# Package 'corehunter'

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Title Multi-Purpose Core Subset Selection

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**Description** Core Hunter is a tool to sample diverse, representative subsets from large germplasm collections, with minimum redundancy. Such so-called core collections have applications in plant breeding and genetic resource management in general. Core Hunter can construct cores based on genetic marker data, phenotypic traits or precomputed distance matrices, optimizing one of many provided evaluation measures depending on the precise purpose of the core (e.g. high diversity, representativeness, or allelic richness). In addition, multiple measures can be simultaneously optimized as part of a weighted index to bring the different perspectives closer together. The Core Hunter library is implemented in Java 8 as an open source project (see <a href="http://www.corehunter.org">http://www.corehunter.org</a>).

**Depends** R (>= 3.2.3), rJava (>= 0.9-8)

**Imports** naturalsort (>= 0.1.2), methods

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Author Herman De Beukelaer [aut, cre], Guy Davenport [aut], Veerle Fack [ths]

Maintainer Herman De Beukelaer < herman.debeukelaer@gmail.com>

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## **Description**

Core Hunter is a tool to sample diverse, representative subsets from large germplasm collections, with minimum redundancy. Such so-called core collections have applications in plant breeding and genetic resource management in general. Core Hunter can construct cores based on genetic marker data, phenotypic traits or precomputed distance matrices, optimizing one of many provided evaluation measures depending on the precise purpose of the core (e.g. maximum diversity, representativeness, or allelic richness). In addition, multiple measures can be simultaneously optimized as part of a weighted index to bring the different perspectives closer together. The Core Hunter library is implemented in Java 8 as an open source project (see http://www.corehunter.org).

#### See Also

coreHunterData, genotypes, phenotypes, distances, sampleCore, evaluateCore, objective

## Examples

```
## Not run:
# sample core based on genetic marker data (default format)
geno.file <- system.file("extdata", "genotypes.csv", package = "corehunter")
geno <- genotypes(file = geno.file, format = "default")
core <- sampleCore(geno)

# sample core based on genetic marker data (biparental format)
geno.file <- system.file("extdata", "genotypes-biparental.csv", package = "corehunter")
geno <- genotypes(file = geno.file, format = "biparental")
core <- sampleCore(geno)</pre>
```

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```
# sample core based on genetic marker data (frequency format)
geno.file <- system.file("extdata", "genotypes-frequency.csv", package = "corehunter")</pre>
geno <- genotypes(file = geno.file, format = "frequency")</pre>
core <- sampleCore(geno)</pre>
# sample core based on phenotypic traits
pheno.file <- system.file("extdata", "phenotypes.csv", package = "corehunter")</pre>
pheno <- phenotypes(file = pheno.file)</pre>
core <- sampleCore(pheno)</pre>
# sample core based on precomputed distance matrix
dist.file <- system.file("extdata", "distances.csv", package = "corehunter")</pre>
dist <- distances(file = dist.file)</pre>
core <- sampleCore(dist)</pre>
# sample core from genotypes with custom objective (allelic richness)
core <- sampleCore(geno, obj = objective("HE"))</pre>
# sample core from genotypes with custom size and objective (representativeness)
core <- sampleCore(geno, obj = objective("AN", "MR"), size = 0.1)</pre>
# sample core from genotypes with custom size and stop condition
core <- sampleCore(geno, size = 0.1, impr.time = 2)</pre>
# sample core based on both genotypes and phenotypes
geno.pheno <- coreHunterData(geno, pheno)</pre>
core <- sampleCore(geno.pheno)</pre>
## End(Not run)
```

coreHunterData

Initialize Core Hunter data.

## **Description**

The data may contain genotypes, phenotypes and/or a precomputed distance matrix. All provided data should describe the same individuals which is verified by comparing the item ids and names.

#### Usage

```
coreHunterData(genotypes, phenotypes, distances)
```

#### **Arguments**

genotypes Genetic marker data (chgeno).

phenotypes Phenotypic trait data (chpheno).

distances Precomputed distance matrix (chdist).

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#### Value

Core Hunter data (chdata) with elements

geno Genotype data of class chgeno if included.

pheno Phenotype data of class chpheno if included.

dist Distance data of class chdist if included.

size Number of individuals in the dataset.

ids Unique item identifiers.

names Item names. Names of individuals to which no explicit name has been assigned are equal to the unique ids.

java Java version of the data object.

Core Hunter data of class chdata.

#### See Also

