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**pyscal** is a python module for the calculation of local atomic structural environments including Steinhardt’s bond orientational order parameters\(^1\) during post-processing of atomistic simulation data. The core functionality of pyscal is written in C++ with python wrappers using **pybind11** which allows for fast calculations and easy extensions in python.

Steinhardt’s order parameters are widely used for the identification of crystal structures\(^3\). They are also used to distinguish if an atom is in a solid or liquid environment\(^4\). pyscal is inspired by the **BondOrderAnalysis** code, but has since incorporated many additional features and modifications. The pyscal module includes the following functionalities:

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• calculation of Steinhart’s order parameters and their averaged version\(^2\).
• links with the Voro++ code, for the calculation of Steinhart parameters weighted using the face areas of Voronoi polyhedra\(^3\).
• classification of atoms as solid or liquid\(^4\).
• clustering of particles based on a user defined property.
• methods for calculating radial distribution functions, Voronoi volumes of particles, number of vertices and face area of Voronoi polyhedra, and coordination numbers.
• calculation of angular parameters to identify diamond structure\(^5\).

CHAPTER 2

Getting started

For general documentation and examples see here.
3.1 pyscal reference

3.1.1 pyscal Reference

pyscal.core module

Main module of pyscal. This module contains definitions of the two major classes in pyscal - the System and Atom. Atom is a pure pybind11 class whereas System is a hybrid class with additional python definitions. For the ease of use, Atom class should be imported from the core module. The original pybind11 definitions of Atom and System can be found in catom and csystem respectively.

class pyscal.core.System
    Bases: pyscal.csystem.System

    A python/pybind11 hybrid class for holding the properties of a system.

    box
        A list containing the dimensions of the simulation box in the format [[x_low, x_high], [y_low, y_high], [z_low, z_high]]
        
        Type  list of list of floats

    atoms

    Type  list of Atom objects

Notes

A System consists of two major components - the simulation box and the atoms. All the associated variables are then calculated using this class.
**Note:** atoms can be accessed or set as *atoms*. However, due to technical reasons individual atoms should be accessed using the *get_atom()* method. An atom can be assigned to the atom using the *set_atom()* method.

**Examples**

```python
>>> sys = System()
>>> sys.read_inputfile('atoms.dat')
```

**atoms**

Atom access

**box**

Wrap for inbuilt box

**calculate_angularcriteria()**

Calculate the angular criteria for each atom :param None:

**Returns**

**Return type** None

**Notes**

Calculates the angular criteria for each atom as defined in [1]. Angular criteria is useful for identification of diamond cubic structures. Angular criteria is defined by,

\[ A = \sum_{i=1}^{6} \left( \cos(\theta_i) + \frac{1}{3} \right)^2 \]

where \( \cos(\theta) \) is the angle size suspended by each pair of neighbors of the central atom. \( A \) will have a value close to 0 for structures if the angles are close to 109 degrees. The calculated \( A \) parameter for each atom is stored in *angular*.

**References**

**calculate_centrosymmetry** (*nmax=12, get_vals=True*)

Calculate the centrosymmetry parameter

**Parameters** *nmax* (*int, optional*) – number of neighbors to be considered for centrosymmetry parameters. Has to be a positive, even integer. Default 12

**Returns**

**Return type** None

**Notes**

Calculate the centrosymmetry parameter for each atom which can be accessed by *centrosymmetry* attribute. It calculates the degree of inversion symmetry of an atomic environment. Centrosymmetry recalculates the neighbor using the number method as specified in *pyscal.core.System.find_neighbors()* method. This is the ensure that the required number of neighbors are found for calculation of the parameter.

References

calculate_chiparams (angles=False)
Calculate the chi param vector for each atom

Parameters angles (bool, optional) – If True, return the list of cosines of all neighbor pairs

Returns angles – list of all cosine values, returned only if angles is True.

Return type array of floats

Notes

This method tries to distinguish between crystal structures by finding the cosines of angles formed by an atom with its neighbors. These cosines are then histogrammed with bins [-1.0, -0.945, -0.915, -0.755, -0.705, -0.195, 0.195, 0.245, 0.795, 1.0] to find a vector for each atom that is indicative of its local coordination. Compared to chi parameters from chi_0 to chi_7 in the associated publication, the vector here is from chi_0 to chi_8. This is due to an additional chi parameter which measures the number of neighbors between cosines -0.705 to -0.195.

Parameter nlimit specifies the number of nearest neighbors to be included in the analysis to find the cutoff. If parameter angles is true, an array of all cosine values is returned. The publication further provides combinations of chi parameters for structural identification which is not implemented here. The calculated chi params can be accessed using chiparams.

References

calculate_cna (lattice_constant=None)
Calculate the Common Neighbor Analysis indices

Parameters lattice_constant (float, optional) – lattice constant to calculate CNA. If not specified, adaptive CNA will be used

Returns cna – dict containing the cna signature of the system

Return type dict

Notes

Performs the common neighbor analysis [1][2] or the adaptive common neighbor analysis [2] and assigns a structure to each atom.

If lattice_constant is specified, a conventional common neighbor analysis is used. If lattice_constant is not specified, adaptive common neighbor analysis is used. The assigned structures can be accessed by structure. The values assigned for structure are 0 Unknown, 1 fcc, 2 hcp, 3 bcc, 4 icosahedral.

References

calculate_disorder (averaged=False, q=6)
Calculate the disorder criteria for each atom
Parameters

- `averaged (bool, optional)` – If True, calculate the averaged disorder. Default False.

- `q (int, optional)` – The Steinhardt parameter value over which the bonds have to be calculated. Default 6.

