Package ‘haplotypes’

March 16, 2019

Type Package
Title Manipulating DNA Sequences and Estimating Unambiguous Haplotype Network with Statistical Parsimony
Version 1.1
Date 2019-03-14
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Description Provides S4 classes and methods for reading and manipulating aligned DNA sequences, supporting an indel coding methods (only simple indel coding method is available in the current version), showing base substitutions and indels, calculating absolute pairwise distances between DNA sequences, and collapses identical DNA sequences into haplotypes or inferring haplotypes using user provided absolute pairwise character difference matrix. This package also includes S4 classes and methods for estimating genealogical relationships among haplotypes using statistical parsimony and plotting parsimony networks.
License GPL-2
URL https://cran.r-project.org,
    https://biolsystematics.wordpress.com/r/
Imports methods, stats, utils, network, sna, ape, phangorn, pegas, plotrix
NeedsCompilation no
Repository CRAN
Date/Publication 2019-03-16 22:50:12 UTC

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Description

This package provides S4 classes and methods for reading and manipulating aligned DNA sequences, supporting an indel coding methods (only simple indel coding method is available in the current version), showing base substitutions and indels, calculating absolute pairwise distances between DNA sequences, and collapses identical DNA sequences into haplotypes or inferring haplotypes using user provided absolute pairwise character difference matrix. This package also includes S4 classes and methods for estimating genealogical relationships among haplotypes using statistical parsimony and plotting parsimony networks.

Author(s)

Caner Aktas, <caktas.aca@gmail.com>

Examples

```r
## Read example FASTA file.
# Example: read example fasta file
f<-system.file("example.fas",package="haplotypes")
# invalid character 'N' was replaced with '?' with a warning message
x<-read.fas(file=f)
# an object of class 'Dna'
x

## or load DNA Sequence data set.
data("dna.obj")
x<-dna.obj
## Not run:
x

## End(Not run)

## Compute an absolute pairwise character difference matrix from DNA sequences.
# coding gaps using simple indel coding method
d<-distance(x,indels="sic")
## Not run:
d

## End(Not run)

## Infer haplotypes using the 'Dna' object.
# coding gaps using simple indel coding method
h<-haplotype(x,indels="s")
## Not run:
h
```
#append-methods

Combines two Dna objects

## Description

Combines two Dna objects.

## Usage

```
## S4 method for signature 'Dna'
append(x, values)
```

## Arguments

- `x` an object of class Dna.
- `values` an object of class Dna.
as.data.frame-methods

Value

an object of class Dna.

Methods

signature(x = "Dna", values = "Dna") combines two Dna objects.

Examples

data("dna.obj")

x <- dna.obj
ty <- dna.obj

nrow(x)

## Combining two 'Dna' objects.
z <- append(x, y)
nrow(z)

as.data.frame-methods Coerces a Dna object to a data.frame

Description

Coerces an object to a data.frame.

Usage

## S4 method for signature 'Dna'

as.data.frame(x)

Arguments

x

an object of class Dna.

Value

returns a data frame.

Methods

signature(x = "Dna") coerces a Dna object to a data.frame.
Examples

data("dna.obj")

x<-dna.obj
x<-as.dna(x[1:4,1:6])

## Coercing a 'Dna' object to a data.frame.
df<-as.data.frame(x)
df

# TRUE
is.data.frame(df)

## Not run:
# gives the same result
df<-as.data.frame(x@sequence)
df

## End(Not run)

as.dna-methods  Coerces an object to a Dna object

Description

Coerces an object that contains DNA sequences to an object of Class Dna.

Usage

## S4 method for signature 'matrix'
as.dna(x)
## S4 method for signature 'data.frame'
as.dna(x)
## S4 method for signature 'list'
as.dna(x)
## S4 method for signature 'character'
as.dna(x)
## S4 method for signature 'Haplotype'
as.dna(x)
## S4 method for signature 'DNAbin'
as.dna(x)
## S4 method for signature 'phyDat'
as.dna(x)
as.dna-methods

Arguments

x               a matrix, a data.frame, a list, a character, an object of class Haplotype, DNABin
                 (ape) or phyDat (phangorn) object containing the DNA sequences.

Details

Elements of the list must be vectors. Each element of the list contains a single DNA sequence. If
the sequence lengths differ, the longest sequence is taken into account and gaps are introduced to
the shorter sequences at the end of the matrix in the slot sequence. Sequence length information is
stored in the slot seqlengths.

Valid characters for the slot sequence are "A","C","G","T","a","c","g","t","-","?". During the con-
version of the object to the class Dna, integers 0,1,2,3,4,5 or characters "0","1","2","3","4","5" are
converted to "?","A","C","G","T","-", respectively. Invalid characters are replaced with "?" with a
warning message.

Value

an object of class Dna.

Methods

signature(x = "matrix") coerces matrix to a Dna object.
signature(x = "data.frame") coerces data.frame to a Dna object.
signature(x = "list") coerces list to a Dna object.
signature(x = "character") coerces characters to a Dna object.
signature(x = "Haplotype") coerces a Haplotype object to a Dna object.
signature(x = "DNABin") coerces a DNABin object to a Dna object.
signature(x = "phyDat") coerces a phyDat object to a Dna object.

Examples

## Coercing a matrix to a 'Dna' object.
# all valid characters
x<-matrix(c("?","A","C","G","t","-","0","1","2","3","4","5"),4,6)
rownames(x)<-c("seq1","seq2","seq3","seq4")
dna.obj<as.dna(x)
dna.obj

## Not run:
# includes invalid characters
x<-matrix(c("x","y","x","?","t","-","0","1","2","3","4","5"),4,6)
rownames(x)<-c("seq1","seq2","seq3","seq4")
dna.obj<as.dna(x)
dna.obj
dna.obj@sequence
# all valid integers
x<-matrix(c(0,1,2,3,4,5,0,1,2,3,4,5),4,6)
rownames(x)<-c("seq1","seq2","seq3","seq4")
dna.obj<-as.dna(x)
dna.obj@dna@sequence

## Coercing a data.frame to a 'Dna' object.
x<-data.frame(matrix(c("?", "A", "C", "G", "T", "$", "0", "1", "2", "3", "4", "5"),4,6))
rownames(x)<-c("seq1","seq2","seq3","seq4")
dna.obj<-as.dna(x)
dna.obj@dna@sequence

## Coercing a list to a 'Dna' object.
seq1<-c("?", "A", "C", "G", "T", "$", "0", "1")
seq2<-c("?", "A", "C", "G", "T", "$", "0", "1", "2")
seq3<-c("?", "A", "C", "G", "T", "$", "0", "1", "2", "3")
x<-list(seq1=seq1,seq2=seq2,seq3=seq3)
dna.obj<-as.dna(x)

## sequence lengths differ
dna.obj@seqlengths
dna.obj@dna@sequence

## Coercing a character vector to a Dna object.
seq<-c("?", "A", "C", "G", "T", "$", "0", "1")
x<-as.dna(seq)

## Coercing a Haplotype object to a Dna object.
data("dna.obj")
x<-dna.obj
h<-haplotype(x)

## DNA Sequences of unique haplotypes
dna.obj<-as.dna(h)
dna.obj
d<-distance(x)

## if 'Haplotype' object does not contain 'DNA' Sequences
h<-haplotype(d)

## returns an error
as.dna(h)

## Coercing a DNABin object to a Dna object.
require(ape)
data(woodmouse)
x<-as.dna(woodmouse)
as.DNAbin-methods

Description
This function coerces Dna object to DNAbin (ape) object.

Usage
## S4 method for signature 'Dna'
as.DNAbin(x, endgaps=TRUE)

Arguments

x an object of class Dna.
endgaps boolean; gaps at the end of the sequences are included if this is TRUE.

Value
an object of class DNAbin.

Methods
signature(x = "Dna") coerces a Dna object to a DNAbin object.

