# **Aqueduct Documentation**

Release 0.2.17

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### **AQUEDUCT INSTALLATION GUIDE**

### 1.1 Overview

This package comprises of two elements:

- 1. aqueduct,
- 2. valve.

Aqueduct is a Python module. It is a collection of tools to trace residues in MD simulation. Valve is a driver Python script. It uses aqueduct to perform such a tracing.

### 1.2 Install

### 1.2.1 Aqueduct

Installation was tested on limited number of POSIX-like systems.

In some specific cases installation is very simple:

- 1. Download aqueduct.tar.gz bundle file..
- 2. Unpack aqueduct bundle file.
- 3. Go to src directory.
- 4. Type:

```
python setup.py install
```

Aqueduct requires several Python modules to work and in particular it requires MDAnalysis with AMBER support. This, on the other hand, requires netCDF4. Installation of this combination is sometimes cumbersome. General procedure is following:

- 1. Install libnetcdf4 and libhdf5 development libraries.
- 2. Install netCDF4:

```
pip install netCDF4
```

3. Try to install aqueduct.

If, by chance, you are on Ubuntu 14.04 you can try helper script ubuntu\_mdanalysis\_install\_helper.sh.

### 1.2.2 Valve

Valve does not need installation per se. Once aqueduct is installed, valve can be run by a following command:

python valve.py --help

Valve script, ie valve.py, is located in apps directory.

### **1.2.3 Extras**

Access to some visualization capabilities of Aqueduct requires additional Python modules:

- 1. matplotlib,
- 2. pymol.

These are usually easy to install.

**CHAPTER** 

**TWO** 

### **VALVE MANUAL**

Valve application is a driver that uses aqueduct module to perform analysis of trajectories of selected residues in MD simulation.

### 2.1 Valve invocation

Once aqueduct module is installed (see Aqueduct installation guide) properly on the machine Valve is available as valve.py command line tool.

### 2.1.1 **Usage**

Basic help of *Valve* usage can be displayed by following command:

```
valve.py --help
```

It should display following information:

```
usage: valve.py [-h] [--dump-template-config] [-t THREADS] [-c CONFIG_FILE]
                [--max-frame MAX_FRAME]
Valve, Aqueduct driver
optional arguments:
                        show this help message and exit
 -h, --help
  --dump-template-config
                        Dumps template config file. Suppress all other output
                        or actions. (default: False)
 -t THREADS
                        Limit Aqueduct calculations to given number of
                       threads. (default: None)
 -c CONFIG_FILE
                        Config file filename. (default: None)
  --max-frame MAX_FRAME
                        Limit number of frames. (default: None)
  --version
                        Prints versions and exits.. (default: False)
```

### 2.1.2 Configuration file template

Configuration file used by *Valve* is of moderate length and complexity. It can be easily prepared with a template file that can be printed by *Valve*. Use following command to print configuration file template on the screen:

```
valve.py --dump-template-config
```

Configuration file template can also be easily saved in to a file with:

```
valve.py --dump-template-config > config.txt
```

Where config.txt is a configuration file template.

For detailed description of configuration file and available options see Configuration file options

#### 2.1.3 Valve calculation run

Once configuration file is ready Valve calculations can be run with a following simple command:

```
valve.py -c config.txt
```

Some of *Valve* calculations can be run in parallel. By default all available CPU cores are used. This is not always desired - limitation of used CPU cores can be done with -t option which limits number of concurrent threads used by *Valve*. If it equals 1 no parallelism is used.

**Note:** Specifying number of threads greater then available CPU cores is generally not optimal.

However, in order to maximize usage of available CPU power it is recommended to set it as number of cores + 1. The reason is that *Valve* uses one thread for the main process and the excess over one for processes for parallel calculations. When parallel calculations are executed the main threads waits for results.

**Note:** Option —max—frame can be used for testing or debugging purposes. It allows to limit number of frames processed by *Valve*. If it set, for example, to 1000 only first 1000 frames will be processed making all calculations very fast.

### 2.2 How does Valve work

Application starts with parsing input options. If --help or --dump-template-config options are provided appropriate messages are printed on the screen and Valve quits with signal 0.

**Note:** In current version *Valve* does not check the validity of the config file.

If config file is provided *Valve* parse it quickly and regular calculations starts according to its content. Calculations performed by *Valve* are done in several stages described in the next sections.

#### 2.2.1 Traceable residues

The first stage finds all residues that should be traced and appends them to the list of *traceable residues*. It is done in a loop over all frames. In each frame residues of interest are searched and appended to the list but only if they are not already present on the list.

The search of the residues is done according to user provided definitions.. Two requirements have to be met to append residue to the list:

- 1. The residue have to be found according to the *Object* definition.
- 2. The residues have to be within the *Scope* of interest.

The *Object* definition encompasses usually the active site of the protein. The *Scope* of interest defines, on the other hand, the boundaries in which residues are traced and is usually defined as protein.

Since aqueduct in its current version uses MDAnalysis Python module for reading, parsing and searching of MD trajectory data, definitions of *Object* and *Scope* have to be given as its *Selection Commands*.

### **Object definition**

Object definition have to comprise of two elements:

- 1. It have to define residues to trace.
- 2. It have to define spatial boundaries of the *Object* site.

For example, proper Object definition could be following:

```
(resname WAT) and (sphzone 6.0 (resnum 99 or resnum 147))
```

It defines WAT as residues that should be traced and defines spatial constrains of the *Object* site as spherical zone within 6 Angstroms of the center of masses of residues with number 99 and 147.

### Scope definition

Scope can be defined in two ways: as Object but with broader boundaries or as the convex hull of selected molecular object.

In the first case definition is very similar to *Object* and it have to follow the same limitations. For example, proper *Scope* definition could be following:

```
resname WAT around 2.0 protein
```

It consequently have to define WAT as residues of interest and defines spatial constrains as all WAT residues that are within 2 Angstroms of the protein.

If the *Scope* is defined as the convex hull of selected molecular object (which is recommended), the definition itself have to comprise of this molecular object only, for example protein. In that case the scope is interpreted as the interior of the convex hull of atoms from the definition. Therefore, *traceable residues* would be in the scope only if they are within the convex hull of atoms of protein.

### 2.2.2 Raw paths

The second stage of calculations uses the list of all traceable residues from the first stage and finds coordinates of center of masses for each residue in each frame. As in the first stage, it is done in a loop over all frames. For each residue in each frame *Valve* calculates or checks two things:

- 1. Is the residue in the *Scope* (this is always calculated according to the Scope definition).
- 2. Is the residue in the *Object*. This information is calculated in the first stage and can be reused in the second. However, it is also possible to recalculate this data according to the new *Object* definition.

For each of the *traceable residues* a special *Path* object is created. If the residue is in the *Scope* its center of mass is added to the appropriate *Path* object together with the information if it is in the *Object* or not.

### 2.2.3 Separate paths

The third stage uses collection of *Path* objects to create *Separate Path* objects. Each *Path* comprise data for one residue. It may happen that the residue enters and leaves the *Scope* and the *Object* many times over the entire MD. Each such an event is considered by *Valve* as a separate path.

Each separate path comprises of three parts:

- 1. *Incoming* Defined as a path that leads from the point in which residue enters the *Scope* and enters the object for the firs time.
- 2. *Object* Defined as a path that leads from the point in which residue enters the *Object* for the first time and leaves it for the last time.
- 3. *Outgoing* Defined as a path that leads from the point in which residue leaves the *Object* for the last lime and leaves the *Scope*.

#### **Auto Barber**

After the initial search of *Separate Path* objects it is possible to run procedure which trims paths down to the surface of macromolecule or other molecular entity defined by the user. This is done by removing parts of raw paths that are inside spheres that originate in the points marking these ends of separate paths that end at the boundary of *Scope*. Recreation of separate paths is run automatically after Auto Barber procedure.

