

# Package: GENLIB (via r-universe)

September 17, 2024

**Type** Package

**Title** Genealogical Data Analysis

**Version** 1.1.10

**Date** 2024-02-16

**Description** Genealogical data analysis including descriptive statistics (e.g., kinship and inbreeding coefficients) and gene-dropping simulations. See: ``GENLIB: an R package for the analysis of genealogical data" Gauvin et al. (2015) <[doi:10.1186/s12859-015-0581-5](https://doi.org/10.1186/s12859-015-0581-5)>.

**License** GPL (>= 2)

**LazyLoad** yes

**Depends** R (>= 3.1.0), Rcpp (>= 0.9.10)

**Imports** kinship2, methods, bootstrap, Matrix, lattice, quadprog, foreach, parallel, doParallel

**LinkingTo** Rcpp, BH

**NeedsCompilation** yes

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**Repository** CRAN

**RoxygenNote** 7.1.2

**Date/Publication** 2024-02-19 19:20:21 UTC

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Classes of basic handling of genealogy data

Classes "GLmultiVector" "GLmultiArray4" "GLmultiMatrix"  
"GLmultiNumber"

---

## Description

Objects created to carry information mostly between invisible functions.

## Objects from the Class

Objects can be created by calls of the form `new("GLmultiVector", 'matrix', depth = 'integer')`  
`new("GLmultiArray4", 'Array', depth = 'integer')`  
`new("GLmultiMatrix", 'Array', depth = 'integer')`  
`new("GLmultiNumber", 'numeric', depth = 'integer', .Names = 'character')`

## Slots

GLmultiVector

Object of class "matrix"

.Data: Object of class "integer"

GLmultiArray4, GLmultiMatrix

.Data: Object of class "array"

depth: Object of class "integer"

GLmultiNumber

.Data: Object of class "numeric" ~~

depth: Object of class "integer" ~~

.Names: Object of class "character" ~~

## Extends

Class "matrix", from data part. Class "array", by class "matrix", distance 2. Class "structure", by class "matrix", distance 3. Class "vector", by class "matrix", distance 4, with explicit coerce.

## Methods

[<- signature(x = "GLmultiVector", i = "ANY", j = "ANY", value = "ANY"): ...

[ signature(x = "GLmultiVector", i = "ANY", j = "ANY", drop = "ANY"): ...

**depth** signature(x = "GLmultiVector"): ...

**Dim** signature(object = "GLmultiVector"): ...

**Examples**

```
showClass("GLmultiVector")
showClass("GLmultiArray4")
showClass("GLmultiMatrix")
showClass("GLmultiNumber")
```

---

**Classes of GLgroup handling**

<i>Classes</i>	"GLmultiPhiGroup"	"GLmultiPhiGroupSingle"
	"GLmultiFGroup"	"GLmultiFGroupSingle"
	"GLCGMatrixGroupSingle"	

---

**Description**

Objects used to carry information mostly between invisible functions.

**Objects from the Class**

Objects can be created by calls of the form

```
new("GLmultiPhiGroup", MatriceCG, group = 'GLgroup', grindex = 'list')
new("GLmultiPhiGroupSingle", MatriceCG, group = 'GLgroup', grindex = 'list')
new("GLmultiFGroup", MatriceCG, group = 'GLgroup', grindex = 'list')
new("GLmultiFGroupSingle", MatriceCG, group = 'GLgroup', grindex = 'list')
new("GLCGMatrixGroupSingle", MatriceCG, group = 'GLgroup', grindex = 'list')
```

**Slots**

GLCGMatrixGroupSingle, GLmultiFGroupSingle, GLmultiPhiGroupSingle

Object of class "matrix"

**group:** Object of class "GLgroup"

**grindex:** Object of class "list" containing the indices of the probands of 'group'

GLmultiPhiGroup :

**.Data:** Object of class "GLmultiMatrix"

**group:** Object of class "GLgroup"

**grindex:** Object of class "list" containing the indices of the probands of 'group'

GLmultiFGroup :

**.Data:** Object of class "GLmultiVector"

**group:** Object of class "GLgroup"

**grindex:** Object of class "list" containing the indices of the probands of 'group'

**Extends**

Class "**matrix**", from data part. Class "**array**", by class "matrix", distance 2. Class "**structure**", by class "matrix", distance 3. Class "**vector**", by class "matrix", distance 4, with explicit coerce.

**Methods**

```
[<- signature(x = "GLCGMatrixGroupSingle", i = "ANY", j = "ANY", value = "ANY"): ...
[ signature(x = "GLCGMatrixGroupSingle", i = "ANY", j = "ANY", drop = "ANY"): ...
group signature(x = "GLCGMatrixGroupSingle"): ...
```

**Examples**

```
showClass("GLCGMatrixGroupSingle")
showClass("GLmultiFGroup")
showClass("GLmultiFGroupSingle")
showClass("GLmultiPhiGroupSingle")
showClass("GLmultiPhiGroup")
```

---

gen.branching

*Genealogy subset*


---

**Description**

Creates an object of class GLgen from an existing GLgen object by selecting specified individuals.

**Usage**

```
gen.branching( gen, pro = 0, ancestors = gen.founder(gen), bflag = 0)
```

**Arguments**

gen	An object of class GLgen obtained with gen.genealogy, gen.lineages or gen.branching. Required.
pro	Vector of proband id numbers to be included. Default is 0, which will select all individuals without children.
ancestors	Vector of ancestors id numbers to be included. Default includes all founders in the original GLgen object.
bflag	If set to 0 (default and recommended), probands and founders are filtered to quicken the calculations.

**Value**

returns a GLgen object

**Note**

Note that if there are no links between some probands and ancestors, these ids will not be included.

**See Also**

[gen.genealogy](#) [gen.graph](#) [gen.lineages](#)

**Examples**

```

data(geneaJi)
genJi<-gen.genealogy(geneaJi)
genJi_part<-gen.branching(genJi,pro=c(2,28))

# Plots of original genealogy and of the branched version
layout(matrix(1:2,c(1,2),byrow=TRUE))
gen.graph(genJi)
mtext("Original", line=2, cex=1.2)
gen.graph(genJi_part)
mtext("Branched tree\nfor individuals 2 and 28", line=1, cex=1.2)

```

---

gen.children	<i>Get id numbers of children</i>
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---

**Description**

Returns the id numbers of the children of specified individuals.

**Usage**

```
gen.children( gen, individuals, ...)
```

**Arguments**

gen	An object of class GLgen obtained with gen.genealogy, gen.lineages or gen.branching. Required.
individuals	Vector of individual id numbers. Required.
...	Option to pass additionnal arguments automaticaly between methods. Internal use only.

**Value**

returns a vector of integer

**See Also**

[gen.genealogy](#) [gen.pro](#) [gen.founder](#) [gen.sibship](#) [gen.parent](#)

**Examples**

```

data(geneaJi)
genJi<-gen.genealogy(geneaJi)
gen.children(genJi, individuals=14)

data(genea140)
gen140<-gen.genealogy(genea140)
children51052<-gen.children(gen140,51052)
gen.graph(gen140, pro=c(children51052))

```

---

gen.climbPAR	<i>Internal function used by gen.findMRCA</i>
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---

**Description**

gen.climbPAR is an internal function used by gen.findMRCA.

**Usage**

```
gen.climbPAR(gen, individuals, founder)
```

**Arguments**

gen	The genealogy to consider.
individuals	probands to consider
founder	the founder by which the distance is calculated

**Value**

returns a list containing the founder ID and the distance.

**See Also**

[gen.findMRCA](#) [gen.getAncestorsPAR](#)

**Examples**

```
data(geneaJi)
genJi<-gen.genealogy(geneaJi)
gen.findMRCA(genJi, individuals=c(1,29), NbProcess = 1)
```

---

gen.completeness	<i>Completeness</i>
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---

**Description**

Returns the completeness of the genealogical data for the specified probands

**Usage**

```
gen.completeness( gen, pro="0", genNo=-1, type="MEAN", ...)
```

**Arguments**

gen	An object of class GLgen obtained with gen.genealogy, gen.lineages or gen.branching. Required.
pro	Vector of proband id numbers to be included. Default is 0, which will select all individuals without children.
genNo	Vector of generation numbers at which completeness should be calculated. Default is -1, which calculates completeness at each generation.
type	If type="MEAN" (default), mean completeness over all specified probands is calculated. If type="IND", completeness is calculated for each specified proband.
...	Option to pass additional arguments automatically between methods. Internal use only.

**Value**

A data frame with each generation at which the completeness is calculated as rows and one column when type is "MEAN". When type is "IND", the number of columns equals the number of probands specified.

**References**

Cazes P, Cazes MH. (1996) Comment mesurer la profondeur genealogique d'une ascendance? Population (French Ed) 51:117-140.

**See Also**

[gen.genealogy](#) [gen.rec](#) [gen.occ](#) [gen.implex](#) [gen.meangendepth](#)

**Examples**

```
data(geneaJi)
genJi<-gen.genealogy(geneaJi)
gen.completeness(genJi, type="IND")
# For the 4th generation
gen.completeness(genJi, type="IND", genNo=4)

data(genea140)
gen140<-gen.genealogy(genea140)
gen.completeness(gen140)
```

---

gen.completenessVar     *Variance of completeness index*

---

**Description**

Returns the variance of the completeness index across probands.



**Usage**

```
gen.completenessVar( gen, pro = "0", genNo = -1, ...)
```

**Arguments**

<code>gen</code>	An object of class <code>GLgen</code> obtained with <code>gen.genealogy</code> , <code>gen.lineages</code> or <code>gen.branching</code> . Required.
<code>pro</code>	Vector of proband id numbers to be included. Default is 0, which will select all individuals without children.
<code>genNo</code>	Vector of generation numbers at which completeness should be calculated. Default is -1, which calculates completeness at each generation.
<code>...</code>	Option to pass additional arguments automatically between methods. Internal use only.

**Value**

A data frame with one column and each generation at which the variance is calculated as rows.

**References**

Cazes P, Cazes MH. (1996) Comment mesurer la profondeur genealogique d'une ascendance? Population (French Ed) 51:117-140.

