### Value

a data frame with one row for each unique family id in the famid argument or the one detected in the Pedigree object. Components of the output are:

- famid: The family id, as entered into the data set
- n: Number of subjects in the family
- unrelated: Number of them that appear to be unrelated to anyone else in the entire Pedigree. This is usually marry-ins with no children (in the Pedigree), and if so are not a problem.
- split: Number of unique 'new' family ids.
  - 0 =no one in this 'family' is related to anyone else (not good)
  - -1 = everythings is fine
  - 2 and + = the family appears to be a set of disjoint trees. Are you missing some of the people?
- join: Number of other families that had a unique family, but are actually joined to this one. 0 is the hope.

### See Also

```
make_famid()
```

```
# use 2 samplepeds
data(sampleped)
pedAll <- Pedigree(sampleped)

## check them giving separate ped ids
fcheck.sep <- family_check(pedAll)
fcheck.sep

## check assigning them same ped id
fcheck.combined <- with(sampleped, family_check(id, dadid, momid,
rep(1, nrow(sampleped))))
fcheck.combined</pre>
```

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family\_infos\_table

Affection and availability information table

### **Description**

This function creates a table with the affection and availability information for all individuals in a pedigree object.

### Usage

```
family_infos_table(pedi, col_val = NA)
```

### **Arguments**

pedi A pedigree object.

col\_val The column name in the fill slot of the pedigree object to use for the table.

#### Value

A cross table dataframe with the affection and availability information.

#### **Examples**

```
data(sampleped)
pedi <- Pedigree(sampleped)
pedi <- generate_colors(pedi, "num_child_tot", threshold = 2)
Pedixplorer:::family_infos_table(pedi, "num_child_tot")
Pedixplorer:::family_infos_table(pedi, "affection")</pre>
```

family\_sel\_ui

Shiny module to select a family in a pedigree

## **Description**

This module allows to select a family in a pedigree object. The function is composed of two parts: the UI and the server. The UI is called with the function family\_sel\_ui() and the server with the function family\_sel\_server().

40 fertility\_to\_factor

#### Usage

```
family_sel_ui(id)

family_sel_server(
   id,
   pedi,
   fam_var = NULL,
   fam_sel = NULL,
   title = "Family selection"
)

family_sel_demo(fam_var = NULL, fam_sel = NULL, title = "Family selection")
```

# Arguments

id A string to identify the module.
 pedi A reactive pedigree object.
 fam\_var The default family variable to use as family indicator.
 fam\_sel The default family to select.

The title of the module.

## Value

title

A reactive list with the subselected pedigree object and the selected family id.

## **Examples**

```
if (interactive()) {
    family_sel_demo()
}
```

fertility\_to\_factor Fertility variable to factor

## **Description**

Transform a fertility variable to a factor variable By default, all other values are transformed to NA and considered as fertile.

#### Usage

```
fertility_to_factor(fertility)
```

findsibs 41

## **Arguments**

fertility

A character, factor or numeric vector corresponding to the fertility status of the individuals. This will be transformed to a factor with the following levels: infertile\_choice\_na, infertile, fertile

The following values are recognized:

- "inferile\_choice\_na" : "infertile\_choice", "infertile\_na"
- "infertile": "infertile", "steril", FALSE, 0
- "fertile": "fertile", TRUE, 1, NA

#### Value

a factor vector containing the transformed variable "infertile\_choice\_na", "infertile", "fertile"

# **Examples**

```
fertility_to_factor(c(
   1, "fertile", TRUE, NA,
   "infertile", "steril", FALSE, 0,
   "infertile_na", "infertile_choice_na", "infertile_choice"
))
```

findsibs

Find the siblings of a subject

## **Description**

Find the siblings of a subject

#### **Usage**

```
findsibs(idpos, plist, lev)
```

## **Arguments**

idpos The position of the subject

plist The alignment structure representing the Pedigree layout. See align() for de-

tails.

lev The generation level of the subject

#### **Details**

This routine is used by auto\_hint(). It finds the siblings of a subject.

#### Value

The positions of the siblings

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## See Also

```
auto_hint()
```

findspouse

Find the spouse of a subject

# Description

Find the spouse of a subject

# Usage

```
findspouse(idpos, plist, lev, obj)
```

# Arguments

idpos	The position of the subject
plist	The alignment structure representing the Pedigree layout. See align() for details.
lev	The generation level of the subject
obj	A Pedigree object

## **Details**

This routine is used by auto\_hint(). It finds the spouse of a subject.

# Value

The position of the spouse

## See Also

```
auto_hint()
```

find\_avail\_affected 43

find_avail_affected Find single affected and available individual from a
--

## **Description**

Finds one subject from among available non-parents with indicated affection status.

# Usage

```
## S4 method for signature 'Ped'
find_avail_affected(obj, avail = NULL, affected = NULL, affstatus = NA)
## S4 method for signature 'Pedigree'
find_avail_affected(obj, avail = NULL, affected = NULL, affstatus = NA)
```

## **Arguments**

obj	A Ped or Pedigree object.
avail	A logical vector with the availability status of the individuals (i.e. $FALSE = not$ available, $TRUE = available$ , $NA = unknown$ ).
affected	A logical vector with the affection status of the individuals (i.e. $FALSE = unaffected$ , $TRUE = affected$ , $NA = unknown$ ).
affstatus	Affection status to search for.

#### **Details**

When used within <a href="https://shrink">shrink</a>(), this function is called with the first affected indicator, if the affected item in the Pedigree is a matrix of multiple affected indicators.

If avail or affected is null, then the function will use the corresponding Ped accessor.

#### Value

A list is returned with the following components

- ped The new Ped object
- newAvail Vector of availability status of trimmed individuals
- idTrimmed Vector of IDs of trimmed individuals
- isTrimmed logical value indicating whether Ped object has been trimmed
- bit\_size Bit size of the trimmed Ped

#### See Also

```
shrink()
```

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### **Examples**

```
data(sampleped)
pedi <- Pedigree(sampleped)
find_avail_affected(pedi, affstatus = 1)</pre>
```

find\_avail\_noninform Find uninformative but available subject

## **Description**

Finds subjects from among available non-parents with all affection equal to 0.

### Usage

```
## S4 method for signature 'Ped'
find_avail_noninform(obj, avail = NULL, affected = NULL)
## S4 method for signature 'Pedigree'
find_avail_noninform(obj, avail = NULL, affected = NULL)
```

#### **Arguments**

obj A Ped or Pedigree object.

avail A logical vector with the availability status of the individuals (i.e. FALSE = not

available, TRUE = available, NA = unknown).

affected A logical vector with the affection status of the individuals (i.e. FALSE = unaf-

fected, TRUE = affected, NA = unknown).

#### **Details**

Identify subjects to remove from a Pedigree who are available but non-informative (unaffected). This is the second step to remove subjects in shrink() if the Pedigree does not meet the desired bit size.

If avail or affected is null, then the function will use the corresponding Ped accessor.

#### Value

Vector of subject ids who can be removed by having lowest informativeness.

## See Also

```
shrink()
```

```
data(sampleped)
pedi <- Pedigree(sampleped)
find_avail_noninform(pedi)</pre>
```

find\_ray\_intersections 45

```
find_ray_intersections
```

Find intersections of a ray with a segment

## **Description**

Given the coordinates of a segment and the angle of a ray from the origin, this function computes the intersection point of the ray with the segment.

#### Usage

```
find_ray_intersections(x0, y0, x1, y1, theta)
```

### **Arguments**

x0	x-coordinate of the segment's starting point
y0	y-coordinate of the segment's starting point
x1	x-coordinate of the segment's ending point
y1	y-coordinate of the segment's ending point
theta	Angle of the ray from the origin (in radians)

#### Value

A vector of the x and y coordinates of the intersection point, or NA if no intersection occurs.

# **Examples**

```
Pedixplorer:::find_ray_intersections(0, 0, 1, 1, pi / 4)
```

find\_unavailable

Find unavailable subjects in a Pedigree

# Description

Find the identifiers of subjects in a Pedigree iteratively, as anyone who is not available and does not have an available descendant by successively removing unavailable terminal nodes.

# Usage

```
## S4 method for signature 'Ped'
find_unavailable(obj, avail = NULL)
## S4 method for signature 'Pedigree'
find_unavailable(obj, avail = NULL)
```

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### Arguments

obj A Ped or Pedigree object.

avail A logical vector with the availability status of the individuals (i.e. FALSE = not

available, TRUE = available, NA = unknown).

#### **Details**

If avail is null, then the function will use the corresponding Ped accessor.

Originally written as pedTrim by Steve Iturria, modified by Dan Schaid 2007, and now split into the two separate functions: find\_unavailable(), and trim() to do the tasks separately. find\_unavailable() calls exclude\_stray\_marryin() to find stray available marry-ins who are isolated after trimming their unavailable offspring, and exclude\_unavail\_founders(). If the subject ids are character, make sure none of the characters in the ids is a colon (":"), which is a special character used to concatenate and split subjects within the utility. The trim() functions is now replaced by the subset() function.

#### Value

Returns a vector of subject ids for who can be removed.

## **Side Effects**

Relation matrix from subsetting is trimmed of any special relations that include the subjects to trim.

#### See Also

```
shrink()
```

#### **Examples**

```
data(sampleped)
ped1 <- Pedigree(sampleped[sampleped$famid == "1",])
find_unavailable(ped1)</pre>
```

fix\_parents

Fix parents relationship and gender

#### **Description**

Fix the sex of parents, add parents that are missing from the data. Can be used with a dataframe or a vector of the different individuals informations.

fix\_parents 47

# Usage

```
## S4 method for signature 'character'
fix_parents(obj, dadid, momid, sex, famid = NULL, missid = NA_character_)
## S4 method for signature 'data.frame'
fix_parents(obj, del_parents = NULL, filter = NULL, missid = NA_character_)
```

#### **Arguments**

obj	A data frame or a vector of the individuals identifiers. If a dataframe is given it must contain the columns id, dadid, momid, sex and famid (optional).
dadid	A vector containing for each subject, the identifiers of the biologicals fathers.
momid	A vector containing for each subject, the identifiers of the biologicals mothers.
sex	A character, factor or numeric vector corresponding to the gender of the individuals. This will be transformed to an ordered factor with the following levels: male < female < unknown
	The following values are recognized:
	• "male": "m", "male", "man", 1
	• "female": "f", "female", "woman", 2
	• "unknown": "unknown", 3
famid	A character vector with the family identifiers of the individuals. If provide, will be aggregated to the individuals identifiers separated by an underscore.
missid	A character vector with the missing values identifiers. All the id, dadid and momid corresponding to those values will be set to NA_character
del_parents	Boolean defining if missing parents needs to be deleted or fixed. If one then if one of the parent is missing, both are removed, if both then if both parents are missing, both are removed. If NULL then no parent is removed and the missing parents are added as new rows.
filter	Filtering column containing $\emptyset$ or 1 for the rows to kept before proceeding.

# **Details**

First look to add parents whose ids are given in momid/dadid. Second, fix sex of parents. Last look to add second parent for children for whom only one parent id is given. If a **famid** vector is given the family id will be added to the ids of all individuals (id, dadid, momid) separated by an underscore before proceeding.

#### Special case for dataframe:

Check for presence of both parents id in the **id** field. If not both presence behaviour depend of **delete** parameter

- If TRUE then use fix\_parents function and merge back the other fields in the dataframe then set availability to 0 for non available parents.
- If FALSE then delete the id of missing parents

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#### Value

A data.frame with id, dadid, momid, sex as columns with the relationships fixed.

#### Author(s)

Jason Sinnwell

#### **Examples**

```
test1char <- data.frame(
    id = paste('fam', 101:111, sep = ''),
    sex = c('male', 'female')[c(1, 2, 1, 2, 1, 1, 2, 2, 1, 2, 1)],
    father = c(
        0, 0, 'fam101', 'fam101', 'fam101', 0, 0,
        'fam106', 'fam106', 'fam109'
    ),
    mother = c(
        0, 0, 'fam102', 'fam102', 'fam102', 0, 0,
        'fam107', 'fam107', 'fam112'
    )
)
test1newmom <- with(test1char, fix_parents(id, father, mother,
        sex,
        missid = NA_character_
))
Pedigree(test1newmom)</pre>
```

generate\_aff\_inds

Process the affection informations

# Description

Perform transformation uppon a vector given as the one containing the affection status to obtain an affected binary state.

#### Usage

```
generate_aff_inds(
  values,
  mods_aff = NULL,
  threshold = NULL,
  sup_thres_aff = NULL,
  is_num = NULL
)
```

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### **Arguments**

values	Vector containing the values of the column to process.
mods_aff	Vector of modality to consider as affected in the case where the values is a factor.
threshold	Numeric value separating the affected and healthy subject in the case where the values is numeric.
sup_thres_aff	Boolean defining if the affected individual are above the threshold or not. If TRUE, the individuals will be considered affected if the value of values is stricly above the threshold. If FALSE, the individuals will be considered affected if the value is stricly under the threshold.
is_num	Boolean defining if the values need to be considered as numeric.

#### **Details**

This function helps to configure a binary state from a character or numeric variable.

#### If the variable is a character or a factor::

In this case the affected state will depend on the modality provided as an affected status. All individuals with a value corresponding to one of the element in the vector **mods\_aff** will be considered as affected.

#### If the variable is numeric::

In this case the affected state will be TRUE if the value of the individual is above the **threshold** if **sup\_thres\_aff** is TRUE and FALSE otherwise.

#### Value

A dataframe with the affected column processed accordingly. The different columns are:

- mods: The different modalities of the column
- labels: The labels of the different modalities
- affected: The column processed to have only TRUE/FALSE values

```
generate\_aff\_inds(c(1, 2, 3, 4, 5), threshold = 3, sup\_thres\_aff = TRUE) \\ generate\_aff\_inds(c("A", "B", "C", "A", "V", "B"), mods\_aff = c("A", "B"))
```

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generate_border	Process the border colors based on availability	

## **Description**

Perform transformation uppon a vector given as the one containing the availability status to compute the border color. The vector given will be transformed using the vect\_to\_binary() function.

## Usage

```
generate_border(values, colors_avail = c("green", "black"), colors_na = "grey")
```

### **Arguments**

values The vector containing the values to process as available.

colors\_avail Set of 2 colors to use for the box's border of an individual. The first color will

be used for available individual (avail == 1) and the second for the unavailable

individual (avail == 0).

colors\_na Color to use for individuals with no informations.

### Value

A list of three elements

- mods: The processed values column as a numeric factor
- avail: A logical vector indicating if the individual is available
- sc\_bord : A dataframe containing the description of each modality of the scale

## **Examples**

```
generate\_border(c(1, 0, 1, 0, NA, 1, 0, 1, 0, NA))
```

generate_colors	Process the filling and border colors based on affection and availability

## Description

Perform transformation uppon a dataframe given to compute the colors for the filling and the border of the individuals based on the affection and availability status.

