Package 'fastreeR'

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Description Calculate distances, build phylogenetic trees or
     perform hierarchical clustering between the samples of a VCF or FASTA file.
     Functions are implemented in Java and called via rJava.
     Parallel implementation that operates directly on the VCF or FASTA file
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2 fastreeR-package

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R topics documented:

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Description

Calculate distances, build phylogenetic trees or perform hierarchical clustering between the samples of a VCF or FASTA file. Functions are implemented in Java and called via rJava. Parallel implementation that operates directly on the VCF or FASTA file for fast execution.

Author(s)

Maintainer: Anestis Gkanogiannis <anestis@gkanogiannis.com> (ORCID)

See Also

Useful links:

- https://github.com/gkanogiannis/fastreeR
- https://github.com/gkanogiannis/BioInfoJava-Utils
- Report bugs at https://github.com/gkanogiannis/fastreeR/issues

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dist2clusters	Perform Hierarchical Clustering and tree pruning on a distance matrix

Description

Performs Hierarchical Clustering on a distance matrix (i.e. calculated with vcf2dist or fasta2dist) and generates a phylogenetic tree with agglomerative Neighbor Joining method (complete linkage) (as in dist2tree). The phylogenetic tree is then pruned with cutreeDynamic to get clusters (as in tree2clusters).

Usage

```
dist2clusters(inputDist, cutHeight = NULL, minClusterSize = 1, extra = TRUE)
```

Arguments

inputDist Input distances file location (generated with vcf2dist or fasta2dist). File can

be gzip compressed. Or a dist distances object.

cutHeight Define at which height to cut tree. Default automatically defined.

minClusterSize Minimum size of clusters. Default 1.

extra Boolean whether to use extra parameters for the cutreeDynamic.

Value

A list of:

- character vector of the generated phylogenetic tree in Newick format
- character vector of the clusters. Each row contains data for a cluster, separated by space. The id of the cluster, the size of the cluster (number of elements) and the names of its elements, Cluster id 0 contains all the objects not assigned to a cluster (singletons). Example clusters output:

```
0
   3
       Sample1
                Sample2
                         Sample3
  3
1
       Sample4
                Sample5
                         Sample6
2 2
       Sample7
                Sample8
3
  2
       Sample9
                Sample0
```

Author(s)

Anestis Gkanogiannis, <anestis@gkanogiannis.com>

References

Java implementation: https://github.com/gkanogiannis/BioInfoJava-Utils

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Examples

```
my.clust <- dist2clusters(
    inputDist =
        system.file("extdata", "samples.vcf.dist.gz", package = "fastreeR")
)</pre>
```

dist2tree

Generate phylogenetic tree from samples of a distance matrix

Description

Performs Hierarchical Clustering on a distance matrix (i.e. calculated with vcf2dist or fasta2dist) and generates a phylogenetic tree with agglomerative Neighbor Joining method (complete linkage).

Usage

```
dist2tree(inputDist)
```

Arguments

inputDist

Input distances file location (generated with vcf2dist or fasta2dist). File can be gzip compressed. Or a dist distances object.

Value

A character vector of the generated phylogenetic tree in Newick format.

Author(s)

Anestis Gkanogiannis, <anestis@gkanogiannis.com>

References

Java implementation: https://github.com/gkanogiannis/BioInfoJava-Utils

```
my.tree <- dist2tree(
    inputDist =
        system.file("extdata", "samples.vcf.dist.gz", package = "fastreeR")
)</pre>
```

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fasta2dist

Calculate distances between sequences of a FASTA file

Description

This function calculates a d2_S type dissimilarity measurement between the n sequences (which can represent samples) of a FASTA file. See doi:10.1186/s1285901611863 for more details.

Usage

```
fasta2dist(
    ...,
    outputFile = NULL,
    threads = 2,
    kmer = 6,
    normalize = FALSE,
    compress = TRUE
)
```

Arguments

... Input fasta files locations (uncompressed or gzip compressed).

outputFile Output distances file location. threads Number of java threads to use.

kmer Kmer length to use for analyzing fasta sequences.

normalize Normalize on sequences length.

compress Compress output (adds .gz extension).

Value

A dist distances object of the calculation.

Author(s)

Anestis Gkanogiannis, <anestis@gkanogiannis.com>

References

Java implementation: https://github.com/gkanogiannis/BioInfoJava-Utils

```
my.dist <- fasta2dist(
    inputfile = system.file("extdata", "samples.fasta.gz",
        package = "fastreeR"
    )
)</pre>
```

6 tree2clusters

_	
tree2clusters	Perform Hierarchical Clustering and tree pruning on a phylogenetic
	tree

Description

The phylogenetic tree is pruned with cutreeDynamic to get clusters.

Usage

```
tree2clusters(
  treeStr,
  treeDistances = NULL,
  treeLabels = NULL,
  cutHeight = NULL,
  minClusterSize = 1,
  extra = TRUE
)
```

Arguments

treeStr A character vector of a phylogenetic tree in Newick format

treeDistances codenumeric matrix of distances, that were used to generate the tree. If NULL,

it will be inferred from tree branch lengths.

treeLabels A character vector of tree leaf labels.

cutHeight Define at which height to cut tree. Default automatically defined.

minClusterSize Minimum size of clusters. Default 1.

extra Boolean whether to use extra parameters for the cutreeDynamic.