Returns

Return type None

Notes

Calculate the disorder criteria as introduced in [1]. The disorder criteria value for each atom is defined by,

\[ D_j = \frac{1}{N_b} \sum_{i=1}^{N_b} [S_{jj} + S_{kk} - 2S_{jk}] \]

where \( S_{ij} = \sum_{m=-6}^{6} q_{6m}(i) q_{6m}^*(i) \)

The keyword `averaged` is True, the disorder value is averaged over the atom and its neighbors. The disorder value can be accessed using `disorder` and the averaged version can be accessed using `avg_disorder`.

For ordered systems, the value of disorder would be zero which would increase and reach one for disordered systems.

References

`calculate_energy (species='Au', pair_style=None, pair_coeff=None, mass=1.0, averaged=False)`

Calculate the potential energy of atom using LAMMPS

Parameters

- `species (str)` – Name of atomic species

- `pair_style (str)` – lammps pair style

- `pair_coeff (str)` – lammps pair coeff

- `mass (float)` – mass of the atoms

- `averaged (bool, optional)` – Average the energy over neighbors if True default False.

Returns

Return type None

Notes

Calculates the potential energy per atom using the given potential through LAMMPS. More documentation coming up...

Values can be accessed through `pyscal.catom.Atom.energy` and `avg_energy`.

If `averaged` is True, the energy is averaged over the neighbors of an atom. If neighbors were calculated before calling this method, those neighbors are used for averaging. Otherwise neighbors are calculated on the fly with an adaptive cutoff method.
**calculate_entropy** *(rm, sigma=0.2, rstart=0.001, h=0.001, local=False, M=12, N=6, ra=None, averaged=False, switching_function=False)*

Calculate the entropy parameter for each atom

**Parameters**

- **rm** *(float)* – cutoff distance for integration of entropy parameter in distance units
- **sigma** *(float)* – broadening parameter
- **rstart** *(float, optional)* – minimum limit for integration, default 0.00001
- **h** *(float, optional)* – width for trapezoidal integration, default 0.0001
- **local** *(bool, optional)* – if True, use the local density instead of global density, default False
- **averaged** *(bool, optional)* – if True find the averaged entropy parameters, default False
- **switching_function** *(bool, optional)* – if True, use the switching function to average, otherwise do a simple average over the neighbors. Default False
- **ra** *(float, optional)* – cutoff length for switching function used only if switching_function is True
- **M** *(int, optional)* – power for switching function, default 12 used only if switching_function is True
- **N** *(int, optional)* – power for switching function, default 6 used only if switching_function is True

**Returns**

**Return type** None

**Notes**

The entropy parameters can be accessed by `entropy` and `avg_entropy`. For a complete description of the entropy parameter, see the documentation.

The `local` keyword can be used to use a local density instead of the global one. This method will only work with neighbor methods that use a cutoff.

**calculate_q** *(q, averaged=False, only_averaged=False, condition=None, clear_condition=False)*

Find the Steinhardt parameter q_l for all atoms.

**Parameters**

- **q** *(int or list of ints)* – A list of all Steinhardt parameters to be found from 2-12.
- **averaged** *(bool, optional)* – If True, return the averaged q values, default False
- **only_averaged** *(bool, optional)* – If True, only calculate the averaged part. Default False
- **condition** *(callable or atom property)* – Either function which should take an Atom object, and give a True/False output or an attribute of atom class which has value 0 or 1.
- **clear_condition** *(bool, optional)* – clear the condition variable for all atoms

**Returns**
Return type  None

Notes

Enables calculation of the Steinhardt parameters [1] q from 2-12. The type of q values depend on the method used to calculate neighbors. See the description find_neighbors() for more details. If the keyword average is set to True, the averaged versions of the bond order parameter [2] is returned. If only the averaged versions need to be calculated, only_averaged keyword can be set to False.

The neighbors over which the q values are calculated can also be filtered. This is done through the argument condition which is passed as a parameter. condition can be of two types. The first type is a function which takes an Atom object and should give a True/False value. condition can also be an Atom attribute or a value from custom values stored in an atom. See cluster_atoms() for more details. If the condition is equal for both host atom and the neighbor, the neighbor is considered for calculation of q parameters. This is slightly different from cluster_atoms() where the condition has to be True for both atoms. condition is only cleared when neighbors are recalculated. Additionally, the keyword clear_condition can also be used to clear the condition and reset it to 0. By default, condition is applied to both unaveraged and averaged q parameter calculation. If condition is needed for only averaged q parameters, this function can be called twice, initially without condition and averaged=False, and then with a condition specified and averaged=True. This way, the condition will only be applied to the averaged q calculation.

References

**calculate_rdf** (histobins=100, histomin=0.0, histomax=None)
Calculate the radial distribution function.

Parameters

- histobins (int) – number of bins in the histogram
- histomin (float, optional) – minimum value of the distance histogram. Default 0.0.
- histomax (float, optional) – maximum value of the distance histogram. Default, the maximum value in all pair distances is used.

Returns

- rdf (array of ints) – Radial distribution function
- r (array of floats) – radius in distance units

**calculate_solidneighbors()**
Find Solid neighbors of all atoms in the system.

Parameters  None –

Returns

Return type  None

Notes

A solid bond is considered between two atoms if the connection between them is greater than 0.6.

**calculate_sro** (reference_type=1, average=True, shells=2)
Calculate short range order

Parameters
• `reference_type (int, optional)` – type of the atom to be used a reference. default 1

• `average (bool, optional)` – if True, average over all atoms of the reference type in the system. default True.