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## Usage

```
## S4 method for signature 'character'
generate_colors(
 obj,
  avail,
 mods_aff = NULL,
  is_num = FALSE,
  keep_full_scale = FALSE,
  colors_aff = c("yellow2", "red"),
  colors_unaff = c("white", "steelblue4"),
  colors_avail = c("green", "black"),
  colors_na = "grey"
)
## S4 method for signature 'numeric'
generate_colors(
 obj,
  avail,
  threshold = 0.5,
  sup_thres_aff = TRUE,
  is_num = TRUE,
  keep_full_scale = FALSE,
  breaks = 3,
  colors_aff = c("yellow2", "red"),
  colors_unaff = c("white", "steelblue4"),
  colors_avail = c("green", "black"),
  colors_na = "grey"
)
## S4 method for signature 'Pedigree'
generate_colors(
 obj,
  col_aff = "affection",
  add_to_scale = TRUE,
  col_avail = "avail",
  is_num = NULL,
 mods_aff = TRUE,
  threshold = 0.5,
  sup_thres_aff = TRUE,
  keep_full_scale = FALSE,
  breaks = 3,
  colors_aff = c("yellow2", "red"),
  colors_unaff = c("white", "steelblue4"),
  colors_avail = c("green", "black"),
 colors_na = "grey",
  reset = TRUE
)
```

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# Arguments

obj	A Pedigree object or a vector containing the affection status for each individuals. The affection status can be numeric or a character.
avail	A logical vector with the availability status of the individuals (i.e. $FALSE = not$ available, $TRUE = available$ , $NA = unknown$ ).
mods_aff	Vector of modality to consider as affected in the case where the values is a factor.
is_num keep_full_scal	Boolean defining if the values need to be considered as numeric.
	Boolean defining if the affection values need to be set as a scale. If values is numeric the filling scale will be calculated based on the values and the number of breaks given. If values isn't numeric then each levels will get it's own color
colors_aff	Set of increasing colors to use for the filling of the affected individuls.
colors_unaff	Set of increasing colors to use for the filling of the unaffected individuls.
colors_avail	Set of 2 colors to use for the box's border of an individual. The first color will be used for available individual (avail == 1) and the second for the unavailable individual (avail == $\emptyset$ ).
colors_na	Color to use for individuals with no informations.
threshold	Numeric value separating the affected and healthy subject in the case where the values is numeric.
sup_thres_aff	Boolean defining if the affected individual are above the threshold or not. If TRUE, the individuals will be considered affected if the value of values is strictly above the threshold. If FALSE, the individuals will be considered affected if the value is strictly under the threshold.
breaks	Number of breaks to use when using full scale with numeric values. The same number of breaks will be done for values from affected individuals and unaffected individuals.
col_aff	A character vector with the name of the column to be used for the affection status.
add_to_scale	Boolean defining if the scales need to be added to the existing scales or if they need to replace the existing scales.
col_avail	A character vector with the name of the column to be used for the availability status.
reset	If TRUE the scale of the specified column will be reset if already present.

# **Details**

The colors will be set using the generate\_fill()] and the <code>generate\_border()</code> functions respectively for the filling and the border.

## Value

## When used with a vector:

A list of two elements

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- The list containing the filling colors processed and their description
- The list containing the border colors processed and their description

#### When used with a Pedigree object:

The Pedigree object with the affected and avail columns processed accordingly as well as the scales slot updated.

### **Examples**

```
generate_colors(
    c("A", "B", "A", "B", NA, "A", "B", "A", "B", NA),
    c(1, 0, 1, 0, NA, 1, 0, 1, 0, NA),
    mods_aff = "A"
)

generate_colors(
    c(10, 0, 5, 7, NA, 6, 2, 1, 3, NA),
    c(1, 0, 1, 0, NA, 1, 0, 1, 0, NA),
    threshold = 3, keep_full_scale = TRUE
)
data("sampleped")
pedi <- Pedigree(sampleped)
pedi <- generate_colors(pedi, "affection", add_to_scale=FALSE)
scales(pedi)</pre>
```

generate\_fill

Process the filling colors based on affection

#### **Description**

Perform transformation uppon a column given as the one containing affection status to compute the filling color.

### Usage

```
generate_fill(
  values,
  affected,
  labels,
  is_num = NULL,
  keep_full_scale = FALSE,
  breaks = 3,
  colors_aff = c("yellow2", "red"),
  colors_unaff = c("white", "steelblue4"),
  colors_na = "grey"
)
```

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## **Arguments**

values The vector containing the values to process as affection.

affected A logical vector with the affection status of the individuals (i.e. FALSE = unaf-

fected, TRUE = affected, NA = unknown).

labels The vector containing the labels to use for the affection.

is\_num Boolean defining if the values need to be considered as numeric.

keep\_full\_scale

Boolean defining if the affection values need to be set as a scale. If values is numeric the filling scale will be calculated based on the values and the number of breaks given. If values isn't numeric then each levels will get it's own color

breaks Number of breaks to use when using full scale with numeric values. The same

number of breaks will be done for values from affected individuals and unaf-

fected individuals.

colors\_aff Set of increasing colors to use for the filling of the affected individuls.

Set of increasing colors to use for the filling of the unaffected individuls.

colors\_na Color to use for individuals with no informations.

#### **Details**

The colors will be set using the grDevices::colorRampPalette() function with the colors given as parameters.

The colors will be set as follow:

- If keep\_full\_scale is FALSE: Then the affected individuals will get the first color of the colors\_aff vector and the unaffected individuals will get the first color of the colors\_unaff vector.
- If keep\_full\_scale is TRUE:
  - If values isn't numeric: Each levels of the affected values vector will get it's own color from the colors\_aff vector using the grDevices::colorRampPalette() and the same will be done for the unaffected individuals using the colors\_unaff.
  - If values is numeric: The mean of the affected individuals will be compared to the mean
    of the unaffected individuals and the colors will be set up such as the color gradient follow
    the direction of the affection.

#### Value

A list of three elements

- mods: The processed values column as a numeric factor
- affected: A logical vector indicating if the individual is affected
- sc\_fill: A dataframe containing the description of each modality of the scale

```
aff <- generate_aff_inds(seq_len(5), threshold = 3, sup_thres_aff = TRUE)
generate_fill(seq_len(5), aff$affected, aff$labels)
generate_fill(seq_len(5), aff$affected, aff$labels, keep_full_scale = TRUE)</pre>
```

get\_dataframe 55

get\_dataframe

Get dataframe name

# Description

Extract the name of the different dataframe present in a file

## Usage

```
get_dataframe(file)
```

# **Arguments**

file

The file path

#### **Details**

This function detect the extension of the file and extract if necessary the different dataframe / sheet names available.

#### Value

A vector of all the dataframe name present.

# **Examples**

```
## Not run:
    get_dataframe('path/to/my/file.txt')
## End(Not run)
```

get\_famid

Get family id

# Description

Get the family id from the individuals identifiers.

# Usage

```
get_famid(obj)
## S4 method for signature 'character'
get_famid(obj)
```

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# **Arguments**

obj

A character vector of individual ids

#### **Details**

The family id is the first part of the individual id, separated by an underscore. If the individual id does not contain an underscore, then the family id is set to NA.

## Value

A character vector of family ids

## **Examples**

```
get_famid(c("A", "1_B", "C_2", "D_", "_E", "F"))
```

get\_families\_table

Summarise the families information for a given variable in a data frame

# Description

This function summarises the families information for a given variable in a data frame. It returns the most numerous modality for each family and the number of individuals in the family.

#### Usage

```
get_families_table(df, var)
```

# **Arguments**

df a data frame

var the variable to summarise

# Value

a data frame with the family information

```
df <- data.frame(
    famid = c(1, 1, 2, 2, 3, 3),
    health = c("A", "B", "A", "A", "B", "B"))
get_families_table(df, "health")</pre>
```

get\_title 57

get\_title

Get the title of the family information table

## **Description**

This function generates the title of the family information table depending on the selected family and subfamily and other parameters.

## Usage

```
get_title(
  family_sel,
  subfamily_sel,
  family_var,
  mod,
  inf_selected,
  kin_max,
  keep_parents,
  nb_rows,
  short_title = FALSE
)
```

## **Arguments**

```
family_sel
                  the selected family
subfamily_sel
                  the selected subfamily
family_var
                  the selected family variable
mod
                  the selected affected modality
                  the selected informative individuals
inf_selected
                  the maximum kinship
kin_max
keep_parents
                  the keep parents option
                  the number of individuals
nb_rows
                  a boolean to generate a short title
short_title
```

#### Value

a string with the title

```
get_title(1, 1, "health", "A", "All", 3, TRUE, 10, FALSE)
get_title(1, 1, "health", "A", "All", 3, TRUE, 10, TRUE)
get_title(1, 1, "health", "A", "All", 3, FALSE, 10, FALSE)
```

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get\_twin\_rel

Get twin relationships

# Description

Get twin relationships

## Usage

```
get_twin_rel(obj)
```

# Arguments

obj

A Pedigree object

#### **Details**

This routine function determine the twin relationships in a Pedigree. It determine the order of the twins in the Pedigree. It is used by auto\_hint().

## Value

A list containing components

- 1. twinset the set of twins
- 2. twinrel the twins relationships
- 3. twinord the order of the twins

#### See Also

```
auto_hint()
```

health\_sel\_ui

Shiny module to select a health variable in a pedigree

## **Description**

This module allows to select health variables in a pedigree object. The function is composed of two parts: the UI and the server. The UI is called with the function health\_sel\_ui() and the server with the function health\_sel\_server().

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#### Usage

```
health_sel_ui(id)
health_sel_server(
  id,
  pedi,
  var = NULL,
  as_num = NULL,
  mods_aff = NULL,
  threshold = NULL,
  sup_threshold = NULL
)
health_sel_demo()
```

### **Arguments**

id A string to identify the module.

pedi A reactive pedigree object.

var A string with the name of the health variable to select.

as\_num A boolean to know if the health variable needs to be considered as numeric.

mods\_aff A character vector of the affected modalities.

threshold A numeric threshold to determine affected individuals.

sup\_threshold A boolean to know if the affected individuals are strickly superior to the thresh-

old.

### Value

A reactive list with the following informations:actions-box

- health\_var: the selected health variable,
- to\_num: a boolean to know if the health variable needs to be considered as numeric,
- mods\_aff: a character vector of the affected modalities,
- threshold: a numeric threshold to determine affected individuals,
- sup\_threshold: a boolean to know if the affected individuals are strickly superior to the threshold.

```
if (interactive()) {
    health_sel_demo()
}
```

60 Hints-class

Hints-class

Hints object

#### **Description**

The hints are used to specify the order of the individuals in the pedigree and to specify the order of the spouses.

#### **Constructor::**

You either need to provide **horder** or **spouse** in the dedicated parameters (together or separately), or inside a list.

### Usage

```
Hints(horder, spouse)
## S4 method for signature 'list,missing_OR_NULL'
Hints(horder, spouse)
## S4 method for signature 'numeric,data.frame'
Hints(horder, spouse)
## S4 method for signature 'numeric,missing_OR_NULL'
Hints(horder, spouse)
```

#### **Arguments**

horder

A named numeric vector with one element per subject in the Pedigree. It determines the relative horizontal order of subjects within a sibship, as well as the relative order of processing for the founder couples. (For this latter, the female founders are ordered as though they were sisters). The names of the vector should be the individual identifiers.

spouse

A data frame with one row per hinted marriage, usually only a few marriages in a pedigree will need an added hint, for instance reverse the plot order of a husband/wife pair. Each row contains the id of the left spouse (i.e. id1), the id of the right hand spouse (i.e. idr), and the anchor (i.e: anchor: 1 = left, 2 = right, 0 = either). Children will preferentially appear under the parents of the anchored spouse.

## Value

A Hints object.

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#### **Slots**

horder A numeric named vector with one element per subject in the Pedigree. It determines the relative horizontal order of subjects within a sibship, as well as the relative order of processing for the founder couples. (For this latter, the female founders are ordered as though they were sisters).

spouse A data frame with one row per hinted marriage, usually only a few marriages in a Pedigree will need an added hint, for instance reverse the plot order of a husband/wife pair. Each row contains the identifiers of the left spouse, the right hand spouse, and the anchor (i.e : 1 = left, 2 = right, 0 = either).

#### Accessors

- horder(x): Get the horder vector
- horder(x) <- value : Set the horder vector
- spouse(x): Get the spouse data.frame
- spouse(x) <- value : Set the spouse data.frame

#### Generics

- as.list(x): Convert a Hints object to a list
- subset(x, i, keep = TRUE): Subset a Hints object based on the individuals identifiers given.
  - i : A vector of individuals identifiers to keep.
  - keep: A logical value indicating if the individuals should be kept or deleted.

#### See Also

```
Pedigree()
```

```
Hints(
    list(
        horder = c("1" = 1, "2" = 2, "3" = 3),
        spouse = data.frame(
            idl = c("1", "2"),
            idr = c("2", "3"),
            anchor = c(1, 2)
        )
    )
}

Hints(
    horder = c("1" = 1, "2" = 2, "3" = 3),
    spouse = data.frame(
    idl = c("1", "2"),
    idr = c("2", "3"),
```

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```
anchor = c(1, 2)
)

Hints(
    horder = c("1" = 1, "2" = 2, "3" = 3)
)
```

ibd\_matrix

IBD matrix

## **Description**

Transform identity by descent (IBD) matrix data from the form produced by external programs such as SOLAR into the compact form used by the coxme and lmekin routines.

### Usage

```
ibd_matrix(id1, id2, ibd, idmap, diagonal)
```

## **Arguments**

id1	A character vector with the id of the first individuals of each pairs or a matrix or data frame with 3 columns: id1, id2, and ibd
id2	A character vector with the id of the second individuals of each pairs
ibd	the IBD value for that pair
idmap	an optional 2 column matrix or data frame whose first element is the internal value (as found in id1 and id2, and whose second element will be used for the dimnames of the result
diagonal	optional value for the diagonal element. If present, any missing diagonal elements in the input data will be set to this value.

### **Details**

The IBD matrix for a set of n subjects will be an n by n symmetric matrix whose i,j element is the contains, for some given genetic location, a 0/1 indicator of whether 0, 1/2 or 2/2 of the alleles for i and j are identical by descent. Fractional values occur if the IBD fraction must be imputed. The diagonal will be 1. Since a large fraction of the values will be zero, programs such as Solar return a data set containing only the non-zero elements. As well, Solar will have renumbered the subjects as seq\_len(n) in such a way that families are grouped together in the matrix; a separate index file contains the mapping between this new id and the original one. The final matrix should be labeled with the original identifiers.

#### Value

a sparse matrix of class dsCMatrix. This is the same form used for kinship matrices.

inf\_sel\_ui 63

#### See Also

```
kinship()
```

#### **Examples**

```
df <- data.frame(
    id1 = c("1", "2", "1"),
    id2 = c("2", "3", "4"),
    ibd = c(0.5, 0.16, 0.27)
)
ibd_matrix(df$id1, df$id2, df$ibd, diagonal = 2)</pre>
```

inf\_sel\_ui

Shiny module to select the informative individuals in a pedigree

# **Description**

This module allows to select informative individuals in a pedigree object. They will be used to subset the pedigree object with the function useful\_inds(). Further filtering options are available (max kinship and keep parents). The function is composed of two parts: the UI and the server. The UI is called with the function inf\_sel\_ui() and the server with the function inf\_sel\_server().

# Usage

```
inf_sel_ui(id)
inf_sel_server(id, pedi)
inf_sel_demo(pedi)
```

## Arguments

id A string to identify the module.pedi A reactive pedigree object.

#### Value

A reactive pedigree object subselected from the informative individuals.

