## Package 'BASiNETEntropy'

August 16, 2023

```
Title Classification of RNA Sequences using Complex Network and Information Theory
```

**Version** 0.99.6

Description It makes the creation of networks from sequences of RNA, with this is done the abstraction of characteristics of these networks with a methodology of maximum entropy for the purpose of making a classification between the classes of the sequences. There are two data present in the 'BASiNET' package, ``mRNA", and ``ncRNA" with 10 sequences. These sequences were taken from the data set used in the article (LI, Aimin; ZHANG, Junying; ZHOU, Zhongyin, 2014) <doi:10.1186/1471-2105-15-311>, these sequences are used to run examples.

```
License GPL-3
Encoding UTF-8
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```

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entropy theories

## Description

Given three or two distinct data sets, one of mRNA, one of lncRNA and one of sncRNA. The classification of the data is done from the structure of the networks formed by the sequences, that is filtered by an entropy methodology. After this is done, the classification starts.

## Usage

```
classify(
 mRNA,
  lncRNA,
  sncRNA = NULL,
  trainingResult,
  save_dataframe = NULL,
  save_model = NULL,
  predict_with_model = NULL
)
```

## **Arguments**

mRNA	Directory where the file .FASTA lies with the mRNA sequences
lncRNA	Directory where the file .FASTA lies with the lncRNA sequences
sncRNA	Directory where the file .FASTA lies with the sncRNA sequences (optional)
trainingResult	The result of the training, (three or two matrices)
save_dataframe	save when set, this parameter saves a .csv file with the features in the current directory. No file is created by default.

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save\_model save when set, this parameter saves a .rds file with the model in the current directory. No file is created by default.

predict\_with\_model

predict the input sequences with the previously generated model.

#### Value

Results

#### Author(s)

Murilo Montanini Breve

## **Examples**

```
library(BASiNETEntropy)
arqSeqMRNA <- system.file("extdata", "mRNA.fasta",package = "BASiNETEntropy")
arqSeqLNCRNA <- system.file("extdata", "ncRNA.fasta", package = "BASiNETEntropy")
load(system.file("extdata", "trainingResult.RData", package = "BASiNETEntropy"))
r_classify <- classify(mRNA=arqSeqMRNA, lncRNA=arqSeqLNCRNA, trainingResult = trainingResult)</pre>
```

createedges

Creates an untargeted graph from a biological sequence

#### Description

A function that from a biological sequence generates a graph not addressed having as words vertices, this being able to have its size parameter set by the' word 'parameter. The connections between words depend of the' step 'parameter that indicates the next connection to be formed

## Usage

```
createedges(sequence, word = 3, step = 1)
```

## **Arguments**

sequence It is a vector that represents the sequence

word This integer parameter decides the size of the word that will be formed

step It is the integer parameter that decides the step that will be taken to make a new

connection

#### Value

Returns the array used to creates the edge list

#### Author(s)

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 ${\tt creatingDataframe}$ 

Creates a feature matrix using complex network topological measures

## **Description**

A function that from the complex network topological measures create the feature matrix.

## Usage

```
creatingDataframe(measures, tamM, tamLNC, tamSNC)
```

## Arguments

measures The complex network topological measures

tamM mRNA sequence size
tamLNC lncRNA sequence size
tamSNC snRNA sequence size

#### Value

Returns the feature matrix in scale 0-1

## Author(s)

Murilo Montanini Breve

curveofentropy

Creates an entropy curve

## Description

A function that from the entropy measures and threshold creates an entropy curve.

## Usage

```
curveofentropy(H, threshold)
```

## **Arguments**

H The 'training' return for the entropy measures

threshold The 'training' return for the threshold

#### Value

Returns a entropy curve

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## Author(s)

Murilo Montanini Breve

entropy

Calculates the entropy

## **Description**

A function that calculates the entropy

## Usage

entropy(x)

## **Arguments**

Χ

The probabilities P0 and P1

## Value

Returns the entropy

#### Author(s)

Murilo Montanini Breve

filtering

Filters the edges

## Description

A function that filters the edges after the maximum entropy is obtained

## Usage

```
filtering(edgestoselect, edgestofilter)
```

## Arguments

```
edgestoselect The selected edges edgestofilter The edges used to filter
```

## Value

Returns the filtered edges

## Author(s)

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matrixmultiplication Compares the matrices

## Description

A function that compares the matrices 'trainingResult' and the adjacency matrix to produce a filtered adjacency matrix.

## Usage

```
matrixmultiplication(data, histodata)
```

## Arguments

data Adjacency matrix histodata 'trainingResult' data

## Value

Returns the filtered adjacency matrix

## Author(s)

Murilo Montanini Breve

maxentropy

Calculates the maximum entropy

## Description

A function that calculates the maximum entropy

## Usage

```
maxentropy(histogram)
```

## Arguments

histogram

The histogram (used in 'training' function)

## Value

Returns the maximum entropy

## Author(s)

preprocessing 7

preprocessing	Rescales the results between values from 0 to 1
pi chi occastilg	Rescutes the results between values from 0 to 1

#### **Description**

Given the results the data is rescaled for values between 0 and 1, so that the length of the sequences does not influence the results. The rescaling of the sequences are made separately

## Usage

```
preprocessing(datah, tamM, tamLNC, tamSNC)
```

#### **Arguments**

datah Array with results numerics

tamM Integer number of mRNA sequences
tamLNC Integer number of lncRNA sequences
tamSNC Integer number of sncRNA sequences

#### Value

Returns the array with the rescaled values

## Author(s)

Murilo Montanini Breve

selectingEdges Selects the edges of the adjacency matrix

## **Description**

A function that selects the edges of the adjacency matrix

## Usage

```
selectingEdges(MAX, data)
```

## **Arguments**

MAX The maximum entropy data The adjacency matrix

#### Value

Returns the selected edges of the adjacency matrix

8 training

## Author(s)

Murilo Montanini Breve

training	Trains the algorithm to select the edges that maximize the entropy

## Description

A function that trains the algorithm to select the edges that maximize the entropy

## Usage

```
training(mRNA, lncRNA, sncRNA = NULL)
```

## Arguments

mRNA Directory where the file .FASTA lies with the mRNA sequences

1ncRNA Directory where the file .FASTA lies with the lncRNA sequences

sncRNA Directory where the file .FASTA lies with the sncRNA sequences (optional)

#### Value

Returns the edge lists and the 'curveofentropy' function inputs

## Author(s)

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