Package 'PdPDB'

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Title Pattern Discovery in PDB Structures of Metalloproteins
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Description Looks for amino acid and/or nucleotide patterns and/or small ligands coordinated to a given prosthetic centre. Files have to be in the local file system and contain proper extension.
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PdPDB

Pattern Discovery in PDB Structures of Metalloproteins

Description

Looks for amino acid and/or nucleotide patterns coordinated to a given prosthetic centre. It also accounts for small molecule ligands. Patterns are aligned, clustered and translated to logo-like sequences to infer coordination motifs.

Usage

PdPDB(path, metal, n, perc, interactive, dropsReplicate)

Arguments

path	A string containing the path to the PDB directory.
metal	A string containing the PDB chemical symbol of the target prosthetic centre; e.g. SF4 for [4Fe-4S] cluster, ZN for zinc. The PDB chemical symbol is case sensitive for macOS.
n	A numerical value that contains the number or residue in following/preceding n positions from the ligated amino acid or nucleotide; if n=1 PdPDB searches for $x(L)x$ motif-like chains, if n=2 for $xx(L)xx$. (L)igand.
perc	A numerical value about the minimum percent of letters in a column otherwise residues are dropped.
interactive	A numerical value. 0 interactive, 1 automated (will not cut dendrogram), 2 user decided cut. In mode 1 and 2 ExPASy amino acid frequencies are used as reference.
dropsReplicate	A numerical value. 0 keeps replicated patterns, 1 drops replicated patterns entry by entry, 2 keeps only unique patterns.

Value

PdPDB generates a list of ".csv" and ".svg" files that will be stored in the same folder of the analyzed pdb/cif files (see "path"), its output is as follows:

frequency.csv	PDB-like patterns (i.e. with PDB chem Ids). "-" and "+" are used for residues out of the n inspecting window or from different monomers, respectively. Patterns come along with their frequency.			
alignment.csv	Ligand-aligned patterns with dashes, plus signs and gaps ("*"). See 'frequency.csv			
following_X_enrichment.csv				
	n files. Each file contains enrichment score, z-score and statistics at up to n following positions. X is the +position from ligated residue.			
ligands_enrichment.csv				
	Enrichment scores and statistics for ligands.			
notLigands_enrichment.csv				
	Enrichment statistics for the whole specimen but ligands.			
<pre>preceeding_X_enrichment.csv</pre>				
	As for "following" but this is meant for residues preceeding ligands. See "fol- lowing_X_enrichment.csv."			
root_enrichment.csv				
	Overall enrichment score.			
logo_Y.csv	Y files. Each file contains the logo and consensus sequence for a cluster. Y is the cluster number.			
dendrogram.svg	The dendrogram along with the user deciced cutoff and clusters.			

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following_X_pro	portions.svg		
	Plot of the enrichment score per each amino acid in following positions.		
ligands_proport	ions.svg		
	Plot of the enrichment score per each amino acid in ligated position.		
notLigands_prop	portions.svg		
	Plot of the enrichment score per each amino acid in non ligated position.		
<pre>preceeding_X_pr</pre>	oportions.svg		
	Plot of the enrichment score per each amino acid in preceeding positions.		
root_proportion	is.svg		
	Plot of the root enrichment score.		
logo_Y.svg	Plot of the logo and consensus sequence of the Yth cluster. The complete aligned cluster is given as homonym '.csv' file. Sequences come along with percentages. If the dendrogram is not cut the root logo is given.		
following_X_standardized.svg			
	Plot of the z-score per each amino acid in following positions.		
ligands_standar	dized.svg		
	Plot of the z-score per each amino acid in ligated position.		
notLigands_stan	dardized.svg		
	Plot of the z-score per each amino acid in non ligated position.		
preceeding_X_standardized.svg			
	Plot of the z-score per each amino acid in preceeding positions.		
<pre>root_standardiz</pre>	red.svg		
	Plot of the root z-score.		
patterns.csv	PDB like extracted patterns along with the PDB ID and metal IDs. Useful for debbugging. Needed for restore.		
PdPDB.log	PdPDB log file. Useful for debbugging. Needed for restore.		

Note

Files have to be in the local file system and contain the ".pdb" or ".cif" extension. Output files use brackets to highlight ligands and/or 'L' in heading line.

Author(s)

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References

Belmonte L, Mansy SS Patterns of Ligands Coordinated to Metallocofactors Extracted from the Protein Data Bank, Journal of Chemical Information and Modeling (accepted)

Examples

############### Research Parameters

```
metal="SF4" # searches for [4fe-4s] coordinating patterns
n=1 # searches for x(L)x patterns, (L) coordinates to SF4
perc=20 # drops residues with less than the 20% of frequency
interactive= 0 # interactive. User decided references and dendrogram cut
dropsReplicate=0 # do not remove replicated patterns
```

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