#### 2.2.4 Clusterization of inlets

Each of the separate paths has beginning and end. If either of them are at the boundaries of the *Scope* they are considered as *Inlets*, i.e. points that mark where the *traceable residues* enters or leaves the *Scope*. Clusters of inlets, on the other hand, mark endings of tunnels or ways in the system which was simulated in the MD.

Clusterization of inlets is performed in following steps:

- 1. Initial clusterization. Depending on the method, some of the inlets might not be arranged to any cluster and are considered as outliers.
- 2. [Optional] Outliers detection. Arrangement of inlets to clusters is sometimes far from optimal. In this step, *inlets* that do not fit to cluster are detected and annotated as outliers. This step can be executed in two modes:
  - (a) Automatic mode. Inlet is considered to be an outlier if its distance from the centroid is greater then mean distance + 4 \* standard deviation of all distances within the cluster.
  - (b) Defined threshold. Inlet is considered to be an outlier if its minimal distance from any other point in the cluster is greater then the threshold.
- 3. [Optional] Reclusterization of outliers. It may happen that the outliers form actually clusters but it was not recognized in initial clusterization. In this step clusterization is executed for outliers only and found clusters are appended to the clusters identified in the first step. Rest of the inlets are marked as outliers.

#### Potentially recursive clusterization

Both *Initial clusterization* and *Reclustarization* can be run in a recursive manner. If in the appropriate sections defining clusterization methods option *recursive\_clusterization* is used appropriate method is run for each cluster separately. Clusters of specific size can be excluded from recursive clusterization (option *recursive\_threshold*). It is also possible to limit maximal number of recursive levels - option *max\_level*. For additional information see *Clusterization sections*.

### 2.2.5 Analysis

Fifth stage of *Valve* calculations analyses results calculated in stages 1 to 4. Results of the analysis is displayed on the screen or can be save to text file and comprise of following parts:

- Tile and data stamp.
- [Optional] Dump of configuration options.
- Basic information on traceable residues and separate paths.
  - Number of traceable residues.
  - Number of separate paths.
- Basic information on inlets.
  - Number of inlets.
  - Number of clusters.
  - Are outliers detected.
- Summary of inlets clusters. Table with 5 columns:

- 1. Nr: Row number, starting from 0.
- 2. Cluster: ID of the cluster. Outliers have 0.
- 3. Size: Size of the cluster.
- 4. **INCOMING**: Number of inlets corresponding to separate paths that enter the scope.
- 5. **OUTGOING**: Number of inlets corresponding to separate paths that leave the scope.

### • Summary of separate paths clusters types. Table with 9 columns.

- 1. Nr: Row number, starting from 0.
- 2. **CType**: Separate path Cluster Type.
- 3. **Size**: Number of separate paths belonging to Cluster type.
- 4. **Inp**: Average length of incoming part of the path. If no incoming part is available it is nan.
- 5. **InpStd**: Standard deviation of length Inp.
- 6. **Obj**: Average length of object part of the path. If no incoming part is available it is nan.
- 7. **ObjStd**: Standard deviation of length Inp.
- 8. Out: Average length of outgoing part of the path. If no incoming part is available it is nan.
- 9. OutStd: Standard deviation of length Inp.

#### • List of separate paths and their properties. Table with 17 columns.

- 1. **Nr**: Row number, starting from 0.
- 2. **ID**: Separate path ID.
- 3. **BeginF**: Number of frame in which the path begins.
- 4. **InpF**: Number of frame in which path begins Incoming part.
- 5. **ObjF**: Number of frame in which path begins Object part.
- 6. OutF: Number of frame in which path begins Outgoing part.
- 7. **EndF**: Number of frame in which the path ends.
- 8. **InpL**: Length of Incoming part. If no incoming part nan is given.
- 9. ObjL: Length of Object part.
- 10. OutL: Length of Outgoing part. If no outgoing part nan is given.
- 11. **InpS**: Average step of Incoming part. If no incoming part nan is given.
- 12. InpStdS: Standard deviation of InpS.
- 13. **ObjS**: Average step of Object part.
- 14. **ObjStdS**: Standard deviation of ObjS.
- 15. OutS: Average step of Outgoing part. If no outgoing part nan is given.
- 16. OutStdS: Standard deviation of OutS.
- 17. **CType**: Cluster type of separate path.

#### Separate path ID

Separate Paths IDs are composed of two numbers separated by colon. First number is the residue number. Second number is consecutive number of the separate path made by the residue. Numeration starts with 0.

#### **Cluster Type of separate path**

Each separate paths has two ends: beginning and end. Both of them either belong to one of the inlets clusters, or are among outliers, or are inside the scope. If an end belongs to one of the clusters (including outliers) it has ID of the cluster. If it is inside the scope it has special ID of N. Cluster type is an ID composed of IDs of both ends of separate path separated by colon charter.

#### 2.2.6 Visualization

Sixth stage of *Valve* calculations visualizes results calculated in stages 1 to 4. Visualization is done with PyMOL. *Valve* automatically starts PyMOL and loads visualizations in to it. Molecule is loaded as PDB file. Other objects like Inlets clusters or paths are loaded as CGO objects.

Following is a list of objects created in PyMOL (all of them are optional). PyMOL object names given in **bold** text or short explanation is given.

- Selected frame of the simulated system. Object name: molecule.
- Inlets clusters, each cluster is a separate object. Object name: **cluster\_** followed by cluster annotation: otliers are annotated as Out; regular clusters by ID.
- List of cluster types, raw paths. Each cluster type is a separate object. Object name composed of cluster type (colon replaced by underline) plus **\_raw**.
- List of cluster types, smooth paths. Each cluster type is a separate object. Object name composed of cluster type (colon replaced by underline) plus **\_smooth**.
- All raw paths. They can be displayed as one object or separated in to Incoming, Object and Outgoing part. Object name: all\_raw\_in, all\_raw\_obj, and all\_raw\_out.
- All raw paths inlets arrows. Object name: all\_raw\_paths\_io.
- All smooth paths. They can be displayed as one object or separated in to Incoming, Object and Outgoing part. Object name: all\_smooth, or all\_smooth\_in, all\_smooth\_obj, and all\_smooth\_out.
- All raw paths inlets arrows. Object name: all\_raw\_paths\_io.
- Raw paths displayed as separate objects or as one object with several states. Object name: **raw\_paths\_** plus number of path or **raw\_paths** if displayed as one object.
- Smooth paths displayed as separate objects or as one object with several states. Object name: **smooth paths** plus number of path or **smooth paths** if displayed as one object.
- Raw paths arrows displayed as separate objects or as one object with several states. Object name: raw\_paths\_io\_ plus number of path or raw\_paths\_io if displayed as one object.
- Smooth paths arrows displayed as separate objects or as one object with several states. Object name: **smooth\_paths\_io**\_ plus number of path or **smooth\_paths\_io** if displayed as one object.

#### **Color schemes**

Inlets clusters are colored automatically. Outliers are gray.

Incoming parts of paths are red, Outgoing parts are blue. Object parts in case of smooth paths are green and in case of raw paths are green if residue is precisely in the object area or yellow if is leaved object area but it is not in the Outgoing part yet.

Arrows are colored in accordance to paths colors.

### **CONFIGURATION FILE OPTIONS**

Valve Configuration file is a simple and plain text file. It has similar structure as INI files commonly used in one of the popular operating systems and is compliant with Python module ConfigParser.

Configuration file comprises of several *sections*. They can be grouped in to three categories. Names of sections given in **bold** text.

- 1. Global settings:
  - global
- 2. Stages options:
  - (a) traceable\_residues
  - (b) raw\_paths
  - (c) separate\_paths
  - (d) inlets\_clusterization
  - (e) analysis
  - (f) visualize
- 3. Methods options:
  - smooth
  - clusterization
  - reclusteriation

### 3.1 Section global

This section allows settings of trajectory data and some other future global options.

Option	Default value	Description
top	None	Path to topology file. Aqueduct supports PDB, PRMTOP, PFS topology files.
trj	None	Path to trajectory file. Aqueduct supports NC and DCD trajectory files.