```
genotypes, phenotypes, distances
```

## **Examples**

```
geno.file <- system.file("extdata", "genotypes.csv", package = "corehunter")
pheno.file <- system.file("extdata", "phenotypes.csv", package = "corehunter")
dist.file <- system.file("extdata", "distances.csv", package = "corehunter")

my.data <- coreHunterData(
    genotypes(file = geno.file, format = "default"),
    phenotypes(file = pheno.file),
    distances(file = dist.file)
)</pre>
```

distances

Create Core Hunter distance data from matrix or file.

## **Description**

Specify either a symmetric distance matrix or the file from which to read the matrix. See www.corehunter.org for documentation and examples of the distance matrix file format used by Core Hunter.

## Usage

```
distances(data, file)
```

evaluateCore 5

## **Arguments**

data Symmetric distance matrix. Unique row and column headers are required, should

be the same and are used as item ids. Can be a numeric matrix or a data frame. The data frame may optionally include a first column NAME used to assign names

to some or all individuals. The remaining columns should be numeric.

file File from which to read the distance matrix.

#### Value

Distance matrix data of class chdist with elements

data Distance matrix (numeric matrix).

size Number of individuals in the dataset.

ids Unique item identifiers.

names Item names. Names of individuals to which no explicit name has been assigned are equal to the unique ids.

java Java version of the data object.

file Normalized path of file from which data was read (if applicable).

## **Examples**

```
# create from distance matrix
m <- matrix(runif(100), nrow = 10, ncol = 10)
diag(m) <- 0
# make symmetric
m[lower.tri(m)] <- t(m)[lower.tri(m)]
# set headers
rownames(m) <- colnames(m) <- paste("i", 1:10, sep = "-")
dist <- distances(m)
# read from file
dist.file <- system.file("extdata", "distances.csv", package = "corehunter")
dist <- distances(file = dist.file)</pre>
```

evaluateCore

Evaluate a core collection using the specified objective.

## Description

Evaluate a core collection using the specified objective.

#### Usage

```
evaluateCore(core, data, objective)
```

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#### **Arguments**

core A core collection of class choore, or a numeric or character vector indicating

the indices or ids, respectively, of the individuals in the evaluated core.

data Core Hunter data (chdata) containing genotypes, phenotypes and/or a precom-

puted distance matrix. Can also be an object of class chdist, chgeno or chpheno

if only one type of data is provided.

objective Objective function (chobj) used to evaluate the core.

#### Value

Value of the core when evaluated with the chosen objective (numeric).

#### See Also

coreHunterData, objective

## **Examples**

```
data <- exampleData()
core <- sampleCore(data, objective("EN", "PD"))
evaluateCore(core, data, objective("EN", "PD"))
evaluateCore(core, data, objective("AN", "MR"))
evaluateCore(core, data, objective("EE", "GD"))
evaluateCore(core, data, objective("CV"))
evaluateCore(core, data, objective("HE"))</pre>
```

exampleData

Small example dataset with 218 individuals.

## **Description**

Data was genotyped using 190 SNP markers and 4 quantitative traits were recorded. Includes a precomputed distance matrix read from "extdata/distances.csv", genotypes read from "extdata/genotypes-biparental. and phenotypes read from "extdata/phenotypes.csv". The distance matrix is computed from the genotypes (Modified Rogers' distance).

## Usage

```
exampleData()
```

#### **Details**

Data was taken from the CIMMYT Research Data Repository (Study Global ID hdl:11529/10199; real data set 5, cycle 0).

genotypes 7

#### Value

Core Hunter data of class chdata

#### Source

Cerón-Rojas, J. Jesús; Crossa, José; Arief, Vivi N.; Kaye Basford; Rutkoski, Jessica; Jarquín, Diego; Alvarado, Gregorio; Beyene, Yoseph; Semagn, Kassa; DeLacy, Ian, 2015-06-04, "Application of a Genomics Selection Index to Real and Simulated Data", http://hdl.handle.net/11529/10199 V10

#### **Examples**

exampleData()

genotypes

Create Core Hunter genotype data from data frame, matrix or file.

## **Description**

Specify either a data frame or matrix, or a file from which to read the genotypes. See www.corehunter.org for documentation and examples of the genotype data file format used by Core Hunter

#### **Usage**