Examples

## Coercing a Dna object to a DNAbin object.
data("dna.obj")
x<-dna.obj
dBin<-as.DNAbin(x)
dBin

#gaps at the end removed
dBin<-as.DNAbin(x, endgaps=FALSE)
dBin
as.list-methods

Methods for function as.list in the Package haplotypes

Description
Coerces an object to a list.

Usage
```r
## S4 method for signature 'Dna'
as.list(x)
## S4 method for signature 'Haplotype'
as.list(x)
## S4 method for signature 'Parsimnet'
as.list(x)
```

Arguments
- `x` an object of class `Dna`, `Haplotype` or `Parsimnet`.

Details
If `x` is a `Dna` object, elements of the list are character vectors that contain the DNA sequences of length equal to corresponding value in the slot `seqlengths`. If `x` is `Haplotype` or `Parsimnet` objects, slots are converted to list elements.

Value
returns a list.

Methods
- `signature(x = "Dna")` coerces an object of class `Dna` to a list.
- `signature(x = "Haplotype")` coerces an object of class `Haplotype` to a list.
- `signature(x = "Parsimnet")` coerces an object of class `Parsimnet` to a list.

Examples
```r
data("dna.obj")

## Coercing a 'Dna' object to a list.
x<-dna.obj[1:3,as.matrix=FALSE]
as.list(x)

## Not run:
## Coercing a 'Haplotype' object to a list.
x<-dna.obj
```
as.matrix-methods

h <- haplotype(x)
as.list(h)

## Coercing a 'Parsimnet' object to a list.
x <- dna.obj
p <- parsimnet(x)
as.list(p)

## End(Not run)

as.matrix-methods

Methods for function as.matrix in the Package haplotypes

Description

Coerces an object to a matrix.

Usage

## S4 method for signature 'Dna'
as.matrix(x)

Arguments

x an object of class Dna.

Value

returns a character matrix.

Methods

signature(x = "Dna") coerces an object of class Dna to a matrix.

Examples

data("dna.obj")

## Coercing a 'Dna' object to a matrix.
x <- dna.obj[1:4,1:6,as.matrix=FALSE]
x
as.matrix(x)

## Not run:
# gives the same result
dna.obj[1:4,1:6,as.matrix=TRUE]
## End(Not run)
as.network-methods

Coerces an object to a network object

Description

This function coerces Parsimnet object to network (network) object.

Usage

```r
## S4 method for signature 'Parsimnet'
as.network(x, net = 1, ...)
```

Arguments

- `x`: an object of class Parsimnet.
- `net`: a numeric vector of length one indicating which network to convert.
- `...`: additional arguments to function network.

Value

an object of class network.

Methods

signature(x = "Parsimnet") coerces a Parsimnet object to a network object.

Examples

```r
## Coercing a Parsimnet object to a network object.
data("dna.obj")
x <- dna.obj
p <- parsimnet(x)
n <- as.network(p)

#Fourth network (with only two edges)
p <- parsimnet(x, prob = .99)
n <- as.network(p, net = 4)
```
as.networx-methods

Coerces an object to a networx object

Description

This function coerces Parsimnet object to networx \{phangorn\} object.

Usage

```r
## S4 method for signature 'Parsimnet'
as.networx(x, net = 1, ...)
```

Arguments

- `x`: an object of class \texttt{Parsimnet}.
- `net`: a numeric vector of length one indicating which network to convert.
- `...`: additional arguments to \texttt{as.splits}.

Value

An object of class \texttt{networx}.

Methods

signature(x = "Parsimnet") coerces a \texttt{Parsimnet} object to a \texttt{networx} object.

Examples

```r
## Coercing a Parsimnet object to a networx object.
data("dna.obj")
x <- dna.obj
p <- parsimnet(x)
x <- as.networx(p)
plot(nx, "2D")
```
as.numeric-methods

Coerces a Dna object to a numeric matrix

Description

Converts a character matrix to a numeric matrix.

Usage

## S4 method for signature 'Dna'
as.numeric(x)

Arguments

x
an object of class Dna.

Details

Function as.numeric() coerces the character matrix in the slot sequence to a numeric matrix. Lower or upper case characters “?”, “A”, “C”, “G”, “T”, “-” are converted to integers 0,1,2,3,4,5, respectively.

Value

returns a numeric matrix.

Methods

signature(x = "Dna") coerces a Dna object to a numeric matrix.

Examples

x<-matrix(c("?","A","C","G","T","-","0","1","2","3","4","5"),4,6)
rownames(x)<-c("seq1","seq2","seq3","seq4")
x<-as.dna(x)
# original character matrix
as.matrix(x)

## Coercing a 'Dna' object to a numeric matrix.
# numeric matrix
as.numeric(x)
as.phyDat-methods

Description
This function coerces a dna object to phyDat (phangorn) object.

Usage

```r
## S4 method for signature 'Dna'
as.phyDat(x, indels="sic", ...)
```

Arguments
- `x`: an object of class `Dna`.
- `indels`: the indel coding method to be used. This must be one of "sic", "5th" or "missing". Any unambiguous substring can be given. See also ‘Details’
- `...`: additional arguments to `as.phyDat`.

Details
Available indel coding methods:

- **sic**: Treating gaps as a missing character and coding them separately following the simple indel coding method.
- **5th**: Treating gaps as a fifth state character.
- **missing**: Treating gaps as a missing character.

Value
an object of class `phyDat`.

Methods

```
signature(x = "Dna") coerces a Dna object to a phyDat object.
```

Examples

```r
data("dna.obj")
x<-dna.obj

## Coercing a Dna object to a phyDat object.
# Simple indel coding.
phyd<-as.phyDat(x)
phyd
```
# Gaps as 5th state characters.
phyd<-as.phyDat(x,indels="5")
phyd

# Gaps as 5th state characters.
phyd<-as.phyDat(x,indels="m")
phyd

---

# basecomp-methods

**Calculates base composition**

### Description

Calculates base composition of `Dna` object.

### Usage

```r
## S4 method for signature 'Dna'
basecomp(x)
```

### Arguments

- `x` an object of class `Dna`.

### Value

A matrix with sequence as rows, DNA bases as columns and frequencies as entries.

### Methods

- `signature(x = "Dna")` calculates base composition of `Dna` object.

### Examples

```r
data("dna.obj")
x <- dna.obj

## Calculating base compositions.
basecomp(x)
```
Generates single bootstrap replicate

Description

Methods for generating a single bootstrap replicate.

Usage

```{r}
## S4 method for signature 'Dna'
boot.dna(x, replacement = TRUE)
```

Arguments

- `x` an object of class `Dna`.
- `replacement` boolean; whether the sampling is done with replacement or without replacement.

Value

an object of class `Dna`.

Methods

`signature(x = "Dna")` generates single bootstrap replicate from a Dna object.

Author(s)

Caner Aktas, <caktas.aca@gmail.com>

Examples

```{r}
data("dna.obj")
x<-dna.obj

## Generating a bootstrap replicate.
# with replacement
bxr<-%boot.dna(x)
image(bxr)

# without replacement
bx<-%boot.dna(x, replacement=FALSE)
image(bx)
```
**distance-methods**

*Calculates absolute pairwise character difference matrix using a Dna object*

**Description**

Computes and returns an absolute pairwise character difference matrix from DNA sequences.

**Usage**

```r
# S4 method for signature 'Dna'
distance(x, subset=NULL, indels="sic")
```

**Arguments**

- `x`: an object of class `Dna`.
- `subset`: a vector of integers in the range [1,nrow(x)], specifying which sequence(s) are used in the distance calculation. Only distance between selected sequence(s) and the rest of the sequences are calculated. If it is NULL, all comparisons are done.
- `indels`: the indel coding method to be used. This must be one of "sic", "5th" or "missing". Any unambiguous substring can be given. See also ‘Details’

**Details**

Available indel coding methods:

- **sic**: Treating gaps as a missing character and coding them separately following the simple indel coding method.
- **5th**: Treating gaps as a fifth state character.
- **missing**: Treating gaps as a missing character.

**Value**

returns an object of class `dist`.