**See Also**

[gen.gc](#)

**Examples**

```
data(geneaJi)
genJi<-gen.genealogy(geneaJi)
gen.completeness(genJi, type="IND")
gen.completenessVar(genJi)
# For the 4th generation
gen.completenessVar(genJi, genNo=4)
```

```
data(genea140)
gen140<-gen.genealogy(genea140)
gen.completeness(gen140)
gen.completenessVar(gen140)
```

---

gen.depth	<i>Genealogy depth</i>
-----------	------------------------

---

**Description**

Returns the number of generations in the genealogy from a GLgen object.

**Usage**

```
gen.depth(gen)
```

**Arguments**

gen	An object of class GLgen obtained with gen.genealogy, gen.lineages or gen.branching. Required.
-----	------------------------------------------------------------------------------------------------

**Value**

returns a vector of integer

**See Also**

[gen.genealogy](#) [gen.min](#) [gen.mean](#) [gen.min](#) [gen.meangendepth](#)

**Examples**

```
data(geneaJi)
genJi<-gen.genealogy(geneaJi)
gen.depth(genJi)
```

```
data(genea140)
gen140<-gen.genealogy(genea140)
gen.depth(gen140)
```

---

gen.f	<i>Inbreeding coefficient</i>
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---

**Description**

Returns the inbreeding coefficients of the specified probands

**Usage**

```
gen.f( gen, pro, depthmin= (gen.depth(gen)-1), depthmax= (gen.depth(gen)-1))
```

**Arguments**

gen	An object of class GLgen obtained with gen.genealogy, gen.lineages or gen.branching. Required.
pro	Vector of proband id numbers to be included.
depthmin	Minimum genealogical depth to consider in the calculation. Default is the whole genealogy.
depthmax	Maximum genealogical depth to consider in the calculation. Default is the whole genealogy.

**Value**

A vector or GLmultiVector object depending on the number of generations treated. GLmultiVector is an array of vectors, one for each depth. Array of size ('depthMax'-'depthMin'), vectors of size 'length(pro)'. A vector of size 'length(pro)' is returned when ('depthMax'-'depthMin') equals 1.

**References**

- Malecot G. (1948) Les mathematiques de l'heredite. Paris: Masson, p 65.
- Thompson EA. (1986) Pedigree Analysis in Human Genetics. Baltimore, MD, USA: Johns Hopkins University Press, p 25.
- Karigl G. (1981) A recursive algorithm for the calculation of identity coefficients. Ann Hum Genet 45:299-305.

**See Also**

[gen.genealogy](#) [gen.phi](#)

**Examples**

```
data(geneaJi)
genJi<-gen.genealogy(geneaJi)
f_values<-gen.f(genJi)
f_values
f_allgen<-gen.f(genJi, depthmin=1)
f_allgen<-unclass(f_allgen)

plot(1:7,f_allgen[1,],type="b",xlab="Generation",ylab="Inbreeding values",ylim=c(0,0.25),pch=0)
points(1:7,f_allgen[3,], type="b", lty=12, pch=1)
legend("topright", legend=c("Individual 1", "Individual 29"),lty=c(1,12), pch=c(0,1))

data(genea140)
gen140<-gen.genealogy(genea140)
inbreeding_val<-gen.f(gen140)
boxplot(inbreeding_val, horizontal=TRUE, xlab="Inbreeding values")
```

---

gen.fCI	<i>Average inbreeding coefficient confidence interval</i>
---------	-----------------------------------------------------------

---

**Description**

Returns the confidence interval of the average inbreeding among specified individuals

**Usage**

```
gen.fCI(vectF, prob=c(0.025,0.05,0.95,0.975), b="5000", print.it=FALSE)
```

**Arguments**

vectF	A vector of inbreeding coefficients obtained for example with gen.f. Required.
prob	Confidence limits probabilities. Default is probs=c(0.025, 0.05, 0.95, 0.975).
b	Number of simulations used to calculate the confidence interval. Default is b="5000".
print.it	If print.it=FALSE (default), progression in the number of simulations performed is not displayed.

**Value**

A vector or a GLmultiVector object depending on the type of 'vectF' entered. If 'vectF' is a GLmultiVector, an array of vectors (GLmultiVector), one for each depth, is returned. Array of size ('depthMax'-'depthMin'), vectors of size 'length(probs)'. If 'vectF' is a GLmultiVector, a vector of size 'length(probs)' is returned.

**See Also**

[gen.f](#)

**Examples**

```
data(genea140)
gen140<-gen.genealogy(genea140)
gen10 <-gen.branching(gen140, pro=gen.pro(gen140)[c(1:10)])
fval<-gen.f(gen10)
gen.fCI(fval)
```

---

`gen.find.Min.Distance.MRCA`

*Finds the minimum distances between pairs of individuals given a set of ancestors.*

---

### Description

Returns the shortest distances (number of meioses) between pairs of probands given the matrix of MRCAs output by the `gen.findMRCA` function.

### Usage

```
gen.find.Min.Distance.MRCA(genMatrix, individuals="ALL", ancestors="ALL")
```

### Arguments

<code>genMatrix</code>	Matrix of most recent common ancestors, MRCAs, obtained with <code>gen.findMRCA</code> . Required.
<code>individuals</code>	Vector of proband id numbers to include. All are included by default.
<code>ancestors</code>	Vector of MRCA id numbers to include. All are included by default.

### Value

returns a matrix

### See Also

[gen.genealogy](#) [gen.founder](#) [gen.findMRCA](#) [gen.findDistance](#) [gen.findFounders](#)

### Examples

```
data(geneaJi)
genJi<-gen.genealogy(geneaJi)
matMRCA<-gen.findMRCA(genJi, individuals=c(1,29), NbProcess = 1)
gen.find.Min.Distance.MRCA(matMRCA)

# Increasing NbProcess will decrease execution time
data(genea140)
gen140<-gen.genealogy(genea140)
matMRCA<-gen.findMRCA(gen140, individuals=c(409033,408728,408828), NbProcess = 1)
gen.find.Min.Distance.MRCA(matMRCA, individuals=c(409033,408728))
```

---

<code>gen.findDistance</code>	<i>Minimum genetic distance between two individuals</i>
-------------------------------	---------------------------------------------------------

---

**Description**

Returns the minimum distance (number of meioses) between the specified individuals through one specified ancestor.

**Usage**

```
gen.findDistance(gen, individuals, ancestor)
```

**Arguments**

<code>gen</code>	An object of class <code>GLgen</code> obtained with <code>gen.genealogy</code> , <code>gen.lineages</code> or <code>gen.branching</code> . Required.
<code>individuals</code>	A vector of individual id numbers between which to calculate the distance. Required.
<code>ancestor</code>	A common ancestor to the specified individuals. Required.

**Value**

returns a numeric value

**See Also**

[gen.genealogy](#) [gen.founder](#) [gen.findMRCA](#) [gen.findFounders](#) [gen.find.Min.Distance.MRCA](#)

**Examples**

```
data(geneaJi)
genJi<-gen.genealogy(geneaJi)
gen.findDistance(genJi, individuals=c(1,29), ancestor=17)
gen.findDistance(genJi, individuals=c(1,29), ancestor=26)

data(genea140)
gen140<-gen.genealogy(genea140)
gen.findDistance(gen140, individuals=c(409033,408728), ancestor=38714)
gen.findDistance(gen140, individuals=c(408728,408828), ancestor=95080)
```

---

gen.findFounders      *Find common founder ancestors*

---

**Description**

Returns all the ancestors that are founders for specified individuals in the genealogy.

**Usage**

```
gen.findFounders(gen, individuals, NbProcess=parallel::detectCores()-1)
```

**Arguments**

gen	An object of class GLgen obtained with gen.genealogy, gen.lineages or gen.branching. Required.
individuals	Vector of individual id numbers for which to find the founders. Required
NbProcess	Number of processes to use when running this function. Default=parallel::detectCores()-1

**Value**

A vector of integers

**Note**

Uses slave processes to make the search faster. Those slave processes launch the function gen.getFoundersMPI.

**See Also**

[gen.genealogy](#) [gen.founder](#) [gen.findMRCA](#) [gen.findDistance](#) [gen.find.Min.Distance.MRCA](#)

**Examples**

```
data(geneaJi)
genJi<-gen.genealogy(geneaJi)
gen.founder(genJi)
gen.findFounders(genJi, individuals=c(1,29), NbProcess = 1)
```

```
data(genea140)
gen140<-gen.genealogy(genea140)
gen2<-gen.branching(gen140,pro=c(409033,408728))
# 615 founders in genealogy of #409033 and #408728
all_founders<-gen.founder(gen2)
length(all_founders)
```

```
all_commonFounders<-gen.findFounders(gen140, individuals=c(409033,408728), NbProcess = 1)
length(all_commonFounders) # 127 founders common to #409033 and #408728
```

---

gen.findMRCA                      *Finding most recent common ancestors, MRCA*

---

### Description

Returns MRCA of pairs of specified individuals and the distance (number of meioses) between individuals through the MRCA.

### Usage

```
gen.findMRCA(gen, individuals, NbProcess=parallel::detectCores()-1)
```

### Arguments

gen	An object of class GLgen obtained with gen.genealogy, gen.lineages or gen.branching. Required.
individuals	Vector of individual id numbers for which to find the MRCA. Required
NbProcess	Number of processes to use when running this function. Default=parallel::detectCores()-1

### Value

returns a matrix

### Note

This function uses the parallel programming functions of foreach, snow and doSNOW for launching several processes.

### See Also

[gen.genealogy](#) [gen.founder](#) [gen.findFounders](#) [gen.findDistance](#) [gen.find.Min.Distance.MRCA](#)

### Examples

```
data(geneaJi)
genJi<-gen.genealogy(geneaJi)
gen.findMRCA(genJi, individuals=c(1,29), NbProcess = 1)

# For a more complex example:
data(genea140)
gen140<-gen.genealogy(genea140)
all_commonFounders<-gen.findFounders(gen140, individuals=c(409033,408728), NbProcess = 1)
# 127 founders common to #409033 and #408728
length(all_commonFounders)
# 22 most recent common ancestors for #409033 and #408728
MRCA_2ind<-gen.findMRCA(gen140, individuals=c(409033,408728), NbProcess = 1)
```



---

gen.founder	<i>Get founder id numbers</i>
-------------	-------------------------------

---

### Description

Returns the id numbers of the founders. Founders are defined as the individuals without parents in the genealogy (i.e., mother id=0, father id=0).