```
if (interactive()) {
    data("sampleped")
    pedi <- shiny::reactive({
        Pedigree(sampleped[sampleped$famid == "1", ])
    })
    inf_sel_demo(pedi)
}</pre>
```

is\_founder

is\_disconnected

Are individuals disconnected

# Description

Check which individuals are disconnected.

## Usage

```
is_disconnected(id, dadid, momid)
```

## **Arguments**

dadid A vector containing for each subject, the identifiers of the biologicals fathers.

Momid A vector containing for each subject, the identifiers of the biologicals mothers.

#### **Details**

An individuals is considered disconnected if the kinship with all the other individuals is 0.

#### Value

A vector of boolean of the same size as **id** with TRUE if the individual is disconnected and FALSE otherwise

## **Examples**

```
is_disconnected(
    c("1", "2", "3", "4", "5"),
    c("3", "3", NA, NA, NA),
    c("4", "4", NA, NA, NA)
)
```

is\_founder

Are individuals founders

## **Description**

Check which individuals are founders.

## Usage

```
is_founder(momid, dadid, missid = NA_character_)
```

 $is\_informative$ 65

# **Arguments**

momid	A vector containing for each subject, the identifiers of the biologicals mothers.
dadid	A vector containing for each subject, the identifiers of the biologicals fathers.
missid	A character vector with the missing values identifiers. All the id, dadid and
	momid corresponding to those values will be set to NA_character

#### Value

A vector of boolean of the same size as dadid and momid with TRUE if the individual has no parents (i.e is a founder) and FALSE otherwise.

### **Examples**

```
is_founder(c("3", "3", NA, NA), c("4", "4", NA, NA))
```

is\_informative

Find informative individuals

# Description

Select the ids of the informative individuals.

# Usage

```
## S4 method for signature 'character_OR_integer'
is_informative(obj, avail, affected, informative = "AvAf")
## S4 method for signature 'Ped'
is_informative(obj, informative = "AvAf", reset = FALSE)
## S4 method for signature 'Pedigree'
is_informative(obj, col_aff = NULL, informative = "AvAf", reset = FALSE)
```

## **Arguments**

obj	A character vector with the id of the individuals or a data. frame with all the informations in corresponding columns.
avail	A logical vector with the availability status of the individuals (i.e. FALSE = not available, TRUE = available, NA = unknown).
affected	A logical vector with the affection status of the individuals (i.e. $FALSE = unafected$ , $TRUE = affected$ , $NA = unknown$ ).
informative	Informative individuals selection can take 5 values:
	<ul> <li>'AvAf' (available and affected),</li> </ul>
	• 'AvOrAf' (available or affected).

- 'AvOrAf' (available or affected),
- 'Av' (available only),

is\_informative

- 'Af' (affected only),
- 'All' (all individuals)
- A numeric/character vector of individuals id
- A boolean

reset If TRUE, the isinf slot is reset

col\_aff A character vector with the name of the column to be used for the affection

status.

#### **Details**

Depending on the **informative** parameter, the function will extract the ids of the informative individuals. In the case of a numeric vector, the function will return the same vector. In the case of a boolean, the function will return the ids of the individuals if TRUE, NA otherwise. In the case of a string, the function will return the ids of the corresponding informative individuals based on the avail and affected columns.

#### Value

## When obj is a vector:

A vector of individuals informative identifiers.

#### When obj is a Pedigree:

The Pedigree object with its isinf slot updated.

```
is_informative(c("A", "B", "C", "D", "E"), informative = c("A", "B"))
is_informative(c("A", "B", "C", "D", "E"), informative = c(1, 2))
is_informative(c("A", "B", "C", "D", "E"), informative = c("A", "B")) is_informative(c("A", "B", "C", "D", "E"), avail = c(1, 0, 0, 1, 1),
    affected = c(0, 1, 0, 1, 1), informative = "AvAf")
is_informative(c("A", "B", "C", "D", "E"), avail = c(1, 0, 0, 1, 1),
    affected = c(0, 1, 0, 1, 1), informative = "AvOrAf")
is_informative(c("A", "B", "C", "D", "E"),
    informative = c(TRUE, FALSE, TRUE, FALSE, TRUE))
data("sampleped")
pedi <- ped(Pedigree(sampleped))</pre>
pedi <- is_informative(pedi, informative = "Av")</pre>
isinf(pedi)
data("sampleped")
pedi <- Pedigree(sampleped)</pre>
pedi <- is_informative(pedi, col_aff = "affection")</pre>
isinf(ped(pedi))
```

is\_parent 67

is_parent Are	e individuals parents
---------------	-----------------------

# Description

Check which individuals are parents.

# Usage

```
## S4 method for signature 'character_OR_integer'
is_parent(obj, dadid, momid, missid = NA_character_)
## S4 method for signature 'Ped'
is_parent(obj, missid = NA_character_)
```

# **Arguments**

obj	A vector of each subjects identifiers or a Ped object
dadid	A vector containing for each subject, the identifiers of the biologicals fathers.
momid	A vector containing for each subject, the identifiers of the biologicals mothers.
missid	A character vector with the missing values identifiers. All the id, dadid and momid corresponding to those values will be set to NA_character

#### Value

A vector of boolean of the same size as  ${\bf obj}$  with TRUE if the individual is a parent and FALSE otherwise

```
is_parent(c("1", "2", "3", "4"), c("3", "3", NA, NA), c("4", "4", NA, NA))
data(sampleped)
pedi <- Pedigree(sampleped)
is_parent(ped(pedi))</pre>
```

68 is\_valid\_ped

### **Description**

Check if horder and spouse slots are valid:

- horder is named numeric vector
- spouse is a data.frame
  - Has the three idr, idl, anchor columns
  - idr and idl are different and doesn't contains NA
  - idr and idl couple are unique
  - anchor column only have right, left or either values
- all ids in spouse needs to be in the names of the horder vector

### Usage

```
is_valid_hints(object)
```

## **Arguments**

object

A Hints object.

#### Value

A character vector with the errors or TRUE if no errors.

is\_valid\_ped

Check if a Ped object is valid

#### **Description**

Multiple checks are done here

## Usage

```
is_valid_ped(object)
```

#### **Arguments**

object

A Ped object.

#### **Details**

- 1. Check that the ped ids slots have the right values
- 2. Check that the sex, fertility, deceased, avail and affected slots have the right values
- 3. Check that dad are male and mom are female
- 4. Check that individuals have both parents or none
- 5. Check that proband are affected
- 6. Check that proband and consultand are mutually exclusive
- 7. Check that asymptomatic individuals are not affected

is\_valid\_pedigree 69

## Value

A character vector with the errors or TRUE if no errors.

is\_valid\_pedigree

Check if a Pedigree object is valid

## **Description**

Multiple checks are done here

## Usage

is\_valid\_pedigree(object)

## **Arguments**

object

A Ped object.

## **Details**

- 1. Check that the all Rel id are in the Ped object
- 2. Check that twins have same parents
- 3. Check that MZ twins have same sex
- 4. Check that all columns used in scales are in the Ped object
- 5. Check that all fill & border modalities are in the Ped object column
- 6. Check that all id used in Hints object are in the Ped object
- 7. Check that all spouse in Hints object are male / female

## Value

A character vector with the errors or TRUE if no errors.

70 is\_valid\_scales

is\_valid\_rel

Check if a Rel object is valid

## **Description**

Multiple checks are done here

## Usage

```
is_valid_rel(object)
```

## **Arguments**

object

A Ped object.

#### **Details**

- 1. Check that the "id1", "id2", "code", "famid" slots exist
- 2. Check that the "code" slots have the right values (i.e. "MZ twin", "DZ twin", "UZ twin", "Spouse")
- 3. Check that all "id1" are different to "id2"
- 4. Check that all "id1" are smaller than "id2"
- 5. Check that no duplicate relation are present

## Value

A character vector with the errors or TRUE if no errors.

is\_valid\_scales

Check if a Scales object is valid

# Description

Check if the fill and border slots are valid:

- fill slot is a data.frame with "order", "column\_values", "column\_mods", "mods", "labels", "affected", "fill", "density", "angle" columns.
  - "affected" is logical.
  - "density", "angle", "order", "mods" are numeric.
  - "column\_values", "column\_mods", "labels", "fill" are character.
- border slot is a data.frame with "column\_values", "column\_mods", "mods", "labels", "border" columns.
  - "column\_values", "column\_mods", "labels", "border" are character.
  - "mods" is numeric.

kindepth 71

#### Usage

```
is_valid_scales(object)
```

#### **Arguments**

object A Scales object.

#### Value

A character vector with the errors or TRUE if no errors.

kindepth

Individual's depth in a pedigree

### **Description**

Computes the depth of each subject in the Pedigree.

### Usage

```
## S4 method for signature 'character_OR_integer'
kindepth(obj, dadid, momid, align_parents = FALSE, force = FALSE)
## S4 method for signature 'Pedigree'
kindepth(obj, align_parents = FALSE, force = FALSE)
## S4 method for signature 'Ped'
kindepth(obj, align_parents = FALSE, force = FALSE)
```

#### **Arguments**

obj	A character vector with the id of the individuals or a data.frame with all the informations in corresponding columns.
dadid	A vector containing for each subject, the identifiers of the biologicals fathers.
momid	A vector containing for each subject, the identifiers of the biologicals mothers.
align_parents	If align_parents = TRUE, go one step further and try to make both parents of each child have the same depth. (This is not always possible). It helps the drawing program by lining up pedigrees that 'join in the middle' via a marriage.
force	If force = TRUE, the function will return the depth minus min(depth) if depth reach a state with no founders is not possible.

### **Details**

Mark each person as to their depth in a Pedigree; 0 for a founder, otherwise:

```
depth = 1 + \max(fatherDepth, motherDepth)
```

In the case of an inbred Pedigree a perfect alignment may not exist.

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#### Value

An integer vector containing the depth for each subject

#### Author(s)

Terry Therneau, updated by Louis Le Nezet

#### See Also

```
align()
```

## **Examples**

```
kindepth(
     c("A", "B", "C", "D", "E"),
     c("C", "D", "0", "0"),
     c("E", "E", "0", "0", "0")
)
data(sampleped)
ped1 <- Pedigree(sampleped[sampleped$famid == "1",])
kindepth(ped1)</pre>
```

kinship

Kinship matrix

## **Description**

Compute the kinship matrix for a set of related autosomal subjects. The function is generic, and can accept a Pedigree, a Ped or a vector as the first argument.

## Usage

```
## S4 method for signature 'Ped'
kinship(obj, chrtype = "autosome")

## S4 method for signature 'character'
kinship(obj, dadid, momid, sex, chrtype = "autosome")

## S4 method for signature 'Pedigree'
kinship(obj, chrtype = "autosome")
```

## Arguments

obj	A Pedigree or Ped object or a vector of subject identifiers.
chrtype	chromosome type. The currently supported types are 'autosome' and 'X' or 'x'.
dadid	A vector containing for each subject, the identifiers of the biologicals fathers.
momid	A vector containing for each subject, the identifiers of the biologicals mothers.

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sex

A character, factor or numeric vector corresponding to the gender of the individuals. This will be transformed to an ordered factor with the following levels: male < female < unknown

The following values are recognized:

```
"male": "m", "male", "man", 1"female": "f", "female", "woman", 2
```

• "unknown": "unknown", 3

#### **Details**

The function will usually be called with a Pedigree. The call with a Ped or a vector is provided for backwards compatibility with an earlier release of the library that was less capable. Note that when using with a Ped or a vector, any information on twins is not available to the function.

Warning: This function does not work with adopted individuals. If you have adopted individuals in your pedigree, you should remove them before calling this function.

When called with a Pedigree, the routine will create a block-diagonal-symmetric sparse matrix object of class dsCMatrix. Since the [i, j] value of the result is 0 for any two unrelated individuals i and j and a Matrix utilizes sparse representation, the resulting object is often orders of magnitude smaller than an ordinary matrix.

Two genes G1 and G2 are identical by descent (IBD) if they are both physical copies of the same ancestral gene; two genes are identical by state if they represent the same allele. So the brown eye gene that I inherited from my mother is IBD with hers; the same gene in an unrelated individual is not

The kinship coefficient between two subjects is the probability that a randomly selected allele from a locus will be IBD between them. It is obviously 0 between unrelated individuals. For an autosomal site and no inbreeding it will be 0.5 for an individual with themselves, .25 between mother and child, .125 between an uncle and neice, etc.

The computation is based on a recursive algorithm described in Lange, which assumes that the founder alleles are all independent.

### Value

#### When obj is a vector:

A matrix of kinship coefficients.

#### When obj is a Pedigree:

A matrix of kinship coefficients ordered by families present in the Pedigree object.

### References

K Lange, Mathematical and Statistical Methods for Genetic Analysis, Springer-Verlag, New York, 1997.

#### See Also

```
make_famid(), kindepth()
```

74 make\_class\_info

## **Examples**

make\_class\_info

Make class information

# Description

Make class information

## Usage

```
make_class_info(x)
```

## **Arguments**

Х

A list of class

### Value

A character vector of class information

```
Pedixplorer:::make_class_info(list(
    1, "a", 1, 2, 3, list(1, 2)
))
```

make\_famid 75

Compute family id
-------------------

## **Description**

Construct a family identifier from pedigree information

## Usage

```
## S4 method for signature 'character'
make_famid(obj, dadid, momid)
## S4 method for signature 'Pedigree'
make_famid(obj)
```

# Arguments

obj	A character vector with the id of the individuals or a data.frame with all the informations in corresponding columns.
dadid	A vector containing for each subject, the identifiers of the biologicals fathers.
momid	A vector containing for each subject, the identifiers of the biologicals mothers.

#### **Details**

Create a vector of length n, giving the family 'tree' number of each subject. If the Pedigree is totally connected, then everyone will end up in tree 1, otherwise the tree numbers represent the disconnected subfamilies. Singleton subjects give a zero for family number.

### Value

### When used with a character vector:

An integer vector giving family groupings

## When used with a Pedigree object:

An updated Pedigree object with the family id added and with all ids updated

## See Also

```
kinship()
```

```
make_famid(
    c("A", "B", "C", "D", "E", "F"),
    c("C", "D", "0", "0", "0", "0"),
    c("E", "E", "0", "0", "0", "0")
)
```

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```
data(sampleped)
ped1 <- Pedigree(sampleped[,-1])
make_famid(ped1)</pre>
```

make\_rownames

Make rownames for rectangular data display

### **Description**

Make rownames for rectangular data display

## Usage

```
make_rownames(x_rownames, nrow, nhead, ntail)
```

### **Arguments**

x\_rownames The rownames of the data

nrow The number of rows in the data

nhead The number of rownames to display at the beginning

ntail The number of rownames to display at the end

### Value

A character vector of rownames

### **Examples**

```
Pedixplorer:::make_rownames(rownames(mtcars), nrow(mtcars), 3, 3)
```

minnbreast

Minnesota Breast Cancer Study

## **Description**

Data from the Minnesota Breast Cancer Family Study. This contains extended pedigrees from 426 families, each identified by a single proband in 1945-1952, with follow up for incident breast cancer.

```
data(minnbreast)
```

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#### **Format**

A data frame with 28081 observations, one line per subject, on the following 14 variables.

- id : Subject identifier
- proband : If 1, this subject is one of the original 426 probands
- fatherid: Identifier of the father, if the father is part of the data set; zero otherwise
- motherid: Identifier of the mother, if the mother is part of the data set; zero otherwise
- famid: Family identifier
- endage : Age at last follow-up or incident cancer
- cancer: 1 = breast cancer (females) or prostate cancer (males), 0 = censored
- · yob: Year of birth
- education: Amount of education: 1-8 years, 9-12 years, high school graduate, vocational education beyond high school, some college but did not graduate, college graduate, post-graduate education, refused to answer on the questionnaire
- marstat: Marital status: married, living with someone in a marriage-like relationship, separated or divorced, widowed, never married, refused to answer the questionaire
- everpreg: Ever pregnant at the time of baseline survey
- parity: Number of births
- nbreast: Number of breast biopsies
- sex: M or F
- bcpc: Part of one of the families in the breast / prostate cancer substudy: θ = no, 1 = yes.
   Note that subjects who were recruited to the overall study after the date of the BP substudy are coded as zero.

### Details

The original study was conducted by Dr. Elving Anderson at the Dight Institute for Human Genetics at the University of Minnesota. From 1944 to 1952, 544 sequential breast cancer cases seen at the University Hospital were enrolled, and information gathered on parents, siblings, offspring, aunts / uncles, and grandparents with the goal of understanding possible familial aspects of brest cancer. In 1991 the study was resurrected by Dr Tom Sellers.

Of the original 544 he excluded 58 prevalent cases, along with another 19 who had less than 2 living relatives at the time of Dr Anderson's survey. Of the remaining 462 families 10 had no living members, 23 could not be located and 8 refused, leaving 426 families on whom updated pedigrees were obtained.

This gave a study with 13351 males and 12699 females (5183 marry-ins). Primary questions were the relationship of early life exposures, breast density, and pharmacogenomics on incident breast cancer risk. For a subset of the families data was gathered on prostate cancer risk for male subjects via questionnaires sent to men over 40. Other than this, data items other than parentage are limited to the female subjects. In 2003 a second phase of the study was instituted. The pedigrees were further extended to the numbers found in this data set, and further data gathered by questionnaire.

