Package 'rSWeeP'

May 15, 2024

Title Functions to creation of low dimensional comparative matrices of Amino Acid Sequence occurrences

Version 1.16.0

- **Description** The SWeeP method was developed to favor the analizes between amino acids sequences and to assist alignment free phylogenetic studies. This method is based on the concept of sparse words, which is applied in the scan of biological sequences and its the conversion in a matrix of ocurrences. Aiming the generation of low dimensional matrices of Amino Acid Sequence occurrences.
- **biocViews** Software, StatisticalMethod, SupportVectorMachine, Technology, Sequencing, Genetics, Alignment

Depends R (>= 4.0)

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Encoding UTF-8

LazyData true

RoxygenNote 7.0.2

Imports pracma, stats

Suggests Biostrings, methods, knitr, rmarkdown, BiocStyle

VignetteBuilder knitr

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orthBase

Generate a orthonormal matrix (lin, col)

Description

Generate a orthonormal matrix in a specified size, lin by col.

Usage

orthBase(lin, col)

Arguments

lin	Number of rows in the desired matrix
col	Number of columns in the desired matrix

Value

A orthonormal matrix in a specified size, lin by col.

Author(s)

Danrley R. Fernandes.

See Also

sWeeP, orth

Examples

orthBase(160000, 10)

lin <- 160000
col <- 10
orthBase(lin = lin, col = col)</pre>

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sWeeP

Description

The "Spaced Words Projection (SWeeP)" is a method for representing biological sequences using compact vectors. SWeeP uses the spacedwords concept by scanning sequences and generating indices to create a higherdimensional matrix of occurrences that is later projected into a smaller randomly oriented orthonormal base (PIERRI, 2019). This way the resulting matrix will conserve the comparational data but will have a selectable size

Usage

```
sWeeP(xfas, baseMatrix)
## S4 method for signature 'character'
sWeeP(xfas, baseMatrix)
## S4 method for signature 'AAStringSet'
sWeeP(xfas, baseMatrix)
```

Arguments

xfas	A AAStringSet or a FASTA format file
baseMatrix	A orthonormal matrix with 160.000 coordinates

Details

The SWeeP method was developed to favor the comparison between complete proteomic sequences and to assist in machine learning analyzes. This method is based on the concept of spaced words, which are used to scan biological sequences and project them into matrix of occurrences, favoring the manipulation of the data. The sWeeP function can project a matrix n by m, where n is the number of sequences in the analized xfas and m is the number of columns in baseMatrix matrix.

Value

A matrix resulted by the projection of the sequences in xfas in the baseMatrix matrix

Author(s)

Danrley R. Fernandes.

References

Pierri, C. R. et al. SWeeP: Representing large biological sequences data sets in compact vectors. Scientific Reports, accepted in December 2019.doi: 10.1038/s41598-019-55627-4.

Examples

```
baseMatrix <- orthBase(160000,10)
path <- system.file(package = "rSWeeP", "extdata", "exdna.fas")
return <- sWeeP(path,baseMatrix)
distancia <- dist(return, method = "euclidean")
tree <- hclust(distancia, method="ward.D")
plot(tree, hang = -1, cex = 1)</pre>
```

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