# **pyvol Documentation**

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PyVOL is a python library packaged into a *PyMOL* GUI for identifying protein binding pockets, partitioning them into sub-pockets, and calculating their volumes. PyVOL can be run as a PyMOL plugin through its GUI or the PyMOL prompt, as an imported python library, or as a commandline program. Visualization of results is exclusively supported through PyMOL though exported surfaces are compatible with standard 3D geometry visualization programs.

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**ONE** 

# **INSTALLATION**

PyVOL can be installed into any python environment; however, for most users direct installation into PyMOL will be easiest.

# 1.1 Installation into PyMOL

PyVOL is distributed as a GUI and a backend. Installation into PyMOL uses PyMOL's plugin manager to install the GUI and then the GUI to install the backend. The GUI is installed through the plugin manager through loading the zipped GUI file:

https://github.com/rhs2132/pyvol/blob/master/pyvolgui.zip

This creates a new PyVOL menu entry under plugins. The third tab of the GUI allows installation of PyVOL from PyPI along with all available dependencies. For more information, especially on Windows, see the installation page.

**TWO** 

# **RUNNING PYVOL**

## 2.1 Quick Start

From within PyMOL, the simplest binding pocket calculation is simply run at the PyMOL prompt with:

```
pocket protein_selection
```

The two parameters that most dramatically affect calculations are the maximum and minimum radii used to respectively define the exterior surface of the protein and the boundary of the binding pocket itself. In practice, the minimum radius does not need to be changed as its default (1.4) is broadly useful. The maximum radius does often need to be adjusted to find a suitable value using the max\_rad parameter:

```
pocket protein_selection, min_rad=1.4, max_rad=3.4
```

# 2.2 Further Examples

For extensive explanations and documentations of the GUI and command line interfaces, see the general usage page.

THREE

### MODULE DOCUMENTATION

For full documentation of the code, see the modules page.

# 3.1 Installation

PyVOL distribution is hosted by PyPI and accessed through pip. PyVOL can consequently be installed into any python environment. For convenience, the PyMOL GUI contains an installer.

# 3.1.1 Detailed Installation into PyMOL

PyVOL is distributed as a GUI and a backend. Installation into PyMOL uses PyMOL's plugin manager to install the GUI and then the GUI to install the backend. The GUI is installed through the plugin manager through loading the zipped GUI file:

```
https://github.com/rhs2132/pyvol/blob/master/pyvolgui.zip
```

This creates a new PyVOL menu entry under plugins. The third tab of the GUI allows installation of PyVOL from PyPI along with all available dependencies. On Linux and MacOS, MSMS is automatically installed from the platform-limited bioconda channel. MSMS installation instructions are otherwise below.

#### 3.1.2 Manual Installation

PyVOL minimally requires biopython, MSMS, numpy, pandas, scipy, scikit-learn, and trimesh in order to run. PyVOL is available for manual installation from github or through PyPI. Most conveniently:

```
pip install bio-pyvol
```

#### 3.1.3 MSMS Installation

MSMS can be installed on MacOS and Linux using the bioconda channel:

```
conda install -c bioconda msms
```

Otherwise MSMS must be installed manually by downloading it from MGLTools and adding it to the path. PyMOL distributions from Schrodinger have MSMS included; however, it must still be added to the path manually. The executable is located at:

<pymol\_root\_dir>/pkgs/msms-2.6.1-2/bin/msms

### 3.1.4 Updating

PyVOL can be updated via the command line:

```
pip update bio-pyvol
```

If using the PyMOL GUI, the third tab has a button labeled Check for Updates that will query PyPI to detect whether an update is available. If one is available, that button changes to Update PyVOL and permits updating with a single click.

#### 3.1.5 Uninstallation

PyVOL can be uninstalled via the command line:

```
pip uninstall bio-pyvol
```

If using the PyMOL GUI, the third tab has a button labeled Uninstall PyVOL that will remove the PyVOL backend. Afterwards, selecting uninstall on the plugin within the PyMOL plugin manager will the GUI.