78 min\_dist\_inf

#### References

Epidemiologic and genetic follow-up study of 544 Minnesota breast cancer families: design and methods. Sellers TA, Anderson VE, Potter JD, Bartow SA, Chen PL, Everson L, King RA, Kuni CC, Kushi LH, McGovern PG, et al. Genetic Epidemiology, 1995; 12(4):417-29.

Evaluation of familial clustering of breast and prostate cancer in the Minnesota Breast Cancer Family Study. Grabrick DM, Cerhan JR, Vierkant RA, Therneau TM, Cheville JC, Tindall DJ, Sellers TA. Cancer Detect Prev. 2003; 27(1):30-6.

Risk of breast cancer with oral contraceptive use in women with a family history of breast cancer. Grabrick DM, Hartmann LC, Cerhan JR, Vierkant RA, Therneau TM, Vachon CM, Olson JE, Couch FJ, Anderson KE, Pankratz VS, Sellers TA. JAMA. 2000; 284(14):1791-8.

### **Examples**

```
data(minnbreast)
breastped <- Pedigree(minnbreast,
    cols_ren_ped = list(
        "dadid" = "fatherid", "momid" = "motherid"
    ), missid = "0", col_aff = "cancer"
)
summary(breastped)
scales(breastped)
#plot family 8, proband is solid, slash for cancers
if (interactive()) {
    plot(breastped[famid(ped(breastped)) == "8"], aff_mark = TRUE)
}</pre>
```

min\_dist\_inf

Minimum distance to the informative individuals

## **Description**

Compute the minimum distance between the informative individuals and all the others. This distance is a transformation of the maximum kinship degree between the informative individuals and all the others. This transformation is done by taking the log2 of the inverse of the maximum kinship degree.

```
minDist = log2(1/\max(kinship))
```

Therefore, the minimum distance is 0 when the maximum kinship is 1 and is infinite when the maximum kinship is 0. For siblings, the kinship value is 0.5 and the minimum distance is 1. Each time the kinship degree is divided by 2, the minimum distance is increased by 1.

```
## S4 method for signature 'character'
min_dist_inf(obj, dadid, momid, sex, id_inf)
## S4 method for signature 'Pedigree'
```

min\_dist\_inf 79

```
min_dist_inf(obj, reset = FALSE, ...)
## S4 method for signature 'Ped'
min_dist_inf(obj, reset = FALSE)
```

### **Arguments**

obj	A character vector with the id of the individuals or a data.frame with all the informations in corresponding columns.
	Additional arguments
dadid	A vector containing for each subject, the identifiers of the biologicals fathers.
momid	A vector containing for each subject, the identifiers of the biologicals mothers.
sex	A character, factor or numeric vector corresponding to the gender of the individuals. This will be transformed to an ordered factor with the following levels: male < female < unknown  The following values are recognized:
	<ul> <li>"male": "m", "male", "man", 1</li> <li>"female": "f", "female", "woman", 2</li> <li>"unknown": "unknown", 3</li> </ul>
id_inf	An identifiers vector of informative individuals.
reset	If TRUE, the kin and if isinf columns is reset

### Value

# When obj is a vector:

A vector of the minimum distance between the informative individuals and all the others corresponding to the order of the individuals in the obj vector.

# When obj is a Pedigree:

The Pedigree object with a new slot named 'kin' containing the minimum distance between each individuals and the informative individuals. The isinf slot is also updated with the informative individuals.

### See Also

```
kinship()
```

```
min_dist_inf(
    c("A", "B", "C", "D", "E"),
    c("C", "D", "0", "0", "0"),
    c("E", "E", "0", "0", "0"),
    sex = c(1, 2, 1, 2, 1),
    id_inf = c("D", "E")
)
```

80 miscarriage\_to\_factor

```
data(sampleped)
pedi <- is_informative(
    Pedigree(sampleped),
    informative = "AvAf", col_aff = "affection"
)
kin(ped(min_dist_inf(pedi, col_aff = "affection")))</pre>
```

miscarriage\_to\_factor Miscarriage variable to factor

## Description

Transform a miscarriage variable to a factor variable By default, all other values are transformed to NA and considered as FALSE. Space and case are ignored.

# Usage

```
miscarriage_to_factor(miscarriage)
```

### **Arguments**

miscarriage

A character, factor or numeric vector corresponding to the miscarriage status of the individuals. This will be transformed to a factor with the following levels: TOP, SAB, ECT, FALSE The following values are recognized:

- "SAB": "spontaneous", "spontaenous abortion"
- "TOP": "termination", "terminated", "termination of pregnancy"
- "ECT": "ectopic", "ectopic pregnancy"
- FALSE: 0, FALSE, "no", NA

### Value

an factor vector containing the transformed variable "SAB", "TOP", "ECT", "FALSE"

```
miscarriage_to_factor(c(
    "spontaneous", "spontaenous abortion",
    "termination", "terminated", "termination of pregnancy",
    "ectopic", "ectopic pregnancy",
    "0", "false", "no", "NA"
))
```

na\_to\_length 81

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NA to specific length

## **Description**

Check if all value in a vector is NA or NULL. If so set all of them to a new value matching the length of the template. If not check that the size of the vector is equal to the template.

### Usage

```
na_to_length(x, temp, value)
```

# Arguments

x The vector to check.

temp A template vector to use to determine the length.

value The value to use to fill the vector.

## Value

A vector with the same length as temp.

## **Examples**

```
Pedixplorer:::na_to_length(NA, rep(0, 4), "NewValue")
Pedixplorer:::na_to_length(c(1, 2, 3, NA), rep(0, 4), "NewValue")
```

norm\_ped

Normalise a Ped object dataframe

# Description

Normalise dataframe for a Ped object

```
norm_ped(
  ped_df,
  na_strings = c("NA", ""),
  missid = NA_character_,
  try_num = FALSE,
  cols_used_del = FALSE,
  date_pattern = "%Y-%m-%d"
)
```

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#### **Arguments**

ped\_df

A data frame with the individuals informations. The minimum columns required are:

- id individual identifiers
- dadid biological fathers identifiers
- momid biological mothers identifiers
- · sex of the individual

The famid column, if provided, will be merged to the *ids* field separated by an underscore using the upd\_famid() function.

The following columns are also recognize and will be transformed with the vect\_to\_binary() function:

- deceased status -> is the individual dead
- avail status -> is the individual available
- evaluated status -> has the individual a documented evaluation
- consultand status -> is the individual the consultand
- proband status -> is the individual the proband
- carrier status -> is the individual a carrier
- asymptomatic status -> is the individual asymptomatic
- adopted status -> is the individual adopted

The values recognized for those columns are 1 or 0, TRUE or FALSE.

The fertility column will be transformed to a factor using the fertility\_to\_factor() function. infertile\_choice\_na, infertile, fertile

 $The \verb| miscarriage column will be transformed to a using the \verb| miscarriage_to_factor()|$ 

function. SAB, TOP, ECT, FALSE

The date of birth and date of death columns will be transformed to a date ob-

ject using the char\_to\_date() function.

na\_strings Vector of strings to be considered as NA values.

missid A character vector with the missing values identifiers. All the id, dadid and

momid corresponding to those values will be set to NA\_character\_.

try\_num Boolean defining if the function should try to convert all the columns to numeric.

cols\_used\_del Boolean defining if the columns that will be used should be deleted.

date\_pattern The pattern of the date

#### Details

Normalise a dataframe and check for columns correspondance to be able to use it as an input to create a Ped object. Multiple test are done and errors are checked.

Will be considered available any individual with no 'NA' values in the available column. Duplicated id will nullify the relationship of the individual. All individuals with errors will be remove from the dataframe and will be transferred to the error dataframe.

A number of checks are done to ensure the dataframe is correct:

### On identifiers::

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- All ids (id, dadid, momid, famid) are not empty (!= "")
- All id are unique (no duplicated)
- All dadid and momid are unique in the id column (no duplicated)
- id is not the same as dadid or momid
- Either have both parents or none

#### On sex::

- All sex code are either male, female, or unknown.
- No parents are infertile or aborted
- All fathers are male
- · All mothers are female

#### Value

A dataframe with different variable correctly standardized and with the errors identified in the error column

#### See Also

```
Ped() Ped Pedigree()
```

```
df <- data.frame(</pre>
   id = c(1, 2, 3, 4, 5, 6, 7, 8, 9, 10),
   dadid = c("A", 0, 1, 3, 0, 4, 1, 0, 6, 6).
   momid = c(0, 0, 2, 2, 0, 5, 2, 0, 8, 8),
   famid = c(1, 1, 1, 1, 1, 1, 1, 2, 2, 2),
   sex = c(1, 2, "m", "man", "f", "male", "m", 3, NA, "f"),
    fertility = c(
      "TRUE", "FALSE", TRUE, FALSE, 1,
      0, "fertile", "infertile", 1, "TRUE"
   ),
   miscarriage = c("TOB", "SAB", NA, FALSE, "ECT", "other", 1, 0, 1, 0),
   deceased = c("TRUE", "FALSE", TRUE, FALSE, 1, 0, 1, 0, 1, 0),
   avail = c("A", "1", 0, NA, 1, 0, 1, 0, 1, 0),
   evalutated = c(
        "TRUE", "FALSE", TRUE, FALSE, 1, 0, NA, "NA", "other", "0"
   ),
   consultand = c(
        "TRUE", "FALSE", TRUE, FALSE, 1, 0, NA, "NA", "other", "0"
   proband = c("TRUE", "FALSE", TRUE, FALSE, 1, 0, NA, "NA", "other", "0"),
   carrier = c("TRUE", "FALSE", TRUE, FALSE, 1, 0, NA, "NA", "other", "0"),
   asymptomatic = c(
        "TRUE", "FALSE", TRUE, FALSE, 1, 0, NA, "NA", "other", "0"
   ),
   adopted = c("TRUE", "FALSE", TRUE, FALSE, 1, 0, NA, "NA", "other", "0"),
   dateofbirth = c(
         "1978-01-01", "1980-01-01", "1982-01-01", "1984-01-01",
```

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norm\_rel

Normalise a Rel object dataframe

### **Description**

Normalise a dataframe and check for columns correspondance to be able to use it as an input to create a Ped object.

### Usage

```
norm_rel(rel_df, na_strings = c("NA", ""), missid = NA_character_)
```

### **Arguments**

rel\_df

A data.frame with the special relationships between individuals. See Rel() for more informations. The minimum columns required are id1, id2 and code. The famid column can also be used to specify the family of the individuals. If a matrix is given, the columns needs to be ordered as id1, id2, code and famid. The code values are:

- 1 = Monozygotic twin
- 2 = Dizygotic twin
- 3 = twin of unknown zygosity
- 4 = Spouse

The value relation code recognized by the function are the one defined by the rel\_code\_to\_factor() function.

na\_strings

Vector of strings to be considered as NA values.

missid

A character vector with the missing values identifiers. All the id, dadid and momid corresponding to those values will be set to NA\_character\_.

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### **Details**

The famid column, if provided, will be merged to the *ids* field separated by an underscore using the upd\_famid() function. The code column will be transformed with the rel\_code\_to\_factor(). Multiple test are done and errors are checked.

A number of checks are done to ensure the dataframe is correct:

#### On identifiers::

- All ids (id1, id2) are not empty (!= "")
- id1 and id2 are not the same

## On code:

• All code are recognised as either "MZ twin", "DZ twin", "UZ twin" or "Spouse"

# Value

A dataframe with the errors identified

### **Examples**

num\_child

Number of childs

## **Description**

Compute the number of childs per individual

```
## S4 method for signature 'character_OR_integer'
num_child(obj, dadid, momid, rel_df = NULL, missid = NA_character_)
## S4 method for signature 'Pedigree'
num_child(obj, reset = FALSE)
```

86 num\_child

#### **Arguments**

obj A character vector with the id of the individuals or a data, frame with all the informations in corresponding columns. A vector containing for each subject, the identifiers of the biologicals fathers. dadid A vector containing for each subject, the identifiers of the biologicals mothers. momid rel\_df A data frame with the special relationships between individuals. See Rel() for more informations. The minimum columns required are id1, id2 and code. The famid column can also be used to specify the family of the individuals. If a matrix is given, the columns needs to be ordered as id1, id2, code and famid. The code values are: • 1 = Monozygotic twin

• 2 = Dizygotic twin

• 3 = twin of unknown zygosity

• 4 = Spouse

The value relation code recognized by the function are the one defined by the rel\_code\_to\_factor() function.

missid A character vector with the missing values identifiers. All the id, dadid and

momid corresponding to those values will be set to NA\_character\_.

If TRUE, the num\_child\_tot, num\_child\_ind and the num\_child\_dir columns reset

are reset.

### **Details**

Compute the number of direct child but also the number of indirect child given by the ones related with the linked spouses. If a relation ship dataframe is given, then even if no children is present between 2 spouses, the indirect childs will still be added.

### Value

### When obj is a vector:

A dataframe with the columns num\_child\_dir, num\_child\_ind and num\_child\_tot giving respectively the direct, indirect and total number of child.

### When obj is a Pedigree object:

An updated Pedigree object with the columns num\_child\_dir, num\_child\_ind and num\_child\_tot added to the Pedigree ped slot.

```
num_child(
  obj = c("1", "2", "3", "4", "5", "6", "7", "8", "9", "10"),
  dadid = c("3", "3", "6", "8", "0", "0", "0", "0", "0", "0"),
  rel_df = data.frame(
     id1 = "10",
     id2 = "3",
```

parent\_of 87

```
code = "Spouse"
)
)

data(sampleped)
ped1 <- Pedigree(sampleped[sampleped$famid == "1",])
ped1 <- num_child(ped1, reset = TRUE)
summary(ped(ped1))</pre>
```

parent\_of

Get parents of individuals

# **Description**

Get the parents of individuals.

### Usage

```
## S4 method for signature 'character_OR_integer'
parent_of(obj, dadid, momid, id2)

## S4 method for signature 'Ped'
parent_of(obj, id2)

## S4 method for signature 'Pedigree'
parent_of(obj, id2)
```

#### **Arguments**

obj	A character vector with the id of the individuals or a data. frame with all the informations in corresponding columns.
dadid	A vector containing for each subject, the identifiers of the biologicals fathers.
momid	A vector containing for each subject, the identifiers of the biologicals mothers.
id2	A vector of individuals identifiers to get the parents from

# Value

A vector of individuals identifiers corresponding to the parents of the individuals in id2

```
data(sampleped)
ped <- Pedigree(sampleped)
parent_of(ped, "1_121")</pre>
```

paste0max

Print0 to max

## **Description**

Print0 the elements inside a vector until a maximum is reached.

## Usage

```
paste0max(x, max = 5, sep = "", ...)
```

### **Arguments**

x A vector.

max The maximum number of elements to print.

... Additional arguments passed to print0

#### Value

The character vector aggregated until the maximum is reached.

## **Examples**

```
x <- seq_len(10)
Pedixplorer:::paste0max(x, 5)</pre>
```

Ped-class

Ped object

## Description

S4 class to represent the identity informations of the individuals in a pedigree.

### **Constructor::**

You either need to provide a vector of the same size for each slot or a data.frame with the corresponding columns.

The metadata will correspond to the columns that do not correspond to the Ped slots.