**Returns vec** – The short range order averaged over the whole system for atom of the reference type. Only returned if `average` is True. First value is SRO of the first neighbor shell and the second value corresponds to the second nearest neighbor shell.

**Return type** list of float

**Notes**

Calculates the short range order for an AB alloy using the approach by Cowley [1]. Short range order is calculated as,

\[ \alpha_i = 1 - \frac{n_i}{m_A c_i} \]

where \( n_i \) is the number of atoms of the non reference type among the \( c_i \) atoms in the \( i \)th shell. \( m_A \) is the concentration of the non reference atom. Please note that the value is calculated for shells 1 and 2 by default. In order for this to be possible, neighbors have to be found first using the `find_neighbors()` method. The selected neighbor method should include the second shell as well. For this purpose `method=cutoff` can be chosen with a cutoff long enough to include the second shell. In order to estimate this cutoff, one can use the `calculate_rdf()` method.

**References**

- `calculate_vorovector (edge_cutoff=0.05, area_cutoff=0.01, edge_length=False)`
  get the voronoi structure identification vector.

  **Parameters**

  - `edge_cutoff (float, optional)` – cutoff for edge length. Default 0.05.
  - `area_cutoff` [float, optional] cutoff for face area. Default 0.01.
  - `edge_length` [bool, optional] if True, a list of unrefined edge lengths are returned. Default false.

  **Returns**

  - `vorovector` – array of the form (n3, n4, n5, n6)
  - **Return type** array like, int

  **Notes**

  Returns a vector of the form \((n_3, n_4, n_5, n_6)\), where \(n_3\) is the number of faces with 3 vertices, \(n_4\) is the number of faces with 4 vertices and so on. This can be used to identify structures [1] [2].

  The keywords `edge_cutoff` and `area_cutoff` can be used to tune the values to minimise the effect of thermal distortions. Edges are only considered in the analysis if the `edge_length/sum(edge_lengths)` is at least `edge_cutoff`. Similarly, faces are only considered in the analysis if the `face_area/sum(face_areas)` is at least `face_cutoff`. 
References

**cluster_atoms** *(condition, largest=True, cutoff=0)*
Cluster atoms based on a property

**Parameters**

- **condition** *(callable or atom property)* – Either function which should take an Atom object, and give a True/False output or an attribute of atom class which has value or 1 or 0.
- **largest** *(bool, optional)* – If True returns the size of the largest cluster. Default False.
- **cutoff** *(float, optional)* – If specified, use this cutoff for calculation of clusters. By default uses the cutoff used for neighbor calculation.

**Returns**

- **lc** – Size of the largest cluster. Returned only if largest is True.
- **Return type** int

**Notes**

This function helps to cluster atoms based on a defined property. This property is defined by the user through the argument *condition* which is passed as a parameter. *condition* can be of two types. The first type is a function which takes an Atom object and should give a True/False value. *condition* can also be an Atom attribute or a value from custom values stored in an atom.

When clustering, the code loops over each atom and its neighbors. If the *condition* is true for both host atom and the neighbor, they are assigned to the same cluster. For example, a condition to cluster solid atoms would be,

```python
def condition(atom):
    # if both atom is solid
    return (atom1.solid)
```

The same can be done by passing “solid” as the condition argument instead of the above function. Passing a function allows to evaluate complex conditions, but is slower than passing an attribute.

**embed_in_cubic_box()**
Embedded the triclinic box in a cubic box

**find_diamond_neighbors()**
Find underlying fcc lattice in diamond

**Parameters** None –

**Returns**

**Return type** None

**Notes**

This method finds in the underlying fcc/hcp lattice in diamond. It works by the method described in this publication. For each atom, 4 atoms closest to it are identified. The neighbors of the its neighbors are further identified and the common neighbors shared with the host atom are selected. These atom will fall in the underlying fcc lattice for cubic diamond or hcp lattice for hexagonal lattice.

If neighbors are previously calculated, they are reset when this method is used.
find_largestcluster()

Find the largest solid cluster of atoms in the system from all the clusters.

Parameters None –

Returns cluster – the size of the largest cluster

Return type int

Notes

pyscal.core.System.find_clusters() has to be used before using this function.

find_neighbors (method='cutoff', cutoff=None, threshold=2, filter=None, voroexp=1, padding=1.2, nlimit=6, cells=False, nmax=12, assign_neighbor=True)

Find neighbors of all atoms in the System.

Parameters method ({'cutoff', 'voronoi', 'number'}) –
cutoff method finds neighbors of an atom within a specified or adaptive cutoff distance from the atom. voronoi method finds atoms that share a Voronoi polyhedra face with the atom. Default, cutoff number method finds a specified number of closest neighbors to the given atom. Number only populates

cutoff [[float, ‘sann’, ‘adaptive’]] the cutoff distance to be used for the cutoff based neighbor calculation method described above. If the value is specified as 0 or adaptive, adaptive method is used. If the value is specified as sann, sann algorithm is used.

threshold [float, optional] only used if cutoff=adaptive. A threshold which is used as safe limit for calculation of cutoff.

filter [{'None', 'type'}, optional] apply a filter to nearest neighbor calculation. If the filter keyword is set to type, only atoms of the same type would be included in the neighbor calculations. Default None.

toroexp [int, optional] only used if method=voronoi. Power of the neighbor weight used to weight the contribution of each atom towards Steinhardt parameter values. Default 1.

padding [double, optional] only used if cutoff=adaptive or cutoff=number. A safe padding value used after an adaptive cutoff is found. Default 1.2.

nlimit [int, optional] only used if cutoff=adaptive. The number of particles to be considered for the calculation of adaptive cutoff. Default 6.

nmax [int, optional] only used if cutoff=number. The number of closest neighbors to be found for each atom. Default 12

Returns

Return type None

Raises

- RuntimeError – raised when threshold value is too low. A low threshold value will lead to ‘sann’ algorithm not converging when finding a neighbor. This function will try to automatically increase threshold and check again.