**Methods**

- `signature(x = "Dna")` Computes and returns an absolute pairwise character difference matrix from Dna objects.

**Author(s)**

Caner Aktas, &lt;caktas.aca@gmail.com&gt;
### References


### See Also

`indelcoder` and `subs`

### Examples

```r
data("dna.obj")
x<-dna.obj[4:7,13:22,as.matrix=FALSE]

## Simple indel coding.
distance(x,indels="s")

## Gaps as 5th state characters.
distance(x,indels="5")

## Gaps as missing characters.
distance(x,indels="m")

## Not run:
## Using 'subset'.
x<-dna.obj[4:10,13:22,as.matrix=FALSE]
distance(x, NULL)
distance(x, subset=c(1))
distance(x, subset=c(2,4))

## End(Not run)
```

---

**Dna-class**  

**Description**  

S4 class to hold DNA sequence data.

**Objects from the Class**  

Objects can be created by calls of the form `new("Dna", sequence, seqlengths, seqnames)`, however reading fasta file using `read.fas` function or coerce matrix, data.frame or list objects to a Dna object using `as.dna` methods is preferable.
Slots

sequence: Object of class "matrix" containing DNA sequence data, rows represent sequences and columns represent sites. See also ‘Note’.

seqlengths: Object of class "numeric" containing the length of each DNA sequence.

seqnames: Object of class "character" containing the name of each DNA sequence.

Methods

[ signature(x = "Dna", i = "ANY", j = "ANY"): extracts part of a DNA sequence as an object of class matrix.

[ signature(x = "Dna", i = "ANY", j = "ANY", value = "ANY"): replaces part of a Dna sequence with an object of class "matrix", "numeric" or "character".

append signature(x = "Dna", value = "ANY"): combines two Dna objects.

as.data.frame signature(x = "Dna"): coerces an object of class Dna to a data.frame.

as.list signature(x = "Dna"): coerces an object of class Dna to a list; elements of the list are character vectors that contains the DNA sequences of length equal to corresponding value in the slot seqlengths.

as.matrix signature(x = "Dna"): coerces an object of class Dna to a matrix.

as.numeric signature(x = "Dna"): coerces an object of class Dna to a numeric matrix.

as.DNAbin signature(x = "Dna"): coerces an object of class Dna to a DNAbin object.

as.phyDat signature(x = "Dna"): coerces an object of class Dna to a phyDat object.

basecomp signature(x = "Dna"): calculates base composition of Dna object.

boot.dna signature(x = "Dna"): generates single bootstrap replicate.

distance signature(x = "Dna"): computes and returns an absolute pairwise character difference matrix from DNA sequences.

haplotype signature(x = "Dna"): infers haplotypes from DNA sequences.

image signature(x = "Dna"): displays DNA sequences

indelcoder signature(x = "Dna"): supports simple indel coding method.

length signature(x = "Dna"): returns the longest sequence length.

append signature(x = "Dna", value = "ANY"): combines two Dna objects.

names signature(x = "Dna"): gets the names of an object Dna.

names<- signature(x = "Dna"): sets the names of an object Dna.

ncol signature(x = "Dna"): returns the longest sequence length.

nrow signature(x = "Dna"): returns the total sequence number.

pairnei signature(x = "Dna"): calculates pairwise Nei’s average number of differences between populations.

pairPhiST signature(x = "Dna"): calculates pairwise PhiST between populations.

parsimnet signature(x = "Dna"): estimates genealogies using statistical parsimony.

polymorph signature(x = "Dna"): displays information about DNA polymorphisms of two sequences; indels and base substitutions, respectively.
range signature(object = "Dna"): returns a vector containing the minimum and maximum length of DNA sequences.

remove.gaps signature(object = "Dna"): removes alignment gaps.

rownames signature(object = "Dna"): retrieve the row names of a DNA sequence matrix.

rownames<- signature(object = "Dna"): set the row names of a DNA sequence matrix.

show signature(object = "Dna"): displays Dna object briefly.

subs signature(x = "Dna"): displays information about base substitutions.

tolower signature(x = "Dna"): Translate characters in DNA sequence matrix from upper to lower case.

toupper signature(x = "Dna"): Translate characters in DNA sequence matrix from lower to upper case.

unique signature(x = "Dna"): returns a list with duplicate DNA sequences removed.

Note


Author(s)

Caner Aktas, <caktas.aca@gmail.com>

dna.obj Example DNA sequence data

Description

An example object of the class Dna.

Usage

data(dna.obj)

Format

dna.obj contains a Dna object.

Examples

data(dna.obj)
dna.obj
grouping-methods

Groups haplotypes according to the grouping variable (populations, species, etc.)

Description

Function for creating a matrix with haplotypes as rows, grouping factor (populations, species, etc.) as columns and abundance as entries.

Usage

```r
## S4 method for signature 'Haplotype'
grouping(x, factors)
```

Arguments

- `x`: an object of class `Haplotype`.
- `factors`: a vector or factor giving the grouping variable (populations, species, etc.), with one element per individual.

Value

A list with two components:

- `hapmat`: a matrix with haplotypes as rows, levels of the grouping factor (populations, species, etc.) as columns and abundance as entries.
- `hapvec`: a vector giving the haplotype identities of individuals.

Methods

`signature(x = "Haplotype")`

Author(s)

Caner Aktas, <caktas.aca@gmail.com>

See Also

- `haplotype`

Examples

```r
data("dna.obj")
x<-dna.obj[1:6,as.matrix=FALSE]
# inferring haplotypes from DNA sequences
h<-haplotype(x)

## Grouping haplotypes.
```
# Haplotype-class

C character vector

# length of the argument 'factor' is equal to the number of sequences

\[ g \]

## Description

S4 class to store haplotype information.

## Objects from the Class

Objects can be created by calls of the form `new("Haplotype", haplist, hapind, uniquehapind, sequence, d, freq, nhap)`, however use function `haplotype` instead.

## Slots

- **haplist**: Object of class "list", containing the names of individuals that share the same haplotype.
- **hapind**: Object of class "list", containing the index of individuals that share the same haplotype.
- **uniquehapind**: Object of class "numeric", containing the index of the first occurrence of unique haplotypes.
- **sequence**: Object of class "matrix" if present, giving the DNA sequence matrix of unique haplotypes.
- **d**: Object of class "matrix", giving the absolute pairwise character difference matrix of unique haplotypes.
- **freq**: Object of class "numeric", giving the haplotype frequencies.
- **nhap**: Object of class "numeric", giving the total number of haplotypes.

## Methods

- **as.dna** signature(x = "Haplotype"): if Haplotype object contains dna sequences, coerces an object of class Haplotype to an object of class Dna, else returns an error message.
- **as.list** signature(x = "Haplotype"): assigns slots of an object Haplotype to list elements.
- **grouping** signature(x = "Haplotype"): creates a matrix with haplotypes as rows, grouping factor (populations, species, etc.) as columns and abundance as entries.
- **hapreord** signature(x = "Haplotype"): reorders haplotypes according to the ordering factor.
- **length** signature(x = "Haplotype"): returns the number of haplotypes.
- **pieplot** signature(x = "Parsimnet", y = "Haplotype"): plot pie charts on statistical parsimony network.
**pielegend** signature(x = "Parsimnet", y = "Haplotype"): add legends to pie charts produced using pieplot.

**show** signature(object = "Haplotype"): displays the object briefly.

**Author(s)**

Caner Aktas, <caktas.aca@gmail.com>

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**haplotype-methods**  *Methods for function haplotype in the package haplotypes*

**Description**

Collapses identical DNA sequences into haplotypes or inferring haplotypes using user provided absolute pairwise character difference matrix.

**Usage**

```r
## S4 method for signature 'Dna'
haplotype(x, indels = "sic")
## S4 method for signature 'dist'
haplotype(x)
## S4 method for signature 'matrix'
haplotype(x)
```

**Arguments**

- `x` an object of class `Dna`, `dist`, or `matrix`.
- `indels` the indel coding method to be used. This must be one of "sic", "5th" or "missing". Any unambiguous substring can be given. See `distance` for details.

