

Package: phylotools (via r-universe)

September 14, 2024

Type Package

Title Phylogenetic Tools for Eco-Phylogenetics

Version 0.2.4

Date 2021-02-09

Author Jinlong Zhang [aut, cre], Nancai Pei [ctb], Xiangcheng Mi [ctb]

Maintainer Jinlong Zhang <jinlongzhang01@gmail.com>

Description A collection of tools for building RAxML supermatrix using PHYLIP or aligned FASTA files. These functions will be useful for building large phylogenies using multiple markers.

Depends ape

Suggests vegan, knitr, rmarkdown

VignetteBuilder knitr

License GPL-2

LazyLoad yes

URL <https://github.com/helixcn/phylotools>

NeedsCompilation no

Repository <https://helixcn.r-universe.dev>

RemoteUrl <https://github.com/helixcn/phylotools>

RemoteRef HEAD

RemoteSha 758d338b2d2004c5b94f931802d269251f4fd7a1

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phylotools-package *Phylogenetic tools for building PHYLIP supermatrix and more*

Description

A collection of a few functions for handling DNA-barcoding sequences, building PHYLIP supermatrix for RAxML etc.

Details

Package: phylotools
Type: Package
Version: 0.2.1
Date: 2017-12-08
License: GLP-2
LazyLoad: yes

Author(s)

Jinlong Zhang
Maintainer: Jinlong Zhang <jinlongzhang01@gmail.com>

clean.fasta.name *Clean the name of a fasta file*

Description

Cleaning the names of sequences for a fasta file. The punctuation characters and the white space will be replaced with "_".

Usage

```
clean.fasta.name(infile = NULL, outfile = "name_cleaned.fasta")
```

Arguments

infile	character string representing the name of the fasta file.
outfile	Character string representing the file name to be generated.

Details

Punctuation characters and white space will be replaced by "_". More information can be found at [regex](#).

Value

This is a subroutine without a return value. A fasta file with all the names of sequences renamed will be saved to the working directory.

Author(s)

Jinlong Zhang <jinlongzhang01@gmail.com>

References

http://www.genomatix.de/online_help/help/sequence_formats.html

See Also

[read.fasta](#)

Examples

```
cat(
  ">seq_1*66", "--TTACAAATTGACTTATTATA",
  ">seq_2()r", "GATTACAAATTGACTTATTATA",
  ">seq_3:test", "GATTACAAATTGACTTATTATA",
  ">seq_588", "GATTACAAATTGACTTATTATA",
  ">seq_8$$yu", "GATTACAAATTGACTTATTATA",
  ">seq_10", "---TACAAATTGAATTATTATA",
  file = "matk.fasta", sep = "\n")

clean.fasta.name(infile = "matk.fasta")
get.fasta.name("name_cleaned.fasta")

# Delete file
unlink("matk.fasta")
unlink("name_cleaned.fasta")
```

`dat2fasta`*Convert and Save sequence data frame to fasta file*

Description

Convert and Save sequence data frame to fasta file.

Usage

```
dat2fasta(dat, outfile = "out.fasta")
```

Arguments

<code>dat</code>	data frame by read.phylip or read.fasta
<code>outfile</code>	a character string, representing the name of the fasta file to be generated

Details

The column of the data frame must be: 1. seq.name, 2. seq.text, represent the name of the sequences, the content of the sequence, eg. ATCGGGAAC.

Value

This is a routine without return value.