# 3.2 General Usage

The GUI and shell command line prompts recapitulate the PyMOL prompt interface with few exceptions. General usage of PyVOL is shown here with specific examples and modifications for the other interfaces in the following pages. Programmatic invocation of internal functions is supported and covered through the module documentation.

### 3.2.1 Pocket Specification

PyVOL by default recognizes the largest pocket and returns the volume and geometry for it. However, manual identification of the pocket of interest is generally preferable. This can be done through specification of a ligand, a residue, or a coordinate. If a specification is given, the mode is changed to specific by default.

#### **Default Behavior:**

```
pocket protein_selection, mode=largest
```

#### **Ligand Specification:**

```
pocket protein_selection, mode=specific, ligand=ligand_selection
pocket protein_selection, ligand=ligand_selection
```

#### **Residue Specification:**

```
pocket protein_selection, mode=specific, resid=A15
pocket protein_selection, resid=A15
pocket protein_selection, mode=specific, residue=residue_selection
pocket protein_selection, residue=residue_selection
```

where the resid is written as <Chain><Residue number> (e.g. B432). If there is only one chain in the selection, the chain ID can be excluded.

#### **Coordinate Specification:**

```
pocket protein_selection, mode=specific, pocket_coordinate="5.0 10.0 15.0"
pocket protein_selection, pocket_coordinate="5.0 10.0 15.0"
```

where the coordinate is provided as three floats separated by spaces and bounded by quotation marks.

#### **Calculation of All Pockets**

Alternatively, PyVOL can return the surfaces and volumes for all pockets above a minimum volume that are identified. By default, this volume cutoff is set at 200 A<sup>3</sup>.

```
pocket protein_selection, mode=all, minimum_volume=200
```

### 3.2.2 Extra Ligand Options

When a ligand is provided, the atoms of the ligand can be used to identify both minimum and maximum extents of the calculated binding pocket.

#### **Ligand Volume Inclusion**

To include the volume of the ligand in the pocket volume (useful for when the ligand extends into bulk solvent), use the lig incl rad parameter:

```
pocket protein_selection, ligand=ligand_selection, lig_incl_rad=0.0
```

where the value of lig\_incl\_rad is added to the Van der Waals radii of each atom in the ligand selection when calculating the exterior surface of the protein.

#### **Ligand-defined Maximum Volume**

The atoms of the ligand can also be used to define a maximum boundary to the calculated pocket by specifying the lig\_excl\_rad parameter:

```
pocket protein_selection, ligand=ligand_selection, lig_excl_rad=2.0
```

where the value of 11lig\_excl\_rad11 is added to the Van der Waals radii of each atom in the ligand selection when calculating the exterior surface of the protein.

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# 3.2.3 Sub-pocket Partitioning

Sub-partitioning is enabled by setting the subdivide parameter to True:

```
pocket protein_selection, subdivide=True
```

Parameters controlling the number of sub-pockets identified generally perform well using defaults; however, they can be easily adjusted as needed. The two most important parameters are the minimum radius of the largest sphere in each sub-pocket (this excludes small sub-pockets) and the maximum number of clusters:

```
pocket protein_selection, subdivide=True, min_subpocket_rad=1.7, max_clusters=10
```

If the number of clusters must be reduced, sub-pockets are merged on the basis of connectivity between the defining sets of tangent spheres. Practically, sub-pockets with a greater surface area boundary are merged first.

# 3.2.4 Display and Output Options

By default, PyVOL simply outputs a log containing volumes and, when invoked through PyMOL, displays pocket boundaries as semi-translucent surfaces. This behavior can be extensively customized.