**Note:** Options **top** and **trj** are mandatory.

### 3.2 Common settings of stage sections

Stages 1-4 which perform calsulations have some common options allowig for execution control and saving/loading data.

Option	Default value	Description
execute	runonce	Option controls stage execution. It
		can have one of three possible val-
		ues: run, runonce, and skip.
		If it is set to run calculations are
		always performed and if <b>dump</b> is
		set dump file is saved. If it is set
		to runonce calculations are per-
		formed if there is no dump file spec-
		ified by <b>dump</b> option. If it is present
		calculations are skiped and data is
		loaded from the file. If it is set to
		skip calculations are skip and if
		<b>dump</b> is set data is loaded from the
		file.
dump	[dump file name]	File name of dump data. It is used
		to save results of calculations or
		to load previously calculated data -
		this depends on <b>execute</b> option. De-
		fault value of this option depends on
		the stage and for stages 1 to 4 is one
		of the following (listed in order):
		•
		1_traceable_residues_data.dump
		• 2_raw_paths_data.dump
		• 3_separate_paths_data.dump
		•
		4_inlets_clusterization_data.dum

Stages 5-6 also uses **execute** option, however, since they do not perform calculations *per se* in stead of **dump** option they use **save**.

Option	Default value	Description
execute	run	Option controls stage execution. It
		can have one of three possible val-
		ues: run, runonce, and skip. If
		it is set to run or runonce stage
		is executed and results is saved ac-
		cording to <b>save</b> option. If it is set to
		skip stage is skipped.
save	[save file name]	File name for saving results. De-
		fault value of this option depends on
		the stage and for stages 1 to 4 is one
		of the following (listed in order):
		• 5_analysis_results.txt
		<ul> <li>6_visualize_results.py</li> </ul>
		Stage 6 can save results in two file
		types:
		1. As Python script - extension
		.py plus companion archive
		.tar.gz,
		2. As PyMOL session - exten-
		sion .pse.

### 3.3 Stage traceable\_residues

Option	Default value	Description
scope	None	Definition of <i>Scope</i> of interest. See also <i>Scope definition</i> .
scope_convexhull	True	Flag to set if the <i>Scope</i> is direct or convex hull definition.
object	None	Definition of <i>Object</i> of interest. See also <i>Object definition</i> .

Note: Options scope and object are mandatory.

## 3.4 Stage raw\_paths

This stage also requires definition of the *Scope* and *Object*. If appropriate settings are not given, settings from the previous stage are used.

Option	De-	Description
	fault	
	value	
scope	None	Definition of <i>Scope</i> of interest. See also <i>Scope definition</i> . If None value form
		previous stage is used.
scope_conve	ex <b>No</b> He	Flag to set if the <i>Scope</i> is direct or convex hull definition. If None value form
		previous stage is used.
object	None	Definition of Object of interest. See also Object definition. If None, value form the
		previous stage is used
clear_in_ob	jeÆt <u>al</u> isæfo	If it is set to True information on occupation of <i>Object</i> site by traceable residues
		calculated in the previous stage is cleared and have to be recalculated. This is useful
		if definition of <i>Object</i> was changed.

### 3.5 Stage separate\_paths

Option	De-	Description
	fault	
	value	
dis-	True	If set to True empty paths are discarded.
card_empty_	paths	
sort_by_id	True	If set to True separate paths are sorted by ID. Otherwise they are sorted in order of
		apparance.
ap-	False	If set to True smooth paths are precalculated according to <b>smooth</b> setting. This
ply_smoothin	ng	speed up access to smooth paths in later stages but makes dump data much bigger.
ap-	True	If set to True raw paths are replaced by smooth paths calculated according to
ply_soft_smo	othing	smooth section.
dis-	1	This option allows to discard paths that are shorter then the threshold.
card_short_p	aths	
auto_barber	None	This option allows to select molecular entity used in Auto Barber procedure. See
		also Auto Barber and barber_with_spheres().

# 3.6 Stage inlets\_clusterization

Option	De-	Description
	fault	
	value	
reclus-	False	If set to True reclusterization of outliers is executed according to the method defined
ter_outliers		in <b>reclusterization</b> section.
de-	False	If set detection of outliers is executed. It could be set as a floating point distance
tect_outlier	s	threshold or set tu Auto. See Clusterization of inlets for more details.
single-	False	Maximal size of cluster to be considered as outliers. If set to number > 0 clusters of
tons_outlie	rs	that size are removed and their objects are moved to outliers. See Clusterization of
		<i>inlets</i> for more details.
max_level	5	Maximal number of recursive clusterization levels.

# 3.7 Stage analysis

Option	Default value	Description
dump_confi	True	If set to True configuration options, as seen by Valve, are added to the head of results.

### 3.8 Stage visualize

Option	De-	Description
	fault	
	value	
sim-	0.05236	If set to float number simplification of smooth paths is applied. Simplification
ply_smooths		removes points which do not (or almost do not) change the shape of smooth path.
		For more details see Recursive Vector Linearization.
all_paths_raw	False	If True produces one object in PyMOL that holds all paths visualized by raw
		coordinates.
all_paths_smo	odfalse	If True produces one object in PyMOL that holds all paths visualized by smooth
		coordinates.
all_paths_spli	it False	If is set True objects produced by all_paths_raw and all_paths_smooth are split
		into Incoming, Object, and Outgoing parts and visualized as three different objects.
all_paths_raw	_False	If set True arrows pointing beginning and end of paths are displayed oriented
		accordingly to raw paths orientation.
all_paths_smo	odfal <u>s</u> io	If set True arrows pointing beginning and end of paths are displayed oriented
		accordingly to smooth paths orientation.
paths_raw	False	If set True raw paths are displayed as separate objects or as one object with states
		corresponding to number of path.
paths_smooth	False	If set True smooth paths are displayed as separate objects or as one object with
		states corresponding to number of path.
paths_raw_io	False	If set True arrows indicating beginning and and of paths, oriented accordingly to raw
		paths, are displayed as separate objects or as one object with states corresponding to
		number of paths.
paths_smooth	_Foalse	If set True arrows indicating beginning and and of paths, oriented accordingly to
		smooth paths, are displayed as separate objects or as one object with states
		corresponding to number of paths.
paths_states	False	If True objects displayed by paths_raw, paths_smooth, paths_raw_io, and
		paths_smooth_io are displayed as one object with with states corresponding to
		number of paths. Otherwise they are displayed as separate objects.
ctypes_raw	False	Displays raw paths in a similar manner as non split <b>all_paths_raw</b> but each cluster
		type is displayed in separate object.
ctypes_smoot	hFalse	Displays smooth paths in a similar manner as non split <b>all_paths_smooth</b> but each
		cluster type is displayed in separate object.
show_molecu	laFalse	If is set to selection of some molecular object in the system, for example to
		protein, this object is displayed.
show_molecu	le_frames	Allows to indicate which frames of object defined by <b>show_molecule</b> should be
		displayed. It is possible to set several frames. In that case frames would be
		displayed as states.
show_chull	False	If is set to selection of some molecular object in the system, for example to
		protein, convex hull of this object is displayed.
show_chull_f	rames	Allows to indicate for which frames of object defined by <b>show_chull</b> convex hull
		should be displayed. It is possible to set several frames. In that case frames would
		be displayed as states.

**Note:** Possibly due to limitations of MDAnalysis only whole molecules can be displayed. If **show\_molecule** is set to backbone complete protein will be displayed any way. This may change in future version of MDAnalysis and or aqueduct.

**Note:** If several frames are selected they are displayed as states which may interfere with other PyMOL objects displayed with several states.

Note: If several states are displayed protein tertiary structure data might be lost. This seems to be limitation of

3.8. Stage visualize

either MDAnalysis or PyMOL.

### 3.9 Clusterization sections

Default section for definition of clusterization method is named **clusterization** and default section for reclusterization method definition is named **reclusterization**. All clusterization sections shares some common options. Other options depends on the method.