Author(s)

Jinlong Zhang <jinlongzhang01@gmail.com>

References

http://www.genomatix.de/online_help/help/sequence_formats.html

See Also

[read.fasta](#), [read.phylip](#)

Examples

```
cat(
">seq_2", "GTCTTATAAGAAAGAATAAGAAAG--AAATACAAA-----AAAAAAGA",
">seq_3", "GTCTTATAAGAAAGAAATAGAAAAGTAAAAAAAA-----AAAAAAG",
">seq_5", "GACATAAGACATAAAATAGAATACTCAATCAGAAACCAACCCATAAAAC",
">seq_8", "ATTCCAAAATAAAATACAAAAAGAAAAAACTAGAAAGTTTTTTTCTTTG",
">seq_9", "ATCCTTGTTCTTTTTTCTTTAATCTTTAAATAAACCTTTTTTTTTTA",
file = "trn1.fasta", sep = "\n")

res <- read.fasta("trn1.fasta")
dat2fasta(res)
```

```
unlink("trn1.fasta")
unlink("out.fasta")
```

dat2phylip*Convert the data frame to sequential PHYLIP format file*

Description

Convert and save a data frame to sequential PHYLIP file.

Usage

```
dat2phylip(dat, outfile = "out.phy")
```

Arguments

dat	the data frame returned by read.phylip , read.fasta .
outfile	character string represents the phylip file to be generated.

Details

The output will be in sequential PHYLIP format.

Value

This is a subroutine, there is no return value.

Note

The names of the sequences should not contain white space or Punctuation characters. See [regex](#) for more details.

Author(s)

Jinlong Zhang <jinlongzhang01@gmail.com>

References

http://www.genomatix.de/online_help/help/sequence_formats.html

See Also

[dat2fasta](#), [read.fasta](#), [read.phylip](#)

Examples

```

cat(
  ">seq_2", "GTCTTATAAGAAAGAATAAGAAAG--AAATACAAA-----AAAAAAGA",
  ">seq_3", "GTCTTATAAGAAAGAAATAGAAAAGTAAAAAAAA-----AAAAAAG",
  ">seq_5", "GACATAAGACATAAAATAGAATACTCAATCAGAAACCAACCCATAAAAAC",
  ">seq_8", "ATTCCAAAATAAAATACAAAAGAAAAAACTAGAAAGTTTTTTTCTTTG",
  ">seq_9", "ATTCTTTGTTCTTTTTTTTCTTTAATCTTTAAATAAACCTTTTTTTTTTA",
  file = "trn1.fasta", sep = "\n")

res <- read.fasta("trn1.fasta")
dat2phylip(res)
unlink("trn1.fasta")
unlink("out.phy")

```

<code>get.fasta.name</code>	<i>get the names of all the sequences of fasta file</i>
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Description

get the names of all the sequences of a fasta file, and perform cleaning of the names of the sequences

Usage

```
get.fasta.name(infile, clean_name = FALSE)
```

Arguments

`infile` character string representing the name of the fasta file.
`clean_name` logical, representing cleaning of the names will be performed.

Value

a character vector containing the names of the sequences

Note

Punctuation characters and white space be replaced by "_". Definition of Punctuation characters can be found at [regex](#).

Author(s)

Jinlong Zhang <jinlongzhang01@gmail.com>

References

http://www.genomatix.de/online_help/help/sequence_formats.html

See Also[read.fasta](#), [regex](#)**Examples**

```

cat(
  ">seq_2", "GTCTTATAAGAAAGAATAAGAAAG--AAATACAAA-----AAAAAAGA",
  ">seq_3", "GTCTTATAAGAAAGAAATAGAAAAGTAAAAAAAA-----AAAAAAG",
  ">seq_5", "GACATAAGACATAAAATAGAATACTCAATCAGAAACCAACCCATAAAAAC",
  ">seq_8", "ATTCCAAAATAAAATACAAAAGAAAAAACTAGAAAGTTTTTTTCTTTG",
  ">seq_9", "ATTCTTTGTTCTTTTTTTCTTTAATCTTTAAATAAACCTTTTTTTTTTA",
  file = "trn1.fasta", sep = "\n")
get.fasta.name("trn1.fasta")
unlink("trn1.fasta")

```

get.phylip.name	<i>get the names of sequences from a PHYLIP file</i>
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Description

get the names of sequences from a PHYLIP file.