The output name for all computed PyMOL objects and the base filename for any output files can be specified using the prefix option:

```
pocket protein_selection, prefix=favprot
```

PyVOL can also write the input and output files to a directory if given an output directory. In this case it writes out the input protein and ligand structures, a csv report of all calcuated volumes, and paired csv/obj files containing tangent sphere collections and 3D triangulated mesh files respectively.

```
pocket protein_selection, output_dir=chosen_out_dir
```

Calculated surfaces can be visualized in three different ways by setting the display\_mode parameter. The following three commands set the output as a solid surface with transparency, a wireframe mesh, and a collection of spheres. Color is set with the color parameter and transparency (when applicable) with the alpha parameter:

```
pocket protein_selection, display_mode=solid, alpha=0.85, color=skyblue
pocket protein_selection, display_mode=mesh, color=red
pocket protein_selection, display_mode=spheres, color=firebrick
```

where alpha is [0, 1.0] and the color is any color defined within PyMOL. The presets should generally be sufficient, but custom colors can be chosen using the commands given on the PyMOL wiki.

# 3.3 GUI Interface

The GUI is divided into three tabs that respectively: 1) run new PyVOL calculations 2) load prior PyVOL calculations from disk and 3) install, update, and uninstall the backend.

#### 3.3.1 Run Tab

test run

#### 3.3.2 Load Tab

test load

#### 3.3.3 Install Tab

test install

# 3.4 Shell Interface

PyVOL can also be run from the system command line using bash or any standard shell. If installed using pip, a pyvol entry point should be automatically installed and made available on the path. Otherwise, manual invocation of pyvol/\_\_main\_\_.py should work.

### 3.4.1 Running from the Shell

From the command-line, PyVOL is run exclusively using a configuration file.

```
python -m pyvol <input_parameters.cfg>
```

### 3.4.2 Template Configuration File Generation

A template configuration file with default values supplied can be generated using:

```
python -m pyvol -t <output_template.cfg>
```

# 3.4.3 Notes on Output

Currently, PyVOL only reports standard log output to stdout when run this way. So if an output directory is not provided, there is no easy way to retrieve the results.

# 3.5 pyvol package

#### 3.5.1 Submodules

### 3.5.2 pyvol.cluster module

```
pyvol.cluster.cluster_within_r (spheres, radius, allow_new=True)

Cluster spheres with the same radius using DBSCAN, modifying input data in situ
```

**Args:** spheres (Spheres): complete set of input spheres radius (float): radius at which clustering is to occur allow new (bool): permit new clusters? (Default value = True)

```
pyvol.cluster.cluster_between_r (spheres, ref_radius, target_radius)
```

Cluster spheres from a target radius to a reference radius, modifying input data in situ

**Args:** spheres (Spheres): complete set of input spheres ref\_radius (float): radius from which cluster identities will be drawn target\_radius (float): radius to which cluster identities will be propagated

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Reassigns improperly clustered spheres to 'proper' clusters, modifying input data in situ

**Args:** spheres (Spheres): complete set of input spheres radius (float): radius at which closest groups are identified min\_cluster\_size (int): minimum number of spheres in a 'proper' cluster (Default value = 1) max clusters (int): maximum number of 'proper' clusters (Default value = None)

pyvol.cluster.extract\_groups (spheres, surf\_radius=None, prefix=None)

Extracts spheres belonging to each cluster from the complete input set and optionally calculates bounded surfaces

**Args:** spheres (Spheres): complete set of input spheres surf\_radius: radius used to calculate bounding spheres for individual groups (Default value = None) prefix: prefix to identify new surfaces (Default value = None)

**Returns:** group\_list ([Spheres]): a list of Spheres objects each corresponding to a different cluster

pyvol.cluster.hierarchically\_cluster\_spheres (spheres,

ordered radii,

min\_new\_radius=None, min\_cluster\_size=10, max\_clusters=None)

Cluster spheres by grouping spheres at large radius and propagating those assignments down to smaller radii