Option	Default	Description
	value	
method	meanshift	Name of clasteriation method. It have to be one of the following: dbscan, affprop,
	or dbscan	meanshift, birch, kmeans. Default value depends if it is <b>clusteriation</b> section
		(meanshift) or <b>reclusterization</b> section (dbscan).
recur-	clusteriza-	If it is set to name of some section that holds clusterization method settings this
sive_clusteriz	za <b>tion</b> or	method will be called in the next recursion of clusteriation. Default value for
	None	reclusterization is None.
recur-	None	Allows to set threshold of that excludes clusters of certain size from
sive_threshol	d	reclusterization. Value of this option comprises of <i>operator</i> and <i>value</i> . Operator
		can be one of the following: >, >=, <=, <. Value have to be expressed as floating
		number and it have to be in the range of 0 to 1.

### 3.9.1 dbscan

For detailed description look at sklearn.cluster.DBSCAN documentation. Following table summarized options available in *Valve* and is a copy of original documentation.

Option	Value type	Description
eps	float	The maximum distance between two samples for them to be considered as in the same neighborhood.
min_samples	int	The number of samples (or total weight) in a neighborhood for a point to be considered as a core point. This includes the point itself.
metric	str	The metric to use when calculating distance between instances in a feature array. Can be one of the following:  • cityblock, • cosine, • euclidean, • manhattan.
algorithm	str	The algorithm to be used by the NearestNeighbors module to compute pointwise distances and find nearest neighbors. Can be one of the following:  • auto, • ball_tree, • kd_tree, • brute.
leaf_size	int	Leaf size passed to BallTree or cK-DTree.

### 3.9.2 affprop

For detailed description look at AffinityPropagation documentation. Following table summarized options available in *Valve* and is a copy of original documentation.

Option	Value	Description
	type	
damping	float	Damping factor between 0.5 and 1.
conver-	int	Number of iterations with no change in the number of estimated clusters that
gence_iter		stops the convergence.
max_iter	int	Maximum number of iterations.
preference	float	Points with larger values of preferences are more likely to be chosen as
		exemplars.

### 3.9.3 meanshift

For detailed description look at MeanShift documentation. Following table summarized options available in *Valve* and is a copy of original documentation.

Ор-	Value	Description
tion	type	
band-	Auto	Bandwidth used in the RBF kernel. If Auto or None automatic method for bandwidth
width	or	estimation is used. See estimate_bandwidth().
	float	
clus-	bool	If true, then all points are clustered, even those orphans that are not within any kernel.
ter_all		
bin_seed	in <b>b</b> ool	If true, initial kernel locations are not locations of all points, but rather the location of the
		discretized version of points, where points are binned onto a grid whose coarseness
		corresponds to the bandwidth.
min_bin_	_f <b>re</b> tq	To speed up the algorithm, accept only those bins with at least min_bin_freq points as
		seeds. If not defined, set to 1.

### 3.9.4 birch

For detailed description look at Birch documentation. Following table summarized options available in *Valve* and is a copy of original documentation.

Option	Value	Description
	type	
thresh-	float	The radius of the subcluster obtained by merging a new sample and the closest
old		subcluster should be lesser than the threshold. Otherwise a new subcluster is started.
branch-	int	Maximum number of CF subclusters in each node.
ing_factor		
n_clusters	int	Number of clusters after the final clustering step, which treats the subclusters from the
		leaves as new samples. By default, this final clustering step is not performed and the
		subclusters are returned as they are.

### **3.9.5 kmeans**

For detailed description look at KMeans documentation. Following table summarized options available in *Valve* and is a copy of original documentation.

Op-	Value	Description
tion	type	
n_clust	enisnt	The number of clusters to form as well as the number of centroids to generate.
max_it	erint	Maximum number of iterations of the k-means algorithm for a single run.
n_init	int	Number of time the k-means algorithm will be run with different centroid seeds. The final
		results will be the best output of n_init consecutive runs in terms of inertia.
init	str	Method for initialization, defaults to k-means++. Can be one of following: k-means++
		or random.
tol	float	Relative tolerance with regards to inertia to declare convergence.

### 3.10 Smooth section

Section **smooth** supports following options:

Option	Value type	Description
method	str	Smoothing method. Can be one of
		the following:
		• window, (see
		WindowSmooth)
		• mss, (see
		MaxStepSmooth)
		• window_mss, (see
		WindowOverMaxStepSmooth)
		• awin, (see
		ActiveWindowSmooth)
		• awin_mss, (see
		ActiveWindowOverMaxStepSmooth)
		• dwin, (see
		DistanceWindowSmooth)
		• dwin_mss. (see
		DistanceWindowOverMaxStepSmoot.
recursive	int	Number of recursive runs of
		smoothing method.
window	int or float	In window based method defines
		window size. In plain window it
		has to be int number.
step	int	In step based method defines size of
		the step.
function	str	In window based methods defines
		averaging function. Can be mean
		or median.

**CHAPTER** 

**FOUR** 

### **VALVE TUTORIAL**

This is a tentative *Valve* manual. Created for the sake of Aqueduct training we have today. Eventually, it will be rewritten to the official version.

This tutorial assumes aqueduct and Valve is already installed - see Aqueduct installation guide. It is also assumed that user is acquainted with Valve manual and Valve Configuration file options.

### 4.1 Valve invocation

Usually Valve is run by:

```
valve.py
```

Due to specific setup in our laboratory *Valve* has to be run through simple wrapper script:

```
valve_run
```

Additionally, to speed up all calculations it is assumed that Valve is run with --max-frame 1000 option:

```
valve_run --max-frame 1000
```

To check is Valve is installed and works properly try to issue following commands:

```
valve_run --help
valve_run --version
```

### 4.2 Test data

#### Mouse!

We will use 10ns Amber MD simulation data of sEH protein (PDBID 1cqz). Necessary files can be downloaded here:

- Go to download server.
- Go to 1cgz directory.
- Download all files and save them in sane location on your machine. Please note, that .nc file is ca. 3.5 GB so it may take a while to download it.

### 4.3 Inspect your system

Before we start any calculations lets have a look at the protein of interest. Start *PyMOL* and get 1cqz PDB structure (for example by typing in *PyMOL* command prompt fetch 1cqz).

To setup *Valve* calculations we need to know active site of the protein. More precisely we need to know IDs or residues that are in the active site. This would allow us to create *Object definition*.

But wait. Is it really the correct structure? How many chains there are? What is the numeration of residues?

### 4.3.1 Create Object definition

Lets load another structure. Open file first\_frame\_1cqz.pdb downloaded from test data repository. It is a first frame of the MD simulation and it is en example of how the frame of MD looks like. In order to create *Object definition* you have to discover following things:

- 1. What is the name of water residue?
- 2. What are numbers of residues in the active site?
- 3. What size the active site is?

**Note:** It is also good idea to open . pdb file in your favorite text editor and look at residue numbers and names.

### 4.3.2 Create Scope definition

Scope definition is easy to create. We will use Convex hull version so the scope definition could be simply backbone.

### 4.4 Prepare config file

Valve performs calculations according to the configuration (aka config) file.

Lets start from dumping config file template to config.txt file. Open it in your favorite editor and fill all options. If you have troubles look at *Configuration file options* (and *Valve manual*).

Things to remember:

- 1. Provide correct paths to topology and trajectory data.
- 2. Enter correct *Object* and *Scope* definitions.
- 3. Provide file name of result in analysis section, for example results.txt (for future reference).
- 4. Make sure visualization is switched on and save option points to session file name (.pse)

### 4.5 Run Valve

Make sure all necessary data is in place. Open terminal, go to your working directory and type in:

```
valve_run --max-frame 1000 -c config.txt
```

Depending on your machine and current load it may take a while (matter of minutes) to complete all calculations.

### 4.5.1 Visual inspection

In the last stage *PyMOL* should pop up and *Valve* should start to feed it with visualization data. This would take a moment and if you set up save option a *PyMOL* session would be saved. Once it is done *Valve* quits and switches off *PyMOL*. Now, you can restart it and read saved session.