Usage

```
get.phylip.name(infile, clean_name = FALSE)
```

Arguments

infile	character representing the name or path of the phylip file.
clean_name	logical, representing cleaning of the names will be performed.

Details

Punctuation characters and white space be replaced by "_". Definition of Punctuation characters can be found at [regex](#).

Value

a character vector of the names of the sequences

Author(s)

Jinlong Zhang <jinlongzhang01@gmail.com>

See Also[read.phylip](#), [regex](#)

Examples

```
cat("6 22",
    "seq_1  --TTACAAATTGACTTATTATA",
    "seq_2  GATTACAAATTGACTTATTATA",
    "seq_3  GATTACAAATTGACTTATTATA",
    "seq_5  GATTACAAATTGACTTATTATA",
    "seq_8  GATTACAAATTGACTTATTATA",
    "seq_10 ---TACAAATTGAATTATTATA",
    file = "matk.phy", sep = "\n")
get.phylip.name("matk.phy")
unlink("matk.phy")
```

read.fasta

Read FASTA file

Description

Read and convert the fasta file to data frame

Usage

```
read.fasta(file = NULL, clean_name = FALSE)
```

Arguments

file character string representing the name of the fasta file.

clean_name logical, representing cleaning of the names will be performed. Punctuation characters and white space be replaced by "_". See [regex](#) for more details.

Details

In this function, names of the sequences are identified by ">", and all the lines before next ">" will be concatenated.

Value

a data frame with two columns: (1) seq.name, the names for all the sequences. (2) seq.text, the raw sequence data.

Note

Punctuation characters and white space in the names of the sequences will be replaced by "_".

Author(s)

Jinlong Zhang <jinlongzhang01@gmail.com>

References

http://www.genomatix.de/online_help/help/sequence_formats.html

See Also

[read.phylip](#), [dat2fasta](#), [dat2phylip](#), [split_dat](#)

Examples

```
cat(
  ">seq_2", "GTCTTATAAGAAAGAATAAGAAAG--AAATACAAA-----AAAAAGA",
  ">seq_3", "GTCTTATAAGAAAGAAATAGAAAAGTAAAAAAAA-----AAAAAAG",
  ">seq_5", "GACATAAGACATAAAATAGAATACTCAATCAGAAACCAACCCATAAAAC",
  ">seq_8", "ATTCCAAAATAAAATACAAAAAGAAAAACTAGAAAGTTTTTTTCTTG",
  ">seq_9", "ATTCTTTGTTCTTTTTTCTTTAATCTTAAATAAACCTTTTTTTT",
  file = "trn1.fasta", sep = "\n")

res <- read.fasta("trn1.fasta")
unlink("trn1.fasta")
```

read.phylip

read phylip file

Description

read the phylip file, and store the sequences and their names in a dataframe.

Usage

```
read.phylip(infile, clean_name = TRUE, seq.name.length = NA)
```

Arguments

<code>infile</code>	character string for the name of the phylip file.
<code>clean_name</code>	logical, representing cleaning of the names will be performed.
<code>seq.name.length</code>	Number of characters for the sequence names, if not specified, the function will use the first white space as the identifier. All the character before the first white space will be treated as sequence names, and the rest will be treated as sequences.

Details

`read.phylip` accepts both interleaved and sequential phylip, the number of sequences is identified by parsing the first line of the file. Sequences and their names will be stored in a data frame.

If `clean_name` is `TRUE`, punctuation characters and white space be replaced by `"_"`. Definition of punctuation characters can be found at [regex](#).

Value

a data frame with two columns: (1) seq.name, the names for all the sequences; (2) seq.text, the raw sequence data.

Note

the Punctuation characters and white space in the names of the sequences will be replaced by "_".