Args: spheres (Spheres): complete set of input spheres ordered\_radii ([float]): list of radii ordered from largest to smallest min\_new\_radius (float): smallest spheres to keep (Default value = None) min\_cluster\_size (int): minimum number of spheres in a cluster (Default value = 10) max\_clusters (int): maximum number of clusters (Default value = None)

pyvol.cluster.identify\_closest\_grouped(spheres, group, radius)

Identifies the closest 'properly' grouped cluster to a specified group

**Args:** spheres (Spheres): complete set of input spheres group (float): group for which to identify the closest clusters radius (float): radius at which to perform the search

**Returns:** group (float): passthrough of input group closest (float): id of the closest cluster magnitude (int): number of pairwise closest connections between the queried group and the closest identified cluster

pyvol.cluster.merge\_sphere\_list(s\_list, r=None, g=None)

**Args:** s\_list ([Spheres]): list of input spheres r (float): radius value to assign to output Spheres (Default value = None) g (float): group value to assign to output Spheres (Default value = None)

**Returns:** merged\_spheres (Spheres): a single Spheres object containing the merged input lists

pyvol.cluster.reassign\_group(spheres, source\_group, target\_group)
Reassign a group in place

**Args:** spheres (Spheres): complete set of input spheres source\_group (float): group to change target\_group (float): new group id

pyvol.cluster.reassign\_groups\_to\_closest (spheres, group\_list, radius, iterations=None, preserve largest=False)

Reassign a group to the closest group as identified by maximum linkage; operates in place

**Args:** spheres (Spheres): complete set of input spheres group\_list ([float]): list of group ids which are to be iteratively reassigned radius (float): radius at which searches are to take place iterations (int): number of times to attempt to reassign groups (Default value = None) preserve\_largest: keep the group id of the group with more members? (Default value = False)

pyvol.cluster.remove\_interior(spheres)

Remove all spheres which are completely enclosed in larger spheres; operates in place

**Args:** spheres (Spheres): complete set of input spheres

pyvol.cluster.remove\_overlap (spheres, radii=None, spacing=0.1, iterations=20, tolerance=0.02)
Remove overlap between groups; operates in place

**Args:** spheres (Spheres): complete set of input spheres radii ([float]): radii at which to perform searches for overlap (Default value = None) spacing (float): binning radius (Default value = 0.1) iterations (int): number of times to attempt overlap removal (Default value = 20) tolerance (float): overlap tolerance (Default value = 0.02)

# 3.5.3 pyvol.identify module

```
pyvol.identify.pocket (prot_file, mode='largest', lig_file=None, coordinate=None, resid=None, residue_coordinates=None, min_rad=1.4, max_rad=3.4, lig_excl_rad=None, lig_incl_rad=None, subdivide=False, minimum_volume=200, min_subpocket_rad=1.7, min_subpocket_surf_rad=1.0, max_clusters=None, prefix=None, output_dir=None, constrain_inputs=False)

Calculates the SES for a binding pocket
```

Args: prot\_file (str): filename for the input pdb file containing the peptidee mode (str): pocket identification mode (can be largest, all, or specific) (Default value = "largest") lig\_file (str): filename for the input pdb file containing a ligand (Default value = None) coordinate ([float]): 3D coordinate used for pocket specification (Default value = None) resid (str): residue identifier for pocket specification (Default value = None) residue coordinates ([float]): 3D coordinate of an atom in a surrounding residue used for pocket specification (Default value = None) min rad (float): radius for SAS calculations (Default value = 1.4) max rad (float): radius used to identify the outer, bulk solvent exposed surface (Default value = 3.4) lig excl rad (float): maximum distance from a provided ligand that can be included in calculated pockets (Default value = None) lig incl rad (float): minimum distance from a provided ligand that should be included in calculated pockets when solvent border is ambiguous (Default value = None) subdivide (bool): calculate subpockets? (Default value = False) minimum\_volume (float): minimum volume of pockets returned when running in 'all' mode (Default value = 200) min\_subpocket\_rad (float): minimum radius that identifies distinct subpockets (Default value = 1.7) min\_subpocket\_surf\_rad (float): radius used to calculate subpocket surfaces (Default value = 1.0) max\_clusters (int): maximum number of clusters (Default value = None) prefix (str): identifying string for output (Default value = None) output\_dir (str): filename of the directory in which to place all output; can be absolute or relative (Default value = None) constrain\_inputs (bool): restrict quantitative input parameters to tested values? (Default value = False)

