

Read me

Command line

```
java -heap -jar antmot.jar names pdb_path graph_build dist
min_size min_intra_freq max_extra_freq minimality arff_path
motif_path
```

Example

```
java -Xmx1024m -jar antmot.jar test.txt ./pdb 1 4 3 0.5 1
false test.arff motifs_test.txt
```

Parameters

Input

heap : java heap memory for example Xmx1024m

names : file containing pdb names in the format illustrated by figure 1

pdb_path : folder where the pdb files are saved

graph_build : method of graph building, values={1, 2}, 1 for *AllAtoms* method and 2 for *CA* method.

dist : distance used by the method of graph building (graph_build)

min_size : minimum number of vertices in motifs

min_intra_freq : minimum frequency of a motif within a given class

max_extra_freq : maximum frequency of a motif in an outer class

minimality : boolean parameter to stop building the motif if it satisfies min_intra_freq and max_extra_freq. This parameter is not yet implemented in our program, so the current and default value is false.

Output

arff_path : arff file used to perform classification with weka workbench.

motif_path: file where motifs are saved (figure 2)

```
#Family_1
1BLX.pdb
1BYG.pdb
1CKI.pdb
1CM8.pdb
1FGK.pdb
#Family_2
1AXI.pdb
1B7Y.pdb
1BH6.pdb
1C7K.pdb
1C9B.pdb
1CF5.pdb
```

Figure1. File containing the names of concerned pdb files belonging to two families

```
t#Motif 321
v0 I
v1 R
v2 L
e 0 1 0
e 0 2 1
t#Motif 322
v0 L
v1 V
v2 K
v3 T
e 0 1 0
e 0 2 1
e 0 3 2
t#Motif 323
```

Figure2. Sample of an output file containing spatial motifs.