Author(s)

Jinlong Zhang <jinlongzhang01@gmail.com>

See Also

[read.fasta](#)

Examples

```
cat("6 22",
    "seq_1  --TTACAAATTGACTTATTATA",
    "seq_2  GATTACAAATTGACTTATTATA",
    "seq_3  GATTACAAATTGACTTATTATA",
    "seq_5  GATTACAAATTGACTTATTATA",
    "seq_8  GATTACAAATTGACTTATTATA",
    "seq_10 ---TACAAATTGAATTATTATA",
    file = "matk.phy", sep = "\n")

res <- read.phylip(infile = "matk.phy")
unlink("matk.phy")
```

rename.fasta

Rename the sequences for a fasta file

Description

Rename the sequences within a fasta file according to a data frame supplied.

Usage

```
rename.fasta(infile = NULL, ref_table, outfile = "renamed.fasta")
```

Arguments

infile	character string containing the name of the fasta file.
ref_table	a data frame with first column for original name, second column for the new name of the sequence.
outfile	The name of the fasta file with sequences renamed.

Details

If the original name was not found in the `ref_table`, the name for the sequence will be changed into "old_name_" + original name.

Value

This is a subroutine without return value.

Note

Since whitespace and punctuation characters will be replaced with "_", name of a sequence might change. It is suggest to obtain the name of the sequences by calling `read.fasta` first, and save the `data.frame` to a csv file to obtain the "original" name for the sequences.

Author(s)

Jinlong Zhang <jinlongzhang01@gmail.com>

References

http://www.genomatix.de/online_help/help/sequence_formats.html

See Also

[read.fasta](#), [split_dat](#)

Examples

```
cat(
  ">seq_1", "--TTACAAATTGACTTATTATA",
  ">seq_2", "GATTACAAATTGACTTATTATA",
  ">seq_3", "GATTACAAATTGACTTATTATA",
  ">seq_5", "GATTACAAATTGACTTATTATA",
  ">seq_8", "GATTACAAATTGACTTATTATA",
  ">seq_10", "---TACAAATTGAATTATTATA",
  file = "matk.fasta", sep = "\n")
old_name <- get.fasta.name("matk.fasta")
new_name <- c("Magnolia", "Ranunculus", "Carex", "Morus", "Ulmus", "Salix")
ref2 <- data.frame(old_name, new_name)
rename.fasta(infile = "matk.fasta", ref_table = ref2, outfile = "renamed.fasta")
unlink("matk.fasta")
unlink("renamed.fasta")
```

rm.sequence.fasta *Delete sequences from fasta file*

Description

Delete sequences from fasta file

Usage

```
rm.sequence.fasta(infile, outfile = "sequence.removed.fasta", to.rm = NULL)
```

Arguments

infile	Character string representing the name of the fasta file.
outfile	Character string representing the name of the output fasta file.
to.rm	Vector of character string containing the names of sequences to be deleted.

Details

Delete sequences from a fasta file.

Value

This is a subroutine without return value.

Author(s)

Jinlong Zhang <jinlongzhang01@gmail.com>

References

http://www.genomatix.de/online_help/help/sequence_formats.html

See Also

[read.fasta](#), [dat2fasta](#)

Examples

```
cat(
">seq_1", "---TCCGCCCCCTACTCTA",
">seq_3", "CTCTCCGCCCCTACTCTA",
">seq_5", "---TCCGCCC-TTACTCTA",
">seq_6", "---TCCGCCCCTACTCTA",
">seq_9", "---TCCGCCC-TCTACTCTA",
">seq_12", "CTCTCCGCCC-TCTACTCTA",
file = "trn2.fasta", sep = "\n")
```

```
rm.sequence.fasta(infile = "trn2.fasta", to.rm = c("seq_1","seq_12"))  
  
unlink("trn2.fasta")  
unlink("sequence.removed.fasta")
```

split_dat	<i>grouping the data frame containing sequences and names and generate fasta file</i>
-----------	---------------------------------------------------------------------------------------

Description

Split the data frame of sequences based on the reference table of grouping.

Usage

```
split_dat(dat, ref_table)
```

Arguments

dat	data frame generated by read.phylip or read.fasta
ref_table	data frame with first column for the name of the sequence, second column for the group the sequence belongs to.