**Returns:** pockets ([Spheres]): a list of Spheres objects each of which contains the geometric information describing a distinct pocket or subpocket

```
pyvol.identify.subpockets (bounding_spheres, ref_spheres, min_rad, max_rad, min_subpocket_rad=1.7, min_subpocket_surf_rad=1.0, max_subpocket_rad=None, sampling=0.1, inclusion_radius_buffer=1.0, min_cluster_size=50, max_clusters=None, prefix=None)
```

Args: bounding\_spheres (Spheres): a Spheres object containing both the peptide and solvent exposed face external spheres ref\_spheres (Spheres): a Spheres object holding the interior spheres that define the pocket to be subdivided min\_rad (float): radius for original SES calculations (Default value = 1.4) max\_rad (float): radius originally used to identify the outer, bulk solvent exposed surface (Default value = 3.4) min\_subpocket\_rad (float): minimum radius that identifies distinct subpockets (Default value = 1.7) min\_subpocket\_surf\_rad (float): radius used to calculate subpocket surfaces (Default value = 1.0) max\_subpocket\_rad (float): maximum spheres radius used for subpocket clustering (Default value = None) sampling (float): radial sampling frequency for clustering (Default value = 0.1) inclusion\_radius\_buffer (float): defines the inclusion distance for nonextraneous spheres in combination with min\_rad and max\_rad (Default value = 1.0) min\_cluster\_size (int): minimum number of spheres that can constitute a proper clusterw (Default value = 50) max\_clusters (int): maximum number of clusters (Default value = None) prefix (str): identifying string for output (Default value = None)

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**Returns:** pockets ([Spheres]): a list of Spheres objects each of which contains the geometric information describing a distinct subpocket

```
pyvol.identify.write_report (all_pockets, output_dir, prefix)
Write a brief report of calculated volumes to file
```

**Args:** all\_pockets ([Spherese]): a list of Spheres objects each of which contains the complete information about a distinct pocket or subpocket output\_dir (str): output directory, relative or absolute prefix (str): identifying prefix for output files

### 3.5.4 pyvol.pymol interface module

**Args:** spheres\_file (str): filename name (str): internal display name (Default value = None) display\_mode (str): display mode (Default value = "solid") color (str): PyMOL color string (Default value = 'marine') alpha (float): transparency value (Default value = 0.85)

```
pyvol.pymol_interface.pocket(protein, mode=None, ligand=None, pocket_coordinate=None,
                                      residue=None,
                                                       resid=None,
                                                                      prefix=None,
                                                                                      min_rad=1.4,
                                      max rad=3.4,
                                                         lig_excl_rad=None,
                                                                                lig_incl_rad=None,
                                      display mode='solid',
                                                             color='marine',
                                                                                alpha=0.85,
                                                        subdivide=None.
                                      put dir=None,
                                                                             minimum volume=200,
                                      min\ subpocket\ rad=1.7,
                                                                       min\ subpocket\ surf\ rad=1.0,
                                      max_clusters=None, excl_org=False, constrain_inputs=True)
```