```
genotypes(data, alleles, file, format)
```

## Arguments

data

Data frame or matrix containing the genotypes (individuals x markers) depending on the chosen format:

default Data frame. One row per individual and one or more columns per marker. Columns contain the names, numbers, references, ... of observed alleles. Unique row names (item ids) are required and columns should be named after the marker to which they belong, optionally extended with an arbitrary suffix starting with a dot (.), dash (-) or underscore (\_) character.

biparental Numeric matrix or data frame. One row per individual and one column per marker. Data consists of 0, 1 and 2 coding for homozygous (AA), heterozygous (AB) and homozygous (BB), respectively. Unique row names (item ids) are required and optionally column (marker) names may be included as well.

frequency Numeric matrix or data frame. One row per individual (or bulk sample) and multiple columns per marker. Data consists of allele frequencies, grouped per marker in consecutive columns named after the corresponding marker, optionally extended with an arbitrary suffix starting with a dot (.), dash (-) or underscore (\_) character.. The allele frequencies of each

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marker should sum to one in each sample. Unique row names (item ids) are required.

In case a data frame is provided, an optional first column NAME may be included to specify item names. The remaining columns should follow the format as described above. See <a href="www.corehunter.org">www.corehunter.org</a> for more details about the supported genotype formats. Note that both the frequency and biparental format syntactically also comply with the default format but with different semantics, meaning that it is very important to specify the correct format. Some checks have been built in that raise warnings in case it seems that the wrong format might have been specified based on an inspection of the data. If you are sure that you have selected the correct format these warnings, if any, can be safely ignored.

alleles Allele names per marker (character vector). Ignored except when creating

frequency data from a matrix or data frame. Allele names should be ordered in

correspondence with the data columns.

file File containing the genotype data.

format Genotype data format, one of default, biparental or frequency.

#### Value

Genotype data of class chgeno with elements

data Genotypes. Data frame for default format, numeric matrix for other formats.

size Number of individuals in the dataset.

ids Unique item identifiers (character).

names Item names (character). Names of individuals to which no explicit name has been assigned are equal to the unique ids.

markers Marker names (character). May contain NA values in case only some or no marker names were specified. Marker names are always included for the default and frequency format but are optional for the biparental format.

alleles List of character vectors with allele names per marker. Vectors may contain NA values in case only some or no allele names were specified. For biparental data the two alleles are name "0" and "1", respectively, for all markers. For the default format allele names are inferred from the provided data. Finally, for frequency data allele names are optional and may be specified either in the file or through the alleles argument when creating this type of data from a matrix or data frame.

java Java version of the data object.

format Genotype data format used.

file Normalized path of file from which data was read (if applicable).

## **Examples**

```
# create from data frame or matrix
# default format
geno.data <- data.frame(</pre>
```

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```
NAME = c("Alice", "Bob", "Carol", "Dave", "Eve"),
M1.1 = c(1,2,1,2,1),
M1.2 = c(3,2,2,3,1),
M2.1 = c("B", "C", "D", "B", NA),
M2.2 = c("B", "A", "D", "B", NA),
M3.1 = c("a1", "a1", "a2", "a2", "a1"),
M3.2 = c("a1", "a2", "a2", "a1", "a1"),
M4.1 = c(NA, "+", "+", "+", "-"),
M4.2 = c(NA, "-", "+", "-", "-"),
row.names = paste("g", 1:5, sep = "-")
geno <- genotypes(geno.data, format = "default")</pre>
# biparental (e.g. SNP)
geno.data <- matrix(</pre>
sample(c(0,1,2), replace = TRUE, size = 1000),
nrow = 10, ncol = 100
)
rownames(geno.data) <- paste("g", 1:10, sep = "-")</pre>
colnames(geno.data) <- paste("m", 1:100, sep = "-")</pre>
geno <- genotypes(geno.data, format = "biparental")</pre>
# frequencies
geno.data <- matrix(</pre>
c(0.0, 0.3, 0.7, 0.5, 0.5, 0.0, 1.0,
   0.4, 0.0, 0.6, 0.1, 0.9, 0.0, 1.0,
   0.3, 0.3, 0.4, 1.0, 0.0, 0.6, 0.4),
byrow = TRUE, nrow = 3, ncol = 7
rownames(geno.data) <- paste("g", 1:3, sep = "-")</pre>
colnames(geno.data) <- c("M1", "M1", "M1", "M2", "M2", "M3", "M3")</pre>
alleles <- c("M1-a", "M1-b", "M1-c", "M2-a", "M2-b", "M3-a", "M3-b")
geno <- genotypes(geno.data, alleles, format = "frequency")</pre>
# read from file
# default format
geno.file <- system.file("extdata", "genotypes.csv", package = "corehunter")</pre>
geno <- genotypes(file = geno.file, format = "default")</pre>
# biparental (e.g. SNP)
geno.file <- system.file("extdata", "genotypes-biparental.csv", package = "corehunter")</pre>
geno <- genotypes(file = geno.file, format = "biparental")</pre>
# frequencies
geno.file <- system.file("extdata", "genotypes-frequency.csv", package = "corehunter")</pre>
geno <- genotypes(file = geno.file, format = "frequency")</pre>
```