- RuntimeWarning – raised when neighbor search was unsuccessful. This is due to a low threshold value.
Notes

This function calculates the neighbors of each particle. There are several ways to do this. A complete
description of the methods can be found here.

Method cutoff and specifying a cutoff radius uses the traditional approach being the one in which the
neighbors of an atom are the ones that lie in the cutoff distance around it.

In order to reduce time during the distance sorting during the adaptive methods, pyscal sets an initial guess
for a cutoff distance. This is calculated as,

\[ r_{\text{initial}} = threshold \times \left( \frac{\text{simulation box volume}}{\text{number of particles}} \right)^{1/3} \]

threshold is a safe multiplier used for the guess value and can be set using the `threshold` keyword.

In Method cutoff, if `cutoff='adaptive'`, an adaptive cutoff is found during runtime for each atom
[1]. Setting the cutoff radius to 0 also uses this algorithm. The cutoff for an atom i is found using,

\[ r_c(i) = padding \times \left( \frac{1}{\text{nlimit}} \right) \sum_{j=1}^{\text{nlimit}} (r_{ij}) \]

padding is a safe multiplier to the cutoff distance that can be set through the keyword `padding`. `nlimit`
keyword sets the limit for the top nlimit atoms to be taken into account to calculate the cutoff radius.

In Method cutoff, if `cutoff='sann'`, sann algorithm is used [2]. There are no parameters to tune sann
algorithm.

The second approach is using Voronoi polyhedra which also assigns a weight to each neighbor in the ratio
of the face area between the two atoms. Higher powers of this weight can also be used [3]. The keyword
`voroexp` can be used to set this weight.

If method os `number`, instead of using a cutoff value for finding neighbors, a specified number of closest
atoms are found. This number can be set through the argument `nmax`.

**Warning:** Adaptive and number cutoff uses a padding over the initial guessed “neighbor distance”. By
default it is 2. In case of a warning that `threshold` is inadequate, this parameter should be further
increased. High/low value of this parameter will correspond to the time taken for finding neighbors.

References

`find_solids` (bonds=0.5, threshold=0.5, avgthreshold=0.6, cluster=True, q=6, cutoff=0, 
right=True)

Distinguish solid and liquid atoms in the system.

**Parameters**

- **bonds** *(int or float, optional)* – Minimum number of solid bonds for an atom
to be identified as a solid if the value is an integer. Minimum fraction of neighbors of an
atom that should be solid for an atom to be solid if the value is float between 0-1. Default
0.5.

- **threshold** *(double, optional)* – Solid bond cutoff value. Default 0.5.

- **avgthreshold** *(double, optional)* – Value required for Averaged solid bond cut-
off for an atom to be identified as solid. Default 0.6.

- **cluster** *(bool, optional)* – If True, cluster the solid atoms and return the number
of atoms in the largest cluster.
• \( q(int, \text{optional}) \) – The Steinhardt parameter value over which the bonds have to be calculated. Default 6.

• \( \text{cutoff}(double, \text{optional}) \) – Separate value used for cluster classification. If not specified, cutoff used for finding neighbors is used.

• \( \text{right}(bool, \text{optional}) \) – If true, greater than comparison is to be used for finding solid particles. default True.

Returns \( \text{solid} \) – Size of the largest solid cluster. Returned only if \( \text{cluster=True} \).

Return type \( \text{int} \)

Notes

The neighbors should be calculated before running this function. Check \( \text{find_neighbors()} \) method.

\( \text{bonds} \) define the number of solid bonds of an atom to be identified as solid. Two particles are said to be 'bonded' if [1],

\[
s_{ij} = \sum_{m=-6}^{6} q_{6m}(i)q_{6m}^{*}(i) \geq \text{threshold}
\]

where \( \text{threshold} \) values is also an optional parameter.

If the value of \( \text{bonds} \) is a fraction between 0 and 1, at least that much of an atom’s neighbors should be solid for the atom to be solid.

An additional parameter \( \text{avgthreshold} \) is an additional parameter to improve solid-liquid distinction. In addition to having a the specified number of \( \text{bonds} \),

\[
\langle s_{ij} \rangle > \text{avgthreshold}
\]

also needs to be satisfied. In case another \( q \) value has to be used for calculation of \( S_{ij} \), it can be set used the \( q \) attribute. In the above formulations, > comparison for \( \text{threshold} \) and \( \text{avgthreshold} \) can be changed to < by setting the keyword \( \text{right} \) to False.

If \( \text{cluster} \) is True, a clustering is done for all solid particles. See \( \text{find_clusters()} \) for more details.

References

\( \text{get_atom}(\text{index}) \)

Get the \( \text{Atom} \) object at the queried position in the list of all atoms in the \( \text{System} \).

Parameters \( \text{index}(int) \) – index of required atom in the list of all atoms.

Returns \( \text{atom} \) – atom object at the queried position.

Return type \( \text{Atom object} \)

\( \text{get_custom}(\text{atom, customkeys}) \)

Get a custom attribute from Atom

Parameters

• \( \text{atom}(\text{Atom object}) \) –

• \( \text{customkeys}(\text{list of strings}) \) – the list of keys to be found

Returns \( \text{vals} \) – array of custom values
Return type  list

get_distance (atom1, atom2, vector=False)
Get the distance between two atoms.

