Package 'COMA'

June 29, 2024

	0000 = 5, = 0 = 1
Title Conve	ex Optimization of Mate Allocation
Version 0.1	17
Author Jef	frey B. Endelman
Maintainer	· Jeffrey Endelman <endelman@wisc.edu></endelman@wisc.edu>
Description	Convex Optimization of Mate Allocation
Depends R	A (>= 4.0)
License GI	PL-3
RoxygenNo	ote 7.2.3
Encoding	UTF-8
Imports gg	gplot2, CVXR, tidyr
Suggests k	nitr, rmarkdown, Rmosek
oc on ple	s documented: s
ocs	Optimal Contribution Selection
Description Optimize Usage	n ze the contribution of each individual to the next generation
ocs(dF	, parents, ploidy, K, tol = 1e-06, dF.adapt = NULL, solver = "ECOS")

2 oma

Arguments

dF inbreeding rate

parents input data frame (see Details)

ploidy ploidy

K kinship matrix

tol tolerance, values below this set to 0

dF.adapt see Details

solver solver for CVXR (default is "ECOS")

Details

The first four columns of parents should be named as follows: id, merit, min, max. Min and max are real numbers between 0 and 1 specifying the minimum and maximum contribution for each parent. An optional fifth column named "female" is a logical TRUE/FALSE variable for species with separate sexes.

Additional columns can be used to specify constraints on linear combinations of the contributions. The values are the coefficients of the contribution variables, and the name of each column specifies the right-hand side of the constraint. Each name must begin with "lt", "gt", or "eq", followed by a non-negative numeric value. For example, "lt0.5" means less than or equal to 0.5.

The average inbreeding coefficient of the current generation is based on all individuals in K, which may exceed the list of individuals in parents.

It is possible that no feasible solution exists for the specified dF. Argument dF. adapt can be used to automatically progressively higher values. The software increases the dF limit by dF. adapt\$step up to the smaller of dF. adapt\$max or the realized value under the original dF, in an attempt to find a solution with less inbreeding.

Value

list containing

response data.frame with realized dF, merit, n.parent

oc data frame of optimal contributions

oma

Optimal Mate Allocation

Description

Optimize the allocation for each mating

Usage

```
oma(
   dF,
   parents,
   matings,
   ploidy,
   K,
```

oma 3

```
tol = 1e-06,
  dF.adapt = NULL,
  solver = "ECOS"
)
```

Arguments

dF inbreeding rate

parents data frame (see Details)

matings matings data frame (see Details)

ploidy ploidy

K kinship matrix

tol tolerance, values below this set to 0

dF.adapt see Details

solver solver for CVXR (default is "ECOS")

Details

The first three columns of parents should be named "id", "min", "max", with an optional fourth column "female" to indicate sex in dioecious species. Additional columns can be used to specify constraints on linear combinations of the contributions. The values are the coefficients of the contribution variables, and the name of each column specifies the right-hand side of the constraint. Each name must begin with "lt", "gt", or "eq", followed by a non-negative numeric value. For example, "lt0.5" means less than or equal to 0.5.

The data.frame matings has five columns: "female, male, merit, min, max" for dioecious species, or else "parent1, parent2, merit, min, max".

The average inbreeding coefficient of the current generation is based on all individuals in K, which may exceed the list of individuals in parents.

It is possible that no feasible solution exists for the specified dF. Argument dF. adapt can be used to automatically progressively higher values. The software increases the dF limit by dF. adapt\$step up to the smaller of dF. adapt\$max or the realized value under the original dF, in an attempt to find a solution with less inbreeding.

Value

list containing

response data.frame with realized dF, merit, n.parent, n.mate

oc data frame of optimal contributions for each individual

om data frame of optimal allocations for each mating

4 read_data

plot_ribbon

Ribbon Plot of Optimal Contributions

Description

Ribbon Plot of Optimal Contributions

Usage

```
plot_ribbon(oc, min.c = 0.001)
```

Arguments

oc named list of optimal contributions
min.c minimum contribution for plotting

Details

The inbreeding rates are taken from the names of oc. Each element of oc should be a data frame with columns "id" and "value".

Value

ggplot2 object

read_data

Read data for OCS/OMA

Description

Predicts merit for OCS/OMA

Usage

```
read_data(
  geno.file,
  kinship.file,
  ploidy,
  sex = NULL,
  matings = "none",
  standardize = FALSE,
  n.core = 1
)
```

read_data 5

Arguments

geno.file file with marker effects and genotypes

kinship.file file with kinship matrix

ploidy even integer

sex optional, data frame with columns id and female (T/F)

matings see Details

standardize T/F, standardize merit in parental candidates

n.core multi-core evaluation

Details

The first column of geno. file is the marker name. The second column contains the additive effects for the breeding value parameterization, and (digenic) dominance effects (when available) should be in the third column with the header "dom". Subsequent columns contain the marker data for the population, coded as allele dosage, from 0 to ploidy. Missing marker data is imputed with the population mean.

The kinship file should contain an N x N kinship matrix with id names in the first column and row.

There are several options for argument matings: (1) "none" = no matings; (2) "all" = all possible matings of the individuals in geno.file (excluding reciprocals); (3) a character vector of genotype ids to calculate all pairs of matings; (4) a 2-column data.frame of desired matings with header "female", "male" for dioecious species (separate sexes) or "parent1", "parent2" for hermaphrodites. Self-matings are included under options (2) and (3) for dioecious species but are easily removed in the output if desired.

Value

list containing

K kinship matrix

parents data frame of individual meritsmatings data frame of mating merits

Index

```
ocs, 1
oma, 2
plot_ribbon, 4
read_data, 4
```