### 4.5.2 Analysis tables

Open results.txt file and look at summaries and tables. See also *Valve manual*.

### 4.6 Feedback

Give us your opinion. Send your questions, inquires, anything to developer(s): Tomasz Magdziarz. This are couple of questions that might be useful to form your opinion.

- 1. What do you like in Valve and Aqueduct?
- 2. What do you do not like in Valve or Aqueduct?
- 3. What is missing?
- 4. Do you find it useful?

4.6. Feedback 21

**CHAPTER** 

**FIVE** 

### **AQUEDUCT**

### 5.1 aqueduct package

### 5.1.1 Subpackages

aqueduct.geom package

### **Submodules**

**aqueduct.geom.cluster module** This module provides functions for clusterization. Clusterization is done by scikit-learn module.

```
MeanShiftBandwidth (X, **kwargs)
class PerformClustering (method, **kwargs)
     Bases: object
     __init__ (method, **kwargs)
     __call__(coords)
     _get_noclusters(n)
     fit (coords)
     centers()
aqueduct.geom.convexhull module
_vertices_ids(convexhull)
_vertices_points(convexhull)
_point_within_convexhull (convexhull, point)
_facets(convexhull)
_edges (*args, **kwargs)
is_point_within_convexhull (point_chull)
aqueduct.geom.master module
fit_trace_to_points (trace, points)
decide_on_type (cont, s2o_treshold=0.5)
simple_types_distribution(types)
get_weights_(spaths, smooth=None)
\mathtt{get\_mean\_coord\_}(\mathit{coords}, l)
```

concatenate (\*args)

```
create_master_spath (spaths, smooth=None, resid=0, ctype=None, bias_long=5, pbar=None)
calculate_master(spaths_resid_ctype_smooth)
class MasterTrace
     Bases: object
     ___init___()
aqueduct.geom.pca module
class Center (X)
     Bases: object
     \_\_\mathtt{init}\_\_(X)
      \underline{\hspace{1cm}}call\underline{\hspace{1cm}}(X)
     {\tt undo}\,(X)
class Normalize (X)
     Bases: object
     \_\_\mathtt{init}\_\_(X)
      \mathtt{call} (X)
     \mathbf{undo}\left( X\right)
class Standartize (X)
     Bases: aqueduct.geom.pca.Center, aqueduct.geom.pca.Normalize
      ___init___(X)
     \_\_\mathtt{call}\_\_(X)
     undo (X)
class PCA (X, prepro=None)
     Bases: object
      ___init___(X, prepro=None)
     P
     preprocess(X)
     {\tt preprocess\_undo}\,(X)
     \_\_\mathtt{call}\_\_(X)
     undo(T)
aqueduct.geom.smooth module Created on Dec 15, 2015
@author: tljm
class Smooth (recursive=None, **kwargs)
     Bases: object
     __init__ (recursive=None, **kwargs)
      smooth (coords)
      __call__(coords)
class GeneralWindow
     static max_window_at_pos (pos, size)
      \verb|check_bounds_at_max_window_at_pos|(lb, ub, pos, size)|
```

```
class WindowSmooth (window=5, function=<function mean>, **kwargs)
    Bases: aqueduct.geom.smooth.Smooth, aqueduct.geom.smooth.GeneralWindow
     ___init___(window=5, function=<function mean>, **kwargs)
     smooth (*args, **kwargs)
class DistanceWindowSmooth (window=5, function=<function mean>, **kwargs)
    Bases: aqueduct.geom.smooth.Smooth, aqueduct.geom.smooth.GeneralWindow
     ___init___(window=5, function=<function mean>, **kwargs)
     smooth (*args, **kwargs)
class ActiveWindowSmooth (window=5, function=<function mean>, **kwargs)
    Bases: aqueduct.geom.smooth.Smooth, aqueduct.geom.smooth.GeneralWindow
     ___init___(window=5, function=<function mean>, **kwargs)
     smooth (*args, **kwargs)
class MaxStepSmooth (step=1.0, **kwargs)
    Bases: aqueduct.geom.smooth.Smooth
     ___init___(step=1.0, **kwargs)
     smooth (*args, **kwargs)
class WindowOverMaxStepSmooth(**kwargs)
    Bases: aqueduct.geom.smooth.Smooth
     ___init___(**kwargs)
     smooth (coords)
class ActiveWindowOverMaxStepSmooth(**kwargs)
    Bases: aqueduct.geom.smooth.Smooth
     ___init___(**kwargs)
     smooth (coords)
class DistanceWindowOverMaxStepSmooth (**kwargs)
    Bases: aqueduct.geom.smooth.Smooth
     ___init___(**kwargs)
     smooth (coords)
aqueduct.geom.test linearizeOneWay module
class TestLinearizeOneWay (methodName='runTest')
    Bases: unittest.case.TestCase
    test_Linearize_One_Way()
aqueduct.geom.test_linearizeRecursive module
class TestLinearizeRecursive (methodName='runTest')
    Bases: unittest.case.TestCase
    test_here()
    test_here_nonlinear()
```

```
aqueduct.geom.test_triangle_height module
class TestTriangle_height (methodName='runTest')
    Bases: unittest.case.TestCase
    test_OutType()
    test_2dim()
    test_3dim()
aqueduct.geom.test_trianlgeLinearize module
class TestTrianlgeLinearize (methodName='runTest')
    Bases: unittest.case.TestCase
    test_is_linear()
    test_false_linear()
    test_low_threshold()
    test_list()
    test_numpyarr()
    test_sillyvalue()
aqueduct.geom.test_vectors_angle_anorm module
class TestVectors_angle_anorm (methodName='runTest')
    Bases: unittest.case.TestCase
    test_output_value()
    test_vectors_angle_anorm()
    test_list_input()
    test_zero()
    test_zero2()
    test_huge_values()
    test_mixed_values()
    test_negative_vector()
    test_one_negative_value()
aqueduct.geom.traces module
\mathtt{vector}\ \mathtt{norm}\,(V)
triangle\_angles(A, B, C)
triangle_angles_last(A, B, C)
triangle_height(A, B, C)
vectors\_angle(A, B)
vectors\_angle\_alt(A, B)
\verb"vectors_angle_alt_anorm" (A, B, A\_norm)
vectors\_angle\_anorm(A, B, A\_norm)
class LinearizeOneWay
    Bases: object
    here (coords)
     __call__(coords)
```

#### class LinearizeHobbit

```
Bases: aqueduct.geom.traces.LinearizeOneWay
and_back_again(coords)
__call__(coords)

class TriangleLinearize(threshold)
Bases: object
__init__(threshold)
```

# is\_linear(coords, \*\*kwargs) class LinearizeRecursive(threshold)

Bases: aqueduct.geom.traces.TriangleLinearize

Base class for linearization methods classes.

It implements recursive algorithm.

```
here (coords, depth=0)
```

Core of recursive linearization argorithm.

It checks if the first, the last and the middle point are linear according to the criterion. The middle point is selected a point that is in the middle of length of the paths made by input coordinates.

If these points are linear their indices are returned. Otherwise, coordinates are split into two parts. First part spans points from the first point to the middle point (inclusive) and the second parth spans points from the middle (inclusive) to the last point. Next, these two parts are submitted recursively to here ().

Results of these recursive calls are joined, redundant indices are removed and sorted result is returned.

#### **Parameters**

- coords (numpy.ndarray) Input coordinates.
- **depth** (*int*) Depth of recurence.

**Returns** Indices of :arg: 'coords' points that can be used instead of all points in visulatization.

Return type list of int

```
__call__(coords)
```

### class VectorLinearize (treshold)

Bases: object

Base class for linearization methods classes.

It implements vectro linearization criterion.

```
__init__(treshold)
```

#### is\_linear\_core (coords, depth=None)

Method checks if input coordinates are linear according to the threshold and depth.

