

Package ‘fibos’

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Title Occlusion Surface Using the Occluded Surface and Fibonacci
Occluded Surface

Version 2.0.1

Maintainer Herson Soares <hersonhebert@hotmail.com>

Description The Occluded Surface (OS) algorithm is a widely used approach for analyzing atomic packing in biomolecules as described by Pattabiraman N, Ward KB, Fleming PJ (1995) <[doi:10.1002/jmr.300080603](https://doi.org/10.1002/jmr.300080603)>. Here, we introduce 'fibos', an 'R' and 'Python' package that extends the 'OS' methodology, as presented in Soares HHM, Romanelli JPR, Fleming PJ, da Silveira CH (2024) <[doi:10.1101/2024.11.01.621530](https://doi.org/10.1101/2024.11.01.621530)>.

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Author Herson Soares [cre, aut],
Carlos Silveira [aut],
João Romanelli [aut],
Patrick Fleming [aut],
Posit Software, PBC [cph, fnd]

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`fibos_config`*Install the 'Python' 'fibos' Module*

Description

This function creates a 'Python' virtual environment and installs the 'Python' module 'fibos' required for the full functionality of this package. It handles different system configurations and ensures that the correct compiler paths are set.

Usage

```
fibos_config()
```

Note

This function will install external software (a 'Python' package) on your system. Administrator/sudo privileges might be required on some systems.

See Also

[occluded_surface\(\)](#)

[osp\(\)](#)

Examples

```
# Set up the 'Python' environment and install the required module
fibos_config()
```

`get_radii`*Load Radii Values*

Description

The 'get_radii' function is responsible for loading the atomic radii values used for surface-occlusion calculations. The values it returns are those currently employed in those calculations.

Usage

```
get_radii()
```

Value

A data frame containing the radii values.

Author(s)

Carlos Henrique da Silveira (carlos.silveira@unifei.edu.br)
Herson Hebert Mendes Soares (hersonhebert@hotmail.com)
Joao Paulo Roquim Romanelli (joaoromanelli@unifei.edu.br)
Patrick Fleming (Pat.Fleming@jhu.edu)

See Also

[set_radii\(\)](#)
[reset_radii\(\)](#)

Examples

```
library(fibos)
fibos_config()
#Loads the radii values that have been configured for code execution.
radii = get_radii()
#Displays the first three lines.
radii |> utils::head(3) |> print()
```

occluded_surface *Occluded Surface (OS)*

Description

The 'Occluded Surface (OS)' algorithm is a widely used approach for analyzing atomic packing in biomolecules. Here, we introduce 'FIBOS', an 'R' and 'Python' package that extends the 'OS' methodology with enhancements. The homonymous function 'occluded_surface' calculates 'OS' per atom.

Usage

```
occluded_surface(pdb, method = "FIBOS", density_dots = 5)
```

Arguments

pdb	4-digit PDB id (will fetch it from the RCSB repository) or the path to a PDB local file.
method	Method to be used: 'OS' (classic) or 'FIBOS'(default).The classic 'OS' covers the surface radially with one of the axes as a reference when allocating the dots. In 'FIBOS', Fibonacci spirals were used to allocate the dots, which is known to produce lower axial anisotropy as well as more evenly spaced points on a sphere.
density_dots	Distribution density of atomic dots for surface occlusion calculation.

Details

'Occluded Surface (OS)' (Pattabiraman et al. 1995) method distributes dots (representing patches of area) across the atom surfaces. Each dot has a normal that extends until it reaches either a van der Waals surface of a neighboring atom (the dot is considered occluded) or covers a distance greater than the diameter of a water molecule (the dot is considered non-occluded and disregarded). Thus, with the summed areas of dots and the lengths of normals, it is possible to compose robust metrics capable of inferring the average packing density of atoms, residues, proteins, as well as any other group of biomolecules.

For more details, see (Fleming et al, 2000) and (Soares, et al, 2024)

Value

A table containing:

ATOM the atomic contacts for each atom.

NUMBER OF POINTS the number of dots (patches of area) on atomic surface.

AREA the summed areas of dots.

RAYLENGTH the average lengths of normals normalized by 2.8 Å (water diameter). So, raylen is a value between 0 and 1. A raylen close to 1 indicates worse packaging.

DISTANCE the average distances of contacts in (Å).

Author(s)

Herson Soares, Joao Romanelli, Patrick Fleming, Carlos Silveira.

