

Package ‘DNAcycP2’

April 2, 2025

Title DNA Cyclizability Prediction

Version 0.99.7

Description This package performs prediction of intrinsic cyclizability of every 50-bp subsequence in a DNA sequence. The input could be a file either in FASTA or text format. The output will be the C-score, the estimated intrinsic cyclizability score for each 50 bp sequences in each entry of the sequence set.

Depends R (>= 4.4.0)

Imports basilisk, reticulate

License Artistic-2.0

Encoding UTF-8

Roxygen list(markdown = TRUE)

RoxygenNote 7.3.2

StagedInstall yes

BinaryFiles inst/extdata/irlstm/saved_model.pb,
inst/extdata/irlstm_smooth/saved_model.pb

biocViews NeuralNetwork, StructuralPrediction

VignetteBuilder knitr

Suggests knitr, rmarkdown, BiocGenerics, RUnit, tinytest, BiocStyle,
Biostrings

LazyData false

URL <https://github.com/jipingw/DNAcycP2>

BugReports <https://github.com/jipingw/DNAcycP2>

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cycle_fasta	<i>Predict Cyclizability</i>
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Description

This predicts cyclizability for all subsequences of length 50bp from a .fasta input file.

Usage

```
cycle_fasta(  
  file_path,  
  smooth,  
  n_cores = 1,  
  chunk_length = 1e+05,  
  save_path_prefix = ""  
)
```

Arguments

<code>file_path</code>	.fasta input file path
<code>smooth</code>	Whether to predict smoothed C0 (DNAcycP2) or original C0 (DNAcycP)
<code>n_cores</code>	Number of cores to use for parallel processing (default=1)
<code>chunk_length</code>	Length of sequence that each core will predict on at a given time. (default=100000)
<code>save_path_prefix</code>	Base path for output files. If it is an empty string, the output files will not be saved (default="")

Details

Optionally, saves output files (use argument 'save_path_prefix')

Value

A list of predictions for each ID in the .fasta file.

Each list item has the following columns: position, c_score_norm (predictions on a normalized scale), and c_score_unnorm (predictions on an unnormalized scale).

Each list item is named "cycle_\${id}" corresponding to the fasta id

Examples

```
# Create a temporary file  
temp_file <- tempfile(fileext = ".fasta")  
writeLines(">1", temp_file)  
writeLines("ACTGCTAGTCACTGCTAGTCACTGCTAGTCACTGCTAGTCACTGCTAGTC", temp_file)  
  
# Example usage of cycle_fasta
```

```
cycle_fasta(temp_file, smooth=TRUE)  
  
# Cleanup  
unlink(temp_file)
```

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