## Usage

```
## S4 method for signature 'data.frame'
Ped(
  obj,
  cols_used_init = FALSE,
  cols_used_del = FALSE,
  date_pattern = "%Y-%m-%d"
)
## S4 method for signature 'character_OR_integer'
Ped(
  obj,
  dadid,
  momid,
  sex,
  famid = NA,
  fertility = NA,
  miscarriage = NA,
  deceased = NA,
  avail = NA,
  evaluated = NA,
  consultand = NA,
  proband = NA,
  affected = NA,
  carrier = NA,
  asymptomatic = NA,
  adopted = NA,
  dateofbirth = NA,
  dateofdeath = NA,
  missid = NA_character_,
  useful = NA,
  isinf = NA,
  kin = NA_real_
)
```

### **Arguments**

obj	A character vector with the id of the individuals or a data. frame with all the informations in corresponding columns.
cols_used_init	Boolean defining if the columns that will be used should be initialised to NA.
cols_used_del	Boolean defining if the columns that will be used should be deleted.
date_pattern	The pattern of the date
dadid	A vector containing for each subject, the identifiers of the biologicals fathers.
momid	A vector containing for each subject, the identifiers of the biologicals mothers.
sex	A character, factor or numeric vector corresponding to the gender of the individuals. This will be transformed to an ordered factor with the following levels: $male < female < unknown$

The following values are recognized:

- "male": "m", "male", "man", 1
- "female": "f", "female", "woman", 2
- "unknown": "unknown", 3

famid A character vector with the family identifiers of the individuals. If provide, will be aggregated to the individuals identifiers separated by an underscore.

A character, factor or numeric vector corresponding to the fertility status of the individuals. This will be transformed to a factor with the following levels:

infertile\_choice\_na, infertile, fertile

• "inferile choice na": "infertile choice", "infertile na"

• "infertile": "infertile", "steril", FALSE, 0

• "fertile": "fertile", TRUE, 1, NA

The following values are recognized:

Miscarriage A character, factor or numeric vector corresponding to the miscarriage status of the individuals. This will be transformed to a factor with the following levels: TOP, SAB, ECT, FALSE The following values are recognized:

• "SAB": "spontaneous", "spontaenous abortion"

• "TOP": "termination", "terminated", "termination of pregnancy"

• "ECT": "ectopic", "ectopic pregnancy"

• FALSE: 0, FALSE, "no", NA

A logical vector with the death status of the individuals (i.e. FALSE = alive, TRUE = dead, NA = unknown).

A logical vector with the availability status of the individuals (i.e. FALSE = not available, TRUE = available, NA = unknown).

A logical vector with the evaluation status of the individuals. (i.e. FALSE = documented evaluation not available, TRUE = documented evaluation available).

A logical vector with the consultand status of the individuals. A consultand being an individual seeking genetic counseling/testing (i.e. FALSE = not a consultand, TRUE = consultand).

A logical vector with the proband status of the individuals. A proband being an affected family member coming to medical attention independent of other family members. (i.e. FALSE = not a proband, TRUE = proband).

A logical vector with the affection status of the individuals (i.e. FALSE = unaffected, TRUE = affected, NA = unknown).

A logical vector with the carrier status of the individuals. A carrier being an individual who has the genetic trait but who is not likely to manifest the disease regardless of inheritance pattern (i.e. FALSE = not carrier, TRUE = carrier, NA = unknown).

A logical vector with the asymptomatic status of the individuals. An asymptomatic individual being an individual clinically unaffected at this time but could later exhibit symptoms. (i.e. FALSE = not asymptomatic, TRUE = asymptomatic, NA = unknown).

fertility

deceased

avail

evaluated

consultand

proband

affected

carrier

asymptomatic

adopted	A logical vector with the adopted status of the individuals. (i.e. FALSE = not adopted, TRUE = adopted, NA = unknown).
dateofbirth	A character vector with the date of birth of the individuals.
dateofdeath	A character vector with the date of death of the individuals.
missid	A character vector with the missing values identifiers. All the id, dadid and momid corresponding to those values will be set to NA_character
useful	A logical vector with the usefulness status of the individuals (i.e. $FALSE = not useful$ , $TRUE = useful$ ).
isinf	A logical vector indicating if the individual is informative or not (i.e. FALSE = not informative, TRUE = informative).
kin	A numeric vector with minimal kinship value between the individuals and the informative individuals.

#### **Details**

The minimal needed informations are id, dadid, momid and sex. The other slots are used to store recognized informations. Additional columns can be added to the Ped object and will be stored in the elementMetadata slot of the Ped object.

### Value

A Ped object.

#### **Slots**

id A character vector with the id of the individuals.

dadid A character vector with the id of the father of the individuals.

momid A character vector with the id of the mother of the individuals.

famid A character vector with the family identifiers of the individuals (optional).

sex An ordered factor vector for the sex of the individuals (i.e. male < female < unknown).

fertility A factor vector with the fertility status of the individuals (optional). (i.e. infertile\_choice\_na = no children by choice or unknown reason, infertile = individual is inferile, fertile = individual is fertile).

miscarriage A factor vector with the miscarriage status of the individuals (optional). (i.e. TOP = Termination of Pregnancy, SAB = Spontaneous Abortion, ECT = Ectopic Pregnancy, FALSE = no miscarriage).

deceased A logical vector with the death status of the individuals (optional). (i.e. FALSE = alive, TRUE = dead, NA = unknown).

avail A logical vector with the availability status of the individuals (optional). (i.e. FALSE = not available, TRUE = available, NA = unknown).

evaluated A logical vector with the evaluation status of the individuals (optional). (i.e. FALSE = documented evaluation not available, TRUE = documented evaluation available).

consultand A logical vector with the consultand status of the individuals (optional). A consultand being an individual seeking genetic counseling/testing (i.e. FALSE = not a consultand, TRUE = consultand).

proband A logical vector with the proband status of the individuals (optional). A proband being an affected family member coming to medical attention independent of other family members. (i.e. FALSE = not a proband, TRUE = proband).

- affected A logical vector with the affection status of the individuals (optional). (i.e. FALSE = not affected, TRUE = affected, NA = unknown).
- carrier A logical vector with the carrier status of the individuals (optional). A carrier being an individual who has the genetic trait but who is not likely to manifest the disease regardless of inheritance pattern (i.e. FALSE = not carrier, TRUE = carrier, NA = unknown).
- asymptomatic A logical vector with the asymptomatic status of the individuals (optional). An asymptomatic individual being an individual clinically unaffected at this time but could later exhibit symptoms. (i.e. FALSE = not asymptomatic, TRUE = asymptomatic, NA = unknown).
- adopted A logical vector with the adopted status of the individuals (optional). (i.e. FALSE = not adopted, TRUE = adopted, NA = unknown).

dateofbirth A date vector with the birth date of the individuals (optional).

dateofdeath A date vector with the death date of the individuals (optional).

- useful A logical vector with the usefulness status of the individuals (computed). (i.e. FALSE = not useful, TRUE = useful).
- isinf A logical vector indicating if the individual is informative or not (computed). (i.e. FALSE = not informative, TRUE = informative).
- kin A numeric vector with minimal kinship value between the individuals and the useful individuals (computed).

num\_child\_tot A numeric vector with the total number of children of the individuals (computed).

num\_child\_dir A numeric vector with the number of children of the individuals (computed).

num\_child\_ind A numeric vector with the number of children of the individuals (computed).

elementMetadata A DataFrame with the additional metadata columns of the Ped object.

metadata Meta informations about the pedigree.

## Accessors

For all the following accessors, the x parameters is a Ped object. Each getters return a vector of the same length as x with the values of the corresponding slot. For each getter, you have a setter with the same name, to be use as  $slot(x) \leftarrow value$ . The value parameter is a vector of the same length as x, except for the mcols() accessors where value is a list or a data frame with each elements with the same length as x.

- id(x): Individuals identifiers
- dadid(x): Individuals' father identifiers
- momid(x): Individuals' mother identifiers
- famid(x): Individuals' family identifiers
- sex(x): Individuals' gender
- fertility(x): Individuals' fertility status

- miscarriage(x): Individuals' miscarriage status
- deceased(x): Individuals' death status
- avail(x): Individuals' availability status
- evaluated(x): Individuals' evaluated status
- consultand(x): Individuals' consultand status
- proband(x): Individuals' proband status
- carrier(x): Individuals' carrier status
- asymptomatic(x): Individuals' asymptomatic status
- adopted(x): Individuals' adopted status
- affected(x): Individuals' affection status
- dateofbirth(x): Individuals' birth dates
- dateofdeath(x): Individuals' death dates
- isinf(x): Individuals' informativeness status
- kin(x): Individuals' kinship distance to the informative individuals
- useful(x): Individuals' usefullness status
- mcols(x): Individuals' metadata

#### Generics

- summary(x): Compute the summary of a Ped object
- show(x): Convert the Ped object to a data.frame and print it with its summary.
- as.list(x): Convert a Ped object to a list with the metadata columns at the end.
- as.data.frame(x): Convert a Ped object to a data.frame with the metadata columns at the end.
- subset(x, i, del\_parents = FALSE, keep = TRUE): Subset a Ped object based on the individuals identifiers given.
  - i : A vector of individuals identifiers to keep.
  - del\_parents : A value indicating if the parents of the individuals should be deleted.
  - keep: A logical value indicating if the individuals should be kept or deleted.

### See Also

Pedigree()

### **Examples**

```
data(sampleped)
Ped(sampleped)

Ped(
    obj = c("1", "2", "3", "4", "5", "6"),
    dadid = c("4", "4", "6", "0", "0", "0"),
    momid = c("5", "5", "5", "0", "0", "0"),
    sex = c(1, 2, 3, 1, 2, 1),
    missid = "0"
)
```

Pedigree-class

Pedigree object

## **Description**

A pedigree is a ensemble of individuals linked to each other into a family tree. A Pedigree object store the informations of the individuals and the special relationships between them. It also permit to store the informations needed to plot the pedigree (i.e. scales and hints).

#### **Constructor::**

Main constructor of the package. This constructor help to create a Pedigree object from different data. frame or a set of vectors.

If any errors are found in the data, the function will return the data.frame with the errors of the Ped object and the Rel object.

```
Pedigree(obj, ...)
## S4 method for signature 'character_OR_integer'
Pedigree(
  obj,
  dadid,
 momid,
  sex,
  famid = NA,
  fertility = NULL,
  miscarriage = NULL,
  deceased = NULL,
  avail = NULL,
  evaluated = NULL,
  consultand = NULL,
  proband = NULL,
  affections = NULL,
  carrier = NULL,
```

```
asymptomatic = NULL,
  adopted = NULL,
  dateofbirth = NULL,
  dateofdeath = NULL,
  rel_df = NULL,
 missid = NA_character_,
  col_aff = "affection",
  date_pattern = "%Y-%m-%d",
 normalize = TRUE.
)
## S4 method for signature 'data.frame'
Pedigree(
 obj = data.frame(id = character(), dadid = character(), momid = character(), famid =
   character(), sex = numeric(), fertility = numeric(), miscarriage = numeric(),
   deceased = numeric(), avail = numeric(), evaluated = logical(), consultand =
    logical(), proband = logical(), affection = logical(), carrier = logical(),
  asymptomatic = logical(), adopted = logical(), dateofbirth = character(), dateofdeath
    = character()),
 rel_df = data.frame(id1 = character(), id2 = character(), code = numeric(), famid =
    character()),
 cols_ren_ped = list(id = "indId", dadid = "fatherId", momid = "motherId", famid =
  "family", sex = "gender", fertility = c("sterilisation", "steril"), miscarriage =
  c("miscarriage", "aborted"), deceased = c("status", "dead", "vitalStatus"), avail =
    "available", evaluated = "evaluation", consultand = "consultant", proband =
    "proband", affection = "affected", carrier = "carrier", asymptomatic =
  "presymptomatic", adopted = "adoption", dateofbirth = c("dob", "birth"), dateofdeath
    = c("dod", "death")),
  cols_ren_rel = list(id1 = "indId1", id2 = "indId2", famid = "family"),
  hints = list(horder = NULL, spouse = NULL),
  normalize = TRUE,
 missid = NA_character_,
  col_aff = "affection",
  date_pattern = "%Y-%m-%d";
 na_strings = c("NA", "N/A", "None", "none", "null", "NULL"),
)
```

# Arguments

obj	A vector of the individuals identifiers or a data.frame with the individuals informations. See Ped() for more informations.
	Arguments passed on to generate_colors

dadid A vector containing for each subject, the identifiers of the biologicals fathers.

Momid A vector containing for each subject, the identifiers of the biologicals mothers.

sex

A character, factor or numeric vector corresponding to the gender of the individuals. This will be transformed to an ordered factor with the following levels: male < female < unknown

The following values are recognized:

- "male": "m", "male", "man", 1
- "female": "f", "female", "woman", 2
- "unknown": "unknown", 3

famid

A character vector with the family identifiers of the individuals. If provide, will be aggregated to the individuals identifiers separated by an underscore.

fertility

A character, factor or numeric vector corresponding to the fertility status of the individuals. This will be transformed to a factor with the following levels: infertile\_choice\_na, infertile, fertile

The following values are recognized:

- "inferile\_choice\_na" : "infertile\_choice", "infertile\_na"
- "infertile": "infertile", "steril", FALSE, 0
- "fertile": "fertile", TRUE, 1, NA

miscarriage

A character, factor or numeric vector corresponding to the miscarriage status of the individuals. This will be transformed to a factor with the following levels: TOP, SAB, ECT, FALSE The following values are recognized:

- "SAB": "spontaneous", "spontaenous abortion"
- "TOP": "termination", "terminated", "termination of pregnancy"
- "ECT": "ectopic", "ectopic pregnancy"
- FALSE: 0, FALSE, "no", NA

deceased

A logical vector with the death status of the individuals (i.e. FALSE = alive, TRUE = dead, NA = unknown).

avail

A logical vector with the availability status of the individuals (i.e. FALSE = not available, TRUE = available, NA = unknown).

evaluated

A logical vector with the evaluation status of the individuals. (i.e. FALSE = documented evaluation not available, TRUE = documented evaluation available).

consultand

A logical vector with the consultand status of the individuals. A consultand being an individual seeking genetic counseling/testing (i.e. FALSE = not a consultand, TRUE = consultand).

proband

A logical vector with the proband status of the individuals. A proband being an affected family member coming to medical attention independent of other family members. (i.e. FALSE = not a proband, TRUE = proband).

affections

A logical vector with the affections status of the individuals (i.e. FALSE = unaffected, TRUE = affected, NA = unknown). Can also be a data.frame with the same length as obj. If it is a matrix, it will be converted to a data.frame and the columns will be named after the col\_aff argument.

carrier

A logical vector with the carrier status of the individuals. A carrier being an individual who has the genetic trait but who is not likely to manifest the disease regardless of inheritance pattern (i.e. FALSE = not carrier, TRUE = carrier, NA = unknown).

asymptomatic A logical vector with the asymptomatic status of the individuals. An asymp-

tomatic individual being an individual clinically unaffected at this time but could later exhibit symptoms. (i.e. FALSE = not asymptomatic, TRUE = asymptomatic,

NA = unknown).

adopted A logical vector with the adopted status of the individuals. (i.e. FALSE = not

adopted, TRUE = adopted, NA = unknown).

dateofbirth A character vector with the date of birth of the individuals.

dateofdeath A character vector with the date of death of the individuals.

rel\_df A data.frame with the special relationships between individuals. See Rel() for more informations. The minimum columns required are id1, id2 and code.

The famid column can also be used to specify the family of the individuals. If a matrix is given, the columns needs to be ordered as id1, id2, code and famid.

The code values are:

• 1 = Monozygotic twin

• 2 = Dizygotic twin

• 3 = twin of unknown zygosity

• 4 = Spouse

The value relation code recognized by the function are the one defined by the

rel\_code\_to\_factor() function.

missid A character vector with the missing values identifiers. All the id, dadid and

momid corresponding to those values will be set to NA\_character\_.

col\_aff A character vector with the name of the column to be used for the affection

status.

date\_pattern The pattern of the date

normalize A logical to know if the data should be normalised.

cols\_ren\_ped A named list with the columns to rename for the pedigree dataframe. This is

useful if you want to use a dataframe with different column names. The names of the list should be the new column names and the values should be the old

column names. The default values are to be used with normalize = TRUE.

cols\_ren\_rel A named list with the columns to rename for the relationship matrix. This is

useful if you want to use a dataframe with different column names. The names of the list should be the new column names and the values should be the old

column names.

hints A Hints object or a named list containing horder and spouse.

na\_strings Vector of strings to be considered as NA values.