Parameters
  • atom1 (Atom object) – first atom
  • atom2 (Atom object) – second atom
  • vector (bool, optional) – If True, the displacement vector connecting the atoms
    is also returned. default false.

Returns distance – distance between the first and second atom.

Return type  double

Notes
Periodic boundary conditions are assumed by default.

get_qvals (q, averaged=False)
Get the required q_l (Steinhardt parameter) values of all atoms.

Parameters
  • q_l (int or list of ints) – required q_l value with l from 2-12
  • averaged (bool, optional) – If True, return the averaged q values, default False

Returns qvals – list of q_l of all atoms.

Return type  list of floats

Notes
The function returns a list of q_l values in the same order as the list of the atoms in the system.

identify_diamond (find_neighbors=True)
Identify diamond structure

Parameters find_neighbors (bool, optional) – If True, find 4 closest neighbors

Returns diamondstructure – dict of structure signature

Return type  dict

Notes
Identify diamond structure using the algorithm mentioned in [1]. It is an extended CNA method. The
integers 5, 6, 7, 8, 9 and 10 are assigned to the structure variable of the atom. 5 stands for cubic diamond,
6 stands for first nearest neighbors of cubic diamond and 7 stands for second nearest neighbors of cubic
diamond. 8 signifies hexagonal diamond, the first nearest neighbors are marked with 9 and second nearest
neighbors with 10.
References

read_inputfile (filename, format='lammps-dump', compressed=False, customkeys=None)
Read input file that contains the information of system configuration.

Parameters

- **filename (string)** – name of the input file.
- **format (list) (default: ['lammps-dump', 'poscar', 'ase', 'mdtraj'])** – format of the input file, in case of *ase* the ASE Atoms object
- **compressed (bool, optional)** – If True, force to read a gz compressed format, default False.
- **customkeys (list)** – A list containing names of headers of extra data that needs to be read in from the input file.

Returns

Return type: None

Notes

format keyword specifies the format of the input file. Currently only a *lammps-dump* and *poscar* files are supported. Additionally, the widely use Atomic Simulation environment (https://wiki.fysik.dtu.dk/ase/ase.html). mdtraj objects (http://mdtraj.org/1.9.3/) are also supported by using the keyword *mdtraj* for format. Please note that triclinic boxes are not yet supported for mdtraj format. Atoms object can also be used directly. This function uses the *traj_process()* module to process a file which is then assigned to system.

compressed keyword is not required if a file ends with .gz extension, it is automatically treated as a compressed file.

Triclinic simulation boxes can also be read in.

If custom_keys are provided, this extra information is read in from input files if available. This information is not passed to the C++ instance of atom, and is stored as a dictionary. It can be accessed directly as atom.custom['customval']

repeat (reps, atoms=None, ghost=False, scale_box=True)
Replicate simulation cell

Parameters

- **reps (list of ints of size 3)** – repetitions in each direction
- **atoms (list of atoms, optional)** – if not provided, use atoms that are assigned
- **ghost (bool, optional)** – If True, assign the new atoms as ghost instead of actual atoms

reset_neighbors ()
Reset the neighbors of all atoms in the system.

Parameters None –

Returns

Return type None
Notes

It is used automatically when neighbors are recalculated.

set_atom(\texttt{atom})

Return the atom to its original location after modification.

Parameters

\texttt{atom} (\texttt{ Atom}) – atom to be replaced

Returns

None

Notes

For example, an \texttt{ Atom} at location \texttt{i} in the list of all atoms in \texttt{ System} can be queried by, \texttt{atom = System.get_atom(i)}, then any kind of modification, for example, the position of the \texttt{Atom} can done by, \texttt{atom.pos = [2.3, 4.5, 4.5]}. After modification, the \texttt{Atom} can be set back to its position in \texttt{ System} by \texttt{set_atom()}. Although the complete list of atoms can be accessed or set using \texttt{atoms = sys.atoms, get_atom} and \texttt{set_atom} functions should be used for accessing individual atoms. If an atom already exists at that index in the list, it will be overwritten and will lead to loss of information.

set_atom_cutoff(\texttt{factor=1.0})

Set cutoff for each atom

Parameters

\texttt{factor} (float, optional) – factor for multiplication of cutoff value. default 1

Returns

None

Notes

Assign cutoffs for each atom based on the nearest neighbor distance. The cutoff assigned is the average nearest neighbor distance multiplied by \texttt{factor}.

show(\texttt{colorby=None, filterby=None})

Plot the system

Parameters

- \texttt{sys} (\texttt{ System object}) –

- \texttt{colorby} (string, optional) – property over which the atoms are to be colored. It can be any attributed of \texttt{Atom}, a custom attribute, or calculated q values which can be accessed as \texttt{qx} or \texttt{aqx} where \texttt{x} stands for the \texttt{q} number.

- \texttt{filterby} (string, optional) – property over which the atoms are to be filtered before plotting. It can be any attribute of atom, or a custom value of atom. It should provide a True or False value.

Returns

None

Notes

to_file(\texttt{outfile, format='lammps-dump', customkeys=None, customvals=None, compressed=False, timestep=0, species=None})

Save the system instance to a trajectory file.
Parameters

- **outfile** *(string)* – name of the output file

- **format** *(string, {'lammps-dump', 'lammps-data', 'poscar'})* – format of the output file, default *lammps-dump* Currently only *lammps-dump* format is supported.

- **customkeys** *(list of strings, optional)* – a list of extra atom wise values to be written in the output file.

- **customvals** *(list or list of lists, optional)* – If customkey is specified, customvals take an array of the same length as number of atoms, which contains the values to be written out.

- **compressed** *(bool, optional)* – If true, the output is written as a compressed file.