It begins with calculation of the threshold. If *depth* is None it is set to 1. Current treshold is calculated with following simple equation:

$$treshold_{current} = treshold_{initial} * (2 - 0.9^{depth})$$

Next, in a loop over all points but the first and the last the angle is calculated between two vectors. The first one made by the point and the first point, and the second vector made by the last and the first point. If any of the calculated angles is bigger the the treshold methods returns False; otherwise method returns True.

#### **Parameters**

- coords (numpy.ndarray) Coodrdinates for which linearization criterion is checked.
- **depth** (*int*) Depth of recurence.

**Returns** True if input coordinates are linear and False otherwise.

Return type bool

```
is_linear (coords, depth=None, **kwargs)
```

For more detail see is\_linear\_core() which is used as the criterion of linearity in this method.

#### **Parameters**

- **coords** (numpy.ndarray) Coodrdinates for which linearizetion criterion is checked.
- **depth** (*int*) Depth of recurence.

**Returns** True if input coordinates are linear and False otherwise. Criterion is checked for coordinates in normal and reverse order.

Return type bool

#### class LinearizeRecursiveVector (threshold)

Bases: aqueduct.geom.traces.LinearizeRecursive, aqueduct.geom.traces.VectorLinearize Class provides recursive linearization of coordinates with LinearizeRecursive algorithm and the criterion of linearity implemented by VectorLinearize.

```
diff(trace)
tracepoints(start, stop, nr)
midpoints(paths)
length_step_std(trace)
derrivative(values)
Module contents
```

### aqueduct.traj package

#### **Submodules**

# aqueduct.traj.dumps module class TmpDumpWriterOfMDA

```
Bases: object
__init__()
dump_frames (reader, frames, selection='protein')
close()
__del__()
```

# aqueduct.traj.inlets module class ProtoInletTypeCodes

```
surface = 'surface'
internal = 'internal'
incoming = 'inin'
```

```
outgoing = 'inout'
class InletTypeCodes
     \textbf{Bases:}\ aqueduct.traj.inlets. \textit{ProtoInletTypeCodes}
     all_surface = [('surface', 'inin'), ('surface', 'inout')]
     all_internal = [('internal', 'inin'), ('internal', 'inout')]
     all_incoming = [('surface', 'inin'), ('internal', 'inin')]
     all_outgoing = [('surface', 'inout'), ('internal', 'inout')]
     surface_incoming = ('surface', 'inin')
     internal_incoming = ('internal', 'inin')
     internal_outgoing = ('internal', 'inout')
     surface_outgoing = ('surface', 'inout')
     itype = 'internal'
{\bf class} \ {\bf InletClusterGenericType} \ (inp, \ out)
     Bases: object
      __init___(inp, out)
     input
     output
     {\tt cluster2str}\,(cl)
     __getitem__(item)
      __len__()
     __str__()
     __repr__()
     make_val(base)
     ___cmp___(other)
      __hash___()
class InletClusterExtendedType (surfin, interin, interout, surfout)
     Bases: aqueduct.traj.inlets.InletClusterGenericType
      ___init___(surfin, interin, interout, surfout)
     generic
class Inlet (coords, type, reference)
     Bases: tuple
     __getnewargs__()
          Return self as a plain tuple. Used by copy and pickle.
     __getstate__()
          Exclude the OrderedDict from pickling
     static __new__ (_cls, coords, type, reference)
          Create new instance of Inlet(coords, type, reference)
          Return a nicely formatted representation string
     __slots__ = ()
     _asdict()
          Return a new OrderedDict which maps field names to their values
```

```
_fields = ('coords', 'type', 'reference')
     classmethod _make (iterable, new=<built-in method __new__ of type object>, len=<built-in func-
                         tion len>)
          Make a new Inlet object from a sequence or iterable
     _replace(_self, **kwds)
          Return a new Inlet object replacing specified fields with new values
     coords
          Alias for field number 0
     reference
          Alias for field number 2
     type
          Alias for field number 1
class Inlets (spaths, onlytype=[('surface', 'inin'), ('surface', 'inout')])
     Bases: object
     __init__ (spaths, onlytype=[('surface', 'inin'), ('surface', 'inout')])
     extend_inlets (spath, onlytype=None)
     add_cluster_annotations (clusters)
     size
     coords
     types
     refs
     perform_clustering(method)
     perform_reclustering (method, skip_outliers=False, skip_size=None)
     recluster_cluster (method, cluster)
     recluster_outliers (method)
     small_clusters_to_outliers (maxsize)
     renumber_clusters()
     sort_clusters()
     clusters list
     clusters_centers
     clusters_size
     clusters std
     spaths2ctypes (*args, **kwargs)
     lim_to(what, towhat)
     lim2spaths (spaths)
     lim2types (types)
     lim2clusters (clusters)
     limspaths2(*args, **kwargs)
```

```
aqueduct.traj.paths module Created on Dec 10, 2015
@author: tljm
union (a, b)
glue(a, b)
xor (*args, **kwargs)
left(a, b)
right(a, b)
class PathTypesCodes
    path_in_code = 'i'
     path_object_code = 'c'
    path_out_code = '0'
class GenericPathTypeCodes
     object_name = 'c'
     scope_name = 's'
     out name = 'n'
class GenericPaths (id, min_pf=None, max_pf=None)
     Bases: object, aqueduct.traj.paths.GenericPathTypeCodes
     __init___(id, min_pf=None, max_pf=None)
     add_coord(coord)
     add_object (frame)
     add_scope (frame)
     add_type (frame, ftype)
    max_frame
    min_frame
     get_paths_in()
     get_paths_out()
     get_paths_for_frames_range(*args, **kwargs)
     find_paths (fullonly=False)
     find_paths_coords (fullonly=False)
     find_paths_types (fullonly=False)
     find_paths_coords_types (fullonly=False)
     get_single_path_coords(spath)
     get_single_path_types (spath)
    barber_with_spheres (spheres)
class SinglePathID (id=None, nr=None)
    Bases: object
     ___init___(id=None, nr=None)
     __str__()
```

```
yield_single_paths (gps, fullonly=False, progress=False)
class SinglePath (id, paths, coords, types)
    Bases:
                                                aqueduct.traj.paths.PathTypesCodes,
    aqueduct.traj.inlets.InletTypeCodes
    empty_coords = array([], shape=(0, 3), dtype=float64)
    ___init___(id, paths, coords, types)
    coords_first_in
    coords_last_out
    coords_filo
    get_inlets()
    coords
    coords_cont
    paths
    paths_cont
    types
    types_cont
    gtypes
    gtypes_cont
    etypes
    etypes_cont
    size
    begins
    ends
    has_in
    has_object
    has_out
    get_coords (*args, **kwargs)
    get_coords_cont (smooth=None)
    _make_smooth_coords (*args, **kwargs)
    apply_smoothing(smooth)
    get_distance_cont (smooth=None, normalize=False)
    get_distance_rev_cont(*args, **kwargs)
    get_distance_both_cont(*args, **kwargs)
    get_velocity_cont (*args, **kwargs)
    get_acceleration_cont(*args, **kwargs)
class MasterPath (sp)
    Bases: aqueduct.traj.paths.SinglePath
    ___init___(sp)
    add_width (width)
```

```
aqueduct.traj.reader module Created on Nov 19, 2015
@author: tljm
class Reader (topology, trajectory)
     Bases: object
     ___init___(topology, trajectory)
     open_trajectory()
    number_of_frames
     set_current_frame (frame)
     next_frame()
     iterate_over_frames()
    parse_selection (selection)
     select_resnum(resnum)
     select_multiple_resnum(resnum_list)
class ReadViaMDA (topology, trajectory)
     Bases: aqueduct.traj.reader.Reader
     number_of_frames
     set_current_frame (frame)
    next_frame()
    parse_selection(selection)
     select_resnum(resnum)
     select_multiple_resnum(resnum_list)
     __enter__()
     __exit__(typ, value, traceback)
     open_trajectory()
class ReadAmberNetCDFviaMDA (topology, trajectory)
     Bases: aqueduct.traj.reader.ReadViaMDA
     open_trajectory()
class ReadDCDviaMDA (topology, trajectory)
     Bases: aqueduct.traj.reader.ReadViaMDA
     open_trajectory()
aqueduct.traj.selections module
class Selection
    Bases: object
     def __init__(self,selection,selection_string=None):
         self.selection_object = selection self.selection_string = selection_string
     center_of_mass()
     iterate_over_residues()
     unique_resids()
     unique_resids_number()
     atom_positions()
```

```
center_of_mass_of_residues()
    get_convexhull_of_atom_positions()
    uniquify()
    ___add___(other)
    first resid()
class SelectionMDA (atoms)
    Bases: MDAnalysis.core.AtomGroup.AtomGroup, aqueduct.traj.selections.Selection
    iterate over residues()
    unique_resids (ikwid=False)
    atom_positions()
    ___add___(other)
    uniquify()
class CompactSelectionMDA (sMDA)
    Bases: object
     __init__(sMDA)
    toSelectionMDA (reader)
Module contents
```

