Package 'odseq'

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Type Package

Title Outlier detection in multiple sequence alignments

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Description Performs outlier detection using bootstrap of distance metrics of the multiple alignment.
License MIT + file LICENSE
LazyData True
Encoding UTF-8
biocViews Alignment, MultipleSequenceAlignment
VignetteBuilder knitr
Depends R (>= 3.2.3), msa (>= 1.2.1), kebabs (>= 1.4.1), mclust (>= 5.1), knitr(>= 1.11)
NeedsCompilation no
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odseq-package	Outlier detection in multiple sequence alignments

Description

Performs outlier detection using bootstrap of distance metrics of the multiple alignment.

Details

The DESCRIPTION file:

Package: odseq Type: Package

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Version: 0.99.2 Date: 2015-12-20 Author: José Jiménez

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odseq Outlier detection in a multiple sequence

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odseq_unaligned Outlier detection provided a

distance/similarity matrix of sequences.

seqs PFAM plus random data.

Author(s)

José Jiménez

Maintainer: José Jiménez <jose@jimenezluna.com>

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References

[1] OD-seq: outlier detection in multiple sequence alignments. *Peter Jehl, Fabian Sievers and Desmond G. Higgins*. BMC Bioinformatics. 2015.

See Also

```
odseq odseq_unaligned
```

Examples

```
data(seqs)
al <- msa(seqs)
odseq(al, distance_metric = "affine", B = 1000, threshold = 0.025)</pre>
```

odmix

Gaussian mixture modelling of distances in a multiple sequence alignment.

Description

This function performs clustering of biological sequences via fitting a Gaussian mixture model of the distances defined by the odseq algorithm

Usage

```
odmix(msa_object, distance_metric, groups)
```

Arguments

msa_object

An object of formal class ${\tt MsaAAMultipleAlignment},$ as provided by the ${\tt msa}$

package.

distance_metric

A string indicating the type of distance metric to be computed. Either 'linear'

and 'affine' is supported at the moment.

groups

Number of groups to fit in the mixture model. If a numeric vector of size n, n models will be fitted and a list of BIC values will be given to choose a single

model.

Value

A list containing the following items:

prob A numeric matrix of size n x groups where the probability of belonging to a

group is provided for each sequence.

class The class assigned according to prob. Returns a numeric vector.

BIC values for the models proposed in groups

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Author(s)

José Jiménez <jose@jimenezluna.com>

See Also

```
odseq_unaligned odseq
```

Examples

```
data(seqs)
al <- msa(seqs)
odmix(al, distance_metric = "affine", groups = 2)</pre>
```

odseq

Outlier detection in a multiple sequence alignment

Description

This function will first compute a distance metric among every sequence in the multiple alignment. Then it will bootstrap an average score of these distance to provide information on the distribution of scores, which is used to distinguish outlier sequences with a certain threshold

Usage

```
odseq(msa_object, distance_metric = "linear", B = 100, threshold = 0.025)
```

Arguments

msa_object An object of formal class MsaAAMultipleAlignment, as provided by the msa

package.

distance_metric

A string indicating the type of distance metric to be computed. Either 'linear'

and 'affine' is supported at the moment.

B Integer indicating the number of bootstrap replicates to be run. The higher the

more robust the detection should be.

threshold Float indicating the probability to be left at the right of the bootstrap scores

distribution when computing outliers. This parameter may need some tuning

depending on each specific problem

Value

Returns a logical vector, where TRUE indicates an outlier.

Author(s)

José Jiménez <jose@jimenezluna.com>

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References

[1] OD-seq: outlier detection in multiple sequence alignments. *Peter Jehl, Fabian Sievers and Desmond G. Higgins*. BMC Bioinformatics. 2015.

See Also

```
odseq_unaligned
```

Examples

```
data(seqs)
al <- msa(seqs)
odseq(al, distance_metric = "affine", B = 1000, threshold = 0.025)</pre>
```

odseq_unaligned

Outlier detection provided a distance/similarity matrix of sequences.

Description

Provided a similarity matrix (like the ones provided using string kernels in **kebabs**). It will then compute a score for each sequence and perform bootstrap to provide information on the distribution of the scores, which is used to distinguish outlier sequences.

Usage

```
odseq_unaligned(distance_matrix, B = 100, threshold = 0.025, type = "similarity")
```

Arguments

distance matrix

A numeric matrix representing either similarity or distance among unaligned

sequences. Package kebabs may be useful for this task.

B Integer indicating the number of bootstrap replicates to be run. The higher the

more robust the detection should be.

threshold Float indicating the probability to be left at the right of the bootstrap scores

distribution when computing outliers. This parameter may need some tuning

depending on each specific problem

type A string indicating the type of distance metric used. Either 'similarity' or

'distance'.

Value

Returns a logical vector, where TRUE indicates an outlier.

Author(s)

José Jiménez <jose@jimenezluna.com>

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References

[1] OD-seq: outlier detection in multiple sequence alignments. *Peter Jehl, Fabian Sievers and Desmond G. Higgins*. BMC Bioinformatics. 2015.

See Also

odseq

Examples

```
data(seqs)
sp <- spectrumKernel(k = 3)
mat <- getKernelMatrix(sp, seqs)
odseq_unaligned(mat, B = 1000, threshold = 0.025, type = "similarity")</pre>
```

seqs

PFAM plus random data.

Description

Sequences from a certain PFAM family plus 100 random sequences.

Usage

```
data("seqs")
```

Value

An object of class AAStringSet.

Examples

```
data(seqs)
```

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