**Value**

`haplotype` returns an object of class `Haplotype, as.list-methods` can be used to coerce the object to a list.

**Methods**

- `signature(x = "Dna")` Inferring haplotypes from DNA sequences.
- `signature(x = "dist")` Inferring haplotypes using an absolute pairwise character difference matrix (dist object).
- `signature(x = "matrix")` Inferring haplotypes using an absolute pairwise character difference matrix.

**Author(s)**

Caner Aktas, <caktas.aca@gmail.com>
Examples

data("dna.obj")
x<-dna.obj[1:6,,as.matrix=FALSE]

### Inferring haplotypes using 'Dna' object.
# coding gaps using simple indel coding method
h<-haplotype(x,indels="sic")
h

# giving DNA sequences of haplotypes
as.dna(h)

## Not run:

## Slots of an object Haplotypes
h@haplist  # haplotype list (names)
h@hapind   # haplotype list (index)
h@uniquehapind  # getting index of the first occurrence of haplotypes
h@sequence  # DNA sequences of haplotypes
h@d         # distance matrix of haplotypes
h@freq      # haplotype frequencies
h@nhap      # total number of haplotypes

## End(Not run)

## Inferring haplotypes using dist object.
d<-distance(x)
h<-haplotype(d)
h
## Not run:
# returns an error message
as.dna(h)

## End(Not run)

## Inferring haplotypes using distance matrix.
d<-as.matrix(distance(x))
h<-haplotype(d)
h
## Not run:
# returns an error message
as.dna(h)

## End(Not run)
Description

Reorders haplotypes according to the ordering factor.

Usage

```r
## S4 method for signature 'Haplotype'
hapreord(x, order=c(1:x@nhap))
```

Arguments

- `x`: an object of class `Haplotype`.
- `order`: a vector giving the order of haplotypes, with one element per haplotype.

Value

returns an object of class `Haplotype`.

Methods

`signature(x = "Haplotype")` Reorders haplotypes.

Author(s)

Caner Aktas, <caktas.aca@gmail.com>

See Also

`haplotype`

Examples

```r
data("dna.obj")
x<-dna.obj[1:6,,as.matrix=FALSE]
# inferring haplotypes from DNA sequences
h<-haplotype(x)

## Reordering haplotypes.

# length of the argument 'order' is equal to the number of haplotypes
rh<-hapreord(h,order=c(4,3,1,2))
rh
```
Description

This function returns the list of homoplastic indels and substitutions.

Usage

```r
## S4 method for signature 'Dna'
homopoly(x, indels, ...) 
```

Arguments

- `x` an object of class `Dna`.
- `indels` the indel coding method to be used. This must be one of "sic", "5th" or "missing". Any unambiguous substring can be given. See `distance` for details.
- `...` additional arguments to `parsimnet`.

Value

A list with following components:

- `indels` a character vector of homoplastic indels sitewise Consistency Index, names of the character vector gives the site of homoplastic indel.
- `subs` a character vector of homoplastic substitutions sitewise Consistency Index, names of the character vector gives the site of substitution.

Methods

```r
signature(x = "Dna")
```

Author(s)

Caner Aktas, <caktas.aca@gmail.com>

Examples

```r
data("dna.obj")

### Method for signature 'Dna'.
x <- dna.obj
homopoly(x)
```
Display DNA Sequence

Description

Display an image of DNA sequences.

Usage

```r
## S4 method for signature 'Dna'
image(x, all=FALSE, fifth=TRUE,
      col=c("#BFBFBF","#B99FD","#FD00B","#11A808","#F5D0B","#F8F8FF"),
      chars=TRUE, cex=1, show.names=TRUE, show.sites=TRUE, xlab="", ylab="", ...)
```

Arguments

- `x`: an object of class `Dna`.
- `all`: boolean; should entire sequence be displayed or only the polymorphic sites?
- `fifth`: boolean; if all==FALSE, should gaps be displayed?
- `col`: an integer or character vector for the colors. By default it is blue for "A", red for "C", green for "G", yellow for "T", white for ".", and grey for "?".
- `chars`: boolean; should characters be displayed on image?
- `cex`: a numeric vector of expansion factor for characters.
- `show.names`: boolean; should sequence names be displayed on the left side.
- `show.sites`: boolean; should site labels be displayed on the bottom side.
- `xlab`: a title for the x axis.
- `ylab`: a title for the y axis.
- `...`: additional arguments to `image.default`.

Methods

- `signature(x = "Dna")` Display an image of Dna objects

Author(s)

Caner Aktas, <caktas.aca@gmail.com>.

See Also

- `image.default`
Examples

data("dna.obj")
x<-dna.obj

## Display only polymorphic sites without gaps
image(x,all=FALSE,fifth=FALSE,show.names=TRUE,cex=0.6)

## Display only polymorphic sites with gaps
image(x,all=FALSE,fifth=TRUE,show.names=TRUE,cex=0.6)

## Not run:
## Display entire sequences
image(x,all=FALSE,show.names=TRUE,cex=0.6)

## End(Not run)

---

Description

Function for coding gaps separately. Only simple indel coding method is available in the current version.

Usage

## S4 method for signature 'Dna'
indelcoder(x)

Arguments

x an object of class Dna.

Value

a list with two components:

indels: a matrix giving the indel positions (beginnings and ends) and lengths.
codematrix: a binary matrix giving the indel codings. Missing values are denoted by -1.

Methods

signature(x = "Dna") Function for coding gaps separately.

Author(s)

Caner Aktas, <caktas.aca@gmail.com>
References


See Also

distance

Examples

data("dna.obj")
x<-dna.obj

## Simple indel coding.
indelcoder(x)

---

**length-methods**

*Methods for function length in the package haplotypes*

Description

Methods for function length.

Usage

```r
library(haplotypes)

## S4 method for signature 'Dna'
length(x)

## S4 method for signature 'Haplotype'
length(x)

## S4 method for signature 'Parsimnet'
length(x)
```

Arguments

- `x` an object of class `Dna`, `Haplotype` or `Parsimnet`.

Value

returns a non-negative integer vector.

Methods

- `signature(x = "Dna")` returns the longest sequence length.
- `signature(x = "Haplotype")` returns the number of haplotypes.
- `signature(x = "Parsimnet")` returns the length of network(s).
names-methods

See Also

ncol-methods

Examples

data("dna.obj")
x<-dna.obj

## Longest sequence length
length(x)

## Total number of haplotypes
h<-haplotype(x)
length(h)

## Length of network(s)
p<-parsimnet(x,prob=.95)
# length of the network
length(p)

p<-parsimnet(x,prob=.99)
# length of the networks
length(p)

---

names-methods  Function to get or set names of a Dna object or Parsimnet object

Description

Function to get or set sequence names of Dna object or names of network in Parsimnet object.

Usage

## S4 method for signature 'Dna'
names(x)
## S4 method for signature 'Parsimnet'
names(x)
## S4 replacement method for signature 'Dna'
names(x)<-value
## S4 replacement method for signature 'Parsimnet'
names(x)<-value

Arguments

x an object of class Dna or Parsimnet.
value a character vector of the same length as number of sequence or networks.
Methods

signature(x = "Dna") Function to get or set names of an object of Dna.
signature(x = "Parsimnet") Function to get or set names of networks in Parsimnet object.

Examples

data("dna.obj")

x<-dna.obj
x<-as.dna(x[1:4,1:6])

## Getting sequence names.
names(x)

## Setting sequence names.
names(x)<-c("u","v","z","y")
names(x)

x<-dna.obj
p<-parsimnet(x,prob=.99)

## Getting network names in parsimnet object
names(p)
## Setting network names
names(p)<-c("a","b","c","d","f","g")
names(p)

ncol-methods

Returns the length of the longest DNA sequence

Description

ncol returns the number of columns present in a matrix.

Usage

## S4 method for signature 'Dna'
ncol(x)

Arguments

x an object of class Dna.
**nrow-methods**

**Value**

an integer of length one.