## **Description**

Get Allele frequency matrix.

## Usage

```
getAlleleFrequencies(data)
```

## **Arguments**

data

Core Hunter data containing genotypes

## Value

allele frequency matrix

getNormalizationRanges

Determine normalization ranges of all objectives in a multi-objective configuration.

## **Description**

Executes an independent stochastic hill-climbing search (random descent) per objective to approximate the optimal solution for each objective, from which a suitable normalization range is inferred based on the Pareto minima/maxima. These normalization searches are executed in parallel.

## Usage

```
getNormalizationRanges(data, obj, size = 0.2, always.selected = integer(0),
never.selected = integer(0), mode = c("default", "fast"), time = NA,
impr.time = NA, steps = NA, impr.steps = NA)
```

## **Arguments**

data	Core Hunter data (chdata) containing genotypes, phenotypes and/or a precomputed distance matrix. Can also be an object of class chdist, chgeno or chpheno if only one type of data is provided.
obj	List of objectives (chobj). If no objectives are specified Core Hunter maximizes a weighted index including the default entry-to-nearest-entry distance (EN) for each available data type. For genotypes, the Modified Roger's distance (MR) is used. For phenotypes, Gower's distance (GD) is applied.
size	Desired core subset size (numeric). If larger than one the value is used as the absolute core size after rounding. Else it is used as the sampling rate and multiplied with the dataset size to determine the size of the core. The default sampling rate is 0.2.

always.selected

vector with indices (integer) or ids (character) of items that should always be

selected in the core collection

never.selected vector with indices (integer) or ids (character) of items that should never be

selected in the core collection

mode Execution mode (default or fast). In default mode, the normalization searches

terminate when no improvement is found for ten seconds. In fast mode, searches terminate as soon as no improvement is made for two seconds. These stop conditions can be overridden using arguments time, impr.time, steps and/or impr.steps. In default mode, the value of the latter two, step-based conditions is multiplied with 500, in line with the behaviour of sampleCore when

executed in default mode.

time Absolute runtime limit in seconds. Not used by default (NA). If used, it should

be a strictly positive value, which is rounded to the nearest integer.

impr.time Maximum time without improvement in seconds. If no explicit stop conditions

are specified, the maximum time without improvement defaults to ten or two seconds, when executing Core Hunter in default or fast mode, respectively. If a custom improvement time is specified, it should be strictly positive and is

rounded to the nearest integer.

steps Maximum number of search steps. Not used by default (NA). If used, it should

be a strictly positive value, which is rounded to the nearest integer. In default mode, the value is multiplied with 500, in line with the behaviour of sampleCore

when executed in default mode.

impr.steps Maximum number of steps without improvement. Not used by default (NA).

If used, it should be a strictly positive value, which is rounded to the nearest integer. In default mode, the value is multiplied with 500, in line with the

behaviour of sampleCore when executed in default mode.

## **Details**

For an objective that is being maximized, the upper bound is set to the value of the best solution for that objective, while the lower bound is set to the Pareto minimum, i.e. the minimum value obtained when evaluating all optimal solutions (for each single objective) with the considered objective. For an objective that is being minimized, the roles of upper and lower bound are interchanged, and the Pareto maximum is used instead.