Details

Each group of sequences will be saved to a fasta file. Sequences not included in the ref_table will be saved in "Ungrouped.fasta"

Value

This is a subroutine, there is no return value.

Author(s)

Jinlong Zhang <jinlongzhang01@gmail.com>

References

http://www.genomatix.de/online_help/help/sequence_formats.html

See Also

[rename.fasta](#)

Examples

```

cat(
  ">seq_1", "--TTACAAATTGACTTATTATA",
  ">seq_2", "GATTACAAATTGACTTATTATA",
  ">seq_3", "GATTACAAATTGACTTATTATA",
  ">seq_5", "GATTACAAATTGACTTATTATA",
  ">seq_8", "GATTACAAATTGACTTATTATA",
  ">seq_10", "---TACAAATTGAATTATTATA",
  ">seq_11", "--TTACAAATTGACTTATTATA",
  ">seq_12", "GATTACAAATTGACTTATTATA",
  ">seq_13", "GATTACAAATTGACTTATTATA",
  ">seq_15", "GATTACAAATTGACTTATTATA",
  ">seq_16", "GATTACAAATTGACTTATTATA",
  ">seq_17", "---TACAAATTGAATTATTATA",
  file = "trnh.fasta", sep = "\n")

sequence_name <- get.fasta.name("trnh.fasta")
sequence_group <- c("group1", "group1", "group1", "group1", "group1",
"group2", "group2", "group2", "group3", "group3", "group3", "group3")
group <- data.frame(sequence_name, sequence_group)

fasta <- read.fasta("trnh.fasta")
split_dat(fasta, group)

unlink("trnh.fasta")
unlink("ungrouped.fasta")
unlink("group1.fasta")
unlink("group2.fasta")
unlink("group3.fasta")

```

sub.taxa.label	<i>Substitute the tip labels of a phylogenetic tree</i>
----------------	---------------------------------------------------------

Description

Substitute the tip labels of a phylogenetic tree according to a reference data table.

Usage

```
sub.taxa.label(tree, dat)
```

Arguments

tree	Phylogenetic tree
dat	A dataframe with the first column the tip labels and the second column the new names.

Value

A Phylogenetic tree with the tip labels substituted

Author(s)

Jinlong Zhang <jinlongzhang01@gmail.com>

See Also

[read.tree](#)

Examples

```
library(ape)
data(bird.families)
tips <- bird.families$tip.label
abr <- paste("fam", 1:length(tips), sep = "")
dat <- data.frame(tips, abr)
ntree <- sub.taxa.label(bird.families, dat)
```

supermat	<i>Build PHYLIP supermatrix and RAxML partition file using aligned FASTA or PHYLIP files.</i>
----------	-----------------------------------------------------------------------------------------------

Description

Build PHYLIP supermatrix and create RAxML partition file using aligned fasta or phylip files.

Usage

```
supermat(infile, outfile = "supermat.out.phy",
          partition.file = "gene_partition.txt")
```

Arguments

`infile` a character string vector for phylip or aligned fasta file.
`outfile` the name of the PHYLIP supermatrix
`partition.file` partition data summary describing the genes.

Details

Supermatrix here means a phylip file with combined aligned sequences. The missing sequences should be replaced with either "?" or "-".

Value

A list containing: (1)supermat.dat:a list containing all the data frames read by read.phylip or read.fasta (2)res.super.dat: a data frame containing the sequences and the names (3)partition.dat: summary for all the fasta or phylip files (4)partition.dat.vector: character string vector for the partition file for RAxML

Note

Punctuation characters and white space in the names of the sequences will be replaced by "_". More information can be found at [regex](#). Type of the sequence in the RAxML partition file should be changed manually according to the manual of RAxML.