**Methods**

signature(x = "Dna") ncol returns the number of columns present in the sequence matrix (length of the longest DNA sequence).

**See Also**

length-methods

**Examples**

data("dna.obj")
x <- dna.obj

## Giving the length of the longest sequence.
col(x)
# gives the same result
length(x)

---

**nrow-methods**

Returns the number of DNA sequences

**Description**

nrow returns the number of rows present in a matrix.

**Usage**

## S4 method for signature 'Dna'
nrow(x)

**Arguments**

x an object of class Dna.

**Value**

an integer of length one.

**Methods**

signature(x = "Dna") nrow returns the number of rows present in the sequence matrix (number of sequences).
Examples

data("dna.obj")
x <- dna.obj

## Giving the number of sequences.
nrow(x)

pairnei-methods

Provides the average number of pairwise Nei's (D) differences between populations

Description

Function provides pairwise Nei's raw number of nucleotide differences between populations.

Usage

## S4 method for signature 'Dna'
pairnei(x, populations, indels="sic", nperm=99, subset=NULL, showprogbar=FALSE)
## S4 method for signature 'dist'
pairnei(x, populations, nperm=99, subset=NULL, showprogbar=FALSE)
## S4 method for signature 'matrix'
pairnei(x, populations, nperm=99, subset=NULL, showprogbar=FALSE)

Arguments

x       an object of class Dna, "dist" or "matrix".
populations a vector giving the populations, with one element per individual.
indels the indel coding method to be used. This must be one of "sic", "5th" or "missing". Any unambiguous substring can be given. See distance for details.
nperm the number of permutations. Set this to 0 to skip the permutation procedure.
subset a vector of integers in the range [1, length(unique(populations))], only distances between selected population(s) and the rest of the populations are calculated. If it is NULL, all comparisons are done.
showprogbar boolean; whether the progress bar is displayed or not displayed.

Details

The null distribution of pairwise Nei's differences under the hypothesis of no difference between the populations is obtained by permuting individuals between populations.
Value

a list with following components:

- `neidist` a matrix giving the average number of pairwise Nei’s (D) differences between populations (below diagonal elements) and average number of pairwise differences within populations (diagonal elements).
- `p` a matrix giving the p-values, or NULL if permutation test is not performed.

Methods

signature(x = "Dna")
signature(x = "dist")
signature(x = "matrix")

Author(s)

Caner Aktas, <caktas.aca@gmail.com>

References


Examples

data("dna.obj")

### Method for signature 'Dna'.
x <- dna.obj
x <- dna.obj[21, 20, 21, 26, 27, 28, 30, 3, 4, 7, 13, 14, 15, 16, 23, 24, 25], as.matrix=FALSE]
populations <- c("pop1", "pop1", "pop1", "pop1", "pop1", "pop1", "pop2", "pop2", "pop2", "pop2", "pop2", "pop2", "pop3", "pop4", "pop4", "pop4")

###skip permutation testing
pn <- pair nei(x, populations, nperm=0)

#Between populations
as.dist(pn$neidist)

#Within populations
diag(pn$neidist)

#Gaps as missing characters.
pn <- pair nei(x, populations, indels="m", nperm=0)

#using subset, third population against others
pairPhiST-methods

Provides the pairwise PhiST between populations

Description

This function calculates pairwise PhiST between populations using the AMOVA framework.
Usage

```r
## S4 method for signature 'Dna'
pairPhiST(x, populations, indels = "sic", nperm = 99, negatives = FALSE, subset = NULL, showprogbar = TRUE)
## S4 method for signature 'dist'
pairPhiST(x, populations, nperm = 99, negatives = FALSE, subset = NULL, showprogbar = TRUE)
## S4 method for signature 'matrix'
pairPhiST(x, populations, nperm = 99, negatives = FALSE, subset = NULL, showprogbar = TRUE)
```

Arguments

- `x`: an object of class `Dna`, "dist" or "matrix".
- `populations`: a vector giving the populations, with one element per individual.
- `indels`: the indel coding method to be used. This must be one of "sic", "5th" or "missing". Any unambiguous substring can be given. See `distance` for details.
- `nperm`: the number of permutations. Set this to 0 to skip the permutation procedure.
- `negatives`: boolean; if it is FALSE all negative PhiST values are replaced with zero.
- `subset`: a vector of integers in the range [1, length(unique(populations))], only distances between selected population(s) and the rest of the populations are calculated. If it is NULL, all comparisons are done.
- `showprogbar`: boolean; whether the progress bar is displayed or not displayed.

Details

The null distribution of pairwise PhiST under the hypothesis of no difference between the populations is obtained by permuting individuals between populations.

Value

A list with following components:

- `PhiST`: a matrix giving the PhiST values between populations.
- `p`: a matrix giving the p-values, or NULL if permutation test is not performed.

Methods

- `signature(x = "Dna")`
- `signature(x = "dist")`
- `signature(x = "matrix")`

Note

An internal code `statphi` is taken from package `ade4` version 1.7-8 without any modification, author Sandrine Pavoine. Function `amova` from package `pegas` is used internally to estimate variance components, author Emmanuel Paradis.
Author(s)

Caner Aktas, <caktas.aca@gmail.com>

References


Examples

data("dna.obj")

```r
### Method for signature 'Dna'.
x<-dna.obj
x<-dna.obj[c(1,2,21,26,27,28,30,3,4,7,13,14,15,16,23,24,25),,as.matrix=FALSE]
populations<-c("pop1", "pop1", "pop1", "pop1", "pop1", "pop1", "pop2", "pop2", "pop2", "pop2", "pop2", "pop3", "pop3", "pop4", "pop4", "pop4")

### skip permutation testing
pst<-pairPhiST(x, populations, nperm=0)
pst

### allow negative PhiST values
pst<-pairPhiST(x, populations, nperm=0, negatives=TRUE)
pst

### Gaps as missing characters.
pst<-pairPhiST(x, populations, indels="m", nperm=0, negatives=TRUE)
pst

### using subset, second population against others
pst<-pairPhiST(x, populations, nperm=0, subset=c(2))
pst

### Not run:
### 999 permutations.
pst<-pairPhiST(x, populations, nperm=999,showprogbar=TRUE)
pst

### random populations
x<-dna.obj
populations<-sample(1:4,nrow(x),replace=TRUE)
pst<-pairPhiST(x, populations, nperm=999,showprogbar=TRUE)
pst

### populations based on clusters
x<-dna.obj
d<-distance(x)
hc<-hclust(d,method="ward.D")
populations<-cutree(hc,4)
```
parsimnet-class

### Description

S4 class to store statistical parsimony networks and additional information.

### Objects from the Class

Objects can be created by calls of the form `new("Parsimnet", d, tempProbs, conlimit, prob, nhap, rowindex)`, however use function `parsimnet` instead.

### Slots

- **d**: Object of class "list" containing the geodesic distance matrix of haplotypes and intermediates for each network.
- **tempProbs**: Object of class "numeric" giving the probabilities of parsimony for mutational steps beyond the connection limit.
- **conlimit**: Object of class "numeric" giving the number of maximum connection steps at connection limit.
- **prob**: Object of class "numeric" giving the user defined connection limit.
- **nhap**: Object of class "numeric" giving the number of haplotypes in each network.
- **rowindex**: Object of class "list" containing vectors giving the index of haplotypes in each network.
### Methods

- **as.list** signature(x = "Parsimnet"): assigns slots of an object Parsimnet to list elements.
- **as.network** signature(x = "Parsimnet"): coerces Parsimnet object to network {network} object.
- **as.networx** signature(x = "Parsimnet"): coerces Parsimnet object to networx {phangorn} object.
- **length** signature(x = "Parsimnet"): returns the length of network(s).
- **names** signature(x = "Parsimnet"): gets names of networks in Parsimnet object.
- **names<-** signature(x = "Parsimnet"): sets names of networks in Parsimnet object.
- **plot** signature(x = "Parsimnet"): plots statistical parsimony networks.
- **pieplot** signature(x = "Parsimnet", y = "Haplotype"): plots pie charts on statistical parsimony networks.
- **pielegend** signature(x = "Parsimnet", y = "Haplotype"): add legends to pie charts produced using pieplot.
- **rownames** signature(x = "Parsimnet"): gets names of vertices in networks.
- **rownames<-** signature(x = "Parsimnet"): gets names of vertices in networks.
- **show** signature(object = "Parsimnet"): displays the object briefly.