# aqueduct.utils package

#### **Submodules**

aqueduct.utils.clui module Module comprises convieniences functions and definitios for different operations related to command line user interface.

```
emit_message_to_file_in_root_logger(mess)
message (mess, cont=False)
```

Prints message to standard error. If FileHandler is present in the :py:'root\_logger' the same message is appended to the log file.

Unknown interpreted text role "py".

#### **Parameters**

- mess (str) message to print
- cont (bool) if set True no new line is printed

```
class fbm (info, cont=True)
     Bases: object
     __init__ (info, cont=True)
     __enter__()
     __exit__(typ, value, traceback)
     __call__(info)
```

gregorian\_year\_in\_days = 365.2425 Length of Gregorian year in days. Average value. Source: https://en.wikipedia.org/wiki/Year  $smart\_time\_string(s, rl=0, t=1.1, maximal\_length=None, maximal\_units=5)$ 

Function transforms time in seconds to nicely formatted string of length defined by maximal\_length. Depending on number of seconds time is represented with one or more of the following units:

Unit name	Unit abbreviation
seconds	S
minutes	m
hours	h
days	d
years	У

Maximal number of units used in time string can be set with maximal\_units.

#### **Parameters**

- **s** (*int*) Input time in seconds.
- **rl** (*int*) Number of units already used for representing time.
- t (float) Exces above standard number of current time units.
- maximal\_length (int) Maximal length of the output string. Must be greater then 0.
- maximal\_units (int) Maximal number of units used in the output string. Must be greater then 0 and lower then 6.

Returns string of nicely formated time

Return type str

**gsep** (*sep='-'*, *times=72*, *length=None*)
Generic separator.

#### **Parameters**

- sep(str) Element(s) of separator.
- times (int) Number of times sep is printed.
- **length** (*int*) Optional maximal length of output.

**Returns** String separator.

Return type str

tsep(line)

Parameters line (str) - Input line.

**Returns** Returns default gsep() of length of line.

underline (line)

**Parameters** line (str) – Input line.

**Returns** String made by concatenation of line, os.linesep, and output of tsep() called with line.

Return type str

thead(line)

**Parameters** line (str) – Input line.

**Returns** String made by concatenation of output of tsep() called with line, line, os.linesep, and again output of tsep() called with line.

Return type str

#### class SimpleProgressBar (maxval=None, mess=None)

Bases: object

Simple progress bar displaying progress with percent indicator, progress bar and ETA. Progress is measured by iterations.

#### Variables

- rotate (str) String comprising characters with frames of a rotating toy.
- barlenght (int) Length of progress bar.
- maxval (int) maximal number of iterations
- **current** (*int*) current number of iterations
- overrun\_notice (bool) if True, overrun above :ivar:'maxval' iterations causes insert of newline
- overrun (bool) flag of overrun
- **begin** (int) time in seconds at the initialization of the SimpleProgressBar class.
- tcurrent (int) time in seconds of current iteration

```
rotate = '\\\/-"
```

#### barlenght = 24

\_\_\_init\_\_\_(maxval=None, mess=None)

#### **Parameters**

- maxval (int) Maximal number of iterations stored to maxval.
- **mess** (str) Optional message displayed at progress bar initialization.

# bar()

# ETA()

Returns ETA calculated on the basis of current number of iterations current and current time tcurrent. If number of iterations is 0 returns?. Time is formated with <code>smart\_time\_string()</code>.

**Returns** ETA as string.

Return type str

### percent()

Returns float number of precent progress calculated in the basis of current number of iterations current. Should return number between 0 and 100.

**Returns** percent progress number

Return type float

#### show()

Shows current progress.

If value returned by percent() is =< 100 then progres is printed as percent indicator leaded by ETA calculated by ETA().

If value returned by percent() is > 100 then progress is printed as number of iterations and total time.

Progress bar is writen to standard error.

# heartbeat()

#### update (step)

Updates number of current iterations current by one if step is > 0. Otherwise number of current iterations is not updated. In boths cases time of current iteration tcurrent is updated and show() is called.

#### **Parameters** step (int) – update step

#### ttime()

Calculates and returns total time string formated with smart\_time\_string().

Returns string of total time

Return type str

#### finish()

Finishes progress bar. First, update() is called with step = 0. Next message of total time is writen to standard error.

#### pbar

alias of SimpleProgressBar

```
get_str_timestamp()
```

**aqueduct.utils.helpers module** Collection of helpers - functions and decorators.

#### combine (seqin)

This is an alien function. It is not extensively used.

Directly taken form http://aspn.activestate.com/ASPN/Cookbook/Python/Recipe/302478/index\_txt

Returns a list of all combinations of argument sequences. For example, following call:

```
combine(((1,2),(3,4)))
```

gives following list of combinations:

```
[[1, 3], [1, 4], [2, 3], [2, 4]]
```

**Parameters** sequin (tuple) – Tuple of sequences to combine.

**Returns** All possible combinations of all input sequences.

Return type list of lists

# $is_number(s)$

# lind(l, ind)

Indexes lists using lists of integers as identificators. For example:

```
lind(['a','b','c','d','e'],[1,4,2])
```

returns:

```
['b', 'e', 'c']
```

# **Parameters**

- 1 (list) List to be indexed.
- ind (list) Integer indexes.

Returns Reindexed list.

Return type list

#### class Auto

Auto type definition. The class is used as an alternative value for options (if particular option supports it). If options (or variables/parameters etc.) have value of Auto it means that an automatic process for parametrization should be performed.

For example, if the input parameter is set to Auto it is supposed that its value is calculated on the basis of input data or other parameters.

```
__repr__()
```

Returns String Auto.

Return type str

#### create tmpfile(ext=None)

Creates temporary file. File is created, closed and its file name is returned.

**Note:** It is responsibility of the caller to delete the file.

**Parameters** ext (str) – Optional extension of the file.

**Returns** File name of created temporary file.

Return type str

# range2int(r, uniq=True)

Transforms a string range in to a list of integers (with added missing elements from given ranges).

For example, a following string:

```
'0:2 4:5 7 9'
```

is transformed into:

```
[0,1,2,4,5,7,9]
```

# **Parameters**

- **r** (str) String of input range.
- uniq (bool) Optional parameter, if set to *True* only unique and sorted integers are returned.

Returns List of integers.

Return type list of int

#### int2range(l)

Transforms a list of integers in to a string of ranges.

For example, a following list:

```
[0,1,2,4,5,7,9]
```

is transformed into:

```
0:2 4:5 7 9
```

Parameters 1 (list) – input list of int

Returns String of ranges.

Return type str

### is iterable(l)

Checks if provided object is iterable. Returns True is it is iterable, otherwise returns False.

Parameters 1 (list) – input object

**Returns** True if submitted object is iterable otherwise returns False.

Return type bool

Warning: Current implementation cannot be used with generators!

#### Todo

Current implementation is primitive and HAVE TO be replaced.

#### sortify (gen)

Decorator to convert functions' outputs into a sorted list. If the output is iterable it is converted in to a list of appropriate length. If the output is not iterable it is converted in to a list of length 1.