Calculates the SAS for a binding pocket and displays it

Args: protein (str): PyMOL selection string for the protein mode (str): pocket identification mode (can be largest, all, or specific) (Default value = None) ligand (str): PyMOL selection string for the ligand (Default value = None) pocket coordinate ([float]): 3D coordinate used for pocket specification (Default value = None) residue (str): PyMOL residue selection string for pocket specification (Default value = None) resid (str): residue identifier for pocket specification (Default value = None) prefix (str): identifying string for output (Default value = None) min\_rad (float): radius for SAS calculations (Default value = 1.4) max\_rad (float): radius used to identify the outer, bulk solvent exposed surface (Default value = 3.4) lig\_excl\_rad (float): maximum distance from a provided ligand that can be included in calculated pockets (Default value = None) lig incl rad (float): minimum distance from a provided ligand that should be included in calculated pockets when solvent border is ambiguous (Default value = None) display\_mode (str): display mode for calculated pockets (Default value = "solid") color (str): PyMOL color string (Default value = 'marine') alpha (float): transparency value (Default value = 0.85) output\_dir (str): filename of the directory in which to place all output; can be absolute or relative (Default value = None) subdivide (bool): calculate subpockets? (Default value = None) minimum\_volume (float): minimum volume of pockets returned when running in 'all' mode (Default value = 200) min subpocket rad (float): minimum radius that identifies distinct subpockets (Default value = 1.7) min subpocket surf rad (float): radius used to calculate subpocket surfaces (Default value = 1.0) max clusters (int): maximum number of clusters (Default value = None) excl\_org (bool): exclude non-peptide atoms from the protein selection? (Default value = False) constrain\_inputs (bool): constrain input quantitative values to tested ranges? (Default value = True)

# 3.5.5 pyvol.pymol\_utilities module

**Args:** color\_list ([str]): list of PyMOL color strings (Default value = None) max\_value (int): max palette index (Default value = 7) min\_value (int): min palette index (Default value = 1)

```
Returns: palette ([str]): list of color definitions
```

pyvol.pymol\_utilities.display\_pseudoatom\_group(spheres, name, color='gray60', palette=None)

Displays a collection of pseudoatoms

**Args:** spheres (Spheres): Spheres object holding pocket geometry name (str): display name color (str): PyMOL color string (Default value = 'gray60') palette ([str]): palette (Default value = None)

pyvol.pymol\_utilities.display\_spheres\_object (spheres, name, state=1, color='marine', alpha=0.7, mode='solid', palette=None)

Loads a mesh object into a cgo list for display in PyMOL

**Args:** spheres (Spheres): Spheres object containing all geometry name (str): display name state (int): model state (Default value = 1) color (str): PyMOL color string (Default value = 'marine') alpha (float): transparency value (Default value = 0.7) mode (str): display mode (Default value = "solid") palette ([str]): palette (Default value = None)

pyvol.pymol\_utilities.mesh\_to\_solid\_CGO (mesh, color='gray60', alpha=1.0)
Creates a solid CGO object for a mesh for display in PyMOL

**Args:** mesh (Trimesh): Trimesh mesh object color (str): PyMOL color string (Default value = 'gray60') alpha (float): transparency value (Default value = 1.0)

Returns: cgobuffer (str): CGO buffer that contains the instruction to load a solid object

pyvol.pymol\_utilities.mesh\_to\_wireframe\_CGO (mesh, color='gray60', alpha=1.0)
Creates a wireframe CGO object for a mesh for display in PyMOL

**Args:** mesh (Trimesh): Trimesh mesh object color (str): PyMOL color string (Default value = 'gray60') alpha (float): transparency value (Default value = 1.0)

**Returns:** cgobuffer (str): CGO buffer that contains the instruction to load a wireframe object

#### 3.5.6 pyvol.spheres module

Args: probe\_radius (float): radius for surface calculations (Default value = 1.4) cavity\_atom (int): id of a single atom which lies on the surface of the interior cavity of interest (Default value = None) coordinate ([float]): 3D coordinate to identify a cavity atom (Default value = None) all\_components (bool): return all pockets? (Default value = False) exclusionary\_radius (float): maximum permissibile distance to the closest identified surface element from the supplied coordinate (Default value = 2.5) largest\_only (bool): return only the largest pocket? (Default value = False) noh (bool): remove waters before surface calculation? (Default value = True) minimum\_volume (int): minimum volume of pockets returned when using 'all\_components' (Default value = 200)