### Author(s)

Caner Aktas, <caktas.aca@gmail.com>

---

**parsimnet-methods** Estimates gene genealogies using statistical parsimony

### Description

Function for estimating gene genealogies from DNA sequences or user provided absolute pairwise character difference matrix using statistical parsimony.

### Usage

```r
## S4 method for signature 'Dna'
parsimnet(x, indels="sic", prob=.95)
## S4 method for signature 'dist'
parsimnet(x, seqlength, prob=.95)
## S4 method for signature 'matrix'
parsimnet(x, seqlength, prob=.95)
```
 Arguments

x  an object of class Dna, dist, or matrix.
indels  the indel coding method to be used. This must be one of "sic", "5th" or "missing". Any unambiguous substring can be given. See distance for details.
seqlength  an integer of length one giving the sequence length information (number of characters).
prob  a numeric vector of length one in the range [0.01, 0.99] giving the probability of parsimony as defined in Templeton et al. (1992). In order to set maximum connection steps to Inf (to connect all the haplotypes in a single network), set the probability to NULL.

Details

The network estimation methods implemented in parsimnet function finds one of the most parsimonious network (or sub-networks if connection between haplotypes exceeds the parsimony limit). This is an implementation of the TCS method proposed in Templeton et al. (1992) and Clement et al. (2002). parsimnet function generates an unambiguous haplotype network without loops. If more than one best networks found (results in ambiguous connections), only a network with the lowest average all-pairs distance is returned. Loops may occur only if they are present in initial haplotype distance matrix when using algorithmic method.

Value

S4 methods for signature 'Dna', 'matrix' or 'dist' returns an object of class Parsimnet.

Methods

signature(x = "Dna") estimating gene genealogies from DNA sequences.
signature(x = "dist") estimating gene genealogies from distance matrix (dist object).
signature(x = "matrix") estimating gene genealogies from distance matrix.

Note

Duplicate names in the final distance matrices in slot d are renamed without warning. An internal function .TempletonProb is taken from package pegas version 0.6 without any modification, authors Emmanuel Paradis, Klaus Schliep.

Author(s)

Caner Aktas, <caktas.ac@gmail.com>.

References


See Also

`network`, `plot-methods` and `pieplot-methods`

Examples

```r
## Not run:
data("dna.obj")
x<-dna.obj

### Method for signature 'Dna'.
### statistical parsimony with 95
p<-parsimnet(x)
p
plot(p)

### statistical parsimony with 99
p<-parsimnet(x,prob=.99)
p
# plot the first network
plot(p,net=1)

### statistical parsimony with 99
#indels are coded as missing
p<-parsimnet(x,indels="m",prob=.99)
p
plot(p)

# statistical parsimony without connection limit.
p<-parsimnet(x,prob=NULL)
p
plot(p)

# plot the first network
plot(p,net=1)

### Method for signature 'dist'.
d<-distance(x)
seqlength<-length(x)
### statistical parsimony with 95
p<-parsimnet(d,seqlength)
p
plot(p)

### Method for signature 'matrix'.
```
pielegend-methods

Add Legends to Plots

Description

This function can be used to add legends to pie charts produced using pieplot.

Usage

## S4 method for signature 'Parsimnet,Haplotype'
pielegend(p,h,net=1,factors,...)

Arguments

- `p`: a Parsimnet object.
- `h`: a Haplotype object.
- `net`: a numeric vector of length one indicating which network to plot.
- `factors`: a vector or factor giving the grouping variable (populations, species, etc.), with one element per individual.
- `...`: arguments to be passed to legend and others. See ‘Details’

Details

This method calls legend (graphics), some default parameters changed:

- `col`: an integer or character vector for the edge colors. By default, it is rainbow.
- `fill`: an integer or character vector for the filling colors. By default, it is rainbow
- `legend`: a numeric or character vector to appear in the legend. By default, it is the levels of the grouping factor for haplotypes.
- `x`: position of the legend. Default is set to “topright”.

Value

See ‘legend (graphics)’
Methods

signature(p = "Parsimnet", h = "Haplotype")

Author(s)

Caner Aktas, <caktas.aca@gmail.com>.

See Also

plot, Parsimnet-method, floating.pie, plot.default, plot.network.default and legend.

Examples

data("dna.obj")
x<-dna.obj
h<-haplotypes::haplotype(x)

### Statistical parsimony with 95% connection limit
p<-parsimnet(x)

#randomly generated populations
pop<-c("pop1","pop2","pop3","pop4","pop5","pop6","pop7","pop8")
set.seed(5)
pops<-sample(pop, nrow(x), replace=TRUE)

## Plotting with default parameters.
pieplot(p, h, 1, pops)

## Add legend with default parameters.
pielegend(p, h, 1, pops)

## Change colors for the populations.
## 8 colors for 8 populations
cols<--colors()[c(30, 369, 552, 558, 538, 642, 142, 91)]
pieplot(p, h, 1, pops, col=cols)
pielegend(p, h, 1, pops, col= cols)

pieplot-methods

Plots pie charts on statistical parsimony network

Description

Plotting pie charts on the statistical parsimony network.

Usage

## S4 method for signature 'Parsimnet,Haplotype'
pieplot(x, y, net=1, factors, coord = NULL, inter.labels=FALSE, interactive=FALSE,rex=1,...)
pieplot-methods

Arguments

- **x**: a *Parsimnet* object.
- **y**: a *Haplotype* object.
- **net**: a numeric vector of length one indicating which network to plot.
- **factors**: a vector or factor giving the grouping variable (populations, species, etc.), with one element per individual.
- **coord**: a matrix that contains user specified coordinates of the vertices, or NULL to generate vertex layouts with *network.layout* function from package *network*.
- **inter.labels**: boolean; should vertex labels of intermediate haplotypes be displayed?
- **interactive**: boolean; should vertices be interactively adjusted?
- **rex**: expansion factor for the pie radius.
- **...**: arguments to be passed to *floating.pie* and others. See ‘Details’

Details

This method calls *floating.pie* (*plotrix*), *network.vertex* (*network*), and *plot.default*, *lines*, and *text* (*graphics*). This method also uses some internal structures of *plot.network.default* from package *network*. The following additional arguments can be passed to these functions:

- **mode**: the vertex placement algorithm. Default is set to "fruchtermanreingold".
- **pad**: amount to pad the plotting range; useful if labels are being clipped. Default is set to 1.
- **displaylabels**: boolean; should vertex labels be displayed?
- **label**: a vector of vertex labels. By default, the rownames of the distance matrix (*rownames(p@d[[net]])*) are used. If *inter.labels*==FALSE only haplotype labels are displayed.
- **label.cex**: character expansion factor for labels. Default is set to 0.75.
- **label.col**: an integer or character vector for the label colors. By default, it is 1 (black).
- **label.pos**: position at which labels should be placed relative to vertices. 0 and 6 result in labels placed away from the center of the plotting region; 1, 2, 3, and 4 result in labels being placed below, to the left of, above, and to the right of vertices, respectively; and label.pos 5 or greater than 6 result in labels which are plotted with no offset (i.e., at the vertex positions). Default is set to 0.
- **label.pad**: amount to pad the labels. This setting is available only if the labels are plotted with offset relative to vertex positions. Default is set to 1.
- **vertex.cex**: a numeric vector of expansion factor for intermediate vertices (only). By default it is \((0.5)\times\text{min(radius)}\). Use ‘radius’ to specify size of pie charts.
- **col**: the colors of the pie sectors (i.e., colors for populations), by default *rainbow*.
- **vertex.col**: an integer or character vector for the intermediate vertex colors. By default, it is 1 (black).
- **edge.col**: an integer or character vector for the edge colors. By default, it is 1 (black).
- **edge.lwd**: a numeric vector, edges line width. By default, it is 1.
- **edge.lty**: a numeric vector of length one, specifies the line type for the edges. By default it is 1.
- **edges**: the number of lines forming a pie circle. By default, it is 200.
radius  a numeric vector of length p@nhap[net] for the radius of drawn pie charts. Useful for specifying the radius independent of the haplotype frequencies. Default is $0.8 \times (\text{haplotype frequencies}) \times \text{rex} / \text{max(haplotype frequencies)}$.

vertex.sides  number of polygon sides for vertices. Default is set to 50.

xlab  x axis label.

xlab  y axis label.

