# Package 'DNAmotif'

# September 27, 2024

Type Package

Version 0.1.1

**Title** DNA Sequence Motifs

<b>Description</b> Motifs within biological sequences show a significant role. This package utilizes a user-defined threshold value (window size and similarity) to create consensus segments or motifs through local alignment of dynamic programming with gap and it calculates the frequency of each identified motif, offering a detailed view of their prevalence within the dataset. It allows for thorough exploration and understanding of sequence patterns and their biological importance.
License GPL-3
Encoding UTF-8
Imports stats, Biostrings, Rcpp
LinkingTo Rcpp
NeedsCompilation yes
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Generation of motifs from DNA sequences

## Description

Using a fasta file as input, the motifs and its corresponding frequencies are generated, considering threshold values (window size and similarity), by making consensus segments via local alignment with gap.

### Usage

```
DNAmotifs(fasta_file, ws, cut_off)
```

#### **Arguments**

fasta\_file Sequence file path (.fasta format)

ws Window size

cut\_off Minimum similarity percentage between the motifs for generating a consensus

motif

#### Value

final\_results A dataframe of motifs and their corresponding frequncies

### **Examples**

```
example_fasta = system.file("exdata/sample.fasta", package = "DNAmotif")
DNAmotifs(fasta_file = example_fasta, ws = 15, cut_off = 0.75)
```

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