### Usage

```
gen.founder( gen, ...)
```

### Arguments

gen	An object of class GLgen obtained with <code>gen.genealogy</code> , <code>gen.lineages</code> or <code>gen.branching</code> . Required.
...	Option to pass additionnal arguments automaticaly between methods. Internal use only.

### Value

A vector of integer

### See Also

[gen.genealogy](#) [gen.pro](#) [gen.half.founder](#) [gen.parent](#)

### Examples

```
data(geneaJi)
genJi<-gen.genealogy(geneaJi)
# There are 6 founders
gen.founder(genJi)

data(genea140)
gen140<-gen.genealogy(genea140)
founder140<-gen.founder(gen140)
# There are 7399 founders
length(founder140)
```

gen.gc

*Genetic contribution of ancestors***Description**

Returns the genetic contribution of ancestors to the gene pool of sepcific probands

**Usage**

```
gen.gc(gen, pro=0, ancestors=0, vctProb=c(0.5,0.5,0.5,0.5), typeCG="IND")
```

**Arguments**

gen	An object of class GLgen obtained with gen.genealogy, gen.lineages or gen.branching. Required.
pro	Vector of proband id numbers to be included. Default is 0, which will select all individuals without children.
ancestors	Vector of ancestors id numbers to be included. Default is 0, which will select all individuals without parents.
vctProb	Vector of transmission probabilities. The first two values indicate the probabilities of transmission of a father to his son and daughter, respectively, and the following two values are the same for the mother. Default is vctProb=c(0.5,0.5,0.5,0.5).
typeCG	IF typeCG="IND" (default), the genetic contribution from each ancestor is calculated for each proband. If typeCG="MEAN", the average (over all probands) genetic contributions of each ancestor is returned. If typeCG="PRODUCT", the product (over all probands) of genetic contributions is returned for each ancestor. If typeCG="TOTAL", the sum (over all probands) of genetic contributions is returned for each ancestor. If typeCG="CUMUL", ancestors are ranked in decreasing order of total contribution and cumulative contribution is returned.

**Value**

A matrix with rows corresponding to probands and columns corresponding to ancestors when typeCG="IND". For the other typeCG values, rows are ancestors and there is one column containing mean, product, total or cumulative values.

**References**

- Roberts DF. (1968) Genetic effects of population size reduction. *Nature*, 220, 1084-1088.
- O'Brien E, Jorde LB, Ronnlof B, Fellman JO, Eriksson AW. (1988) Founder effect and genetic disease in Sottunga, Finland. *American Journal of Physical Anthropology*, 77, 335-346.

**See Also**

[gen.genealogy](#) [gen.rec](#) [gen.occ](#) [gen.implex](#) [gen.meangendepth](#) [gen.completeness](#)

**Examples**

```

data(geneaJi)
genJi<-gen.genealogy(geneaJi)
gc<-gen.gc(genJi, pro=c(1,29), ancestors=c(3,6,10,12,14,16,18,20,26,28))
gc
gc_cum<-gen.gc(genJi, pro=c(1,29), ancestors=c(3,6,10,12,14,16,18,20,26,28), type="CUMUL")
gc_cum

data(genea140)
gen140<-gen.genealogy(genea140)
gc<-gen.gc(gen140, pro=c(454422,676521,677273,717537,717634,717709,868572),
           ancestors=c(18305,18528,31114,18325))

gc
# Mother-daughter transmission only with probability=0.5
gc_MoLi<-gen.gc(gen140, pro=c(217891,302718,409282,802424,409682,443151),
               ancestors=c(18321,218231,296200,39066,18679,442607),vctProb=c(0,0,0,0.5))
gc_MoLi

```

---

<code>gen.genealogy</code>	<i>Create object of class GLgen</i>
----------------------------	-------------------------------------

---

**Description**

Creates an object of class GLgen that contains the ascending genealogies derived from input data in pedigree format

**Usage**

```
gen.genealogy(ped, autoComplete=FALSE, ...)
```

**Arguments**

<code>ped</code>	A matrix or data frame with the following columns: individual id, father id, mother id, sex. Required. Individual id numbers must be numeric and unique. If an individual does not have a father and/or mother, the father and/or mother id numbers must be set to 0. Sex must be equal to M or 1 for males and F or 2 for females. The sex column is optional for this function but necessary for some other functions using GLgen objects.
<code>autoComplete</code>	If TRUE, any non-zero father and mother id numbers not appearing in the individual id column, will be added in the individual column as having no father or mother (both set to 0). Default to FALSE.
<code>...</code>	Option to pass additional arguments automatically between methods. Internal use only.

**Value**

An object of class GLgen.

**See Also**

[gen.graph](#) [gen.genout](#)

**Examples**

```
ind<-c(1,2,3,11:21,101:108,201:202)
father<-c(11,15,15,102,0,102,0,103,103,103,105,105,107,107,0,202,0,202,202,0,202,0,0,0)
mother<-c(12,14,14,101,0,101,0,104,104,104,106,106,108,108,0,201,0,201,201,0,201,0,0,0)
sex<-c(1,2,2,1,2,1,2,1,2,1,2,2,2,1,2,1,1,2,1,2,1,2,2,1)
gen.df<-data.frame(ind, father, mother, sex)
gen.genealogy(gen.df)

data(geneaJi)
geneaJi[1:5,]
genJi<-gen.genealogy(geneaJi)
# Print basic information about the genealogy
genJi
```

---

gen.genout

*Create pedigree data*

---

**Description**

Creates data frame in pedigree format from an object of class GLgen

**Usage**

```
gen.genout( gen, sorted = FALSE)
```

**Arguments**

gen	An object of class GLgen obtained with <code>gen.genealogy</code> , <code>gen.lineages</code> or <code>gen.branching</code> . Required.
sorted	If <code>sorted=FALSE</code> (default), individual id number are not sorted in output data. Id numbers for parents will be placed before their children ids. If <code>sorted=TRUE</code> , individual id numbers will be sorted.

**Value**

returns a data.frame containing the following: **ind father mother sex**

**See Also**

[gen.genealogy](#) [gen.graph](#) [gen.branching](#) [gen.lineages](#)

### Examples

```
data(geneaJi)
# Original data is a data.frame
geneaJi[1:12,]

genJi<-gen.genealogy(geneaJi)
# As a genealogy object
genJi

# Genealogy as a data.frame
genJi_df<-gen.genout(genJi)
genJi_df[1:12,]

# Maternal lineage
genJi_MaLi<-gen.lineages(geneaJi, maternal = TRUE)
# Maternal lineage as a data.frame
genJi_MaLi_df<-gen.genout(genJi_MaLi)
genJi_MaLi_df
```

---

gen.getAncestorsPAR     *Internal function used by gen.findMRCA*

---

### Description

gen.getAncestorsPAR is an internal function used by gen.findMRCA.

### Usage

```
gen.getAncestorsPAR(gen, pro)
```

### Arguments

gen	The genealogy to consider.
pro	probands to consider

### Value

Vector of the founders identity.

### See Also

[gen.findMRCA](#) [gen.climbPAR](#)

### Examples

```
data(geneaJi)
genJi<-gen.genealogy(geneaJi)
gen.findMRCA(genJi, individuals=c(1,29), NbProcess = 1)
```

---

gen.getFoundersPAR      *Internal function used by gen.findFounders*

---

### Description

gen.getFoundersPAR is an internal function used by gen.findFounders.

### Usage

```
gen.getFoundersPAR(gen, pro)
```

### Arguments

gen	The genealogy to consider.
pro	probands to consider

### Value

A vector of the founders identity.

### See Also

[gen.findFounders](#) [gen.findDistance](#)

### Examples

```
data(geneaJi)
genJi<-gen.genealogy(geneaJi)
gen.founder(genJi, individuals=c(1,29))
gen.findFounders(genJi, individuals=c(1,29), NbProcess = 1)
```

---

gen.graph      *Pedigree graphical tool*

---

### Description

Function that plots pedigrees of GLgen objects.

### Usage

```
gen.graph( gen, pro=gen.pro(gen), ancestors=gen.founder(gen),
  indVarAffected=gen.genout(gen)$ind, varAffected=gen.genout(gen)$ind, cex="1",
  col="0", symbolsize="1", width="1", packed=FALSE, align=TRUE, ...)
```

**Arguments**

gen	An object of class GLgen obtained with gen.genealogy, gen.lineages or gen.branching. Required.
pro	Vector of proband id numbers to be included. Default is 'gen.pro(gen)', which will select all individuals without children.
ancestors	Vector of ancestors id numbers to be included. Default is 'gen.founder(gen)', which will select all individuals without parents.
indVarAffected	Vector of individuals id numbers used for labeling. Default is 'gen.genout(gen)\$ind', which is all the individuals of the genealogy.
varAffected	Vector of individuals labels. Default is 'gen.genout(gen)\$ind', meaning that the individuals labels are their ids.
cex	Controls text size (same as kinship2::plot.pedigree). Default is 1.
col	Color for each id (same as kinship2::plot.pedigree). Default is 0, which assigns the same color to everyone.
symbolsize	Controls symbolsize (same as kinship2::plot.pedigree). Default is 1.
width	For a packed pedigree, the minimum width allowed in the realignment of pedigrees (same as kinship2::plot.pedigree). Default is 1.
packed	If TRUE, uniform distance between all individuals at a given level (same as kinship2::plot.pedigree). Default is FALSE.
align	Controls the extra effort spent trying to align children underneath parents, but without making the pedigree too wide (same as kinship2::plot.pedigree). Default is TRUE.
...	Other arguments that can be passed to the kinship2::plot.pedigree.

**Value**

Returns the same invisible list as that returned by kinship2::plot.pedigree, which contains the following: **plist** list containing the information about the pedigree (n, id, pos, fam, spouse) **x** x-axis position **y** y-axis position **boxw** box width **boxh** box height **call** the call made to plot.pedigree() (kinship2 package)

**See Also**

[gen.genealogy](#) [gen.genout](#) [gen.branching](#)

**Examples**

```
ind<-c(1,2,3,11:21,101:108,201:202)
father<-c(11,15,15,102,0,102,0,103,103,0,105,105,107,107,0,202,0,202,202,0,202,0,0,0)
mother<-c(12,14,14,101,0,101,0,104,104,0,106,106,108,108,0,201,0,201,201,0,201,0,0,0)
sex<-c(1,2,2,1,2,1,2,1,2,1,2,2,2,1,2,1,1,2,1,2,1,2,2,1)
gen.df<-data.frame(ind, father, mother, sex)
genEx<-gen.genealogy(gen.df)
gen.graph(genEx)

data(geneaJi)
```

```
geneaJi[1:5,]  
genJi<-gen.genealogy(geneaJi)  
gen.graph(genJi)
```

---

`gen.half.founder`      *Get half-founder id numbers*

---

### **Description**

Returns the id numbers of the half-founders. Half-founders are defined as the individuals with only one known parent in the genealogy (i.e., either mother id=0 or father id=0).