Value

A two-column matrix containing the vertex positions as x,y coordinates.

Methods

signature(x = "Parsimnet", y = "Haplotype")

Note

Some internal structures of plot.network.default is taken from package network with modifications, author Carter T. Butts.

Author(s)

Caner Aktas, <caktas.aca@gmail.com>.

See Also

plot, Parsimnet-method, floating.pie, plot.default and plot.network.default

Examples

```r
data("dna.obj")
x<-dna.obj
h<-(haplotypes::haplotype(x)

### Statistical parsimony with 95% connection limit
p<-parsimnet(x)

# randomly generated populations
pop<-c("pop1","pop2","pop3","pop4","pop5","pop6","pop7","pop8")
set.seed(5)
pops<-sample(pop,nrow(x),replace=TRUE)

### Plotting with default parameters.
pieplot(p,h,1, pops)

### Change colors for the populations.
# 8 colors for 8 populations
cols<-colors()[c(30,369,552,558,538,642,142,91)]
pieplot(p,h,1, pops,col=cols)

### Expanding pie charts and intermediate vertices.
pieplot(p,h,1, pops,rex=2)
```
## Adjusting intermediate vertex sizes.
pieplot(p,h,1, pops, vertex.cex=rep(0.2, nrow(p@d[[1]])-p@nhap))

## Expanding pie charts and intermediate vertices, adjusting intermediate vertex sizes.
pieplot(p,h,1, pops,rex=2, vertex.cex=rep(0.1, nrow(p@d[[1]])-p@nhap))

## Adjusting radius of pie charts.
pieplot(p,h,1, pops,radius=rep(1, p@nhap))

## Not run:
## Interactively adjusting vertex positions.
pieplot(p,h,1, pops,interactive=TRUE)

## End(Not run)

## Multiple networks with 99% connection limit.
p<-.parsimnet(x,prob=.99)

## Plotting first network with default parameters.
pieplot(p,h,1, pops)

## Change colors for the populations.
## 8 colors for 8 populations
cols<-colors()[c(30,369,552,558,538,642,142,91)]
pieplot(p,h,1, pops,col=cols)

---

### plot-methods

*Methods for function plot in the package haplotypes*

### Description

Plots statistical parsimony networks.

### Usage

```r
## S4 method for signature 'Parsimnet'
plot(x,y,net=1,inter.labels=FALSE,...)
```
Arguments

- **x**: an object of class `Parsimnet`.
- **y**: not used (needed for compatibility with generic plot function).
- **net**: a numeric vector of length one indicating which network to plot.
- **inter.labels**: boolean; should vertex labels of intermediate haplotypes to be displayed?
- **...**: additional arguments to `plot.default`, `plot.network.default` and arguments to be passed to plot method for `Parsimnet` objects. See ‘Details’

Details

These methods call `plot.network.default` from package `network`. Some default parameters are changed:

- **label**: a vector of vertex labels. By default the row names of the distance matrices in slot `d` are used. If `inter.labels`==FALSE only haplotype labels are displayed.
- **usearrows**: boolean; should arrows (rather than line segments) be used to indicate edges? Default is set to FALSE.
- **mode**: the vertex placement algorithm. Default is set to "kamadakawai".
- **pad**: amount to pad the plotting range; useful if labels are being clipped. Default is set to 1.
- **label.cex**: character expansion factor for label text. Default is set to 0.75.
- **vertex.cex**: a numeric vector of expansion factor for vertices. By default it is 0.8 for haplotypes and 0.5 for intermediates.
- **vertex.col**: an integer or character vector for the vertex colors. By default it is 2 (red) for haplotypes and 4 (blue) for intermediates.

Value

A two-column matrix containing the vertex positions as x,y coordinates.

Methods

```r
signature(x = "Parsimnet", y = "missing") Plots Parsimnet objects.
```

Author(s)

Caner Aktas, <caktas.aca@gmail.com>.

See Also

`parsimnet`, `plot.default` and `plot.network.default`
Examples

```r
## Not run:

data("dna.obj")
x<-dna.obj

### Method for signature 'Parsimnet'.

## Statistical parsimony with 95
p<-parsimnet(x)
p
## Plotting with default parameters.
plot(p)

## Displaying vertex labels of intermediate haplotypes.
plot(p, inter.labels=TRUE)

## Interactively adjusting vertex positions.
plot(p, interactive=TRUE)

## Interactively adjusting and saving vertex positions.
p<-parsimnet(x)
#saving vertex positions as x,y coordinates.
coo<-plot(p,interactive=TRUE)
#reuse saved coordinates.
plot(p,coord=coo)

## Adjusting vertex sizes.
plot(p, vertex.cex=c(rep(3,nrow(p@d[[1]]))))
# different sizes for haplotypes and intermediates
plot(p, vertex.cex=c(rep(3,p@nhap),rep(1,c(nrow(p@d[[1]])-p@nhap))))

## Adjusting vertex colors
# different color for haplotypes and intermediates
plot(p, vertex.col=c(rep("magenta",p@nhap),rep("deepskyblue3",c(nrow(p@d[[1]])-p@nhap))))

## Statistical parsimony with 98
p<-parsimnet(x,prob=.99)
p
#plot the first network
plot(p,net=1)
#plot the second network
plot(p,net=2)
#plot the third network. It is a single vertex.
plot(p,net=3)
```
polymorp-methods

Displays polymorphic sites (base substitutions and indels) between two sequences

Description

This function displays the polymorphic sites (base substitutions and indels) between the two sequences.

Usage

## S4 method for signature 'Dna'
polymorp(x, pair, indels = "sic")

Arguments

- **x**: an object of class `Dna`.
- **pair**: a vector of integers in the range [1, row(x)] of length two, specifying sequence pair.
- **indels**: the indel coding method to be used. This must be one of "sic", "5th" or "missing". Any unambiguous substring can be given. See `distance` for details.

Value

a list with two components:

- **indels**: a list of matrices of the indel regions if `indels=="sic"`. The component names of the list gives the position of the indels.
- **subst**: a list of matrices of the base substitutions. If `indels=="5th"`, each gap is treated as a base substitution. The component names of the list gives the position of the base substitutions.

Methods

signature(x = "Dna") Showing base substitutions and indels between the two sequences.

Author(s)

Caner Aktas, <caktas.aca@gmail.com>

See Also

`indelcoder` and `subs`
Examples

data("dna.obj")
x<-dna.obj

## Showing base substitutions and indels between seq1 and seq6.

# gaps are coded following the simple indel coding method
polymorp(x,c(1,5),indels="s")

# gaps are coded as a fifth state character
polymorp(x,c(1,5),indels="5")

# gaps are treated as missing character
polymorp(x,c(1,5),indels="m")

---

range-methods

Returns the minimum and maximum lengths of the DNA sequences

Description

range returns the lengths of shortest and longest DNA sequences.

Usage

## S4 method for signature 'Dna'
range(x)

Arguments

x an object of class Dna.

Value

an integer of length two.

Methods

signature(x = "Dna") range

See Also

length-methods
Examples

data("dna.obj")
x <- dna.obj

## shortest and longest DNA sequence lengths
range(x)

---

read.fas | Read sequences from a file in FASTA format

Description

Read DNA sequences from a file in FASTA Format.