Because Core Hunter uses stochastic algorithms, repeated runs may produce different results. To eliminate randomness, you may set a random number generation seed using set.seed prior to executing Core Hunter. In addition, when reproducible results are desired, it is advised to use step-based stop conditions instead of the (default) time-based criteria, because runtimes may be affected by external factors, and, therefore, a different number of steps may have been performed in repeated runs when using time-based stop conditions.

#### Value

Numeric matrix with one row per objective and two columns:

lower Lower bound of normalization range.

upper Upper bound of normalization range.

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#### See Also

coreHunterData, objective

#### **Examples**

```
data <- exampleData()

# maximize entry-to-nearest-entry distance between genotypes and phenotypes (equal weight)
objectives <- list(objective("EN", "MR"), objective("EN", "GD"))
# get normalization ranges for default size (20%)
ranges <- getNormalizationRanges(data, obj = objectives, mode = "fast")

# set normalization ranges and sample core
objectives <- lapply(1:2, function(o){setRange(objectives[[o]], ranges[o,])})
core <- sampleCore(data, obj = objectives)</pre>
```

objective

Create Core Hunter objective.

#### **Description**

The following optimization objectives are supported by Core Hunter:

- EN Average entry-to-nearest-entry distance (default). Maximizes the average distance between each selected individual and the closest other selected item in the core. Favors diverse cores in which each individual is sufficiently different from the most similar other selected item (low redundancy). Multiple distance measures are provided to be used with this objective (see below).
- AN Average accession-to-nearest-entry distance. Minimizes the average distance between each individual (from the full dataset) and the closest selected item in the core (which can be the individual itself). Favors representative cores in which all items from the original dataset are represented by similar individuals in the selected subset. Multiple distance measures are provided to be used with this objective (see below).
- EE Average entry-to-entry distance. Maximizes the average distance between each pair of selected individuals in the core. This objective is related to the entry-to-nearest-entry (EN) distance but less effectively avoids redundant, similar individuals in the core. In general, use of EN is preferred. Multiple distance measures are provided to be used with this objective (see below).
- SH Shannon's allelic diversity index. Maximizes the entropy, as used in information theory, of the selected core. Independently takes into account all allele frequencies, regardless of the locus (marker) where to which the allele belongs. Requires genotypes.
- HE Expected proportion of heterozygous loci. Maximizes the expected proportion of heterozygous loci in offspring produced from random crossings within the selected core. In contrast to Shannon's index (SH) this objective treats each marker (locus) with equal importance, regardless of the number of possible alleles for that marker. Requires genotypes.

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CV Allele coverage. Maximizes the proportion of alleles observed in the full dataset that are retained in the selected core. Requires genotypes.

The first three objective types (EN, AN and EE) aggregate pairwise distances between individuals. These distances can be computed using various measures:

- MR Modified Rogers distance (default). Requires genotypes.
- CE Cavalli-Sforza and Edwards distance. Requires genotypes.
- GD Gower distance. Requires phenotypes.
- PD Precomputed distances. Uses the precomputed distance matrix of the dataset.

## Usage

```
objective(type = c("EN", "AN", "EE", "SH", "HE", "CV"), measure = c("MR", "CE", "GD", "PD"), weight = 1, range = NULL)
```

## **Arguments**

type	Objective type, one of EN (default), AN, EE, SH, HE or CV (see description). The
	former three objectives are distance based and require to choose a distance
	mangura By default Modified Roger's distance is used computed from the

measure. By default, Modified Roger's distance is used, computed from the

genotypes.

measure Distance measure used to compute the distance between two individuals, one of

MR (default), CE, GD or PD (see description). Ignored when type is SH, HE or CV.

weight Weight assigned to the objective when maximizing a weighted index. Defaults

to 1.0.

range Normalization range [l,u] of the objective when maximizing a weighted index.

By default the range is not set (NULL) and will be determined automatically prior to execution, if normalization is enabled (default). Values are rescaled to [0,1] with the linear formula v'=(v-l)/(u-l). When an explicit normalization range is set, it overrides the automatically inferred range. Also, setting the range for all included objectives reduces the computation time when sampling a multi-objective core collection. In case of repeated sampling from the same dataset with the same objectives and size, it is therefore advised to determine the normalization ranges only once using <code>getNormalizationRanges</code> so that they can

be reused for all executions.