#### **Details**

If the normalization is set to TRUE, then the data will be standardized using the function norm\_ped() and norm\_rel().

If a data.frame is given, the columns names needed are as follow:

- id: the individual identifier
- · dadid: the identifier of the biological father

- momid: the identifier of the biological mother
- famid: the family identifier of the individual
- · sex: the sex of the individual
- fertility: the fertility status of the individual
- miscarriage: the miscarriage status of the individual
- · deceased: the death status of the individual
- avail: the availability status of the individual
- evaluated: the evaluation status of the individual
- consultand: the consultand status of the individual
- proband: the proband status of the individual
- affection: the affection status of the individual
- carrier: the carrier status of the individual
- asymptomatic: the asymptomatic status of the individual
- adopted: the adopted status of the individual
- dateofbirth: the date of birth of the individual
- dateofdeath: the date of death of the individual
- ...: other columns that will be stored in the elementMetadata slot

The minimum columns required are:

- id
- dadid
- momid
- sex

The famid column can also be used to specify the family of the individuals and will be merge to the id field separated by an underscore.

The columns deceased, avail, evaluated, consultand, proband, carrier, asymptomatic, adopted will be transformed with the vect\_to\_binary() function when the normalisation is selected.

The fertility column will be transformed with the fertility\_to\_factor() function.

The miscarriage column will be transformed with the miscarriage\_to\_factor() function.

The dateofbirth and dateofdeath columns will be transformed with the char\_to\_date() function.

If affections is a data.frame, col\_aff will be overwritten by the column names of the data.frame.

#### Value

A Pedigree object.

#### Slots

ped A Ped object for the identity informations. See Ped() for more informations.

rel A Rel object for the special relationships. See Rel() for more informations.

scales A Scales object for the filling and bordering colors used in the plot. See Scales() for more informations.

hints A Hints object for the ordering of the individuals in the plot. See Hints() for more informations.

#### Accessors

- ped(x, slot): Get the value of a specific slot of the Ped object
- ped(x): Get the Ped object
- ped(x, slot) <- value: Set the value of a specific slot of the Ped object Wrapper of slot(ped(x)) <- value
- ped(x) <- value : Set the Ped object
- mcols(x): Get the metadata of a Pedigree object. This function is a wrapper around mcols(ped(x)).
- mcols(x) <- value: Set the metadata of a Pedigree object. This function is a wrapper around mcols(ped(x)) <- value.</li>
- rel(x, slot): Get the value of a specific slot of the Rel object
- rel(x): Get the Rel object
- rel(x, slot) <- value: Set the value of a specific slot of the Rel object Wrapper of slot(rel(x)) <- value
- rel(x) <- value : Set the Rel object
- scales(x): Get the Scales object
- scales(x) <- value : Set the Scales object
- fill(x): Get the fill data.frame from the Scales object. Wrapper of fill(scales(x))
- fill(x) <- value: Set the fill data.frame from the Scales object. Wrapper of fill(scales(x))</li>
   value
- border(x): Get the border data.frame from the Scales object. Wrapper of border(scales(x))
- border(x) <- value: Set the border data.frame from the Scales object. Wrapper of border(scales(x)) <- value
- hints(x): Get the Hints object
- hints(x) <- value : Set the Hints object

- horder(x): Get the horder vector from the Hints object. Wrapper of horder(hints(x))
- horder(x) <- value: Set the horder vector from the Hints object. Wrapper of horder(hints(x))</li>
   value
- spouse(x): Get the spouse data.frame from the Hints object. Wrapper of spouse(hints(x)).
- spouse(x) <- value: Set the spouse data.frame from the Hints object. Wrapper of spouse(hints(x)) <- value.

#### Generics

- length(x): Get the length of a Pedigree object. Wrapper of length(ped(x)).
- show(x): Print the information of the Ped and Rel object inside the Pedigree object.
- summary(x): Compute the summary of the Ped and Rel object inside the Pedigree object.
- as.list(x): Convert a Pedigree object to a list
- subset(x, i, keep = TRUE): Subset a Pedigree object based on the individuals identifiers given.
  - i : A vector of individuals identifiers to keep.
  - del\_parents: A logical value indicating if the parents of the individuals should be deleted.
  - keep: A logical value indicating if the individuals should be kept or deleted.
- x[i, del\_parents, keep]: Subset a Pedigree object based on the individuals identifiers given.

#### See Also

```
Pedigree() Ped() Rel() Scales() Hints()
Ped() Rel() Scales()
```

```
Pedigree(
    obj = c("1", "2", "3", "4", "5", "6"),
    dadid = c("4", "4", "6", "0", "0", "0"),
    momid = c("5", "5", "5", "0", "0", "0"),
    sex = c(1, 2, 3, 1, 2, 1),
    avail = c(0, 1, 0, 1, 0, 1),
    affections = matrix(c(
        0, 1, 0, 1, 0, 1,
        1, 1, 1, 1, 1, 1
), ncol = 2),
    col_aff = c("aff1", "aff2"),
    missid = "0",
    rel_df = matrix(c(
        "1", "2", 2
```

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```
), ncol = 3, byrow = TRUE),
)
data(sampleped)
Pedigree(sampleped)
```

ped\_avaf\_infos\_ui

Shiny modules to display family information

# Description

This module allows to display the health and availability data for all individuals in a pedigree object. The output is a datatable. The function is composed of two parts: the UI and the server. The UI is called with the function ped\_avaf\_infos\_ui() and the server with the function ped\_avaf\_infos\_server().

## Usage

```
ped_avaf_infos_ui(id)
ped_avaf_infos_server(id, pedi, title = "Family informations", height = "auto")
ped_avaf_infos_demo(height = "auto")
```

# **Arguments**

id A string to identify the module.
 pedi A reactive pedigree object.
 title The title of the module.
 height The height of the datatable.

#### Value

A reactive dataframe with the selected columns renamed to the names of cols\_needed and cols\_supl.

```
if (interactive()) {
    ped_avaf_infos_demo()
}
```

ped\_shiny

ned	serve	r

Create the server logic for the ped\_shiny application

# Description

Create the server logic for the ped\_shiny application

# Usage

```
ped_server(input, output, session, precision = 6)
```

# Arguments

input The input object from a Shiny app.
output The output object from a Shiny app.
session The session object from a Shiny app.

precision Number of decimal for the position of the boxes in the plot.

#### Value

```
shiny::shinyServer()
```

## **Examples**

```
if (interactive()) {
    ped_shiny()
}
```

ped\_shiny

Run Pedixplorer Shiny application

# Description

This function creates a shiny application to manage and visualize pedigree data using the ped\_ui() and ped\_server() functions.

```
ped_shiny(
  port = getOption("shiny.port"),
  host = getOption("shiny.host", "127.0.0.1"),
  precision = 6
)
```

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## **Arguments**

port (optional) Specify port the application should list to.

host (optional) The IPv4 address that the application should listen on.

precision Number of decimal for the position of the boxes in the plot.

## **Details**

The application is composed of several modules:

- Data import
- Data column selection
- · Data download
- · Family selection
- · Health selection
- Informative selection
- Subfamily selection
- Plotting pedigree
- Family information

### Value

**Running Shiny Application** 

## **Examples**

```
if (interactive()) {
    ped_shiny()
}
```

ped\_to\_legdf

Create plotting legend data frame from a Pedigree

### **Description**

Convert a Pedigree to a legend data frame for it to be plotted afterwards with plot\_fromdf().

```
## S4 method for signature 'Pedigree'
ped_to_legdf(
   obj,
   boxh = 1,
   boxw = 1,
   cex = 1,
   adjx = 0,
```

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```
adjy = 0,
lwd = par("lwd"),
precision = 4
)
```

## **Arguments**

obj	A Pedigree object
boxh	Height of the polygons elements
boxw	Width of the polygons elements
cex	Character expansion of the text
adjx	default=0. Controls the horizontal text adjustment of the labels in the legend.
adjy	default=0. Controls the vertical text adjustment of the labels in the legend.
lwd	default=par("lwd"). Controls the bordering line width of the elements in the legend.
precision	The number of significatif numbers to round the numbers to.

## **Details**

The data frame contains the following columns:

- x0, y0, x1, y1: coordinates of the elements
- type: type of the elements
- fill: fill color of the elements
- border: border color of the elements
- angle: angle of the shading of the elements
- density: density of the shading of the elements
- cex: size of the elements
- label: label of the elements
- tips: tips of the elements (used for the tooltips)
- adjx: horizontal text adjustment of the labels
- adjy: vertical text adjustment of the labels

All those columns are used by plot\_fromdf() to plot the graph.

### Value

A list containing the legend data frame and the user coordinates.

```
data("sampleped")
pedi <- Pedigree(sampleped)
leg_df <- ped_to_legdf(pedi)
summary(leg_df$df)
plot_fromdf(leg_df$df, usr = c(-1,15,0,7))</pre>
```

ped\_to\_plotdf

ped\_to\_plotdf

Create plotting data frame from a Pedigree

## Description

Convert a Pedigree to a data frame with all the elements and their characteristic for them to be plotted afterwards with plot\_fromdf().

# Usage

```
## S4 method for signature 'Pedigree'
ped_to_plotdf(
  obj,
  packed = TRUE,
 width = 6,
  align = c(1.5, 2),
  align_parents = TRUE,
  force = FALSE,
  cex = 1,
  symbolsize = cex,
  pconnect = 0.5,
  branch = 0.6,
  aff_mark = TRUE,
  id_lab = "id",
  label = NULL,
  precision = 4,
  lwd = par("lwd"),
  tips = NULL,
)
```

## **Arguments**

obj	A Pedigree object
	Other arguments passed to par()
packed	Should the Pedigree be compressed. (i.e. allow diagonal lines connecting parents to children in order to have a smaller overall width for the plot.)
width	For a packed output, the minimum width of the plot, in inches.
align	For a packed Pedigree, align children under parents TRUE, to the extent possible given the page width, or align to to the left margin FALSE. This argument can be a two element vector, giving the alignment parameters, or a logical value. If TRUE, the default is c(1.5, 2), or if numeric the routine alignped4() will be called.
align_parents	If align_parents = TRUE, go one step further and try to make both parents of each child have the same depth. (This is not always possible). It helps the drawing program by lining up pedigrees that 'join in the middle' via a marriage.

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If force = TRUE, the function will return the depth minus min(depth) if depth force

reach a state with no founders is not possible.

Character expansion of the text cex

Size of the symbols symbolsize

When connecting parent to children the program will try to make the connecting pconnect

> line as close to vertical as possible, subject to it lying inside the endpoints of the line that connects the children by at least pconnect people. Setting this option to a large number will force the line to connect at the midpoint of the children.

branch defines how much angle is used to connect various levels of nuclear families.

aff\_mark If TRUE, add a aff\_mark to each box corresponding to the value of the affection

column for each filling scale.

id\_lab The column name of the id for each individuals.

label If not NULL, add a label to each box under the id corresponding to the value of

the column given.

The number of significatif numbers to round the solution to. precision

lwd default=par("lwd"). Controls the line width of the segments, arcs and polygons. tips

A character vector of the column names of the data frame to use as tooltips. If

NULL, no tooltips are added.

#### **Details**

The data frame contains the following columns:

• x0, y0, x1, y1: coordinates of the elements

• type: type of the elements

• fill: fill color of the elements

• border: border color of the elements

• angle: angle of the shading of the elements

• density: density of the shading of the elements

• cex: size of the elements

• label: label of the elements

• tips: tips of the elements (used for the tooltips)

• adjx: horizontal text adjustment of the labels

• adjy: vertical text adjustment of the labels

All those columns are used by plot\_fromdf() to plot the graph.

#### Value

A list containing the data frame and the user coordinates.

### See Also

```
plot_fromdf() ped_to_legdf()
```

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# **Examples**

ped\_ui

Create the user interface for the ped\_shiny application

## **Description**

Create the user interface for the ped\_shiny application

### Value

```
shiny::shinyUI()
```

## **Examples**

```
if (interactive()) {
    ped_shiny()
}
```

permute

Generate all possible permutation

## Description

Given a vector of length  $\mathbf{n}$ , generate all possible permutations of the numbers 1 to  $\mathbf{n}$ . This is a recursive routine, and is not very efficient.

### Usage

```
permute(x)
```

### **Arguments**

Х

A vector of length n

#### Value

A matrix with **n** cols and **n!** rows

```
Pedixplorer:::permute(seq_len(3))
```

```
{\tt plot,Pedigree,missing-method} \\ {\tt Plot\ Pedigrees}
```

## **Description**

This function is used to plot a Pedigree object.

It is a wrapper for plot\_fromdf() and ped\_to\_plotdf() as well as ped\_to\_legdf() if legend = TRUE.

```
## S4 method for signature 'Pedigree, missing'
plot(
  Х,
  aff_mark = TRUE,
  id_lab = "id",
  label = NULL,
  ggplot_gen = FALSE,
  cex = 1,
  symbolsize = 1,
  branch = 0.6,
  packed = TRUE,
  align = c(1.5, 2),
  align_parents = TRUE,
  force = FALSE,
 width = 6,
  title = NULL,
  subreg = NULL,
  pconnect = 0.5,
  fam_to_plot = 1,
  legend = FALSE,
  leg_cex = 0.8,
  leg_symbolsize = 0.5,
  leg_loc = NULL,
  leg_adjx = 0,
  leg_adjy = 0,
  precision = 4,
  lwd = par("lwd"),
  ped_par = list(),
  leg_par = list(),
  tips = NULL,
  title_cex = 2,
  leg_usr = NULL,
  add_{to}=FALSE
)
```

#### **Arguments**

packed

x A Pedigree object.

aff\_mark If TRUE, add a aff\_mark to each box corresponding to the value of the affection

column for each filling scale.

id\_lab The column name of the id for each individuals.

label If not NULL, add a label to each box under the id corresponding to the value of

the column given.

ggplot\_gen If TRUE add the segments to the ggplot object

cex Character expansion of the text

symbolsize Size of the symbols

branch defines how much angle is used to connect various levels of nuclear families.

Should the Pedigree be compressed. (i.e. allow diagonal lines connecting par-

ents to children in order to have a smaller overall width for the plot.)

align For a packed Pedigree, align children under parents TRUE, to the extent possible

given the page width, or align to to the left margin FALSE. This argument can be a two element vector, giving the alignment parameters, or a logical value. If TRUE, the default is c(1.5, 2), or if numeric the routine alignped4() will be

called.

align\_parents If align\_parents = TRUE, go one step further and try to make both parents of

each child have the same depth. (This is not always possible). It helps the drawing program by lining up pedigrees that 'join in the middle' via a marriage.

force If force = TRUE, the function will return the depth minus min(depth) if depth

reach a state with no founders is not possible.

width For a packed output, the minimum width of the plot, in inches.

title The title of the plot.

subreg A 4-element vector for (min x, max x, min depth, max depth), used to edit away

portions of the plot coordinates returned by ped\_to\_plotdf(). This is useful

for zooming in on a particular region of the Pedigree.

pconnect When connecting parent to children the program will try to make the connecting

line as close to vertical as possible, subject to it lying inside the endpoints of the line that connects the children by at least pronnect people. Setting this option to a large number will force the line to connect at the midpoint of the children.

fam\_to\_plot default=1. If the Pedigree contains multiple families, this parameter can be used

to select which family to plot. It can be a numeric value or a character value. If numeric, it is the index of the family to plot returned by unique(x\$ped\$famid).