Value

• character vector of the clusters. Each row contains data for a cluster, separated by space. The id of the cluster, the size of the cluster (number of elements) and the names of its elements, Cluster id 0 contains all the objects not assigned to a cluster (singletons). Example clusters output:

```
0
   3
       Sample 1
                Sample2
                         Sample3
1
   3
       Sample4
                Sample5
                         Sample6
2
  2
       Sample7
                Sample8
       Sample9
                Sample0
```

Author(s)

Anestis Gkanogiannis, <anestis@gkanogiannis.com>

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References

Java implementation: https://github.com/gkanogiannis/BioInfoJava-Utils

Examples

```
my.clust <- tree2clusters(
    treeStr = dist2tree(
        inputDist = system.file("extdata", "samples.vcf.dist.gz",
            package = "fastreeR"
        )
    )
)</pre>
```

vcf2clusters

Perform Hierarchical Clustering and tree pruning on samples of VCF file

Description

Performs Hierarchical Clustering on a distance matrix calculated as in vcf2dist and generates a phylogenetic tree with agglomerative Neighbor Joining method (complete linkage) (as in dist2tree). The phylogenetic tree is then pruned with cutreeDynamic to get clusters (as in tree2clusters).

Usage

```
vcf2clusters(
  inputFile,
  threads = 2,
  ignoreMissing = FALSE,
  onlyHets = FALSE,
  ignoreHets = FALSE,
  cutHeight = NULL,
  minClusterSize = 1,
  extra = TRUE
)
```

Arguments

inputFile Input vcf file location (uncompressed or gzip compressed). threads Number of java threads to use.

ignoreMissing Ignore variants with missing data (./. or .|.)
onlyHets Only calculate on variants with heterozygous calls.
ignoreHets Only calculate on variants with homozygous calls.

cutHeight Define at which height to cut tree. Default automatically defined.

minClusterSize Minimum size of clusters. Default 1.

extra Boolean whether to use extra parameters for the cutreeDynamic.

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Details

Biallelic or multiallelic (maximum 7 alternate alleles) SNP and/or INDEL variants are considered, phased or not. Some VCF encoding examples are:

- heterozygous variants: 1/0 or 0/1 or 0/2 or 1 | 0 or 0 | 1 or 0 | 2
- homozygous to the reference allele variants: 0/0 or 0 | 0
- homozygous to the first alternate allele variants: 1/1 or 1|1

If there are n samples and m variants, an nxn zero-diagonal symmetric distance matrix is calculated. The calculated cosine type distance (1-cosine_similarity)/2 is in the range [0,1] where value 0 means completely identical samples (cosine is 1), value 0.5 means perpendicular samples (cosine is 0) and value 1 means completely opposite samples (cosine is -1).

The calculation is performed by a Java back-end implementation, that supports multi-core CPU utilization and can be demanding in terms of memory resources. By default a JVM is launched with a maximum memory allocation of 512 MB. When this amount is not sufficient, the user needs to reserve additional memory resources, before loading the package, by updating the value of the java.parameters option. For example in order to allocate 4GB of RAM, the user needs to issue options(java.parameters="-Xmx4g") before library(fastreeR).

Value

A list of:

- dist distances object.
- character vector of the generated phylogenetic tree in Newick format
- character vector of the clusters. Each row contains data for a cluster, separated by space. The id of the cluster, the size of the cluster (number of elements) and the names of its elements, Cluster id 0 contains all the objects not assigned to a cluster (singletons). Example clusters output:

```
0
   3
       Sample 1
                 Sample2
                           Sample3
   3
                           Sample6
       Sample4
                 Sample5
   2
2
       Sample7
                 Sample8
3
   2
       Sample9
                 Sample0
```

Author(s)

Anestis Gkanogiannis, <anestis@gkanogiannis.com>

References

Java implementation: https://github.com/gkanogiannis/BioInfoJava-Utils

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Examples

vcf2dist

Calculate distances between samples of a VCF file

Description

This function calculates a cosine type dissimilarity measurement between the n samples of a VCF file.

Usage

```
vcf2dist(
  inputFile,
  outputFile = NULL,
  threads = 2,
  ignoreMissing = FALSE,
  onlyHets = FALSE,
  ignoreHets = FALSE,
  compress = FALSE
)
```

Arguments

inputFile Input vcf file location (uncompressed or gzip compressed).

outputFile Output distances file location.

threads Number of java threads to use.

ignoreMissing Ignore variants with missing data (./. or .|.)

onlyHets Only calculate on variants with heterozygous calls.

ignoreHets Only calculate on variants with homozygous calls.

Compress output (adds .gz extension).