#### Value

Core Hunter objective of class chobj with elements

type Objective type.

meas Distance measure (if applicable).

weight Assigned weight.

range Normalization range (if specified).

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#### See Also

getNormalizationRanges, setRange

## **Examples**

```
objective()
objective(meas = "PD")
objective("EE", "GD")
objective("HE")
objective("EN", "MR", range = c(0.150, 0.300))
objective("AN", "MR", weight = 0.5, range = c(0.150, 0.300))
```

phenotypes

Create Core Hunter phenotype data from data frame or file.

### **Description**

Specify either a data frame containing the phenotypic trait observations or a file from which to read the data. See <a href="www.corehunter.org">www.corehunter.org</a> for documentation and examples of the phenotype data format used by Core Hunter.

#### Usage

```
phenotypes(data, types, min, max, file)
```

## **Arguments**

data

Data frame containing one row per individual and one column per trait. Unique row and column names are required and used as item and trait ids, respectively. The data frame may optionally include a first column NAME used to assign names to some or all individuals.

types

Variable types (optional). Vector of characters, each of length one or two. Ignored when reading from file.

The first letter indicates the scale type and should be one of N (nominal), 0 (ordinal), I (interval) or R (ratio).

The second letter optionally indicates the variable encoding (in Java) and should be one of B (boolean), T (short), I (integer), L (long), R (big integer), F (float), D (double), M (big decimal), A (date) or S (string). The default encoding is S (string) for nominal variables, I (integer) for ordinal and interval variables and D (double) for ratio variables. Interval and ratio variables are limited to numeric encodings.

If no explicit variable types are specified these are automatically inferred from the data frame column types and classes, whenever possible. Columns of type character are treated as nominal string encoded variables (N). Unordered factor columns are converted to character and also treated as string encoded nominals. Ordered factors are converted to integer encoded interval variables (I) as phenotypes 15

described below. Columns of type logical are taken to be asymmetric binary variables (NB). Finally, integer and more broadly numeric columns are treated as integer encoded interval variables (I) and double encoded ratio variables (R), respectively.

Boolean encoded nominals (NB) are treated as asymmetric binary variables. For symmetric binary variables just use the default string encoding (N or NS). Other nominal variables are converted to factors.

Ordinal variables of class ordered are converted to integers respecting the order and range of the factor levels and subsequently treated as integer encoded interval variables (I). This conversion allows to model the full range of factor levels also when some might not occur in the data. For other ordinal variables it is assumed that each value occurs at least once and that values follow the natural ordering of the chosen data type (in Java).

If explicit types are given for some variables others can still be automatically inferred by setting their type to NA.

min

Minimum values of interval or ratio variables (optional). Numeric vector. Ignored when reading from file. If undefined for some variables the respective minimum is inferred from the data. If the data exceeds the minimum it is also updated accordingly. For nominal and ordinal variables just put NA.

max

Maximum values of interval or ratio variables (optional). Numeric vector. Ignored when reading from file. If undefined for some variables the respective maximum is inferred from the data. If the data exceeds the maximum it is also updated accordingly. For nominal and ordinal variables just put NA.

file

File containing the phenotype data.

#### Value

Phenotype data of class chpheno with elements

data Phenotypes (data frame).

size Number of individuals in the dataset.

ids Unique item identifiers.

names Item names. Names of individuals to which no explicit name has been assigned are equal to the unique ids.

types Variable types and encodings.

ranges Variable ranges, when applicable (NA elsewhere).

java Java version of the data object.

file Normalized path of file from which the data was read (if applicable).