### **Usage**

```
gen.half.founder( gen, ...)
```

### **Arguments**

<code>gen</code>	An object of class GLgen obtained with <code>gen.genealogy</code> , <code>gen.lineages</code> or <code>gen.branching</code> . Required.
<code>...</code>	Option to pass additional arguments automatically between methods. Internal use only.

### **Value**

returns a vector of integer

### **See Also**

[gen.genealogy](#) [gen.pro](#) [gen.founder](#) [gen.parent](#)

### **Examples**

```
data(geneaJi)  
genJi<-gen.genealogy(geneaJi)  
# There are 2 half-founders  
gen.half.founder(genJi)
```



---

gen.implex	<i>Genealogical implex</i>
------------	----------------------------

---

**Description**

Returns the genealogical implex index (a measure of pedigree collapsing) for the specified probands.

**Usage**

```
gen.implex( gen, pro = "0", genNo = -1, type = "MEAN", onlyNewAnc = FALSE, ...)
```

**Arguments**

gen	An object of class GLgen obtained with gen.genealogy, gen.lineages or gen.branching. Required.
pro	Vector of proband id numbers to be included. Default is 0, which will select all individuals without children.
genNo	Vector of generation numbers at which the implex should be calculated. Default is -1, which calculates the implex at each generation.
type	If type="MEAN" (default), implex index values are averaged over all specified probands. If type="IND", the implex index is calculated for all specified proband together. If type="ALL", the implex index is calculated for each specified proband.
onlyNewAnc	If onlyNewAnc=FALSE (default), all ancestors will be considered. If onlyNewAnc=TRUE, only new ancestors will be counted (i.e., an ancestor is not counted again if it has already been counted in another generation).
...	Option to pass additionnal arguments automatically between methods. Internal use only.

**Value**

A data frame with each generation at which the completeness is calculated as rows and one column when type is "MEAN". When type is "IND", the number of columns equals the number of probands specified.

**References**

Cazes P, Cazes MH. (1996) Comment mesurer la profondeur genealogique d'une ascendance? Population (French Ed) 51:117-140.

**See Also**

[gen.genealogy](#) [gen.occ](#) [gen.rec](#) [gen.meangendepth](#) [gen.completeness](#) [gen.gc](#)

**Examples**

```

data(geneaJi)
genJi<-gen.genealogy(geneaJi)
gen.implex(genJi)
gen.implex(genJi, type="IND")
# For the 5th generation
gen.implex(genJi, type="IND", genNo=5)

data(genea140)
gen140<-gen.genealogy(genea140)
gen.implex(gen140)
gen.implex(gen140, pro=c(454422, 676521, 677273, 717537, 717634, 717709, 868572))
gen.implex(gen140, pro=c(454422, 676521, 677273, 717537, 717634, 717709, 868572), type="IND")

```

---

gen.implexVar	<i>Variance of genealogical implex</i>
---------------	----------------------------------------

---

**Description**

Returns the variance of the implex index across probands.

**Usage**

```
gen.implexVar( gen, pro = "0", onlyNewAnc = FALSE, genNo = -1, ...)
```

**Arguments**

gen	An object of class GLgen obtained with gen.genealogy, gen.lineages or gen.branching. Required.
pro	Vector of proband id numbers to be included. Default is 0, which will select all individuals without children.
onlyNewAnc	If onlyNewAnc=FALSE (default), all ancestors will be considered. If onlyNewAnc=TRUE, only new ancestors will be counted (i.e., an ancestor is not counted again if it has already been counted in another generation).
genNo	Vector of generation numbers at which the implex should be calculated. Default is -1, which calculates the implex at each generation.
...	Additional arguments to be passed to methods.

**Value**

A data frame with one column and each generation at which the variance is calculated as rows

**References**

Cazes P, Cazes MH. (1996) Comment mesurer la profondeur genealogique d'une ascendance? Population (French Ed) 51:117-140.

**See Also**[gen.gc](#)**Examples**

```
data(geneaJi)
genJi<-gen.genealogy(geneaJi)
gen.implex(genJi, type="IND")
gen.implexVar(genJi)
# For the 5th generation
gen.implexVar(genJi, genNo=5)
```

```
data(genea140)
gen140<-gen.genealogy(genea140)
gen.implex(gen140)
gen.implex(gen140, pro=c(454422, 676521, 677273, 717537, 717634, 717709, 868572), type="IND")
gen.implexVar(gen140, pro=c(454422, 676521, 677273, 717537, 717634, 717709, 868572), type="IND")
```

---

`gen.lineages`*Create object of class GLgen for maternal or paternal lineages*

---

**Description**

Creates an object of class GLgen that contains maternal or paternal lineages selected from input data in pedigree format

**Usage**

```
gen.lineages(ped, pro = "0", maternal = TRUE, ...)
```

**Arguments**

ped	A matrix or data frame with the following columns: individual id, father id, mother id, sex. Required. Individual id numbers must be numeric and unique. If an individual does not have a father and/or mother, the father and/or mother id numbers must be set to 0. All non-zero father and mother id numbers must also appear in the individual id column. Sex must be equal to M or 1 for males and F or 2 for females. The sex column is optional for this function but necessary for some other functions using GLgen objects.
pro	Vector of individual id numbers for which lineages should be included. Optional.
maternal	If mat=TRUE (default), maternal lineages are selected. mat=FALSE returns paternal lineages.
...	Option to pass additional arguments automatically between methods. Internal use only.

**Value**

returns a GLgen object

**See Also**

[gen.genealogy](#) [gen.graph](#) [gen.branching](#) [gen.genout](#)

**Examples**

```
data(geneaJi)
genJi <- gen.genealogy(geneaJi)
genJi_MaLi<-gen.lineages(geneaJi, maternal = TRUE)
genJi_FaLi<-gen.lineages(geneaJi, maternal = FALSE)

# Plots of original genealogy and maternal and paternal lineages
layout(matrix(1:3,c(1,3),byrow=TRUE), widths =c(3,1,1), heights = 1)
gen.graph(genJi)
mtext("Original", line=2)
gen.graph(genJi_MaLi)
mtext("Maternal\nlineages", line=1)
gen.graph(genJi_FaLi)
mtext("Paternal\nlineages", line=1)
```

---

gen.max

*Maximum number of generations*

---

**Description**

Returns the maximum number of generations between all probands and the individuals specified. Probands are defined as the individuals without children in the genealogy.

**Usage**

```
gen.max( gen, individuals)
```

**Arguments**

**gen** An object of class GLgen obtained with `gen.genealogy`, `gen.lineages` or `gen.branching`. Required.

**individuals** A vector of ids specifying the individuals to include in the calculation. Required.

**Value**

returns a vector of integer

**See Also**

[gen.genealogy](#) [gen.mean](#) [gen.min](#) [gen.depth](#) [gen.meangendepth](#)

**Examples**

```
data(geneaJi)
genJi<-gen.genealogy(geneaJi)
gen.min(genJi,c(17,26))
gen.mean(genJi,c(17,26))
gen.max(genJi,c(17,26))
```

```
data(genea140)
gen140<-gen.genealogy(genea140)
gen.min(gen140,c(18311,18430))
gen.mean(gen140,c(18311,18430))
gen.max(gen140,c(18311,18430))
```

---

gen.mean	<i>Mean number of generations</i>
----------	-----------------------------------

---

**Description**

Returns the average number of generations between all probands and the individuals specified. Probands are defined as the individuals without children in the genealogy.

**Usage**

```
gen.mean( gen, individuals)
```

**Arguments**

gen	An object of class GLgen obtained with gen.genealogy, gen.lineages or gen.branching. Required.
individuals	A vector of ids specifying the individuals to include in the calculation. Required.

**Value**

returns a numeric value

**See Also**

[gen.genealogy](#) [gen.min](#) [gen.max](#) [gen.depth](#) [gen.meangendepth](#)

**Examples**

```
data(geneaJi)
genJi<-gen.genealogy(geneaJi)
gen.min(genJi,c(17,26))
gen.mean(genJi,c(17,26))
gen.max(genJi,c(17,26))
```

```

data(genea140)
gen140<-gen.genealogy(genea140)
gen.min(gen140,c(18311,18430))
gen.mean(gen140,c(18311,18430))
gen.max(gen140,c(18311,18430))

```

---

gen.meangendepth	<i>Expected Genealogical Depth</i>
------------------	------------------------------------

---

### Description

Returns the expected genealogical depth.

### Usage

```
gen.meangendepth( gen, pro = "0", type = "MEAN", ...)
```

### Arguments

gen	An object of class GLgen obtained with gen.genealogy, gen.lineages or gen.branching. Required.
pro	Vector of proband id numbers to be included. Default is 0, which will select all individuals without children.
type	If type="MEAN" (default), mean genealogical depth over all specified probands is calculated. If type="IND", mean genealogical depth is calculated for each specified proband.
...	Option to pass additionnal arguments automaticaly between methods. Internal use only.

### Value

A data frame with only one numeric value when type is "MEAN". When type is "IND", the number of rows equals the number of probands specified.

### References

Cazes P, Cazes MH. (1996) Comment mesurer la profondeur genealogique d'une ascendance? Population (French Ed) 51:117-140.

Kouladjian K. (1986) Une mesure d'entropie genealogique. Chicoutimi, SOREP, Document III-C-43.

De Brakaeleer M, Bellis G. (1994) Genealogies et reconstitutions de familles en genetique humaine. Dossiers et Recherches, no 43, INED, Paris.