If character, it is the family id to plot.

legend default=FALSE. If TRUE, a legend will be added to the plot.

leg\_cex default=0.8. Controls the size of the legend text.
leg\_symbolsize default=0.5. Controls the size of the legend symbols.

leg\_loc default=NULL. If NULL, the legend will be placed in the upper right corner of

the plot. Otherwise, a 4-element vector of the form (x0, x1, y0, y1) can be used to specify the location of the legend. The legend will be fitted to the specified and might be distorted if the aspect ratio of the legend is different from the aspect

ratio of the specified location.

leg_adjx	default=0. Controls the horizontal labels adjustment of the legend.
leg_adjy	default=0. Controls the vertical labels adjustment of the legend.
precision	The number of significatif numbers to round the solution to.
lwd	default=par("lwd"). Controls the line width of the segments, arcs and polygons.
ped_par	default=list(). A list of parameters to use as graphical parameteres for the main plot.
leg_par	default=list(). A list of parameters to use as graphical parameters for the legend.
tips	A character vector of the column names of the data frame to use as tooltips. If NULL, no tooltips are added.
title_cex	The size of the title.
leg_usr	default=NULL. A vector of user coordinates to use for the legend.

If TRUE, the plot will be added to the current plot.

#### **Details**

add\_to\_existing

Two important parameters control the looks of the result. One is the user specified maximum width. The smallest possible width is the maximum number of subjects on a line, if the user's suggestion is too low it is increased to 1 + that amount (to give just a little wiggle room).

To make a Pedigree where all children are centered under parents simply make the width large enough, however, the symbols may get very small.

The second is align, a vector of 2 alignment parameters a and b. For each set of siblings at a set of locations x and with parents at p=c(p1,p2) the alignment penalty is

$$(1/k^{a})\sum_{i} i = 1k[(x_{i} - (p1 + p2)/2)]^{2}$$
$$\sum_{i} (x - \overline{(p)})^{2}/(k^{a})$$

Where k is the number of siblings in the set.

When a = 1 moving a sibship with k sibs one unit to the left or right of optimal will incur the same cost as moving one with only 1 or two sibs out of place.

If a = 0 then large sibships are harder to move than small ones, with the default value a = 1.5 they are slightly easier to move than small ones. The rationale for the default is as long as the parents are somewhere between the first and last siblings the result looks fairly good, so we are more flexible with the spacing of a large family. By tethering all the sibs to a single spot they are kept close to each other. The alignment penalty for spouses is  $b(x_1 - x_2)^2$ , which tends to keep them together. The size of b controls the relative importance of sib-parent and spouse-spouse closeness.

## Value

an invisible list containing

- df: the data.frame used to plot the Pedigree
- par usr: the user coordinates used to plot the Pedigree
- ggplot : the ggplot object if ggplot\_gen = TRUE

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## **Side Effects**

Creates plot on current plotting device.

#### See Also

```
Pedigree()
```

## **Examples**

```
data(sampleped)
pedAll <- Pedigree(sampleped)
if (interactive()) { plot(pedAll) }</pre>
```

plot\_download\_ui

Shiny module to export plot

## **Description**

This module allow to export multiple type of plot from a reactive object. The file type currently supported are png, pdf and html. The function is composed of two parts: the UI and the server. The UI is called with the function plot\_download\_ui() and the server with the function plot\_download\_server().

# Usage

```
plot_download_ui(id)

plot_download_server(
   id,
   my_plot,
   filename = "saveplot",
   label = "Download",
   width = 500,
   height = 500,
   ext = "png"
)
```

## **Arguments**

id A string.

my\_plot Reactive object containing the plot.

filename A string to name the file.

label A string to name the download button.

plot\_fromdf

width	A numeric to set the width of the plot.
height	A numeric to set the height of the plot.
ext	A string to set the extension of the file.

## Value

A shiny module to export a plot.

# **Examples**

```
if (interactive()) {
    plot_download_demo()
}
```

plot\_fromdf

Create a plot from a data.frame

# Description

This function is used to create a plot from a data.frame.

If ggplot\_gen = TRUE, the plot will be generated with ggplot2 and will be returned invisibly.

# Usage

```
plot_fromdf(
    df,
    usr = NULL,
    title = NULL,
    ggplot_gen = FALSE,
    boxw = 1,
    boxh = 1,
    add_to_existing = FALSE,
    title_cex = 2
)
```

## **Arguments**

df

A data.frame with the following columns:

- type: The type of element to plot. Can be text, segments, arc or other polygons. For polygons, the name of the polygon must be in the form poly\_\*\_\* where poly is one of the type given by polygons(), the first \* is the number of slice in the polygon and the second \* is the position of the division of the polygon.
- x0: The x coordinate of the center of the element.
- y0: The y coordinate of the center of the element.

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- x1: The x coordinate of the end of the element. Only used for segments and arc.
- y1: The y coordinate of the end of the element. Only used for segments and arc.
- fill: The fill color of the element.
- border: The border color of the element.
- density: The density of the element.
- angle: The angle of the element.
- label: The label of the element. Only used for text.
- cex: The size of the element.
- adjx: The x adjustment of the element. Only used for text.
- adjy: The y adjustment of the element. Only used for text.

usr The user coordinates of the plot.

title The title of the plot.

boxw Width of the polygons elements
boxh Height of the polygons elements

add\_to\_existing

If TRUE, the plot will be added to the current plot.

title\_cex The size of the title.

#### Value

an invisible ggplot object and a plot on the current plotting device

# **Examples**

plot\_legend

Plot legend

## Description

Small internal function to be used for plotting a Pedigree object legend

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## Usage

```
plot_legend(
  obj,
  cex = 1,
  boxw = 0.1,
  boxh = 0.1,
  adjx = 0,
  adjy = 0,
  leg_loc = c(0, 1, 0, 1),
  add_to_existing = FALSE,
  usr = NULL,
  lwd = par("lwd"),
  precision = 4
)
```

# Arguments

obj	A Pedigree object
cex	Character expansion of the text
boxw	Width of the polygons elements
boxh	Height of the polygons elements
adjx	default=0. Controls the horizontal text adjustment of the labels in the legend.
adjy	default=0. Controls the vertical text adjustment of the labels in the legend.
lwd	default=par("lwd"). Controls the bordering line width of the elements in the legend.
precision	The number of significatif numbers to round the numbers to.

## Value

an invisible list containing

- df: the data.frame used to plot the Pedigree
- par\_usr: the user coordinates used to plot the Pedigree

# **Side Effects**

Creates plot on current plotting device.

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plot\_legend\_ui

Shiny module to generate pedigree graph legend.

## **Description**

This module allows to plot the legend of a pedigree object. The function is composed of two parts: the UI and the server. The UI is called with the function plot\_legend\_ui() and the server with the function plot\_legend\_server().

# Usage

```
plot_legend_ui(id, height = "400px")

plot_legend_server(
   id,
   pedi,
   leg_loc = c(0, 1, 0, 1),
   lwd = par("lwd"),
   boxw = 0.1,
   boxh = 0.1,
   adjx = 0,
   adjy = 0
)

plot_legend_demo(height = "400px", leg_loc = c(0.2, 0.8, 0.2, 0.6))
```

# Arguments

id	A string.
pedi	A reactive pedigree object.
lwd	default=par("lwd"). Controls the bordering line width of the elements in the legend.
boxw	Width of the polygons elements
boxh	Height of the polygons elements
adjx	default=0. Controls the horizontal text adjustment of the labels in the legend.
adjy	default=0. Controls the vertical text adjustment of the labels in the legend.

## Value

A static UI with the legend.

```
if (interactive()) {
    plot_legend_demo()
}
```

plot\_ped\_ui

plot\_ped\_ui

Shiny module to generate pedigree graph.

## **Description**

This module allows to plot a pedigree object. The plot can be interactive. The function is composed of two parts: the UI and the server. The UI is called with the function plot\_ped\_ui() and the server with the function plot\_ped\_server().

## Usage

```
plot_ped_ui(id)

plot_ped_server(
   id,
   pedi,
   title = NA,
   precision = 2,
   max_ind = 500,
   tips = NULL,
   lwd = par("lwd")
)

plot_ped_demo(pedi, precision = 2, max_ind = 500, tips = NULL)
```

## **Arguments**

id A string.

pedi A reactive pedigree object. title A string to name the plot.

precision An integer to set the precision of the plot.

max\_ind An integer to set the maximum number of individuals to plot.

tips A character vector of the column names of the data frame to use as tooltips. If

NULL, no tooltips are added.

1wd A numeric to set the line width of the plot.

## Value

A reactive ggplot or the pedigree object.

```
if (interactive()) {
    data("sampleped")
    pedi <- shiny::reactive({
        Pedigree(sampleped[sampleped$famid == "1", ])</pre>
```

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```
})
plot_ped_demo(pedi)
}
```

polyfun

Polygonal element

## **Description**

Create a list of x and y coordinates for a polygon with a given number of slices and a list of coordinates for the polygon.

# Usage

```
polyfun(nslice, coor, start = 90)
```

## **Arguments**

nslice Number of slices in the polygon

coor Element form which to generate the polygon containing x and y coordinates

start Starting angle in degree

#### Value

a list of x and y coordinates

## **Examples**

```
Pedixplorer:::polyfun(2, list(
    x = c(-0.5, -0.5, 0.5, 0.5),
    y = c(-0.5, 0.5, 0.5, -0.5)
), start = 45)
```

polygons

List of polygonal elements

# Description

Create a list of polygonal elements with x, y coordinates and theta for the square, circle, diamond and triangle. The number of slices in each element can be specified.

# Usage

```
polygons(nslice = 1, start = 90)
```

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## **Arguments**

nslice Number of slices in each element If nslice > 1, the elements are created with

polyfun().

## Value

a list of polygonal elements with x, y coordinates and theta by slice.

# **Examples**

```
Pedixplorer:::polygons()
Pedixplorer:::polygons(4)
```

read\_data

Read data from file path

## **Description**

Read dataframe based on the extension of the file

## Usage

```
read_data(
   file,
   sep = ";",
   quote = "'",
   header = TRUE,
   df_name = NA,
   strings_as_factors = FALSE,
   to_char = TRUE,
   na_values = c("", "NA", "NULL", "None")
)
```

# **Arguments**

file The file path

sep A string defining the separator to use for the file

quote A string defining the quote to use

header A boolean defining if the dataframe contain a header or not df\_name A string defining the name of the dataframe / sheet to use

strings\_as\_factors

A boolean defining if all the strings should be interpreted ad factor

to\_char A boolean defining if all the dataset should be read as character.

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## **Details**

This function detect the extension of the file and proceed to use the according function to read it with the parameters given by the user.

### Value

A dataframe.

## **Examples**

```
## Not run:
    read_data('path/to/my/file.txt', sep=',', header=FALSE)
## End(Not run)
```

Rel-class

Rel object

## **Description**

S4 class to represent the special relationships in a Pedigree.

## **Constructor::**

You either need to provide a vector of the same size for each slot or a data.frame with the corresponding columns.

## Usage

```
## S4 method for signature 'data.frame'
Rel(obj)
## S4 method for signature 'character_OR_integer'
Rel(obj, id2, code, famid = NA_character_)
```

# Arguments

obj	A character vector with the id of the first individuals of each pairs or a data. frame with all the informations in corresponding columns.
id2	A character vector with the id of the second individuals of each pairs
code	A character, factor or numeric vector corresponding to the relation code of the

individuals:MZ twin = Monozygotic twin

- DZ twin = Dizygotic twin
- UZ twin = twin of unknown zygosity
- Spouse = Spouse The following values are recognized:

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- character() or factor(): "MZ twin", "DZ twin", "UZ twin", "Spouse" with of without space between the words. The case is not important.
- numeric(): 1 = "MZ twin", 2 = "DZ twin", 3 = "UZ twin", 4 = "Spouse"

famid

A character vector with the family identifiers of the individuals. If provide, will be aggregated to the individuals identifiers separated by an underscore.

#### **Details**

A Rel object is a list of special relationships between individuals in the pedigree. It is used to create a Pedigree object. The minimal needed informations are id1, id2 and code.

If a famid is provided, the individuals id will be aggregated to the famid character to ensure the uniqueness of the id.

#### Value

A Rel object.

## **Slots**

id1 A character vector with the id of the first individual.

id2 A character vector with the id of the second individual.

code An ordered factor vector with the code of the special relationship.

(i.e. MZ twin < DZ twin < UZ twin < Spouse).

famid A character vector with the famid of the individuals.

#### Accessors

For all the following accessors, the x parameters is a Rel object. Each getters return a vector of the same length as x with the values of the corresponding slot.

- code(x): Relationships' code
- id1(x): Relationships' first individuals' identifier
- id2(x): Relationships' second individuals' identifier
- famid(x): Relationships' individuals' family identifier
- famid(x) <- value : Set the relationships' individuals' family identifier
  - value : A character or integer vector of the same length as x with the family identifiers

#### Generics

- summary (x): Compute the summary of a Rel object
- show(x): Convert the Rel object to a data.frame and print it with its summary.
- as.list(x): Convert a Rel object to a list

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- as.data.frame(x): Convert a Rel object to a data.frame
- subset(x, i, keep = TRUE): Subset a Rel object based on the individuals identifiers given.
  - i : A vector of individuals identifiers to keep.
  - keep: A logical value indicating if the individuals should be kept or deleted.

## See Also

```
Pedigree()
```

## **Examples**

```
rel_df <- data.frame(
    id1 = c("1", "2", "3"),
    id2 = c("2", "3", "4"),
    code = c(1, 2, 3)
)
Rel(rel_df)

Rel(
    obj = c("1", "2", "3"),
    id2 = c("2", "3", "4"),
    code = c(1, 2, 3)
)</pre>
```

relped

Relped data

# Description

Small set of related individuals for testing purposes.

## Usage

```
data("relped")
```

#### **Format**

The dataframe is composed of 4 columns:

- id1: the first individual identifier,
- id2: the second individual identifier,
- code: the relationship between the two individuals,
- famid: the family identifier. The relationship codes are:
- 1 for Monozygotic twin
- 2 for Dizygotic twin
- 3 for Twin of unknown zygosity
- 4 for Spouse relationship

rel\_code\_to\_factor

#### **Details**

This is a small fictive data set of relation that accompanies the sampleped data set. The aim was to create a data set with a variety of relationships. There is 8 relations with 4 different types of relationships.

# Examples

```
data("relped")
data("sampleped")
pedi <- Pedigree(sampleped, relped)
summary(pedi)
if (interactive()) { plot(pedi) }</pre>
```

rel\_code\_to\_factor

Relationship code variable to ordered factor

## **Description**

Relationship code variable to ordered factor

#### **Usage**

```
rel_code_to_factor(code)
```

## **Arguments**

code

A character, factor or numeric vector corresponding to the relation code of the individuals:

- MZ twin = Monozygotic twin
- DZ twin = Dizygotic twin
- UZ twin = twin of unknown zygosity
- Spouse = Spouse The following values are recognized:
- character() or factor(): "MZ twin", "DZ twin", "UZ twin", "Spouse" with of without space between the words. The case is not important.
- numeric(): 1 = "MZ twin", 2 = "DZ twin", 3 = "UZ twin", 4 = "Spouse"

#### Value

an ordered factor vector containing the transformed variable "MZ twin" < "DZ twin" < "UZ twin" < "Spouse"

```
rel_code_to_factor(c(1, 2, 3, 4, "MZ twin", "DZ twin", "UZ twin", "Spouse"))
```

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sampleped

Sampleped data

## **Description**

Small sample pedigree data set for testing purposes.

## Usage

```
data("sampleped")
```

## **Format**

A data frame with 55 observations, one line per subject, on the following 7 variables.

- famid: Family identifier
- id: Subject identifier
- dadid: Identifier of the father, if the father is part of the data set; zero otherwise
- momid: Identifier of the mother, if the mother is part of the data set; zero otherwise
- sex: 1 for male or 2 for female
- affection: 1 or 0
- avail: 1 or 0
- num: Numerical test variable from 0 to 6 randomly distributed

# **Details**

This is a small fictive pedigree data set, with 55 individuals in 2 families. The aim was to create a data set with a variety of pedigree structures.