Usage

read.fas(file)

Arguments

file the name of the file, which the sequence in the FASTA format is to be read from. If it does not contain an absolute path, the file name is relative to the current working directory, getwd().

Value

read.fas returns an object of class DNA.

Note

By default, valid characters are "A","C","G","T","a","c","g","t","-","?" for the class DNA. Numeric entries (integers) between 0-5 will be converted to "?","A","C","G","T","-". Invalid characters will be replaced with "?" with a warning message.

Author(s)

Caner Aktas, <caktas.ac@gmail.com>

See Also

DNA
Examples

```r
# Reading example file.
f <- system.file("example.fas",package="haplotypes")

# invalid character 'N' was replaced with '?' with a warning message
x <- read.fas(file = f)

# an object of class 'Dna'
x
```

---

remove.gaps-methods  Removing gaps from Dna object

Description

Removing gaps(“-“) from Dna object

Usage

```r
## S4 method for signature 'Dna'
remove.gaps(x, entire.col = FALSE)
```

Arguments

- `x`:
  - an object of class `Dna`.
- `entire.col`:
  - boolean; entire columns with gaps are removed if this is TRUE. See also ‘Details’.

Details

If `entire.col`==TRUE, alignment is preserved. If it is FALSE, end gaps are introduced to sequence matrix.

Value

- an object of class `Dna`.

Methods

- `signature(x = "Dna")`

Author(s)

Caner Aktas, <caktas.aca@gmail.com>
## Examples

```r
data("dna.obj")

## original data
x<-dna.obj
range(x)
x@seqlengths

## Only gaps '-' are removed from sequences.
range(x)
x@seqlengths

## entire columns with gaps are removed.
range(x)
x@seqlengths
```

---

### rownames-methods

Retrieve or set the row names

---

### Description

Function to get or set row names of a sequence matrix in a Dna object or distance matrix (or matrices) in a Parsimnet object.

### Usage

```r
## S4 method for signature 'Dna'
rownames(x)

## S4 method for signature 'Parsimnet'
rownames(x)

## S4 replacement method for signature 'Dna'
rownames(x)<-value

## S4 replacement method for signature 'Parsimnet'
rownames(x)<-value
```

### Arguments

- `x` an object of class Dna or Parsimnet.
- `value` a character vector of the same length as number of sequences or a list of the same length as number of networks including vertex names for each network. See ‘Examples’
Methods

signature(x = "Dna")
signature(x = "Parsimnet")

Examples

data("dna.obj")
x<-dna.obj

### Method for signature 'Dna'.

## Getting sequence names.
rownames(x)

## Setting sequence names.
rownames(x)<-c(1:nrow(x))
rownames(x)

### Method for signature 'Parsimnet'.
x<-dna.obj

##single network
p<-parsimnet(x)

##Getting vertex names
rownames(p)

## Setting vertex names.
rownames(p)<-list(c(1:nrow(p@d[[1]])))
rownames(p)
plot(p)

## Multiple networks with 99% connection limit.
p<-parsimnet(x,prob=.99)

## Getting vertex names
rownames(p)

## Setting vertex names.
rownames(p)<-list(1:9, 10, 11,12:13,14,15:16)
rownames(p)
**Description**

Show objects of classes Dna, Haplotype, and Parsimnet

**Methods**

\[
\text{signature(object = "Dna") displays Dna object briefly: The total number of DNA sequences, names of the first six sequences (if nrow(x)>6), length of the shortest and longest sequences and the names of the slots.}
\]

\[
\text{signature(object = "Haplotype") displays Haplotype object briefly: The list of individuals that share the same haplotypes, the total number of haplotypes and the names of the slots.}
\]

\[
\text{signature(object = "Parsimnet") displays Parsimnet object briefly: The total number of networks, the maximum connection steps at chosen probability, the total number of haplotypes in each network, the total number of intermediates in each network, total network lengths (scores) of each network and the names of the slots.}
\]

---

**subs-methods**

*Displays base substitutions*

**Description**

This function displays all base substitutions. If fifth="TRUE", each gap is treated as a fifth state character.

**Usage**

```r
## S4 method for signature 'Dna'
subs(x,fifth=FALSE)
```

**Arguments**

- `x`: an object of class Dna.
- `fifth`: boolean; should gaps be treated as a fifth state character?

**Value**

a list with three components:

- `subsmat`: a sequence matrix showing substitutions.
- `subs`: a list of matrices of the substitutions.
- `subsmnum`: total number of substitutions.

**Methods**

\[
\text{signature(x = "Dna")}
\]
Author(s)

Caner Aktas, <caktas.aca@gmail.com>.

Examples

data("dna.obj")
x<-dna.obj

## Base substitutions.
subs(x)

## Gaps are treated as a fifth state character.
subs(x,fifth=TRUE)

tolower-methods, toupper-methods

Convert sequence characters from upper to lower case or vice versa

Description

Convert sequence characters in a Dna object from upper to lower case or vice versa.

Usage

## S4 method for signature 'Dna'
tolower(x)
## S4 method for signature 'Dna'
toupper(x)

Arguments

x an object of class Dna.

Value

an object of class Dna.

Methods

signature(x = "Dna")
Examples

## Coercing a list to a 'Dna' object.
seq1<-c("?", "A", "C", "G", "T", "-", "0", "1")
seq2<-c("?", "A", "C", "G", "T", "-", "0", "1", "2")
seq3<-c("?", "A", "C", "G", "T", "-", "0", "1", "2", "3")
x<-list(seq1=seq1,seq2=seq2,seq3=seq3)
dna.obj<-as.dna(x)

#characters in Dna object
table(as.matrix(dna.obj))

## all lower case
lowc<-tolower(dna.obj)
#characters
table(as.matrix(lowc))

## all upper case
upc<-toupper(dna.obj)
#characters
table(as.matrix(upc))

---

**unique-methods**

### Extract Unique Sequences

**Description**

unique returns a list with duplicate sequences removed.

**Usage**

```r
## S4 method for signature 'Dna'
unique(x,gaps=FALSE)
```

**Arguments**

- `x` an object of class `Dna`.
- `gaps` boolean; gaps are removed if this is `FALSE`.

**Details**

This function behaves somehow similar to `haplotype`, however indels and missing characters are not taken into account.

**Methods**

`signature(x = "Dna")`
Methods

Examples

data("dna.obj")
x<-dna.obj[1:6,as.matrix=FALSE]

## gaps removed.
unique(x)

## gaps not removed.
unique(x,gaps=TRUE)

## unique vs. haplotype
# unique returns 5 unique sequences.
unique(x)
length(unique(x))

## haplotype returns 4 unique haplotypes with simple indel coding.
h<-haplotype(x)
as.list(as.dna(h))
length(h)

## haplotype returns 3 unique haplotypes with gaps as missing.
h<-haplotype(x, indels="m")
as.list(as.dna(h))
length(h)

[-methods

Extract or replace parts of an object of class Dna

Description

Operators acting on sequence matrix to extract or replace parts.

Usage

## S4 method for signature 'Dna'
x[i, j, as.matrix = TRUE]
## S4 replacement method for signature 'Dna'
x[i, j]<- value

Arguments

x an object of class Dna
i, j elements to extract or replace.
as.matrix boolean; If TRUE, function returns a matrix, else, function returns an object of
value a character vector or a character matrix.
Value
returns an object of class matrix or Dna.

Methods
signature(x = "Dna", i = "ANY", j = "ANY", drop = "ANY")

Author(s)
Caner Aktas, <caktas.aca@gmail.com>

See Also
Dna

Examples

data("dna.obj")
x<-dna.obj

## Extract parts.
# a matrix object
x[1:2,1:3]

# a Dna object, as.dna(x[1:2,1:3]) gives the same result
x[1:2,1:3,as.matrix=FALSE]

## Replace parts.
# "G" "C"
x[1,1:2]
x[1,1:2]<-c("A","T")
x[1,1:2]
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