### See Also

[gen.genealogy](#) [gen.occ](#) [gen.implex](#) [gen.rec](#) [gen.completeness](#) [gen.gc](#)

**Examples**

```
data(geneaJi)
genJi<-gen.genealogy(geneaJi)
gen.meangendepth(genJi)
gen.meangendepth(genJi, type="IND")
```

```
data(genea140)
gen140<-gen.genealogy(genea140)
gen.meangendepth(gen140)
gen.meangendepth(gen140, pro=c(454422, 676521, 677273, 717537, 717634, 717709, 868572))
gen.meangendepth(gen140, pro=c(454422, 676521, 677273, 717537, 717634, 717709, 868572), type="IND")
```

---

gen.meangendepthVar     *Variance of genealogical depth*

---

**Description**

Returns the variance of the genealogical depth

**Usage**

```
gen.meangendepthVar( gen, pro = "0", type = "MEAN", ...)
```

**Arguments**

gen	An object of class GLgen obtained with gen.genealogy, gen.lineages or gen.branching. Required.
pro	Vector of proband id numbers to be included. Default is 0, which will select all individuals without children.
type	If type="MEAN" (default), the average of genealogical depth variances (over all probands) is returned. If type="IND", the variance of the genealogical depth is calculated for each specified proband.
...	Option to pass additionnal arguments automaticaly between methods. Internal use only.

**Value**

A data frame with only one numeric value when type is "MEAN". When type is "IND", the number of rows equals the number of probands specified.

## References

- Cazes P, Cazes MH. (1996) Comment mesurer la profondeur genealogique d'une ascendance? Population (French Ed) 51:117-140.
- Kouladjian K. (1986) Une mesure d'entropie genealogique. Chicoutimi, SOREP, Document III-C-43.
- De Brakaeleer M, Bellis G. (1994) Genealogies et reconstitutions de familles en genetique humaine. Dossiers et Recherches, no 43, INED, Paris.

## See Also

[gen.gc](#)

## Examples

```
data(geneaJi)
genJi<-gen.genealogy(geneaJi)
gen.meangendepth(genJi, type="IND")
gen.meangendepthVar(genJi, type="IND")
gen.meangendepthVar(genJi, type="MEAN")

data(genea140)
gen140<-gen.genealogy(genea140)
gen.meangendepth(gen140)
probands <- c(454422,676521,677273,717537,717634,717709,868572)
gen.meangendepth(gen140, pro=probands)
gen.meangendepthVar(gen140, pro=probands)
gen.meangendepthVar(gen140, pro=probands, type="MEAN")

gen.meangendepth(gen140, pro=probands, type="IND")
gen.meangendepthVar(gen140, pro=probands, type="IND")
```

---

gen.min

*Minimum number of generations*

---

## Description

Returns the minimum number of generations between all probands and the individuals specified. Probands are defined as the individuals without children in the genealogy.

## Usage

```
gen.min( gen, individuals)
```



**Arguments**

- gen                    An object of class GLgen obtained with gen.genealogy, gen.lineages or gen.branching. Required.
- individuals          A vector of ids specifying the individuals to include in the calculation. Required.

**Value**

returns a vector of integer

**See Also**

[gen.genealogy](#) [gen.mean](#) [gen.max](#) [gen.depth](#) [gen.meangendepth](#)

**Examples**

```
data(geneaJi)
genJi<-gen.genealogy(geneaJi)
gen.min(genJi,c(17,26))
gen.mean(genJi,c(17,26))
gen.max(genJi,c(17,26))
```

```
data(genea140)
gen140<-gen.genealogy(genea140)
gen.min(gen140,c(18311,18430))
gen.mean(gen140,c(18311,18430))
gen.max(gen140,c(18311,18430))
```

---

gen.nochildren	<i>Number of children</i>
----------------	---------------------------

---

**Description**

Returns the number of children for specified individuals

**Usage**

```
gen.nochildren( gen, individuals)
```

**Arguments**

- gen                    An object of class GLgen obtained with gen.genealogy, gen.lineages or gen.branching. Required.
- individuals          A vector of ids of the individuals on each of whom the number of children is to be returned. Required.

**Value**

A vector containing the number of children for each individual specified.

**See Also**

[gen.genealogy](#) [gen.nowomen](#) [gen.noind](#) [gen.nomen](#) [gen.children](#)

**Examples**

```
data(geneaJi)
genJi<-gen.genealogy(geneaJi)
gen.children(genJi,14)
gen.nochildren(genJi,individuals=c(1,12,14,20))
```

```
data(genea140)
gen140<-gen.genealogy(genea140)
children51052<-gen.children(gen140,51052)
gen.nochildren(gen140,51052)
gen.graph(gen140, pro=children51052)
```

---

gen.noind	<i>Number of individuals</i>
-----------	------------------------------

---

**Description**

Returns the number of individuals included in the genealogy

**Usage**

```
gen.noind( gen)
```

**Arguments**

gen                    An object of class GLgen obtained with [gen.genealogy](#), [gen.lineages](#) or [gen.branching](#).  
Required.

**Value**

returns a vector of integer

**See Also**

[gen.genealogy](#) [gen.nowomen](#) [gen.nochildren](#) [gen.nomen](#)

**Examples**

```
data(geneaJi)
genJi<-gen.genealogy(geneaJi)
gen.noind(genJi)
```

---

gen.nomen	<i>Number of men</i>
-----------	----------------------

---

**Description**

Returns the number of men included in the genealogy

**Usage**

```
gen.nomen( gen)
```

**Arguments**

gen	An object of class GLgen obtained with gen.genealogy, gen.lineages or gen.branching. Required.
-----	------------------------------------------------------------------------------------------------

**Value**

returns a vector of integer

**See Also**

[gen.genealogy](#) [gen.nowomen](#) [gen.nochildren](#) [gen.noind](#)

**Examples**

```
data(geneaJi)
genJi<-gen.genealogy(geneaJi)
gen.nomen(genJi)
```

---

gen.nowomen	<i>Number of women</i>
-------------	------------------------

---

**Description**

Returns the number of women included in the genealogy

**Usage**

```
gen.nowomen( gen)
```

**Arguments**

gen	An object of class GLgen obtained with gen.genealogy, gen.lineages or gen.branching. Required.
-----	------------------------------------------------------------------------------------------------

**Value**

returns a vector of integer

**See Also**

[gen.genealogy](#) [gen.nomen](#) [gen.nochildren](#) [gen.noind](#)

**Examples**

```
data(geneaJi)
genJi<-gen.genealogy(geneaJi)
gen.nowomen(genJi)
```

---

gen.occ

*Ancestor occurrences*

---

**Description**

Returns the number of times that the specified ancestors are present in the genealogies of the specified probands.

**Usage**

```
gen.occ( gen, pro = "0", ancestors = "0", typeOcc = "IND", ...)
```

**Arguments**

gen	An object of class GLgen obtained with gen.genealogy, gen.lineages or gen.branching. Required.
pro	Vector of proband id numbers to be included. Default is 0, which will select all individuals without children.
ancestors	Vector of ancestors id numbers to be included. Default is 0, which will select all individuals without parents.
typeOcc	If typeOcc="IND" (default), the number of occurrences for each proband will be returned. If typeOcc="TOTAL", the sum of all occurrences over all probands will be returned.
...	Option to pass additional arguments automatically between methods. Internal use only.

**Value**

A matrix with number of lines equal to the number of ancestors included and the number of columns equal to the number of probands included if typeOcc="BRUT" or only one column if typeOcc="TOTAL"

**See Also**

[gen.genealogy](#) [gen.rec](#) [gen.implex](#) [gen.meangendepth](#) [gen.gc](#)

**Examples**

```
data(geneaJi)
genJi<-gen.genealogy(geneaJi)
# Number of occurrences of ancestors in the specified proband's genealogy
gen.occ(genJi, pro=c(1,29), ancestors=c(3,6,10,12,14,16,18,20,26,28))

data(genea140)
gen140<-gen.genealogy(genea140)
gen.occ(gen140, pro=c(454422,677273,717537,717634,717709,868572),
        ancestors=c(18305,18528,31114,18325))
```

---

gen.parent	<i>Get id numbers of parents</i>
------------	----------------------------------

---

**Description**

Returns the id numbers of the parents of specified individuals.

**Usage**

```
gen.parent( gen, individuals, output = "FaMo", ...)
```

**Arguments**

gen	An object of class GLgen obtained with <code>gen.genealogy</code> , <code>gen.lineages</code> or <code>gen.branching</code> . Required.
individuals	Vector of individual id numbers. Required.
output	If <code>output="FaMo"</code> (default) then both mothers and fathers are included. "Mo" outputs mothers only and "Fa", fathers only.
...	Option to pass additional arguments automatically between methods. Internal use only.

**Value**

returns a list containing the following: **Fathers Mothers**

**See Also**

[gen.genealogy](#) [gen.pro](#) [gen.founder](#) [gen.children](#) [gen.sibship](#)

**Examples**

```
data(geneaJi)
genJi<-gen.genealogy(geneaJi)
gen.parent(genJi, individuals=c(3,21,29))
```

---

gen.phi	<i>Kinship coefficient</i>
---------	----------------------------

---

**Description**

Returns the kinship coefficients between pairs of individuals

**Usage**

```
gen.phi(gen, pro, depthmin=(gen.depth(gen)-1), depthmax=(gen.depth(gen)-1), MT=FALSE)
```

**Arguments**

gen	An object of class GLgen obtained with gen.genealogy, gen.lineages or gen.branching. Required.
pro	Vector of proband id numbers to be included. Required.
depthmin	Minimum genealogical depth to consider in the calculation. Default is the whole genealogy.
depthmax	Maximum genealogical depth to consider in the calculation. Default is the whole genealogy.
MT	Allows parallel computing when set to TRUE. Default is MT=FALSE.

**Value**

A matrix or a GLmultiMatrix object depending on the number of generations treated. GLmultiMatrix is an array of matrices, one for each depth. Array of size ('depthMax'-'depthMin') and matrices of size 'length(pro)' \* 'length(pro)'. The matrix object, also of size 'length(pro)' \* 'length(pro)', is returned when ('depthMax'-'depthMin') equals 1.

**References**

- Malecot G. (1948) Les mathematiques de l'heredite. Paris: Masson, p 65.
- Thompson EA. (1986) Pedigree Analysis in Human Genetics. Baltimore, MD, USA: Johns Hopkins University Press, p 25.
- Karigl G. (1981) A recursive algorithm for the calculation of identity coefficients. Ann Hum Genet 45:299-305.