Author(s)

Jinlong Zhang <jinlongzhang01@gmail.com>

References

Kress, W. J., Erickson, D. L., Jones, F. A., Swenson, N. G., Perez, R., Sanjur, O., & Bermingham, E. (2009). Plant DNA barcodes and a community phylogeny of a tropical forest dynamics plot in Panama. *Proceedings of the National Academy of Sciences*, 106(44), 18621-18626.

de Queiroz, A. and Gatesy, J. (2007). The supermatrix approach to systematics. *Trends in Ecology & Evolution*, 22(1), 34-41.

<https://github.com/stamatak/standard-RAxML>

See Also

[read.fasta](#),[read.phylip](#),[dat2phylip](#),

Examples

```
cat("6 22",
    "seq_1  --TTACAAATTGACTTATTATA",
    "seq_2  GATTACAAATTGACTTATTATA",
    "seq_3  GATTACAAATTGACTTATTATA",
    "seq_5  GATTACAAATTGACTTATTATA",
    "seq_8  GATTACAAATTGACTTATTATA",
    "seq_10 ---TACAAATTGAATTATTATA",
    file = "matk.phy", sep = "\n")
```

```
cat("5 15",
    "seq_1  GATTACAAATTGACT",
    "seq_3  GATTACAAATTGACT",
    "seq_4  GATTACAAATTGACT",
    "seq_5  GATTACAAATTGACT",
    "seq_8  GATTACAAATTGACT",
    file = "rbcla.phy", sep = "\n")
```

```
cat("5 50",
    "seq_2  GTCTTATAAGAAAGAATAAGAAAG--AAATACAAA-----AAAAAAGA",
```

```

"seq_3      GTCTTATAAGAAAGAATAGAAAAGTAAAAAAAA-----AAAAAAG",
"seq_5      GACATAAGACATAAAATAGAATACTCAATCAGAAACCAACCCATAAAAC",
"seq_8      ATTCCAAAATAAAATACAAAAGAAAAAACTAGAAAGTTTTTTTCTTTG",
"seq_9      ATTCTTTGTTCTTTTTTTCTTTAATCTTTAAATAAACCTTTTTTTTTTA",
file = "trn1.phy", sep = "\n")

supermat(infiles = c("matk.phy", "rbcla.phy", "trn1.phy"))
unlink(c("matk.phy", "rbcla.phy", "trn1.phy"))
unlink(c("supermat.out.phy", "gene_partition.txt"))

cat(
">seq_1", "--TTACAAATTGACTTATTATA",
">seq_2", "GATTACAAATTGACTTATTATA",
">seq_3", "GATTACAAATTGACTTATTATA",
">seq_5", "GATTACAAATTGACTTATTATA",
">seq_8", "GATTACAAATTGACTTATTATA",
">seq_10", "---TACAAATTGAATTATTATA",
file = "matk.fasta", sep = "\n")

cat(
">seq_1", "GATTACAAATTGACT",
">seq_3", "GATTACAAATTGACT",
">seq_4", "GATTACAAATTGACT",
">seq_5", "GATTACAAATTGACT",
">seq_8", "GATTACAAATTGACT",
file = "rbcla.fasta", sep = "\n")

cat(
">seq_2", "GTCTTATAAGAAAGAATAAGAAAG--AAATACAAA-----AAAAAAGA",
">seq_3", "GTCTTATAAGAAAGAATAGAAAAGTAAAAAAAA-----AAAAAAG",
">seq_5", "GACATAAGACATAAAATAGAATACTCAATCAGAAACCAACCCATAAAAC",
">seq_8", "ATTCCAAAATAAAATACAAAAGAAAAAACTAGAAAGTTTTTTTCTTTG",
">seq_9", "ATTCTTTGTTCTTTTTTTCTTTAATCTTTAAATAAACCTTTTTTTTTTA",
file = "trn1.fasta", sep = "\n")

supermat(infiles = c("matk.fasta", "rbcla.fasta", "trn1.fasta"))
unlink(c("matk.fasta", "rbcla.fasta", "trn1.fasta"))

unlink(c("supermat.out.phy", "gene_partition.txt"))

```

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