```
data("sampleped")
pedi <- Pedigree(sampleped)
summary(pedi)
if (interactive()) { plot(pedi) }</pre>
```

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Scales-class

Scales object

## **Description**

A Scales object is a list of two data frame. The first one is used to represent the affection status of the individuals and therefore the filling of the individuals in the pedigree plot. The second one is used to represent the availability status of the individuals and therefore the border color of the individuals in the pedigree plot.

#### **Constructor::**

You need to provide both **fill** and **border** in the dedicated parameters. However this is usually done using the <code>generate\_colors()</code> function with a Pedigree object.

# Usage

```
Scales(fill, border)
## S4 method for signature 'data.frame,data.frame'
Scales(fill, border)
```

#### **Arguments**

fill

A data.frame with the informations for the affection status. The columns needed are:

- 'order': the order of the affection to be used
- 'column\_values': name of the column containing the raw values in the Ped object
- 'column\_mods': name of the column containing the mods of the transformed values in the Ped object
- 'mods': all the different mods
- 'labels': the corresponding labels of each mods
- 'affected': a logical value indicating if the mod correspond to an affected individuals
- 'fill': the color to use for this mods
- 'density': the density of the shading
- 'angle': the angle of the shading

border

A data frame with the informations for the availability status. The columns needed are:

- 'column\_values': name of the column containing the raw values in the Ped object
- 'column\_mods': name of the column containing the mods of the transformed values in the Ped object
- 'mods': all the different mods
- 'labels': the corresponding labels of each mods
- 'border': the color to use for this mods

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#### Value

A Scales object.

#### Slots

fill A data frame with the informations for the affection status. The columns needed are:

- 'order': the order of the affection to be used
- 'column\_values': name of the column containing the raw values in the Ped object
- 'column\_mods': name of the column containing the mods of the transformed values in the Ped object
- 'mods': all the different mods
- 'labels': the corresponding labels of each mods
- 'affected': a logical value indicating if the mod correspond to an affected individuals
- 'fill': the color to use for this mods
- 'density': the density of the shading
- 'angle': the angle of the shading

border A data.frame with the informations for the availability status. The columns needed are:

- 'column\_values': name of the column containing the raw values in the Ped object
- 'column\_mods': name of the column containing the mods of the transformed values in the Ped object
- 'mods': all the different mods
- 'labels': the corresponding labels of each mods
- 'border': the color to use for this mods

## Accessors

- fill(x): Get the fill data.frame
- $fill(x) \leftarrow value : Set the fill data.frame$
- border(x): Get the border data.frame
- border(x) <- value : Set the border data.frame from the Scales object.

#### Generics

• as.list(x): Convert a Scales object to a list

# See Also

```
Pedigree()
generate_colors()
```

set\_plot\_area

## **Examples**

```
Scales(
   fill = data.frame(
       order = 1,
       column_values = "affected",
       column_mods = "affected_mods",
       mods = c(0, 1),
       labels = c("unaffected", "affected"),
       affected = c(FALSE, TRUE),
       fill = c("white", "red"),
       density = c(NA, 20),
       angle = c(NA, 45)
   ),
   border = data.frame(
       column_values = "avail",
       column_mods = "avail_mods",
       mods = c(0, 1),
       labels = c("not available", "available"),
       border = c("black", "blue")
   )
)
```

set\_plot\_area

Set plotting area

# Description

Set plotting area

## Usage

```
set_plot_area(cex, id, maxlev, xrange, symbolsize, precision = 4, ...)
```

## **Arguments**

cex	Character expansion of the text
id	A character vector with the identifiers of each individuals
maxlev	Maximum level
xrange	Range of x values
symbolsize	Size of the symbols
precision	The number of significant digits to round the solution to.
	Other arguments passed to par()

#### Value

List of user coordinates, old par, box width, box height, label height and leg height

sex\_to\_factor 127

sex\_to\_factor

Gender variable to ordered factor

## **Description**

Gender variable to ordered factor

## Usage

```
sex_to_factor(sex)
```

## **Arguments**

sex

A character, factor or numeric vector corresponding to the gender of the individuals. This will be transformed to an ordered factor with the following levels: male < female < unknown

The following values are recognized:

- "male": "m", "male", "man", 1
- "female": "f", "female", "woman", 2
- "unknown": "unknown", 3

## Value

an ordered factor vector containing the transformed variable "male" < "female" < "unknown"

## **Examples**

```
sex_to_factor(c(1, 2, 3, "f", "m", "man", "female"))
```

shift

Shift set of siblings to the left or right

# Description

Shift set of siblings to the left or right

## Usage

```
shift(id, sibs, goleft, hint, twinrel, twinset)
```

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## **Arguments**

twinrel

id The id of the subject to be shifted

sibs The ids of the siblings

If TRUE, shift to the left, otherwise to the right goleft

hint The current hint vector

The twin relationship matrix twinset The twinset vector

#### **Details**

This routine is used by auto\_hint(). It shifts a set of siblings to the left or right, so that the marriage is on the edge of the set of siblings, closest to the spouse. It also shifts the subject himself, so that he is on the edge of the set of siblings, closest to the spouse. It also shifts the monozygotic twins, if any, so that they are together within the set of twins.

#### Value

The updated hint vector

## See Also

```
auto_hint()
```

shrink

Shrink Pedigree object

# **Description**

Shrink Pedigree object to specified bit size with priority placed on trimming uninformative subjects. The algorithm is useful for getting a Pedigree condensed to a minimally informative size for algorithms or testing that are limited by size of the Pedigree.

If avail or affected are NULL, they are extracted with their corresponding accessors from the Ped object.

## Usage

```
## S4 method for signature 'Pedigree'
shrink(obj, avail = NULL, affected = NULL, max_bits = 16)
## S4 method for signature 'Ped'
shrink(obj, avail = NULL, affected = NULL, max_bits = 16)
```

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## **Arguments**

obj	A Pedigree or Ped object.
avail	A logical vector with the availability status of the individuals (i.e. $FALSE = not$ available, $TRUE = available$ , $NA = unknown$ ).
affected	A logical vector with the affection status of the individuals (i.e. FALSE = unaffected, TRUE = affected, NA = unknown).
max_bits	Optional, the bit size for which to shrink the Pedigree

#### **Details**

Iteratively remove subjects from the Pedigree. The random removal of members was previously controlled by a seed argument, but we remove this, forcing users to control randomness outside the function. First remove uninformative subjects, i.e., unavailable (not genotyped) with no available descendants. Next, available terminal subjects with unknown phenotype if both parents available. Last, iteratively shrinks Pedigrees by preferentially removing individuals (chosen at random if there are multiple of the same status):

- 1. Subjects with unknown affected status
- 2. Subjects with unaffected affected status
- 3. Affected subjects.

#### Value

A list containing the following elements:

- pedObj: Pedigree object after trimming
- id\_trim: Vector of ids trimmed from Pedigree
- id\_lst: List of ids trimmed by category
- bit\_size: Vector of bit sizes after each trimming step
- avail: Vector of availability status after trimming
- pedSizeOriginal: Number of subjects in original Pedigree
- pedSizeIntermed: Number of subjects after initial trimming
- pedSizeFinal: Number of subjects after final trimming

## Author(s)

Original by Dan Schaid, updated by Jason Sinnwell and Louis Le Nezet

#### See Also

```
Pedigree(), bit_size()
```

```
data(sampleped)
ped1 <- Pedigree(sampleped[sampleped$famid == '1',])
shrink(ped1, max_bits = 12)</pre>
```

subregion

sketch\_family\_table

Sketch of the family information table

# Description

Simple function to create a sketch of the family information table.

## Usage

```
sketch_family_table(var_name)
```

## **Arguments**

var\_name

the name of the health variable

## Value

An html sketch of the family information table

subregion

Subset a region of a Pedigree

# Description

Subset a region of a Pedigree

## Usage

```
subregion(df, subreg = NULL)
```

# Arguments

df A data frame with all the plot coordinates

subreg A 4-element vector for (min x, max x, min depth, max depth), used to edit away

portions of the plot coordinates returned by ped\_to\_plotdf(). This is useful

for zooming in on a particular region of the Pedigree.

## Value

A subset of the plot coordinates

unrelated 131

unrelated

Find Unrelated subjects

## **Description**

Determine set of maximum number of unrelated available subjects from a Pedigree.

## Usage

```
## S4 method for signature 'Ped'
unrelated(obj, avail = NULL)
## S4 method for signature 'Pedigree'
unrelated(obj, avail = NULL)
```

#### **Arguments**

obj A Pedigree or Ped object.

avail A logical vector with the availability status of the individuals (i.e. FALSE = not

available, TRUE = available, NA = unknown).

## **Details**

Determine set of maximum number of unrelated available subjects from a Pedigree, given vectors id, father, and mother for a Pedigree structure, and status vector of TRUE / FALSE for whether each subject is available (e.g. has DNA).

This is a greedy algorithm that uses the kinship matrix, sequentially removing rows/cols that are non-zero for subjects that have the most number of zero kinship coefficients (greedy by choosing a row of kinship matrix that has the most number of zeros, and then remove any cols and their corresponding rows that are non-zero. To account for ties of the count of zeros for rows, a random choice is made. Hence, running this function multiple times can return different sets of unrelated subjects.

If avail is NULL, it is extracted with its corresponding accessor from the Ped object.

#### Value

A vector of the ids of subjects that are unrelated.

## Author(s)

Dan Schaid and Shannon McDonnell updated by Jason Sinnwell

132 upd\_famid

## **Examples**

```
data(sampleped)
fam1 <- sampleped[sampleped$famid == 1, -16]
ped1 <- Pedigree(fam1)
unrelated(ped1)
## some possible vectors
## [1] '110' '113' '133' '109'
## [1] '113' '118' '141' '109'
## [1] '113' '118' '140' '109'
## [1] '110' '113' '116' '109'
## [1] '113' '133' '141' '109'</pre>
```

upd\_famid

Update family prefix in individuals id

# Description

Update the family prefix in the individuals identifiers. Individuals identifiers are constructed as follow **famid\_id**. Therefore to update their family prefix the ids are split by the first underscore and the first part is overwritten by **famid**.

# Usage

```
## S4 method for signature 'character,ANY'
upd_famid(obj, famid, missid = NA_character_)

## S4 method for signature 'Ped,character_OR_integer'
upd_famid(obj, famid)

## S4 method for signature 'Ped,missing'
upd_famid(obj)

## S4 method for signature 'Rel,character_OR_integer'
upd_famid(obj, famid)

## S4 method for signature 'Rel,missing'
upd_famid(obj)

## S4 method for signature 'Pedigree,character_OR_integer'
upd_famid(obj)

## S4 method for signature 'Pedigree,character_OR_integer'
upd_famid(obj, famid)

## S4 method for signature 'Pedigree,missing'
upd_famid(obj)
```

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# **Arguments**

obj	Ped or Pedigree object or a character vector of individual ids
famid	A character vector with the family identifiers of the individuals. If provide, will be aggregated to the individuals identifiers separated by an underscore.
missid	A character vector with the missing values identifiers. All the id, dadid and momid corresponding to those values will be set to NA_character

## **Details**

If famid is missing, then the famid() function will be called on the object.

## Value

A character vector of individual ids with family prefix updated

## **Examples**

```
upd_famid(c("1", "2", "B_3"), c("A", "B", "A"))
upd_famid(c("1", "B_2", "C_3", "4"), c("A", NA, "A", NA))

data(sampleped)
ped1 <- Pedigree(sampleped[,-1])
id(ped(ped1))
new_fam <- make_famid(id(ped(ped1)), dadid(ped(ped1)), momid(ped(ped1)))
id(ped(upd_famid(ped1, new_fam)))

data(sampleped)
ped1 <- Pedigree(sampleped[,-1])
make_famid(ped1)</pre>
```

useful\_inds

Usefulness of individuals

## **Description**

Compute the usefulness of individuals

# Usage

```
## S4 method for signature 'character'
useful_inds(
  obj,
  dadid,
  momid,
  avail,
  affected,
  num_child_tot,
```

134 useful\_inds

```
id_inf,
  keep_infos = FALSE
)

## S4 method for signature 'Pedigree'
useful_inds(obj, keep_infos = FALSE, reset = FALSE, max_dist = NULL)

## S4 method for signature 'Ped'
useful_inds(obj, keep_infos = FALSE, reset = FALSE, max_dist = NULL)
```

## Arguments

obj	A character vector with the id of the individuals or a data.frame with all the informations in corresponding columns.
dadid	A vector containing for each subject, the identifiers of the biologicals fathers.
momid	A vector containing for each subject, the identifiers of the biologicals mothers.
avail	A logical vector with the availability status of the individuals (i.e. $FALSE = not$ available, $TRUE = available$ , $NA = unknown$ ).
affected	A logical vector with the affection status of the individuals (i.e. $FALSE = unaf$ -fected, $TRUE = affected$ , $NA = unknown$ ).
num_child_tot	A numeric vector of the number of children of each individuals
id_inf	An identifiers vector of informative individuals.
keep_infos	Boolean to indicate if parents with unknown status but available or reverse should be kept
reset	Boolean to indicate if the useful column should be reset
max_dist	The maximum distance to informative individuals

## **Details**

Check for the informativeness of the individuals based on the informative parameter given, the number of children and the usefulness of their parents. A useful slot is added to the Ped object with the usefulness of the individual.

# Value

## When obj is a vector:

A vector of useful individuals identifiers

# When obj is a Pedigree or Ped object:

The Pedigree or Ped object with the slot 'useful' containing TRUE for useful individuals and FALSE otherwise.

```
data(sampleped)
ped1 <- Pedigree(sampleped[sampleped$famid == "1",])
ped1 <- is_informative(ped1, informative = "AvAf", col_aff = "affection")
ped(useful_inds(ped1))</pre>
```

```
validate_and_rename_df
```

Rename columns in a dataframe

## **Description**

This function renames the columns of a dataframe based on the user-selected columns. It also validates the user-selected columns and ensures that the mandatory columns are selected.

# Usage

```
validate_and_rename_df(
   df,
   selections,
   col_config,
   others_cols = TRUE,
   na_strings = c("", "NA", "NULL")
)
```

# **Arguments**

df	A dataframe to be modified.
selections	A named list of user-selected columns, the names of the list must correspond to the new column names.
col_config	A list of columns configuration see check_col_config() for more details.
others_cols	A boolean to authorize other columns to be present in the output datatable.

## Value

A dataframe with the selected columns renamed.

```
df <- data.frame(
    ColN1 = c(1, 2), ColN2 = 4, ColTU1 = "A", ColTU2 = 3
)
Pedixplorer:::validate_and_rename_df(
    df, list(Need1 = "ColN1", Sup2 = "ColTU1"),
    list(
        Need1 = list(alternate = c("ColN1", "ColN3"), mandatory = TRUE),
        Sup2 = list(alternate = c("ColTU1", "ColTU3"), mandatory = FALSE)
    )
)</pre>
```

vect\_to\_binary

vect\_to\_binary

Vector variable to binary vector

## **Description**

Transform a vector to a binary vector. All values that are not 0, 1, TRUE, FALSE, or NA are transformed to NA

#### **Usage**

```
vect_to_binary(vect, logical = FALSE, default = NA)
```

## **Arguments**

vect

A character, factor, logical or numeric vector corresponding to a binary variable (i.e. 0 or 1). The following values are recognized:

- character() or factor(): "TRUE", "FALSE", "0", "1", "NA" will be respectively transformed to 1, 0, 0, 1, NA. Spaces and case are ignored. All other values will be transformed to NA.
- numeric(): 0 and 1 are kept, all other values are transformed to NA.
- logical(): TRUE and FALSE are tansformed to 1 and0.

logical

Boolean defining if the output should be a logical vector instead of a numeric vector (i.e. 0 and 1 becomes FALSE and 'TRUE).

default

The default value to use for the values that are not recognized. By default, NA is used, but it can be  $\emptyset$  or 1.

## Value

numeric binary vector of the same size as vect with 0 and 1

```
vect_to_binary(
    c(0, 1, 2, 3.6, "TRUE", "FALSE", "0", "1", "NA", "B", TRUE, FALSE, NA)
)
```

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