Details

compress

Biallelic or multiallelic (maximum 7 alternate alleles) SNP and/or INDEL variants are considered, phased or not. Some VCF encoding examples are:

- heterozygous variants : 1/0 or 0/1 or 0/2 or 1 | 0 or 0 | 1 or 0 | 2
- homozygous to the reference allele variants: 0/0 or 0|0
- homozygous to the first alternate allele variants: 1/1 or 1|1

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If there are n samples and m variants, an nxn zero-diagonal symmetric distance matrix is calculated. The calculated cosine type distance (1-cosine_similarity)/2 is in the range [0,1] where value 0 means completely identical samples (cosine is 1), value 0.5 means perpendicular samples (cosine is 0) and value 1 means completely opposite samples (cosine is -1).

The calculation is performed by a Java backend implementation, that supports multi-core CPU utilization and can be demanding in terms of memory resources. By default a JVM is launched with a maximum memory allocation of 512 MB. When this amount is not sufficient, the user needs to reserve additional memory resources, before loading the package, by updating the value of the java.parameters option. For example in order to allocate 4GB of RAM, the user needs to issue options(java.parameters="-Xmx4g") before library(fastreeR).

Output file, if provided, will contain n+1 lines. The first line contains the number n of samples and number m of variants, separated by space. Each of the subsequent n lines contains n+1 values, separated by space. The first value of each line is a sample name and the rest n values are the calculated distances of this sample to all the samples. Example output file of the distances of 3 samples calculated from 1000 variants:

```
3 1000
Sample1 0.0 0.5 0.2
Sample2 0.5 0.0 0.9
Sample3 0.2 0.9 0.0
```

Value

A dist distances object of the calculation.

Author(s)

Anestis Gkanogiannis, <anestis@gkanogiannis.com>

References

Java implementation: https://github.com/gkanogiannis/BioInfoJava-Utils

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vcf2istats

Calculate various per sample statistics from a VCF file

Description

Only biallelic SNPs are considered. For each sample, the following statistics are calculated:

- INDIV : Sample name
- N_SITES: Total number of SNPs
- N_HET: Number of SNPs with heterozygous call (0/1 or 0 | 1 or 1/0 or 1 | 0)
- N_ALT : Number of SNPs with alternate homozygous call (1/1 or 1|1)
- N_REF: Number of SNPs with reference homozygous call (0/0 or 0|0)
- N_MISS: Number of SNPs with missing call (./. or .|.)
- P_HET : Percentage of heterozygous calls
- P_ALT : Percentage of alternate homozygous calls
- P_REF: Percentage of reference homozygous calls
- P_MISS : Percentage of missing calls (missing rate)

Usage

```
vcf2istats(inputFile, outputFile = NULL)
```

Arguments

```
inputFile Input vcf file location (uncompressed or gzip compressed).

outputFile Output samples statistics file location.
```

Value

A data. frame of sample statistics.

Author(s)

Anestis Gkanogiannis, <anestis@gkanogiannis.com>

References

Java implementation: https://github.com/gkanogiannis/BioInfoJava-Utils

```
my.istats <- vcf2istats(
    inputFile =
        system.file("extdata", "samples.vcf.gz", package = "fastreeR")
)</pre>
```

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vcf2tree

Generate phylogenetic tree from samples of a VCF file

Description

This function calculates a distance matrix between the samples of a VCF file as in vcf2dist and performs Hierarchical Clustering on this distance matrix as in dist2tree. A phylogenetic tree is calculated with agglomerative Neighbor Joining method (complete linkage).

Usage

```
vcf2tree(
  inputFile,
  threads = 2,
  ignoreMissing = FALSE,
  onlyHets = FALSE,
  ignoreHets = FALSE
)
```

Arguments

inputFile Input vcf file location (uncompressed or gzip compressed).

threads Number of java threads to use.

ignoreMissing Ignore variants with missing data (./. or .|.)
onlyHets Only calculate on variants with heterozygous calls.
ignoreHets Only calculate on variants with homozygous calls.

Details

Biallelic or multiallelic (maximum 7 alternate alleles) SNP and/or INDEL variants are considered, phased or not. Some VCF encoding examples are:

- heterozygous variants: 1/0 or 0/1 or 0/2 or 1 | 0 or 0 | 1 or 0 | 2
- homozygous to the reference allele variants: 0/0 or 0 | 0
- homozygous to the first alternate allele variants: 1/1 or 1|1

If there are n samples and m variants, an nxn zero-diagonal symmetric distance matrix is calculated. The calculated cosine type distance (1-cosine_similarity)/2 is in the range [0,1] where value 0 means completely identical samples (cosine is 1), value 0.5 means perpendicular samples (cosine is 0) and value 1 means completely opposite samples (cosine is -1).

The calculation is performed by a Java backend implementation, that supports multi-core CPU utilization and can be demanding in terms of memory resources. By default a JVM is launched with a maximum memory allocation of 512 MB. When this amount is not sufficient, the user needs to reserve additional memory resources, before loading the package, by updating the value of the java.parameters option. For example in order to allocate 4GB of RAM, the user needs to issue options(java.parameters="-Xmx4g") before library(fastreeR).

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Value

A character vector of the generated phylogenetic tree in Newick format.

Author(s)

Anestis Gkanogiannis, <anestis@gkanogiannis.com>

References

Java implementation: https://github.com/gkanogiannis/BioInfoJava-Utils

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