- **timestep** *(int, optional)* – timestep to be written to file. default 0

- **species** *(None, optional)* – species of the atoms. Required if any format other than ‘lammps-dump’ is used. Required for conversion to ase object.

Returns

Return type None

Notes

to_file method can handle a number of file formats. The most customizable format is the *lammps-dump* which can take a custom options using customkeys and customvals. customkeys will be the header written to the dump file. It can be any Atom attribute, any property stored in custom variable of the Atom, or calculated q values which can be given by q4, aq4 etc. External values can also be provided using customvals option. customvals array should be of the same length as the number of atoms in the system.

For all other formats, ASE is used to write out the file, and hence the species keyword needs to be specified. If initially, an ASE object was used to create the System, species keyword will already be saved, and need not be specified. In other cases, species should be a list of atomic species in the System. For example [“Cu”] or [“Cu”, “Al”], depending on the number of species in the System. In the above case, atoms of type 1 will be mapped to Cu and of type 2 will be mapped to Al. For a complete list of formats that ASE can handle, see here.

    pyscal.core.test()
    A simple function to test if the module works

Parameters None –

Returns works – True if the module works and could create a System and Atom object False otherwise.

Return type bool

class pyscal.catom.Atom
    Bases: pybind11_builtins.pybind11_object

Class to store atom details.

Parameters

- **pos** *(list of floats of length 3)* – position of the Atom, default [0,0,0]

- **id** *(int)* – id of the Atom, default 0

- **type** *(int)* – type of the Atom, default 1

3.1. pyscal reference
Notes

A pybind11 class for holding the properties of a single atom. Various properties of the atom can be accessed through the attributes and member functions which are described below in detail. Atoms can be created individually or directly by reading a file. Check the examples for more details on how atoms are created. For creating atoms directly from an input file check the documentation of `System` class.

Although an `Atom` object can be created independently, `Atom` should be thought of inherently as members of the `System` class. All the properties that define an atom are relative to the parent class. `System` has a list of all atoms. All the properties of an atom, hence should be calculated through `System`.

Examples

```python
>>> #method 1 - individually
>>> atom = Atom()
>>> #now set positions of the atoms
>>> atom.pos = [23.0, 45.2, 34.2]
>>> #now set id
>>> atom.id = 23
>>> #now set type
>>> atom.type = 1
>>> #Setting through constructor
>>> atom = Atom([23.0, 45.2, 34.2], 23, 1)
```

References

Creation of atoms.

`allaq`

`list of floats`. list of all averaged q values of the atom.

`allq`

`list of floats`. list of all q values of the atom.

`angular`

`Float`. The value of angular parameter A of an atom. The angular parameter measures the tetrahedral coordination of an atom. Meaningful values are only returned if the property is calculated using `calculate_angularcriteria()`.

`avg_angular`

`Float`. The average angular parameter value. Not used currently.

`avg_disorder`

`Float`. The value of averaged disorder parameter.

`avg_energy`

`Float`. Value of averaged energy.

`avg_entropy`

`Float`. Value of averaged entropy parameter.

`avg_sij`

`float`. Value of averaged s_ij which is used for identification of solid atoms. s_ij is defined by

\[
s_{ij} = \sum_{m=-l}^{l} q_{lm}(i) q_{lm}^*(i)
\]
avg_volume
    float. Averaged version of the Voronoi volume which is calculated as an average over itself and its neighbors. Only calculated when the find_neighbors() using the method=’voronoi’ option is used.

bonds

centrosymmetry
    Float. The value of centrosymmetry parameter.

chiparams
    Float. The value of chiparameter of an atom. The return value is a vector of length 8. Meaningful values are only returned if chi params are calculated using calculate_chiparams().

cluster
    int. identification number of the cluster that the atom belongs to.

cna

common

condition
    int. condition that specifies if an atom is included in the clustering algorithm or not. Only atoms with the value of condition=1 will be used for clustering in cluster_atoms().

coordination
    int. coordination number of the atom. Coordination will only be updated after neighbors are calculated using find_neighbors().

custom
    dict. dictionary specifying custom values for an atom. The module only stores the id, type and position of the atom. If any extra values need to be stored, they can be stored in custom using atom.custom = {“velocity”:12}. read_inputfile() can also read in extra atom information. By default, custom values are treated as string.

cutoff
    double. cutoff used for finding neighbors for each atom.

disorder
    Float. The value of disorder parameter.

edge_lengths
    list of floats. For each face, this vector contains the lengths of edges that make up the Voronoi polyhedra of the atom. Only calculated when the find_neighbors() using the method=’voronoi’ option is used.

energy
    Float. Value of energy.

entropy
    Float. Value of entropy parameter.

face_perimeters
    list of floats. List consisting of the perimeters of each Voronoi face of an atom. Only calculated when the find_neighbors() using the method=’voronoi’ option is used.

face_vertices
    list of floats. A list of the number of vertices shared between an atom and its neighbors. Only calculated when the find_neighbors() using the method=’voronoi’ option is used.

get_q()
    Calculate the steinhardt parameter q_l value.

Parameters
• \( \mathbf{q} \) (int or list of ints) – number of the required \( q_l \) - from 2-12
• \textbf{averaged} (bool, optional) – If True, return the averaged \( q \) values, If False, return the non averaged ones default False

Returns \( q_l \) – the value(s) of the queried Steinhardt parameter(s).

Return type  float or list of floats

Notes

Please check this \textbf{link} for more details about Steinhardts parameters and the averaged versions.

Meaningful values are only returned if \textit{calculate_q()} is used.

\textbf{get_qlm()}  
Get the \( q_{lm} \) values.