Written on the basis of listify().

**Returns** Output of decorated function converted to a sorted list.

Return type list

#### uniqify (gen)

Decorator to convert functions' outputs into a sorted list of unique objects. If the output is iterable it is converted in to a list of appropriate length. If the output is not iterable it is converted in to a list of length 1.

Written on the basis of listify().

**Returns** Output of decorated function converted to a sorted list of unique objects.

Return type list

#### listify(gen)

Decorator to convert functions' outputs into a list. If the output is iterable it is converted in to a list of appropriate length. If the output is not iterable it is converted in to a list of length 1.

This function was copied from:

http://argandgahandapandpa.wordpress.com/2009/03/29/python-generator-to-list-decorator/

and further improved by tljm@wp.pl.

**Returns** Output of decorated function converted to a list.

Return type list

# tupleify(gen)

Decorator to convert functions' outputs into a tuple. If the output is iterable it is converted in to a tuple of appropriate length. If the output is not iterable it is converted in to a tuple of length 1.

Written on the basis of listify().

**Returns** Output of decorated function converted to a tuple.

Return type tuple

# arrayify(gen)

Decorator to convert functions' outputs into a 2D numpy array. If the output is iterable it is converted in to a 2D numpy array of appropriate shape. If the output is not iterable it is converted in to a 2D numpy array of shape 1x1.

Written on the basis of listify().

**Returns** Output of decorated function converted to a 2D numpy array.

**Return type** numpy.ndarray

### arrayify1 (gen)

Decorator to convert functions' outputs into a 1D numpy array. If the output is iterable it is converted in to a 2D numpy array of appropriate shape. If the output is not iterable it is converted in to a 2D numpy array of shape 1x1.

Written on the basis of listify().

**Returns** Output of decorated function converted to a 1D numpy array.

Return type numpy.ndarray

```
list\_blocks\_to\_slices(l)
```

Slices list in to block according to its elements identity. Resulting slices correspond to blocks of identical elements.

**Parameters 1** (*list*) – List of any objects.

Returns Generator of slices.

Return type generator

```
what2what (what, towhat)
```

This function search if elements of the one list (:attr: 'what') are present in the other list (:attr: 'towhat') and returns indices of elements form :attr:'what' list as a tuple. If elements from the first list are not present in the second list the tuple is empty. :param list what: Input list for which indices of elements present in towhat are returned. :param list towhat: List of elements which input list is indexed to. :return: Indices of what list that are present in towhat list. :rtype: tuple

# make\_iterable(something)

If input object is not iterable returns it as one element list. Otherwise returns the object.

Parameters something (object) - Input object.

Returns Iterable object.

Return type iterable or list

```
strech_zip(*args)
compress_zip(*args)
zip_zip(*args, **kwargs)
```

aqueduct.utils.multip module Created on Feb 3, 2016

@author: tljm

**Module contents** 

aqueduct.visual package

**Submodules** 

aqueduct.visual.cmaps module

\_\_call\_\_\_(node)

```
aqueduct.visual.helpers module
cc_safe(c)
cc(c)
color_codes(code, custom_codes=None)
get_cmap(size)
class ColorMapDistMap(name='hsv', size=None)
    Bases: object
    default_cm_size = 256
    grey = (0.5, 0.5, 0.5, 1)
    __init__(name='hsv', size=None)
```

```
f_like(n)
aqueduct.visual.pymol_cgo module
aqueduct.visual.pymol_connector module
class BasicPymolCGO
    Bases: object
     cgo_entity_begin = []
     cgo_entity_end = []
     ___init___()
     clean()
    new()
    get()
class BasicPymolCGOLines
    Bases: aqueduct.visual.pymol_connector.BasicPymolCGO
     cgo_entity_begin = [2.0, 1.0]
     cgo_entity_end = [3.0]
     add (coords=None, color=None)
class BasicPymolCGOSpheres
    Bases: aqueduct.visual.pymol_connector.BasicPymolCGO
     cgo_entity_begin = []
     cgo_entity_end = []
     add (coords=None, radius=None, color=None)
class BasicPymolCGOPointers
    Bases: aqueduct.visual.pymol_connector.BasicPymolCGO
     cgo_entity_begin = []
     cgo_entity_end = []
     add_cone (coords1=None, coords2=None, radius1=None, radius2=None, color1=None,
               color2=None)
     add_pointer (point=None, direction=None, length=None, color=None, reverse=False)
class SimpleTarWriteHelper
    Bases: object
    __init__()
     open (filename)
     save_object2tar(obj, name)
     save_file2tar (filename, name)
     __del__()
class ConnectToPymol
    Bases: object
     cgo_line_width = 2.0
    ct_pymol = 'pymol'
     ct_file = 'file'
     ___init___()
```

```
init_pymol()
     init_script (filename)
     add_cgo_object (name, cgo_object, state=None)
     del_cgo_object (name, state=None)
     load_pdb (name, filename, state=None)
     orient on(name)
     ___del___()
class SinglePathPlotter (pymol connector, linearize=None)
     Bases: object
     ___init___(pymol_connector, linearize=None)
     add_single_path_continous_trace(spath,
                                                         smooth=None,
                                                                             plot_in=True,
                                            plot_object=True, plot_out=True, **kwargs)
     paths_trace (spaths, smooth=None, name='paths', state=None, **kwargs)
     paths_inlets (spaths, smooth=None, color=None, plot_in=True, plot_out=True, name='in-out-
                     let', state=None, **kwargs)
     scatter(coords, radius=0.4, color='r', name='scatter', state=None)
     convexhull (chull, color='m', name='convexhull', state=None)
aqueduct.visual.quickplot module
yield_spath_len_and_smooth_diff_in_types_slices(sp,
                                                                           smooth=None,
                                                           smooth_len=None,
                                                           smooth_diff=None,
                                                           types='etypes')
plot_colorful_lines (x, y, c, **kwargs)
spaths_spectra(spaths, **kwargs)
plot_spath_spectrum(sp, **kwargs)
spath_spectrum(sp, **kwargs)
showit (gen)
get_ax3d(fig, sub=111)
class SimpleTracePlotter
     Bases: object
     plot_line (coords, color, **kwargs)
     single_trace (coords, color='r', **kwargs)
     path_trace(path, color=('r', 'g', 'b'), plot_in=True, plot_object=True, plot_out=True,
                   **kwargs)
class SimpleProteinPlotter
     Bases: aqueduct.visual.quickplot.SimpleTracePlotter
     protein_trace (protein, smooth=None, color=('c', 'm', 'y'), **kwargs)
class SimplePathPlotter
     Bases: aqueduct.visual.quickplot.SimpleTracePlotter
     single_path_traces (spaths, smooth=None, color=('r', 'g', 'b'), **kwargs)
class MPLTracePlotter
                                       aqueduct.visual.quickplot.SimplePathPlotter,
     Bases:
     aqueduct.visual.quickplot.SimpleProteinPlotter
```

```
init_ax (*args, **kwargs)
plot_line (*args, **kwargs)
scatter (*args, **kwargs)
```

#### **Module contents**

# 5.1.2 Module contents

Aqueduct - a collection of tools to trace residues in MD simulation.

#### version()

Returns aqueduct version number.

**Returns** 3 element tuple of int numbers

Return type tuple

#### version\_nice()

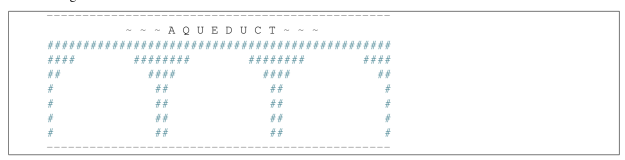
Returns aqueduct version number as nicely formatted string.

**Returns** string composed on the basis of the number returned by version().

Return type str

#### greetings()

Returns fancy greetings of aqueduct. It has a form of ASCII-like graphic. Currently it returns following string:



Returns aqueduct fancy greetings.

Return type str

# **CHAPTER**

# SIX

# **INDICES AND TABLES**

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