References

Fleming PJ, Richards FM (2000). "Protein packing: Dependence on protein size, secondary structure and amino acid composition." doi:10.1006/jmbi.2000.3750

Pattabiraman N, Ward KB, Fleming PJ (1995). "Occluded molecular surface: Analysis of protein packing." doi:10.1002/jmr.300080603

Soares HHM, Romanelli JPR, Fleming PJ, da Silveira CH (2024). "bioRxiv, 2024.11.01.621530." doi:10.1101/2024.11.01.621530

See Also

[osp\(\)](#)

Examples

```
library(fibos)

#Configure the environment
fibos_config()

# Calculate FIBOS per atom and create .srf files in fibos_files folder
pdb_fibos <- occluded_surface("1ptx", method = "FIBOS", density_dots = 5.0)
```

```
# Calculate OSP metric per residue from .srf file in fibos_files folder
pdb_osp <- osp(fs::path("fibos_files", "prot_1ptx.srf"))
```

osp	<i>Ocluded Surface Packing (OSP)</i>
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Description

Implements the 'occluded surface' packing density metric (OSP) averaged by residue, as described in (Fleming and Richards 2000).

Usage

```
osp(file)
```

Arguments

file a SRF File (.srf) generated by 'occluded_surface' in fibos_files folder.

Value

A table containing:

Resnum residue id.

Resname residue name.

OS the summed areas of dots in residue.

'os*[1-raylen]' 'OS' areas weighted by (1-raylen). Raylen is the average lengths of normals normalized by 2.8 Å (water diameter). So, raylen is a value between 0 and 1. A raylen close to 1 indicates worse packaging, and the 'OS' will be reduced.

OSP average occluded surface packing value (OSP) by residue.

Author(s)

Herson Soares

Joao Romanelli

Patrick Fleming

Carlos Silveira.

References

Fleming PJ, Richards FM (2000). "Protein packing: Dependence on protein size, secondary structure and amino acid composition." doi:10.1006/jmbi.2000.3750

Pattabiraman N, Ward KB, Fleming PJ (1995). "Occluded molecular surface: Analysis of protein packing." doi:10.1002/jmr.300080603

Soares HHM, Romanelli JPR, Fleming PJ, da Silveira CH (2024). "bioRxiv, 2024.11.01.621530." doi:10.1101/2024.11.01.621530

See Also

[occluded_surface\(\)](#)

Examples

```
library(fibos)

#Configure the Environment
fibos_config()

# Calculate FIBOS per atom and create .srf files in fibos_files folder
pdb_fibos <- occluded_surface("1ptx", method = "FIBOS", density_dots = 5.0)
# Calculate OSP metric per residue from .srf file in fibos_files folder
pdb_osp <- osp(fs::path("fibos_files", "prot_1ptx.srf"))
```

reset_radii

Reset Radii Values

Description

This function reloads the 'OS' default atomic radii values.

Usage

```
reset_radii()
```

Author(s)

Carlos Henrique da Silveira (carlos.silveira@unifei.edu.br)
Herson Hebert Mendes Soares (hersonhebert@hotmail.com)
Joao Paulo Roquim Romanelli (joaoromanelli@unifei.edu.br)
Patrick Fleming (Pat.Fleming@jhu.edu)

See Also

[get_radii\(\)](#)

[set_radii\(\)](#)

Examples

```
library(fibos)
fibos_config()
#Loads the radius values that have been configured for code execution.
radii = get_radii()
#Displays the first three lines.
radii |> utils::head(3) |> print()
```

```
#Modifies the value of a specific radius.
radii$RAY[1] = 2.15
#Sets the radius value from a tibble.
set_radii(radii)
#Displays the first three lines.
get_radii() |> utils::head(3) |> print()
#Loads the default radius values.
reset_radii()
#Displays the first three lines.
get_radii() |> utils::head(3) |> print()
```

set_radii

Change Radii Values

Description

This function enables modification of the radius values by passing a 'data.frame' as an argument.

Usage

```
set_radii(radii_values)
```

Arguments

radii_values A 'data.frame' containing atomic radii values.

Author(s)

Carlos Henrique da Silveira (carlos.silveira@unifei.edu.br)

Herson Hebert Mendes Soares (hersonhebert@hotmail.com)

Joao Paulo Roquim Romanelli (joaoromanelli@unifei.edu.br)

Patrick Fleming (Pat.Fleming@jhu.edu)

See Also

[get_radii\(\)](#)

[reset_radii\(\)](#)

Examples

```
library(fibos)
fibos_config()
#Loads the radius values that have been configured for code execution.
radii = get_radii()
#Displays the first three lines.
radii |> utils::head(3) |> print()
#Modifies the value of a specific radius.
```

```
radii$RAY[1] = 2.15
#Sets the radius value from a tibble.
set_radii(radii)
#Displays the first three lines.
get_radii() |> utils::head(3) |> print()
```

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