Parameters

• \( \mathbf{q} \) (int) – number of the required \( q_l \) - from 2-12
• \textbf{averaged} (bool, optional) – If True, return the averaged \( q_{lm} \) values, If False, return the non averaged ones default False

Returns

• \( q_{lm} \) (complex vector) – vector of complex numbers.
• Meaningful values are only returned if \textit{calculate_q()} is used.

\textbf{ghost}  
int. int specifying ghost status of the atom.

\textbf{id}  
int. Id of the atom.

\textbf{largest_cluster}  
bool. True if the atom belongs to the largest cluster, False otherwise. Largest cluster is only identified after using the \textit{cluster_atoms()} function.

\textbf{loc}  
int. indicates the position of the atom in the list of all atoms.

\textbf{local_angles}  
List of floats of length 2. List of longitude and colatitude of an atom to its neighbors.

\textbf{mask}  
bool. Mask variable for atom. If mask is true, the atom is ignored from calculations.

\textbf{neighbor_distance}  
List of floats. List of neighbor distances of the atom.

\textbf{neighbor_vector}  
List of floats of length 3. List of vectors connecting an atom to its neighbors.

\textbf{neighbor_weights}  
List of floats. Used to weight the contribution of each neighbor atom towards the value of Steinhardt's parameters. By default, each atom has a weight of 1 each. However, if \textit{find_neighbors()} is used with \textit{method=‘voronoi’}, each neighbor gets a weight proportional to the area shared between the neighboring atom and host atom.
neighbors

*List of ints.* List of neighbors of the atom. The list contains indices of neighbor atoms which indicate their position in the list of all atoms.

**next_neighbor_distances**

double. cutoff used for finding neighbors for each atom.

**next_neighbors**

double. cutoff used for finding neighbors for each atom.

**pos**

*List of floats of the type [x, y, z].* default [0, 0, 0]. Position of the atom.

**set_q()**

Set the value of steinhardt parameter q_l.

**Parameters**

- **q**(int or list of ints) – number of the required q_l - from 2-12
- **val**(float or list of floats) – value(s) of Steinhardt parameter(s).
- **averaged**(bool, optional) – If True, return the averaged q values, If False, return the non averaged ones default False

**Returns**

**Return type** None

**sij**

*float.* Value of s_ij which is used for identification of solid atoms. s_ij is defined by

\[
s_{ij} = \sum_{m=-l}^{l} q_{lm}(i) q_{lm}^*(i)
\]

**solid**

*bool.* True if the atom is solid, False otherwise. Solid atoms are only identified after using the `find_solids()` function.

**sro**

*Float.* The value of short range order parameter.

**structure**

*int.* Indicates the structure of atom. Not used currently.

**surface**

*bool.* True if the atom has at least one liquid neighbor, False otherwise. Surface atoms are only identified after using the `find_solids()` function.

**type**

*int.* int specifying type of the atom.

**vertex_numbers**

*list of floats.* For each Voronoi face of the atom, this values includes a List of vertices that constitute the face. Only calculated when the `find_neighbors()` using the `method='voronoi'` option is used.

**vertex_vectors**

*list of floats.* A list of positions of each vertex of the Voronoi polyhedra of the atom. Only calculated when the `find_neighbors()` using the `method='voronoi'` option is used.

**volume**

*float.* Voronoi volume of the atom. The Voronoi volume is only calculated if neighbors are found using the `find_neighbors()` using the `method='voronoi'` option.
**vorovector**

A vector of the form \((n_3, n_4, n_5, n_6)\) where \(n_3\) is the number of faces with 3 vertices, \(n_4\) is the number of faces with 4 vertices and so on. This can be used to identify structures [1][2]. Vorovector is calculated if the `calculate_vorovector()` method is used.

**References**

**pyscal.crystal_structures module**

pyscal module for creating crystal structures.

`pyscal.crystal_structures.make_crystal(structure, lattice_constant=1.0, repetitions=None, ca_ratio=1.633, noise=0)`

Create a basic crystal structure and return it as a list of `Atom` objects and box dimensions.

**Parameters**

- **structure** (str) – type of the crystal structure
- **lattice_constant** (float, optional) – lattice constant of the crystal structure, default 1
- **repetitions** (list of ints of len 3, optional) – repetitions of the unit cell in x, y and z directions. default \([1, 1, 1]\).
- **ca_ratio** (float, optional) – ratio of c/a for hcp structures, default 1.633
- **noise** (float, optional) – If provided add normally distributed noise with standard deviation `noise` to the atomic positions.

**Returns**

- **atoms** (list of `Atom` objects) – list of all atoms as created by user input
- **box** (list of list of floats) – list of the type \([x_{low}, x_{high}], [y_{low}, y_{high}], [z_{low}, z_{high}]\) where each of them are the lower and upper limits of the simulation box in x, y and z directions respectively.

**Examples**

```python
>>> atoms, box = make_crystal('bcc', lattice_constant=3.48, repetitions=[2,2,2])
>>> sys = System()
>>> sys.assign_atoms(atoms, box)
```

**pyscal.trajectory module**

`class pyscal.trajectory.Timeslice(trajectory, blocklist)`

Timeslice containing info about a single time slice

`to_file(outfile, mode='w')`

Get block as outputfile

**Parameters**

- **outfile** (str) – name of output file
• mode (string) – write mode to be used, optional default “w” write also can be “a” to append.