## **Examples**

```
# create from data frame
pheno.data <- data.frame(
    season = c("winter", "summer", "summer", "winter", "summer"),
    yield = c(34.5, 32.6, 22.1, 54.12, 43.33),
    size = ordered(c("1", "s", "s", "m", "1"), levels = c("s", "m", "1")),</pre>
```

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```
resistant = c(FALSE, TRUE, TRUE, FALSE, TRUE)
)
pheno <- phenotypes(pheno.data)

# explicit types
pheno <- phenotypes(pheno.data, types = c("N", "R", "O", "NB"))
# treat last column as symmetric binary, auto infer others
pheno <- phenotypes(pheno.data, types = c(NA, NA, NA, "NS"))

# explicit ranges
pheno <- phenotypes(pheno.data, min = c(NA, 20.0, NA, NA), max = c(NA, 60.0, NA, NA))

# read from file
pheno.file <- system.file("extdata", "phenotypes.csv", package = "corehunter")
pheno <- phenotypes(file = pheno.file)</pre>
```

read.autodelim

Read delimited file.

## **Description**

Delegates to read.delim where the separator is inferred from the file extension (CSV or TXT). For CSV files the delimiter is set to "," while for TXT file "\t" is used. Also sets some default argument values as used by Core Hunter.

## Usage

```
read.autodelim(file, quote = "'\"", row.names = 1, na.strings = "",
   check.names = FALSE, strip.white = TRUE, stringsAsFactors = FALSE, ...)
```

#### **Arguments**

file File path.

quote the set of quoting characters. To disable quoting altogether, use quote = "".

See scan for the behaviour on quotes embedded in quotes. Quoting is only considered for columns read as character, which is all of them unless colClasses

is specified.

row.names a vector of row names. This can be a vector giving the actual row names, or a

single number giving the column of the table which contains the row names, or character string giving the name of the table column containing the row names.

If there is a header and the first row contains one fewer field than the number of columns, the first column in the input is used for the row names. Otherwise if

row.names is missing, the rows are numbered.

Using row.names = NULL forces row numbering. Missing or NULL row.names generate row names that are considered to be 'automatic' (and not preserved by as.matrix).

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na.strings

a character vector of strings which are to be interpreted as NA values. Blank fields are also considered to be missing values in logical, integer, numeric and complex fields. Note that the test happens *after* white space is stripped from the input, so na.strings values may need their own white space stripped in advance.

check.names

logical. If TRUE then the names of the variables in the data frame are checked to ensure that they are syntactically valid variable names. If necessary they are adjusted (by make.names) so that they are, and also to ensure that there are no duplicates.

strip.white

logical. Used only when sep has been specified, and allows the stripping of leading and trailing white space from unquoted character fields (numeric fields are always stripped). See scan for further details (including the exact meaning of 'white space'), remembering that the columns may include the row names.

stringsAsFactors

logical: should character vectors be converted to factors? Note that this is over-ridden by as.is and colClasses, both of which allow finer control.

... Further arguments to be passed to read.delim.

#### Value

Data frame.

sampleCore

Sample a core collection.

## **Description**

Sample a core collection from the given data.

#### Usage

