Package 'nuCpos'

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Title An R package for prediction of nucleosome positions

Version 1.18.0

Description nuCpos, a derivative of NuPoP, is an R package for prediction of nucleosome positions. nuCpos calculates local and whole nucleosomal histone binding affinity (HBA) scores for a given 147-bp sequence. Note: This package was designed to demonstrate the use of chemical maps in prediction. As the parental package NuPoP now provides chemical-map-based prediction, the function for dHMM-based prediction was removed from this package. nuCpos continues to provide functions for HBA calculation.

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Depends R (>= 4.2.0) Imports graphics, methods Suggests NuPoP, Biostrings, testthat biocViews Genetics, Epigenetics, NucleosomePositioning Date 2023-02-16 NeedsCompilation yes git_url https://git.bioconductor.org/packages/nuCpos git_branch RELEASE_3_17 git_last_commit_3aa150e git_last_commit_date 2023-04-25 Date/Publication 2023-10-15 Maintainer Hiroaki Kato <hkato@med.shimane-u.ac.jp>

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nuCpos-package

Description

nuCpos, a derivative of **NuPoP**, is an R package for prediction of nucleosome positions. **nuCpos** calculates local and whole nucleosomal histone binding affinity (HBA) scores for a given 147-bp sequence. Note: This package was designed to demonstrate the use of chemical maps in prediction. As the parental package **NuPoP** now provides chemical-map-based prediction, the function for dHMM-based prediction was removed from this package. Please refer to Xi et al. (2010) and Wang et al. (2008) for technical details of **NuPoP**. nuCpos continues to provide functions for HBA calculation. The models are based on chemical maps of nucleosomes from budding yeast (Brogaard et al. (2012)), fission yeast (Moyle-Heyrman et al. (2012)), or mouse embryonic stem cells (Voong et al. (2016)).

Details

Package:	nuCpos
Type:	Package
Version:	1.17.4
Date:	2023-02-16
License:	GPL-2

HBA: R function for calculation of the histone binding affinity score of a whole nucleosome.

localHBA: R function for calculation of the local histone binding affinity.

Author(s)

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References

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Examples

```
load(system.file("extdata","inseq.RData",package="nuCpos"))
HBA(inseq, species = "sc")
localHBA(inseq, species = "sc")
```

R function for calculating the histone binding affinity score of a given
147-bp sequence.

Description

HBA

This function calculates the histone binding score for a given 147-bp sequence. Nucleosomal and linker models built upon the chemical maps are used for the calculation.

Usage

HBA(inseq, species = "mm", silent = FALSE)

Arguments

inseq	a character or DNAString object. The length of the character string must be 147 bp.
species	a character = mm, sc or sp; "mm" for mouse, "sc" for <i>S. cerevisiae</i> and "sp" for <i>S. pombe</i> .
silent	a logical value indicating whether messages are printed in the console.

Value

HBA outputs one numeric value: histone binding affinity for a whole nucleosome.

Examples

```
load(system.file("extdata","inseq.RData",package="nuCpos"))
HBA(inseq, species = "sc")
```

localHBA

R function for calculating the local histone binding score of a given 147-bp sequence.

Description

This function calculates local histone binding scores for 13 nucleosomal subsegments. Nucleosomal and linker models built upon the chemical maps are used for the calculation.

Usage

```
localHBA(inseq, species = "mm", silent = FALSE)
```

Arguments

inseq	a character or DNAString object. The length of the character string must be 147 bp.
species	a character = mm, sc or sp; "mm" for mouse, "sc" for <i>S. cerevisiae</i> and "sp" for <i>S. pombe</i> .
silent	a logical value indicating whether messages are printed in the console.

Value

localHBA outputs a numeric vector of length 13: local histone binding affinity scores for specific regions in a nucleosome.

Examples

```
load(system.file("extdata","inseq.RData",package="nuCpos"))
localHBA(inseq, species = "sc")
```

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