Cgenarris Documentation

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1 Features

- Support for special positions.
- $\bullet\,$ Parallelized using MPI and OpenMP.
- $\bullet\,$ Fast and efficient structure checking.
- \bullet Can generate in all possible Z (or NMPC) with Z' <= 1.

2 Installation

2.1 Cgenarris

Requirements

Any C compiler which supports ANSI C99 / GNU99 standard.

Using Makefile

- 1. Uncompress the tar file.
- 2. Execute 'make cgenarris'.
- 3. This will create cgenarris.x which is the desired executable.

NOTE:

- 1. You may change the C compiler using the environment variable CC. You may also uncomment the first line of the makefile and set the compiler.
- 2. Remove object files using 'make clean'

2.2 Pygenarris_mpi

Requirements

- 1. Any C compiler which supports ANSI C99 / GNU99 standard.
- 2. SWIG (Simplified Wrapper code and Interface Generator).
- 3. Numpy and mpi4py
- 4. Distutils for installation through setup.py

Method 1: Using Makefile

- 1. Uncompress the tar file.
- 2. Paste the location of Python.h headerfile in the Makefile. (for Anaconda v3.7 it should be 'anaconda/include/python3.7m/ ')
- 3. Execute 'make pygenarris_mpi'.
- 4. This will create pygenarris_mpi.so library from which you can import pygenarris functions.

Method 2: Using Distutils

- 1. Uncompress the tar file.
- 2. Execute 'python setup.py build_ext --inplace'.
- 3. This will create pygenarris_mpi.so library from which you can import pygenarris functions.

NOTE

- 1. You may change the C compiler using the environment variable CC.
- 2. Pygenarris can be built with both python2 and python3.

3 Cgenarris

Cgenarris is the structure generator written in pure C. It can be compiled into a binary and can run without installing the full genarris python package. If you are interested in only random crystal structures, you can compile and run Cgenarris. It is parallelised using OpenMP and therefore can run only in one node. (An MPI version is under development). The settings for generation are read from control.in file and molecule geometry is read from geometry.in file found in the working directory. The output is printed in file named geometry.out file.

First, the generator first identifies space groups that are compatible with molecular symmetry and given number of molecules in the unit cell. Structures are generated sequentially from lowest space group to the highest. Cell volumes are sampled from a normal distribution. The attempted structures are checked for closeness of molecules. If an atom of a molecule is too close to its own periodic image or another atom of a different molecule in a cell, the structure is discarded. The closeness checks are controlled by the specific radius proportion (sr). If the generation of a space group fails after max_attempts times, the generator moves to the next higher space group. The generated structures are printed to the file in FHI-aims geometry.in. format.

Input

Geometry of the molecule is read from the file *geometry.in* from the working directory.

- 1. num_structures is the number of structures from each space group. Type: integer
- 2. Z number of molecules in the conventional cell. Type: integer
- 3. volume_mean is the mean of the normal distribution from which volume is sampled. Type: float
- 4. volume_std is the standard deviation of the volume distribution. Type: float
- $5. \ sr$ is specific radius proportion. See Genarris paper for definition. Type: float
- 6. tol is the tolerance for special position generation and space group detection. Type: float
- 7. max_attempts is the maximum number of attempts before moving to the next space group. Type: integer
- 8. The number of threads can be set by using the environment variable OMP_NUM_THREADS.

Output

1. A file named *qeometry.out* with all the generated structures in FHI-aims geometry format.

4 Pygenarris_mpi

Pygenarris_mpi is a python API for C structure generator and associated functions. Import the module by:

import pygenarris_mpi

4.1 Generate a pool of random molecular crystals

qenerate_molecular_crystals(filename, num_structures, Z, volume_mean, volume_std, sr, tol, max_attempts)

Description

Generate random molecular crystals by space groups. First, the generator first identifies space groups that are compatible with molecular symmetry and given number of molecules in the unit cell. Structures are generated sequentially from lowest space group to the highest. Cell volumes are sampled from a normal distribution. The attempted structures are checked for closeness of molecules. If an atom of a molecule is too close to its own periodic image or another atom of a different molecule in a cell, the structure is discarded. The closeness checks are controlled by the specific radius proportion (sr). If the generation of a space group fails after max_attempts times, the generator moves to the next higher space group. The generated structures are printed to the file in FHI-aims geometry.in. format.

Input

- 1. Geometry of the molecule is read from the file geometry.in from the working directory.
- 2. filename is the name of the file to which generated structures are printed. Type: string
- 3. num_structures is the number of structures from each space group. Type: integer
- 4. Z number of molecules in the conventional cell. Type: integer
- 5. volume_mean is the mean of the normal distribution from which volume is sampled. Type: float
- 6. volume_std is the standard deviation of the volume distribution. Type: float
- 7. sr is specific radius proportion. See Genarris paper for definition. Type: float
- 8. tol is the tolerance for special position generation and space group detection. Type: float
- 9. max_attempts is the maximum number of attempts before moving to the next space group. Type: integer
- 10. The number of threads can be set by using the environment variable OMP_NUM_THREADS.