Returns

Return type  None

to_system (customkeys=None)
Get block as pyscal system

Parameters  blockno (int) – number of the block to be read, starts from 0

Returns  sys – pyscal System

Return type  System

class  pyscal.trajectory.Trajectory (filename)
Bases: object
A Trajectory class for LAMMPS

convert_data_to_lines (blockno)
Create lines from loaded data

Parameters  blockno (int) – number of the block to be read, starts from 0

Returns  data – list of lines

Return type  list of strs

get_block (blockno)
Get a block from the file as raw data

Parameters  blockno (int) – number of the block to be read, starts from 0

Returns  data – list of strings containing data

Return type  list

get_block_as_system (blockno, customkeys=None)
Get block as pyscal system

Parameters  blockno (int) – number of the block to be read, starts from 0

Returns  sys – pyscal System

Return type  System

get_blocks_to_file (fout, blocklist)
Get a series of blocks from the file as raw data

Parameters  blockno (int) – number of the block to be read, starts from 0

Returns  data – list of strings containing data

Return type  list

get_natom()  
Get number of atoms in the system

Parameters  None –

Returns  

Return type  None

get_nbblocks()  
Get number of blocks in the trajectory file
Parameters None –
Returns
Return type None

**get_nlines()**
Get total number of lines in the file

Parameters None –
Returns nlines – number of lines
Return type int

**load(blockno)**
Load the data of a block into memory as a dictionary of numpy arrays

Parameters blockno (int) – number of the block to be read, starts from 0
Returns
Return type None

**Notes**

When the data of a block is loaded, it is accessible through `Trajectory.data[x]`. This data can then be modified. When the block is written out, the modified data is written instead of existing one. But, loaded data is kept in memory until unloaded using `unload` method.

**unload(blockno)**
Unload the data that is loaded to memory using `load` method

Parameters blockno (int) – number of the block to be read, starts from 0
Returns
Return type None

**pyscal.traj_process module**

pyscal module containing methods for processing of a trajectory. Methods for reading of input files formats, writing of output files etc are provided in this module.

**pyscal.traj_process.read_file(filename, format='lammps-dump', compressed=False, customkeys=None)**

Read input file

Parameters

- filename (string) – name of the input file.
- format (['lammps-dump', 'poscar', 'ase', 'mdtraj']) – format of the input file, in case of ase the ASE Atoms object
- compressed (bool, optional) – If True, force to read a gz compressed format, default False.
- customkeys (list) – A list containing names of headers of extra data that needs to be read in from the input file.

Returns

Return type None
pyscal.traj_process.split_trajectory(infile, format='lammps-dump', compressed=False)

Read in a trajectory file and convert it to individual time slices.

**Parameters**

- `filename (string)` – name of input file
- `format (format of the input file)` – only lammps-dump is supported now.
- `compressed (bool, optional)` – force to read a gz zipped file. If the filename ends with .gz, use of this keyword is not necessary.

**Returns**

- `snaps` – a list of filenames which contain individual frames from the main trajectory.

**Return type**

list of strings

**Notes**

This is a wrapper function around split_traj_lammps_dump function.

pyscal.traj_process.write_file(sys, outfile, format='lammps-dump', compressed=False, customkeys=None, customvals=None, timestep=0, species=None)

Write the state of the system to a trajectory file.

**Parameters**

- `sys (System object)` – the system object to be written out
- `outfile (string)` – name of the output file
- `format (string, optional)` – format of the output file
- `compressed (bool, default false)` – write a gz format
- `customkey (string or list of strings, optional)` – If specified, it adds this custom column to the dump file. Default None.
- `customvals (list or list of lists, optional)` – If customkey is specified, customvals take an array of the same length as number of atoms, which contains the values to be written out.
- `timestep (int, optional)` – Specify the timestep value, default 0
- `species (None, optional)` – species of the atoms. Required if any format other than ‘lammps-dump’ is used. Required for conversion to ase object.

**Returns**

- `Return type` None

pyscal.misc module

pyscal.misc.compare_atomic_env(infile, atomtype=2, precision=2, format='poscar', print_results=True, return_system=False)

Compare the atomic environment of given types of atoms in the input file. The comparison is made in terms of Voronoi volume and Voronoi fingerprint.

**Parameters**

- `infile (string)` – name of the input file
- `atomtype (int, optional)` – type of the atom default 2
• **`precision`** *(float, optional)* – precision for comparing Voronoi volumes default 3
• **`format`** *(string, optional)* – format of the input file default poscar
• **`print_results`** *(bool, optional)* – if True, print the results. If False, return the data instead. default True
• **`return_system`** *(bool, optional)* – if True, return the system object. default False

**Returns**

• **`vvx`** *(list of floats)* – unique Voronoi volumes. Returned only if print results is False
• **`vrx`** *(list of strings)* – unique Voronoi polyhedra. Returned only if print results is False
• **`vvc`** *(list of ints)* – number of unique quantities specified above. Returned only if print results is False

```python
pyscal.misc.find_tetrahedral_voids(infile, format='poscar', print_results=True, return_system=False, direct_coordinates=True, precision=0.1)
```

Check for tetrahedral voids in the system

**Parameters**

• **`infile`** *(string)* – name of the input file
• **`format`** *(string)* – format of the input file, optional default poscar
• **`print_results`** *(bool, optional)* – if True, print the results. If False, return the data instead. default True
• **`return_system`** *(bool, optional)* – if True, return the system object. default False
• **`direct_coordinates`** *(bool, optional)* – if True, results are provided in direct coordinates default False
• **`precision`** *(int, optional)* – the number of digits to check for distances. default 1

**Returns**

• **`types`** *(list of atom types)*
• **`volumes`** *(list of atom volumes)*
• **`pos`** *(list of atom positions)*
• **`sys`** *(system object, returns only if return_sys is True)*
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