**See Also**

[gen.genealogy](#) [gen.f](#)

**Examples**

```

data(geneaJi)
genJi<-gen.genealogy(geneaJi)
kinship<-gen.phi(genJi)
kinship

kinship_allgenerations<-gen.phi(genJi, depthmin =1)
kinship_allgenerations
# 7th generations back in time is equivalent to considering all generations
kinship_allgenerations <- unclass(kinship_allgenerations)
kinship_allgenerations[,7]==kinship

kinship_allgenerations[1,2,]

# Plot of kinship varying according to number of generations considered
plot(1:7,kinship_allgenerations[1,2,], type="b", xlab="Generation", ylab="Kinship value",
     ylim=c(0,0.6), pch=0)
points(1:7,kinship_allgenerations[1,3,], type="b", lty=12, pch=1)
legend("topright", legend=c("Individuals 1 and 2", "Individuals 2 and 29"),lty=c(1,12), pch=c(0,1))

```

---

gen.phiCI

*Average kinship confidence interval*


---

**Description**

Returns the confidence interval of the average kinship among pairs of specified individuals

**Usage**

```
gen.phiCI(phiMatrix, prob=c(0.025,0.05,0.95,0.975), b=5000, print.it=FALSE)
```

**Arguments**

phiMatrix	A square matrix of kinship coefficients obtained for example with gen.phi, or an array of square matrices (GLmultiMatrix object) of kinship coefficients obtained with gen.phi. Required.
prob	Confidence limits probabilities. Default is probs=c(0.025, 0.05, 0.95, 0.975).
b	Number of simulations used to calculate the confidence interval. Default is b="5000".
print.it	If print.it=FALSE (default), progression in the number of simulations performed is not displayed.

**Value**

A vector or a GLmultiVector object depending on the type of 'phiMatrix' entered. If 'phiMatrix' is a GLmultiMatrix, an array of vectors (GLmultiVector), one for each depth, is returned. Array of size ('depthMax'-'depthMin'), vectors of size 'length(probs)'. If 'phiMatrix' is a matrix, a vector of size 'length(probs)' is returned.

**See Also**

[gen.genealogy](#) [gen.phi](#) [gen.phiOver](#) [gen.phiMean](#)

**Examples**

```
data(genea140)
gen140<-gen.genealogy(genea140)
ge07<-gen.branching(gen140, pro=gen.pro(gen140)[c(1:7)])
phiMat<-gen.phi(ge07)
gen.phiCI(phiMat)
```

---

gen.phiMean

*Average kinship*

---

**Description**

Returns the average kinship among pairs of specified individuals

**Usage**

```
gen.phiMean( phiMatrix)
```

**Arguments**

phiMatrix      A square matrix of kinship coefficients obtained for example with gen.phi. Required.

**Value**

returns a numeric value

**See Also**

[gen.genealogy](#) [gen.phi](#) [gen.phiOver](#) [gen.phiCI](#)

**Examples**

```
data(geneaJi)
genJi<-gen.genealogy(geneaJi)
kinship<-gen.phi(genJi)
gen.phiMean(kinship)
```

```
data(genea140)
gen140<-gen.genealogy(genea140)
phi6subjects<-gen.phi(gen140, pro=c(454422, 676521, 677273, 717537, 717634, 717709, 868572))
gen.phiMean(phi6subjects)
```



---

gen.phiOver	<i>Kinship above threshold</i>
-------------	--------------------------------

---

**Description**

Returns the pairs of individuals with kinship coefficient values greater than specified threshold.

**Usage**

```
gen.phiOver( phiMatrix, threshold)
```

**Arguments**

phiMatrix	A square matrix of kinship coefficients obtained for example with gen.phi. Required.
threshold	Threshold of kinship values to return.

**Value**

A data frame containing the probands and their kinship.

**See Also**

[gen.genealogy](#) [gen.phi](#) [gen.phiMean](#) [gen.phiCI](#)

**Examples**

```
data(geneaJi)
genJi<-gen.genealogy(geneaJi)
kinship<-gen.phi(genJi)
gen.phiOver(kinship, 0.1)

data(genea140)
gen140<-gen.genealogy(genea140)
phi9subj<-gen.phi(gen140,pro=c(408758,408950,409082,409111,676521,717537,717634,717709,868572))
gen.phiOver(phi9subj,0.025)
```

---

gen.pro	<i>Get proband id numbers</i>
---------	-------------------------------

---

### Description

Returns the id numbers of the probands. Probands are defined as the individuals without children in the genealogy.

### Usage

```
gen.pro( gen, ...)
```

### Arguments

gen	An object of class GLgen obtained with <code>gen.genealogy</code> , <code>gen.lineages</code> or <code>gen.branching</code> . Required.
...	Option to pass additionnal arguments automaticaly between methods. Internal use only.

### Value

returns a vector of integer

### See Also

[gen.genealogy](#) [gen.founder](#) [gen.half.founder](#) [gen.parent](#) [gen.children](#)

### Examples

```
data(geneaJi)
genJi<-gen.genealogy(geneaJi)
# There are 3 probands
gen.pro(genJi)
```

```
data(genea140)
gen140<-gen.genealogy(genea140)
gen.pro(gen140)
# There are 140 probands
```

---

gen.rec                      *Ancestors coverage*

---

### Description

Returns the number of specified probands genealogically related to specified ancestors (i.e., ancestor occurs in the proband's genealogy).

### Usage

```
gen.rec( gen, pro = "0", ancestors = "0", ...)
```

### Arguments

gen	An object of class GLgen obtained with gen.genealogy, gen.lineages or gen.branching. Required.
pro	Vector of proband id numbers to be included. Default is 0, which will select all individuals without children.
ancestors	Vector of ancestors id numbers to be included. Default is 0, which will select all individuals without parents.
...	Option to pass additional arguments automatically between methods. Internal use only.

### Value

A matrix with number of lines equal to the number of ancestors specified and one column.

### Note

If an ancestor is also a proband, he/she will be counted in his/her coverage.

### See Also

[gen.genealogy](#) [gen.occ](#) [gen.implex](#) [gen.meangendepth](#) [gen.gc](#)

### Examples

```
data(geneaJi)
genJi<-gen.genealogy(geneaJi)
# Number of probands which are descendants of an ancestor
gen.rec(genJi)
```

```
data(genea140)
gen140<-gen.genealogy(genea140)
gen.rec(gen140, ancestors=c(18305, 18528, 31114, 18325))
```

---

gen.sibship                      *Get id numbers of siblings*

---

### Description

Returns the id numbers of the siblings of specified individuals.

### Usage

```
gen.sibship( gen, individuals, halfSibling = TRUE, ...)
```

### Arguments

gen	An object of class GLgen obtained with gen.genealogy, gen.lineages or gen.branching. Required.
individuals	Vector of individual id numbers. Required.
halfSibling	If halfSibling=TRUE (default) then ids of halfsiblings are also returned. halfSibling=FALSE returns only full sibling ids.
...	Option to pass additionnal arguments automaticaly between methods. Internal use only.

### Value

returns a vector of integer

### See Also

[gen.genealogy](#) [gen.pro](#) [gen.founder](#) [gen.children](#) [gen.parent](#)

### Examples

```
data(geneaJi)
genJi<-gen.genealogy(geneaJi)
gen.sibship(genJi, individuals=21,halfSibling=TRUE)
```

```
data(genea140)
gen140<-gen.genealogy(genea140)
sibs<-gen.sibship(gen140,individuals=10174, halfSibling=FALSE)
gen.graph(gen140, pro=c(10174,sibs))
```

---

gen.simuHaplo                      *Gene dropping simulations - haplotypes*

---

### Description

Gene dropping from founders to specified probands according to specified model of meiosis. Output .txt files will have simulated haplotypes, labelled w.r.t. the ancestor of origin

### Usage

```
gen.simuHaplo(gen, pro = NULL, ancestors = NULL, simulNo = 1, model
              = 1, model_params, cM_len, BP_len, physical_map_Mo =
              NULL, physical_map_Fa = NULL, seed = 0, all_nodes = 0,
              outDir = getwd())
```

### Arguments

gen	An object of class GLgen obtained with gen.genealogy, gen.lineages or gen.branching. Required.
pro	Vector of proband id numbers to be included. Will default to all probands in the provided genealogy.
ancestors	Vector of ancestors id numbers to be included. Will default to all ancestors of probands
simulNo	Number of simulations to run. Defaults to 1. Results from all simulations are contained in the same output text file
model	model=1 is Poisson model, model=2 is zero-truncated Poisson, model=3 is Gamma model
model_params	Numeric vector of length 2. Specifies sex-specific parameters for the model of meiosis. Should be in order of male, female
cM_len	Length of chromosome (or subregion) to be simulated in centiMorgans. Integer vector of length 2, first element is cM length for males, second for females
BP_len	Length of chromosome (or subregion) to be simulated in Base pairs. Single integer value (male and female chromosome have same BP length)
physical_map_Mo	Optional map for females (Mother) that specifies relationship between genetic length and physical length. A dataframe with a column named "BP" and another named "cM". The map is the linear piecewise function specified by the points. The order of the pairs should be strictly ascending, the first row should be (0,0) and the last row should be (BP_len, cM_len) If a map is not specified then genetic length is assumed to be constant along the physical length of the region
physical_map_Fa	Optional map for males, should be specified in the same way
seed	integer seed for the random number generator (C++ std library Mersenne twister) for the simulation. Default is the system time

`all_nodes`      `all_nodes=1` will generate Optional "All\_nodes\_haplotypes.txt" output file. Default is `all_nodes=0`.

`outDir`          The directory to write output files to. Default is the current working directory.

**Value**

Will generate output files "Proband\_Haplotypes.txt" and "All\_nodes\_haplotypes.txt" in the specified output directory.

