Package 'ConCat'

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1,000
Title Test your phylogeny
Version 1.0
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Description Perform alignment concatenation and analysis
Depends seqinr
License GPL (>= 2)
URL https://github.com/Ambuj-UF/ConCat-R
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concat Alignment concatenation function
Description Performs sequence alignment concatenation. It adds "?" for missing taxon sequence data.

Usage

```
concat(ext, form, outform, writeData = TRUE)
```

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Arguments

ext Input file extension. example: ".nex"

form Input sequence alignment format. It can be nexus, phylip, fasta and clustal

outform Output sequence alignment format. It can be nexus, phylip-relaxed, phylip-

interleived, phylip-sequential, fasta and clustal

writeData If TRUE (default = TRUE) the program creates a concatenated alignment file.

Otherwise it return the concatenated dataframe element

Author(s)

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entcal Class "entropy"

Description

Calculates entropy value for the given input alignment file.

Usage

```
entcal(file, type = "DNA")
```

Arguments

file Sequence alignment file name specified by either a variable of mode character,

or a double-quoted string; if file = "" (the default) then the program terminates

with message

type Alignment type. Can be DNA or Protein

Author(s)

Ambuj Kumar <ambuj@ufl.edu>

fevol Detects fast evolving sites

Description

To calculate observed variability (OV), all sequences for a given position are compared in a pairwise fashion. Mismatches are scored as 1 and matches as 0; the mean value amongst all the comparisons for a given position is used as the measure of character variability in the subsequent data sorting. Check reference for details.

Usage

```
fevol(file, cutoff = 0.9)
```

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Arguments

file Sequence alignment file name specified by either a variable of mode character,

or a double-quoted string; if file = "" (the default) then the program terminates

with message

cutoff OV value cutoff for fast evolving site detection. cutoff = 0.9 is set as default.

References

Goremykin VV1, Nikiforova SV, Bininda-Emonds OR. 2010. Automated removal of noisy data in phylogenomic analyses. J Mol Evol. 71:319-31. http://www.ncbi.nlm.nih.gov/pubmed/20976444

rcv

Relative confidence value

Description

Calculates RCV scores for the input alignment file

Usage

```
rcv(file, type = "DNA")
```

Arguments

file Sequence alignment file name specified by either a variable of mode character,

or a double-quoted string; if file = "" (the default) then the program terminates

with message

type Alignment type. Can be DNA or Protein

Author(s)

Ambuj Kumar <ambuj@ufl.edu>

read.nexus.data

Read Character Data In NEXUS Format

Description

This function reads a file with sequences in the NEXUS format.

Usage

```
read.nexus.data(file)
```

Arguments

file

a file name specified by either a variable of mode character, or a double-quoted string.

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Details

This parser tries to read data from a file written in a *restricted* NEXUS format (see examples below). Please see files 'data.nex' and 'taxacharacters.nex' for examples of formats that will work. Some noticeable exceptions from the NEXUS standard (non-exhaustive list):

• IComments must be either on separate lines or at the end of lines. Examples:

```
[Comment] — OK
Taxon ACGTACG [Comment] — OK
[Comment line 1
Comment line 2] — NOT OK!
Tax[Comment] on ACG[Comment] T — NOT OK!
```

• IINo spaces (or comments) are allowed in the sequences. Examples:

```
name ACGT — OK
name AC GT — NOT OK!
```

• IIINo spaces are allowed in taxon names, not even if names are in single quotes. That is, single-quoted names are not treated as such by the parser. Examples:

```
Genus_species — OK
'Genus_species' — OK
'Genus species' — NOT OK!
```

• IVThe trailing end that closes the matrix must be on a separate line. Examples:

```
taxon AACCGGT
end; — OK
taxon AACCGGT;
end; — OK
taxon AACCCGT; end; — NOT OK!
```

• VMultistate characters are not allowed. That is, NEXUS allows you to specify multiple character states at a character position either as an uncertainty, (XY), or as an actual appearance of multiple states, {XY}. This is information is not handled by the parser. Examples:

```
taxon 0011?110 — OK
taxon 0011{01}110 — NOT OK!
taxon 0011(01)110 — NOT OK!
```

• VIThe number of taxa must be on the same line as ntax. The same applies to nchar. Examples:

```
ntax = 12 — OK
ntax =
12 — NOT OK!
```

 VIIThe word "matrix" can not occur anywhere in the file before the actual matrix command, unless it is in a comment. Examples:

```
BEGIN CHARACTERS;
TITLE 'Data in file "03a-cytochromeB.nex"';
DIMENSIONS NCHAR=382;
FORMAT DATATYPE=Protein GAP=- MISSING=?;
["This is The Matrix"] — OK
MATRIX

BEGIN CHARACTERS;
TITLE 'Matrix in file "03a-cytochromeB.nex"'; — NOT OK!
DIMENSIONS NCHAR=382;
FORMAT DATATYPE=Protein GAP=- MISSING=?;
MATRIX
```

rycode 5

Value

A list of sequences each made of a single vector of mode character where each element is a (phylogenetic) character state.

Author(s)

Johan Nylander <nylander@scs.fsu.edu>

References

Maddison, D. R., Swofford, D. L. and Maddison, W. P. (1997) NEXUS: an extensible file format for systematic information. *Systematic Biology*, **46**, 590–621.

Examples

```
## Use read.nexus.data to read a file in NEXUS format into object x
## Not run: x <- read.nexus.data("file.nex")</pre>
```

rycode

"RY-coding"

Description

```
Class "RY-coding"
```

Usage

```
rycode(file, type = 3)
```

Arguments

file Sequence alignment file name specified by either a variable of mode character,

or a double-quoted string; if file = "" (the default) then the program terminates

with message

type RY coding position. If type=3 (Default=3), program performs RY coding at the

third codon position whereas if type="all" then program performs RY coding for

all the positions

Author(s)

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