Output

1. A file with all the generated structures in FHI-aims geometry format

4.2 Generate a pool of random molecular crystals with VdW cutoff matrix

generate_molecular_crystals_with_vdw_cutoff_matrix(filename, seedstate, vdw_matrix, num_structures, Z, volume_mean, volume_std, tol, max_attempts)

Description

Almost the same as the previous function. Uses a van der Waal cutoff matrix instead of sr.

Input

- 1. All keywords same as before.
- 2. seedstate is the seed for random number generator. Type: int.
- 3. vdw_matrix is a numpy array. Type: 2D numpy array of type "float32" (single precision) and size $total_atoms \ge total_atoms$.

4.3 Identification of compatible space groups given molecular symmetry

 $find_allowed_positions_using_molecular_symmetry(point_group, Z, Z")$

Description

This function finds the compatible space group positions using molecule's point group.

Input

- 1. point_group is the point_group of the molecule. Eg: "mmm" for tetracene. Type: String.
- 2. Z is the number of molecules in the conventional cell. Type: integer
- 3. Z" is the number of inequivalent molecules in the cell. Not implemented! Set any integer.

Output

1. Compatible Wyckoff positions and space groups are printed.

Example

```
find allowed positions_using_molecular_symmetry("m", ", ")
molecular symmetry = m
spg:2 wyckoff position:2i site symmetry:1 allowed
spg:3 wyckoff position:2e site symmetry:1 allowed
spg:4 wyckoff position:2a site symmetry:1 allowed
spg:6 wyckoff position:2c site symmetry:1 allowed
spg:7 wyckoff position:2a site symmetry:1 allowed
spg:8 wyckoff position:2a site symmetry:m allowed
spg:10 wyckoff position:2n site symmetry:m allowed
spg:10 wyckoff position:2m site symmetry:m allowed
spg:11 wyckoff position:2e site symmetry:m allowed
spg:25 wyckoff position:2h site symmetry:m allowed
spg:25 wyckoff position:2g site symmetry:m allowed
spg:25 wyckoff position:2f site symmetry:m allowed
spg:25 wyckoff position:2e site symmetry:m allowed
spg:26 wýckoff position:2b site sýmmetrý:m allowed
spg:26 wyckoff position:2a site symmetry:m allowed
spg:28 wyckoff position:2c site symmetry:m allowed
spg:31 wyckoff position:2a site symmetry:m allowed
Total allowed spacegroup types : 12
Total allowed positions: 17
```

4.4 Molecule closeness check using specific radius proportion

 $int \ c_check_structure(xtal, \ sr)$

Description

This function uses the built-in cgenarris structure checking function to see if the molecules are unphysically close. The treshold is controlled by specific radius proportion (sr). See the genarris paper for its definition.

Input

First, you need to generate a SWIG object (C structure called crystal internally) of type crystal by:

- 1. sr is the specific radius proportion.
- 2. create an object by xtal = crystal(). [Or use the constructor generated automatically by SWIG; $xtal = new_crystal()$.]
- 3. Then assign values using the function
- create_crystal_from_array(xtal, lattice_vector, X, Y, Z, atoms, total_atoms, Z, spg)
- *lattice vector* is a 3x3 numpy array which has the lattice vectors of the crystal in a row-wise form. Type: numpy 2D array.
- X, Y, Z are numpy arrays which the X, Y, and Z coordinates respectively. Type: numpy 1D array; length total_atoms.

- atoms contain the atom type. This is a string which defines the element at each coordinate. If an atom is represented by a single character, add a trailing space. Eg: for Carbon it should be "C", for Bromine it should be "Br". Type: String of length 2 x total_atoms
- $total_atoms$ is the number of atoms in the molecule. This should be the length of X, Y, Z. Type: integer.
- Z is the number of molecules in the unit cell. Type: integer
- spg is the space group (Not used by structure checker!). Type: integer
- IMPORTANT: It is assumed that coordinates are specified in molecule blocks. i.e, first molecule is first N coordinates, second molecule is from N+1 to 2N etc.
- IMPORTANT: Don't forget to free the swig object. This can be done by delete_crystal(xtal).

Output

Returns 0 if the structure is unphysical, returns 1 if the structure passes the test.

4.5 Molecule closeness check using vdW distance matrix

 $int\ check_structure_with_vdw_matrix(xtal,\ vdw_matrix)$

Description

The vdW distance matrix defines the shortest distance between two atoms belonging to different molecules (which includes periodic images). The matrix should be symmetric and should be of size total_atoms x total_atoms where total_atoms is the number of atoms in a cell.

Input

- 1. xtal is an object of type crystal. See the previous function for more details.
- 2. vdw_matrix is a numpy array. Type: 2D numpy array of type "float32" (single precision) and size $total_atoms \times total_atoms$.

Output

Returns 0 if the structure is unphysical, returns 1 if the structure passes the test.

4.6 Number of space groups compatible with molecule's symmetry

 $int\ num_compatible_spacegroups(Z,\ tolerance)$

Description

Function to get information about the allowed space groups and Wyckoff positions for a given molecule.

Input

- 1. Z is the number of molecules in the unit cell. Type: int.
- 2. tolerance is the tolerance for compatibility check. Type: float.
- 3. Molecule geometry is read from geometry in file from the present working directory.

Output

- 1. Returns the allowed space groups as an integer.
- 2. Prints the detailed output to st dout which includes Wyckoff positions available for given ${\cal Z}$.