**See Also**

[gen.genealogy](#) [gen.simuHaplo\\_traceback](#) [gen.simuHaplo\\_IBD\\_compare](#) [gen.simuSampleFreq](#)

**Examples**

```
gen_table <- data.frame(ind=c(1,2,3,4,5,6,7),
                       mother=c(0,0,2,0,0,5,3),
                       father=c(0,0,1,0,0,4,6),
                       sex=c(1,2,2,1,2,1,1)
)

#convert the genealogy table into a GENLIB object
sample_gen<-gen.genealogy(gen_table)

gen.graph(sample_gen)
# Simulate the gene dropping from founders to probands

gen.simuHaplo(sample_gen, simulNo=5, model = 1, model_params = c(1,1), cM_len=c(100,100),
              BP_len = 100000000, outDir=tempdir())

f<-file(paste(tempdir(),"Proband_Haplotypes.txt",sep="/"),"r")
print(readLines(f,n=10))
```

---

`gen.simuHaplo_convert`    *Convert proband simulation results into sequence data given founder haplotypes*

---

**Description**

Convert the "Proband\_Haplotypes.txt" output file produced by `gen.simuHaplo` into sequence data according to input sequence data for founders.

**Usage**

```
gen.simuHaplo_convert(dir)
```

## Arguments

`dir` path to the directory containing all the required input files. The function expects to find the "Proband\_Haplotypes.txt" file, a file named "founders.hap" which contains the sequence data for all founder chromosomes, and another file called "founders.map" which contains the BP position of all included markers in the specified founder sequences.

## Details

See [https://github.com/R-GENLIB/simuhaplo\\_functions](https://github.com/R-GENLIB/simuhaplo_functions) for examples on how to format the founder sequence input files.

## Value

creates a new file in the specified directory "Proband\_Genotypes.txt"

## See Also

[gen.genealogy](#) [gen.simuHaplo](#)

## Examples

```
#gen_table <- data.frame(ind=c(1,2,3,4,5,6,7,8),
#                         mother=c(0,0,2,0,0,5,3,3),
#                         father=c(0,0,1,0,0,4,6,6),
#                         sex=c(1,2,2,1,2,1,1,2))

#convert the genealogy table into a GENLIB object
#sample_gen<-gen.genealogy(gen_table)

# Simulate the gene dropping from founders to probands
#gen.simuHaplo(sample_gen, simulNo=5, model = 1, model_params = c(1,1), cM_len=c(100,100),
#             BP_len = 100000000, outDir=tempdir())

#after creating a "founders.map" and a "founders.hap" file in the same directory then can ran
#gen.simuHaplo_convert(dir=tempdir())
```

---

gen.simuHaplo\_IBD\_compare

*Compare proband haplotypes for IBD sharing*

---

## Description

Analyze the simulated haplotypes produced by `gen.simuHaplo` and identify segments of IBD sharing between the diploid haplotypes of a pair of probands

**Usage**

```
gen.simuHaplo_IBD_compare(proID_1, proID_2, BP_len, proband_haplotypes_path)
```

**Arguments**

proID_1	integer ID of a proband
proID_2	integer ID of the second proband
BP_len	The chromosome BP length used in the simulation
proband_haplotypes_path	String that gives the location of the "Proband_haplotypes.txt" output file produced by gen.simuHaplo

**Value**

returns a dataframe with the following columns: "simulNo", "n\_seg", "pIBD", "mean\_seg\_len". Each row is a simulation wherein the pair of probands had some IBD sharing. The "simulNo" columns gives the simulation number, "n\_seg" is the number of non-contiguous segments shared between the pair (divide this value by 2 for an average value for the pair), "pIBD" is the percent of the simulated chromosome shared IBD between the pair, and "mean\_seg\_len" is the mean length of IBD segments. Additionally, the function will print the exact location of each IBD segment to the R console. This can be toggled by disabling messages.

**See Also**

[gen.genealogy](#) [gen.simuHaplo](#) [gen.simuHaplo\\_traceback](#)

**Examples**

```
gen_table <- data.frame(ind=c(1,2,3,4,5,6,7,8),
                      mother=c(0,0,2,0,0,5,3,3),
                      father=c(0,0,1,0,0,4,6,6),
                      sex=c(1,2,2,1,2,1,1,2)
)

#convert the genealogy table into a GENLIB object
sample_gen<-gen.genealogy(gen_table)

gen.graph(sample_gen)
# Simulate the gene dropping from founders to probamds

gen.simuHaplo(sample_gen, simulNo=5, model = 1, model_params = c(1,1), cM_len=c(100,100),
              BP_len = 100000000, outDir=tempdir())

IBD_compare <- gen.simuHaplo_IBD_compare(7, 8, 100000000,
                                         paste(tempdir(), "Proband_Haplotypes.txt", sep="/"))
```



---

 gen.simuHaplo\_traceback

*Trace inheritance path for results from gene dropping simulation*


---

## Description

Given the output of a gene dropping simulation using `gen.simuHaplo`, this function can trace the inheritance path of a proband segment back to the founder it is inherited from

## Usage

```
gen.simuHaplo_traceback(gen, proID, ancestorID, all_nodes_path, proband_haplotypes_path)
```

## Arguments

<code>gen</code>	An object of class <code>GLgen</code> obtained with <code>gen.genealogy</code> , <code>gen.lineages</code> or <code>gen.branching</code> . Required.
<code>proID</code>	Integer ID of the proband to trace back from.
<code>ancestorID</code>	Integer ID of the founder, any segment in the proband haplotype inherited from this founder will be traced back
<code>all_nodes_path</code>	String of the path to the "All_nodes_haplotypes.txt" output file generated by <code>gen.simuHaplo</code>
<code>proband_haplotypes_path</code>	String of the path to the "Proband_Haplotypes.txt" output file generated by <code>gen.simuHaplo</code>

## Value

returns a dataframe with the following columns: "simulNo", "seg\_length", "path\_no". For every simulation in the output files where the specified proband inherits a segment from the specified founder. "seg\_length" is the length of the inherited segment in BP. "path\_no" is an integer that specifies the path of inheritance of the segment Unique paths of inheritance are simply identified by the order they appear in the outputs file, and the full paths (ID of every individual along the path) are printed to the R console. If this is not appearing check that messages are not disabled.

## See Also

[gen.genealogy](#) [gen.simuHaplo](#) [gen.simuHaplo\\_IBD\\_compare](#)

## Examples

```
gen_table <- data.frame(ind=c(1,2,3,4,5,6,7),
                        mother=c(0,0,2,0,0,5,3),
                        father=c(0,0,1,0,0,4,6),
                        sex=c(1,2,2,1,2,1,1)
                        )
```

```
#convert the genealogy table into a GENLIB object
sample_gen<-gen.genealogy(gen_table)

gen.graph(sample_gen)
# Simulate the gene dropping from founders to probamds

gen.simuHaplo(sample_gen, simulNo=5, model = 1, model_params = c(1,1), cM_len=c(100,100),
              BP_len = 100000000, all_nodes = 1, outDir=tempdir())

traceback <-gen.simuHaplo_traceback(sample_gen, 7, 1,
                                   paste(tempdir(), "All_nodes_haplotypes.txt", sep = "/"),
                                   paste(tempdir(), "Proband_Haplotypes.txt", sep="/"))
```

---

gen.simuProb

*Gene dropping simulations - Probabilities*


---

### Description

Returns the probabilities that specified probands inherit disease alleles from ancestors.

### Usage

```
gen.simuProb(gen, pro, statePro, ancestors, stateAncestors, simulNo=5000,
             probRecomb=c(0,0), probSurvival=1.0)
```

### Arguments

gen	An object of class GLgen obtained with gen.genealogy, gen.lineages or gen.branching. Required.
pro	Vector of proband id numbers to be included. Required.
statePro	Required vector indicating, for each proband in pro, the probability to be calculated: 0 = no disease allele is transmitted 1 = 1 disease allele is transmitted 2 = 2 disease alleles are transmitted 3 = 1 or 2 disease alleles are transmitted
ancestors	Vector of ancestors id numbers to be included. Required.
stateAncestors	Required vector indicating, for each ancestor in ancestors, the genotype state: 0 = no disease allele present 1 = 1 disease allele present 2 = 2 disease alleles present
simulNo	Number of simulations to perform. Default is 5000.
probRecomb	Recombination probabilities for males and females. Default is no recombination.
probSurvival	Survival probability for homozygotes. Default is 1.

**Value**

A list containing the following:

- the joint probability of specified statePro for all probands,
- the probability of specified statePro for each proband,
- the probability that, 0, 1, ..., and all probands inherit the specified number of disease alleles.

**See Also**

[gen.genealogy](#) [gen.simuSample](#) [gen.simuSet](#) [gen.simuSampleFreq](#)

**Examples**

```
data(geneaJi)
genJi<-gen.genealogy(geneaJi)
# Probability that subjects 1 and 29 get 1 and 2 alleles from ancestors 20 and 25,
# that have themselves 2 and 1.
gen.simuProb(genJi, pro=c(1,29), statePro=c(1,2), ancestors=c(20,25), stateAncestors=c(2,1),
             simulNo=10000)
# Probability that subjects 1 and 29 get 1 segment from ancestors 25,
# knowing the segment has a male
# recombination rate of 0.02 and a female recombination rate of 0.04.
gen.simuProb(genJi, pro=c(1,29), statePro=c(1,1), ancestors=c(25), stateAncestors=c(1),
             simulNo=10000, probRecomb = c(0.02, 0.04))
# Probability that subjects 1 and 29 get 1 and 2 alleles from ancestors 20 and 25,
# that have themselves 2 and 1 and knowing that homozygous people have a survival rate
# of 0.50.
gen.simuProb(genJi, pro=c(1,29), statePro=c(1,2), ancestors=c(20,25), stateAncestors=c(2,1),
             simulNo=10000, probSurvival=0.5)
```

---

gen.simuSample

*Gene dropping simulations - Sample*

---

**Description**

Returns the number of alleles transmitted to specified probands from ancestors

**Usage**

```
gen.simuSample(gen, pro, ancestors, stateAncestors, simulNo = 5000)
```

**Arguments**

gen	An object of class GLgen obtained with gen.genealogy, gen.lineages or gen.branching. Required.
pro	Vector of proband id numbers to be included. Required.
ancestors	Vector of ancestors id numbers to be included. Required.

stateAncestors Required vector indicating, for each ancestor in ancestors, the genotype state:  
 0 = no disease allele present  
 1 = 1 disease allele present  
 2 = 2 disease alleles present

simulNo Number of simulations to perform. Default is 5000.

**Value**

A matrix with number of columns equal to the number of simulations and number of rows equal to the number of probands.