#### identify\_nonextraneous (ref\_spheres, radius)

Calculate the SAS for a given probe radius

Returns all spheres less than radius away from any center in ref\_spheres using cKDTree search built on the non-reference set

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**Args:** ref\_spheres (Spheres): object that defines the pocket of interest radius (float): maximum distance to sphere centers to be considered nonextraneous

Returns: nonextraneous (Spheres): a filtered Spheres object

```
nearest (coordinate, max_radius=None)
```

Returns the index of the sphere closest to a coordinate; if max\_radius is specified, the sphere returned must have a radius <= max\_radius

**Args:** coordinate (float nx3): 3D input coordinate max\_radius (float): maximum permissibile distance to the nearest sphere (Default value = None)

Returns: nearest\_index: index of the closest sphere

#### nearest\_coord\_to\_external (coordinates)

Returns the coordinate of the sphere closest to the supplied coordinates

Args: coordinates (float nx3): set of coordinates

**Returns:** coordinate (float 1x3): coordinate of internal sphere closest to the supplied coordinates

#### remove\_duplicates(eps=0.01)

Remove duplicate spheres by identifying centers closer together than eps using DBSCAN

**Args:** eps (float): DBSCAN input parameter (Default value = 0.01)

#### remove\_ungrouped()

Remove all spheres that did not adequately cluster with the remainder of the set

#### remove\_groups (groups)

Remove all spheres with specified group affiliations

**Args:** groups ([float]): list of groups to remove

#### write (filename, contents='xyzrg', output\_mesh=True)

Writes the contents of \_xyzrg to a space delimited file

**Args:** filename (str): filename to write the report and mesh if indicated contents (str): string describing which columns to write to file (Default value = "xyzrg") output\_mesh (bool): write mesh to file? (Default value = True)

#### xyzrg

Retrieve the coordinates, radii, and group ids

#### xyzr

Retrieve coordinates and radii

#### xyz

Retrieve the coordinates

r

Retrieve the radii

g

Retrieve the group indices

#### 3.5.7 pyvol.utilities module

```
pyvol.utilities.check_dir(location)
```

Ensure that a specified directory exists

**Args:** location (str): target directory

pyvol.utilities.coordinates\_for\_resid(pdb\_file, resid, chain=None, model=0)

Extract the 3D coordinates for all atoms in a specified residue from a pdb file

**Args:** pdb\_file (str): filename of the specified pdb file resid (int): residue number chain (str): chain identifier (Default value = None) model (int): model identifier (Default value = 0)

**Returns:** coordinates ([[float]]): 3xN array containing all atomic positions

pyvol.utilities.run\_cmd (options, in\_directory=None)

Run a program using the command line

**Args:** options ([str]): list of command line options in\_directory (str): directory in which to run the command (Default value = None)

pyvol.utilities.surface\_multiprocessing(args)

A single surface calculation designed to be run in parallel

**Args:** 

**args:** a tuple containing: spheres (Spheres): a Spheres object containing all surface producing objects probe\_radius (float): radius to use for probe calculations kwargs (dict): all remaining arguments accepted by the surface calculation algorithm

Returns: surface (Spheres): the input Spheres object but with calculated surface parameters

pyvol.utilities.sphere\_multiprocessing(spheres, radii, workers=None, \*\*kwargs)

A wrapper function to calculate multiple surfaces using multiprocessing

**Args:** spheres (Spheres): input Spheres object radii ([float]): list of radii at which surfaces will be calculated workers (int): number of workers (Default value = None) \*\*kwargs (dict): all remaining arguments accepted by surface calculation that are constant across parallel calculations

**Returns:** surfaces ([Spheres]): a list of Spheres object each with its surface calculated

#### 3.5.8 Module contents

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