```
sampleCore(data, obj, size = 0.2, always.selected = integer(0),
never.selected = integer(0), mode = c("default", "fast"),
normalize = TRUE, time = NA, impr.time = NA, steps = NA,
impr.steps = NA, indices = FALSE, verbose = FALSE)
```

## **Arguments**

data

Core Hunter data (chdata) containing genotypes, phenotypes and/or a precomputed distance matrix. Typically the data is obtained with coreHunterData. Can also be an object of class chdist, chgeno or chpheno if only one type of data is provided.

obj

Objective or list of objectives (chobj). If no objectives are specified Core Hunter maximizes a weighted index including the default entry-to-nearest-entry distance (EN) for each available data type, with equal weight. For genotypes, the Modified Roger's distance (MR) is used. For phenotypes, Gower's distance (GD) is applied.

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size

Desired core subset size (numeric). If larger than one the value is used as the absolute core size after rounding. Else it is used as the sampling rate and multiplied with the dataset size to determine the size of the core. The default sampling rate is 0.2.

always.selected

vector with indices (integer) or ids (character) of items that should always be selected in the core collection

never.selected vector with indices (integer) or ids (character) of items that should never be selected in the core collection

mode

Execution mode (default or fast). In default mode, Core Hunter uses an advanced parallel tempering search algorithm and terminates when no improvement is found for ten seconds. In fast mode, a simple stochastic hill-climbing algorithm is applied and Core Hunter terminates as soon as no improvement is made for two seconds. Stop conditions can be overridden with arguments time and impr.time.

normalize

If TRUE (default), the applied objectives in a multi-objective configuration (two or more objectives) are automatically normalized prior to execution. For singleobjective configurations, this argument is ignored.

Normalization requires an independent preliminary search per objective (fast stochastic hill-climber, executed in parallel for all objectives). The same stop conditions, as specified for the main search, are also applied to each normalization search. In default execution mode, however, any step-based stop conditions are multiplied by 500 for the normalization searches, because in that case the main search (parallel tempering) executes 500 stochastic hill-climbing steps per replica, in a single step of the main search.

Normalization ranges can also be precomputed (see getNormalizationRanges) or manually specified in the objectives to save computation time when sampling core collections. This is especially useful when multiple cores are sampled for the same objectives, with possibly varying weights.

Absolute runtime limit in seconds. Not used by default (NA). If used, it should be a strictly positive value, which is rounded to the nearest integer.

impr.time

Maximum time without improvement in seconds. If no explicit stop conditions are specified, the maximum time without improvement defaults to ten or two seconds, when executing Core Hunter in default or fast mode, respectively. If a custom improvement time is specified, it should be strictly positive and is rounded to the nearest integer.

steps

Maximum number of search steps. Not used by default (NA). If used, it should be a strictly positive value, which is rounded to the nearest integer. The number of steps applies to the main search. Details of how this stop condition is transferred to normalization searches, in a multi-objective configuration, are provided in the description of the argument normalize.

impr.steps

Maximum number of steps without improvement. Not used by default (NA). If used, it should be a strictly positive value, which is rounded to the nearest integer. The maximum number of steps without improvement applies to the main search. Details of how this stop condition is transferred to normalization searches, in a multi-objective configuration, are provided in the description of the argument normalize.

time

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indices If TRUE, the result contains the indices instead of ids (default) of the selected

individuals.

verbose If TRUE, search progress messages are printed to the console. Defaults to FALSE.

#### **Details**

Because Core Hunter uses stochastic algorithms, repeated runs may produce different results. To eliminate randomness, you may set a random number generation seed using set.seed prior to executing Core Hunter. In addition, when reproducible results are desired, it is advised to use step-based stop conditions instead of the (default) time-based criteria, because runtimes may be affected by external factors, and, therefore, a different number of steps may have been performed in repeated runs when using time-based stop conditions.

#### Value

Core subset (chcore). It has an element sel which is a character or numeric vector containing the sorted ids or indices, respectively, of the selected individuals (see argument indices). In addition the result has one or more elements that indicate the value of each objective function that was included in the optimization.

#### See Also

coreHunterData, objective, getNormalizationRanges

#### **Examples**

```
data <- exampleData()</pre>
# default size, maximize entry-to-nearest-entry Modified Rogers distance
obj <- objective("EN", "MR")</pre>
core <- sampleCore(data, obj)</pre>
# fast mode
core <- sampleCore(data, obj, mode = "f")</pre>
# absolute size
core <- sampleCore(data, obj, size = 25)</pre>
# relative size
core <- sampleCore(data, obj, size = 0.1)</pre>
# other objective: minimize accession-to-nearest-entry precomputed distance
core <- sampleCore(data, obj = objective(type = "AN", measure = "PD"))</pre>
# multiple objectives (equal weight)
core <- sampleCore(data, obj = list(</pre>
objective("EN", "PD"),
objective("AN", "GD")
))
# multiple objectives (custom weight)
core <- sampleCore(data, obj = list(</pre>
objective("EN", "PD", weight = 0.3),
 objective("AN", "GD", weight = 0.7)
```

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```
# custom stop conditions
core <- sampleCore(data, obj, time = 5, impr.time = 2)
core <- sampleCore(data, obj, steps = 300)
# print progress messages
core <- sampleCore(data, obj, verbose = TRUE)</pre>
```

setRange

Set the normalization range of the given objective.

## Description

See argument range of objective for details.

#### Usage

```
setRange(obj, range)
```

## **Arguments**

obj Core Hunter objective of class chobj.

range Normalization range [l,u]. See argument range of objective for details.

#### Value

Objective including normalization range.

#### See Also

objective

wrapData

Wrap distances, genotypes or phenotypes in Core Hunter data.

## Description

If the given data does not match any of these three classes it is returned unchanged.

#### Usage

```
wrapData(data)
```

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# Arguments

data of class chgeno, chpheno or chdist

## Value

Core Hunter data of class chdata

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