**See Also**

[gen.genealogy](#) [gen.simuProb](#) [gen.simuSet](#) [gen.simuSampleFreq](#)

**Examples**

```
data(geneaJi)
genJi<-gen.genealogy(geneaJi)
simu_1000<-gen.simuSample(genJi, pro=c(1,29), ancestors=c(20,25), stateAncestors=c(2,1),
                          simulNo=1000)
# Number of alleles received by probands
table(simu_1000)
# Number of alleles received by simulation
table(colSums(simu_1000))
# Number of alleles received by each probant
table(simu_1000[1,],simu_1000[2,])

data(geneaJi)
genJi<-gen.genealogy(geneaJi)
simu_5000<-gen.simuSample(genJi, pro=c(1,29), ancestors=c(20,25), stateAncestors=c(2,1),
                          simulNo=5000)
# Number of alleles received by probands
table(simu_5000)
# Number of alleles received by simulation
table(colSums(simu_5000))
# Number of alleles received by each probant
table(simu_5000[1,],simu_5000[2,])
```

---

gen.simuSampleFreq      *Gene dropping simulations - Frequencies*

---

**Description**

Returns the number of alleles transmitted to specified probands from ancestors

**Usage**

```
gen.simuSampleFreq( gen, pro, ancestors, stateAncestors, simulNo = "5000")
```

**Arguments**

gen	An object of class GLgen obtained with gen.genealogy, gen.lineages or gen.branching. Required.
pro	Vector of proband id numbers to be included. Required.
ancestors	Vector of ancestors id numbers to be included. Required.
stateAncestors	Required vector indicating, for each ancestor in ancestors, the genotype state: 0 = no disease allele present 1 = 1 disease allele present 2 = 2 disease alleles present
simulNo	Number of simulations to perform. Default is 5000.

**Value**

A data frame containing for each row (one row per proband): **Alleles.transmitted.0 Alleles.transmitted.1 Alleles.transmitted.2**

**See Also**

[gen.genealogy](#) [gen.simuProb](#) [gen.simuSet](#) [gen.simuSample](#)

**Examples**

```
data(geneaJi)
genJi<-gen.genealogy(geneaJi)
gen.simuSampleFreq(genJi, pro=c(1,29), ancestors=c(20,25),stateAncestors=c(2,1), simulNo = 1000)
```

---

gen.simuSet

*Gene dropping simulations with specified transmission probabilities*

---

**Description**

Returns the number of alleles transmitted to specified probands from ancestors considering specified transmission probabilities.

**Usage**

```
gen.simuSet(gen, pro, ancestors, stateAncestors,
            probMatrix=matrix(c(
                c(1,0.5,0,0.5,0.25,0,0,0,0,1,1,1,1,0.75,0.5,1,0.5,0),
                c(1,0.5,0,0.5,0.25,0,0,0,0,1,1,1,1,0.75,0.5,1,0.5,0)),
                nrow=3, ncol=12),
            simulNo = 5000)
```

**Arguments**

gen	An object of class GLgen obtained with gen.genealogy, gen.lineages or gen.branching. Required.
pro	Vector of proband id numbers to be included. Required.
ancestors	Vector of ancestors id numbers to be included. Required.
stateAncestors	Required vector indicating, for each ancestor in ancestors, the genotype state: 0 = no disease allele present 1 = 1 disease allele present 2 = 2 disease alleles present
probMatrix	Matrix of transmission probabilities of 0, 1, or 2 disease alleles. Default is: [,1] [,2] [,3] [,4] [,5] [,6] [,7] [,8] [,9] [,10] [,11] [,12] [1,] 1.0 0.50 0 1 1.00 1.0 1.0 0.50 0 1 1.00 1.0 [2,] 0.5 0.25 0 1 0.75 0.5 0.5 0.25 0 1 0.75 0.5 [3,] 0.0 0.00 0 1 0.50 0.0 0.0 0.00 0 1 0.50 0.0 where rows are genotype states (0,1,2) for the father and columns are states for the mother (in blocks of 3). 1st block (col 1-3) are the probabilities of transmitting 0 allele to a son. 2nd block (col 4-6) are the probabilities of transmitting 1 allele to a son. 3rd block (col 7-9) are the probabilities of transmitting 0 allele to a daughter. 4th block (col 10-12) are the probabilities of transmitting 1 allele to a daughter.
simulNo	Number of simulations to perform. Default is 5000.

**Value**

A matrix with number of columns equal to the number of simulations and number of rows equal to the number of probands.

**See Also**

[gen.genealogy](#) [gen.simuProb](#) [gen.simuSample](#)

**Examples**

```
# A case where only male subjects can receive alleles
data(geneaJi)
genJi<-gen.genealogy(geneaJi)
onlyThroughMale<-matrix(c(c(1,0.5,0,0.5,0.25,0,0,0,0,1,1,1,1,0.75,0.5,1,0.5,0), rep(1,18)),
  nrow=3, ncol=12)
gen.graph(genJi, indVarAffected=c(28,27,25,17,12,8,4,1), varAffected=c(28,27,25,17,12,8,4,1))
simu_1000a<-gen.simuSet(genJi, pro=c(1,28), ancestors=c(17,25), stateAncestors=c(1,1),
  simulNo = 1000,probMatrix=onlyThroughMale)
# Number of alleles received by probands
table(simu_1000a)
# Number of alleles received by all probands at each simulation
table(colSums(simu_1000a))
# Number of alleles received by each probant
```

```

table(simu_1000a[1,],simu_1000a[2,])

# A case where subjects are limited to one copy compared to what is normally expected
max1Allreceived<-matrix(rep(c(1,0.5,0,0.5,0.25,0,0,0,0,1,1,1,1,1,1,1,1), 2) , nrow=3, ncol=12)
gen.graph(genJi)
simu_1000b<-gen.simuSet(genJi, pro=c(1,29), ancestors=25, stateAncestors=1, simulNo=10000,
                        probMatrix=max1Allreceived)
# Normal case matrix not changed
simu_1000original<-gen.simuSet(genJi, pro=c(1,29), ancestors=25, stateAncestors=1, simulNo=10000)
# Number of alleles received by each probant
table(simu_1000b)
table(simu_1000original)
# Number of alleles received by all probants at each simulation
table(colSums(simu_1000b))
table(colSums(simu_1000original))
# Number of alleles received by each probant
table(simu_1000b[1,],simu_1000b[2,])
table(simu_1000original[1,],simu_1000original[2,])

```

---

genea140

*Genealogical information for 140 individuals from the Quebec Reference Sample*

---

## Description

A genealogical corpus made of 41523 individuals from the province of Quebec, Canada. A total of 140 individuals have been sampled in seven sub-populations, listed in pop140, and their genealogies were reconstructed as far back as possible using the BALSAC population register and the Early Quebec Population Register.

## Usage

```
data(gene140)
```

## Format

A data frame with 41523 observations on 11 variables.

[,1] ind: An individual's ID number

[,2] father: This individual's father ID number

[,3] mother: This individual's mother ID number

[,4] sex: Individual's sex coded 1/2 for male/female

## Source

Balsac

**References**

<http://www.quebecgenpop.ca>

Roy-Gagnon, M.-H., Moreau, C., Bherer, C., St-Onge, P., Sinnett, D., Laprise, C., Vezina, H., Labuda, D. (2011). Genomic and genealogical investigation of the French Canadian founder population structure. *Human Genetics*, 129(5), 521-31.

---

geneaJi

*Highly inbred pedigree*

---

**Description**

A modified version of a pedigree of two Jicaque Indians studied by Chapman & Jacquard (1971).

**Usage**

`data(geneaJi)`

**Format**

A data frame with 29 observations on 4 variables.

[, 1] ind An individual's ID number

[, 2] father This individual's father ID number

[, 3] mother This individual's mother ID number

[, 4] sex Individual's sex coded 1/2 for male/female

**Source**

CHAPMAN & JACQUART (1971)

**References**

Chapman, A.M., & Jacquard, A. (1971). Un isolat d'Amérique centrale: Les indiens Jicaques du Honduras. Paris: Presses universitaires de France.



---

 GLgen-class

 Class "GLgen"
 

---

**Description**

Object containing a genealogy.

**Objects from the Class**

Objects can be created by calls of the form `gen.genealogy('data.frame')`.

**Extends**

Class "GLgroup", directly.

**Methods**

**initialize** signature(.Data="integer",Date="character"): ...

**depth** signature(x = "GLgen") Gives the depth of the genealogy.

**length** signature(x = "GLgen") Gives the number of individuals in the genealogy.

**Examples**

```
showClass("GLgen")
```

---

 GLgroup-class

 Class "GLgroup"
 

---

**Description**

Object representing a set of proband in different groups.

Each element of this list is a group named with the proband number.

**Objects from the Class**

Objects can be created by calls of the form `new("GLgroup", 'list')`.

**Extends**

Class "GLgen", directly.

**Methods**

**initialize** signature(.Data=list): ...

[ signature(object = "GLgroup", ANY, ANY, ANY):...

**Examples**

```
showClass("GLgroup")
```

---

```
GLmultiList-class      Class "GLmultiList"
```

---

**Description**

Objects created to carry information mostly between invisible functions.

**Objects from the Class**

Objects can be created by calls of the form `new("GLmultiList", 'Array')`.

**Slots**

.Data: Object of class "list" ~~

liste: Object of class "list" ~~

**Extends**

Class "[list](#)", from data part. Class "[vector](#)", by class "list", distance 2.

**Methods**

No methods defined with class "GLmultiList" in the signature.

**Examples**

```
showClass("GLmultiList")
```

---

```
pop140                Population of origin of the 140 Quebec samples
```

---

**Description**

The 140 individuals from the genealogical corpus from Quebec were sampled from 7 different populations from 5 regions; Quebec City, Montreal, Saguenay, North Shore, Gaspesia. In Gaspesia we find 3 different populations: French-Canadians, Acadians and Loyalists.

**Usage**

```
data(pop140)
```

**Format**

[, 1] ind: An individual's ID number

[, 2] pop: This individual's population

**Source**

Balsac

**References**

<http://www.quebecgenpop.ca/home.html>

Roy-Gagnon, M.-H., Moreau, C., Bherer, C., St-Onge, P., Sinnett, D., Laprise, C., Vezina, H., Labuda, D. (2011). Genomic and genealogical investigation of the French Canadian founder population structure. *Human Genetics*, 129(5), 521-31.

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