Icc Documentation

Release 1.0.0

Christian F. A. Negre

CONTENTS:

1	LCC		3
	1.1	About	3
	1.2	License	3
	1.3	Requirements	4
	1.4	Quick installation using <i>spack</i>	4
	1.5	Download and installation	5
	1.6	Compiling PROGRESS and BML libraries	5
	1.7	Step-by-step installation	5
	1.8	Testing the code	6
	1.9	Quick example run	6
	1.10	Contributors	7
	1.11	Contributing	7
	1.12	Citing	7
	1.12		,
2	Build	ling LCC documentation	9
	2.1	Prerequisites	9
	2.2	Build the full documentation	9
	2.3	Documenting	10
3	Deve	loping	11
4	Innu	t file choices	13
7	4.1		13
	4.2	Jobivanie	10
		Varhasa	12
		Verbose=	13
	4.3	CoordsOutFile=	13
	4.3	CoordsOutFile=	13 13
	4.3 4.4 4.5	CoordsOutFile= PrintCml= ClusterType=	13 13 14
	4.3 4.4 4.5 4.6	CoordsOutFile= PrintCml= ClusterType= ClusterType= Bulk	13 13 14 14
	4.3 4.4 4.5 4.6 4.7	CoordsOutFile= PrintCml= ClusterType= ClusterType= Bulk ClusterType= Spheroid	13 13 14 14 14
	4.3 4.4 4.5 4.6 4.7 4.8	CoordsOutFile= PrintCml= ClusterType= ClusterType= Bulk ClusterType= Spheroid ClusterType= Planes	13 14 14 14 14
	4.3 4.4 4.5 4.6 4.7 4.8 4.9	CoordsOutFile= PrintCml= ClusterType= ClusterType= Bulk ClusterType= Spheroid ClusterType= Planes CenterAtBox=	13 14 14 14 14 15
	4.3 4.4 4.5 4.6 4.7 4.8 4.9 4.10	CoordsOutFile= PrintCml= ClusterType= ClusterType= Bulk ClusterType= Spheroid ClusterType= Planes CenterAtBox= Reorient=	13 13 14 14 14 15 15
	4.3 4.4 4.5 4.6 4.7 4.8 4.9 4.10 4.11	CoordsOutFile= PrintCml= ClusterType= ClusterType= Bulk ClusterType= Spheroid ClusterType= Planes CenterAtBox= Reorient= AtomType=	13 13 14 14 14 14 15 15
	4.3 4.4 4.5 4.6 4.7 4.8 4.9 4.10 4.11 4.12	CoordsOutFile= PrintCml= ClusterType= ClusterType= Bulk ClusterType= Spheroid ClusterType= Planes CenterAtBox= Reorient= AtomType= TypeOfLattice=	13 14 14 14 15 15 15
	4.3 4.4 4.5 4.6 4.7 4.8 4.9 4.10 4.11 4.12 4.13	CoordsOutFile= PrintCml= ClusterType= ClusterType= Bulk ClusterType= Spheroid ClusterType= Planes CenterAtBox= Reorient= AtomType= TypeOfLattice= RandomSeed=	13 13 14 14 14 15 15 15 15
	4.3 4.4 4.5 4.6 4.7 4.8 4.9 4.10 4.11 4.12 4.13 4.14	CoordsOutFile= PrintCml= ClusterType= ClusterType= Bulk ClusterType= Spheroid ClusterType= Planes CenterAtBox= Reorient= AtomType= TypeOfLattice= RandomSeed= PrimitiveFormat=	13 13 14 14 14 15 15 15 15 16
	4.3 4.4 4.5 4.6 4.7 4.8 4.9 4.10 4.11 4.12 4.13 4.14 4.15	CoordsOutFile= PrintCml= ClusterType= ClusterType= Bulk ClusterType= Spheroid ClusterType= Planes CenterAtBox= Reorient= AtomType= TypeOfLattice= RandomSeed= PrimitiveFormat= UseLatticeBase=	13 14 14 14 15 15 15 15 16 16
	4.3 4.4 4.5 4.6 4.7 4.8 4.9 4.10 4.11 4.12 4.13 4.14	CoordsOutFile= PrintCml= ClusterType= ClusterType= Bulk ClusterType= Spheroid ClusterType= Planes CenterAtBox= Reorient= AtomType= TypeOfLattice= RandomSeed= PrimitiveFormat=	13 13 14 14 14 15 15 15 15 16

6	Building/cutting a shape	23
	6.1 Growing shapes from a seed file	23
	6.2 Cutting using planes	24
7	Building a slab	27
	7.1 Building a slab from three PBC vectors	31
8	Building regular shapes	33
9	Computing Roughness	35
10	Tools	4 1
	10.1 Kinetic Monte Carlo Crystal Growth Tool	41
	10.1.1 About	41
	10.1.2 Requirements	41
	10.1.3 Testing and running the code	41
	10.1.4 Notes	42
	10.2 Encapsulating orthogonal vectors	42
11	Indices and tables	43

Issues	Pull Requests	CI

CONTENTS: 1

2 CONTENTS:

CHAPTER

ONE

LCC

1.1 About

Los Alamos Crystal Cut (LCC) is simple crystal builder. It is an easy-to-use and easy-to-develop code to make crystal solid/shape and slabs from a crystal lattice. Provided you have a .pdb file containing your lattice basis you can create a solid or slab from command line. The core developer of this code is Christian Negre (cnegre@lanl.gov). This documentation has been approved for unlimited release with LA-UR-23-28084.

1.2 License

© 2022. Triad National Security, LLC. All rights reserved. This program was produced under U.S. Government contract 89233218CNA000001 for Los Alamos National Laboratory (LANL), which is operated by Triad National Security, LLC for the U.S. Department of Energy/National Nuclear Security Administration. All rights in the program are reserved by Triad National Security, LLC, and the U.S. Department of Energy/National Nuclear Security Administration. The Government is granted for itself and others acting on its behalf a nonexclusive, paid-up, irrevocable worldwide license in this material to reproduce, prepare derivative works, distribute copies to the public, perform publicly and display publicly, and to permit others to do so.

This program is open source under the BSD-3 License.

Redistribution and use in source and binary forms, with or without modification, are permitted provided that the following conditions are met:

- 1. Redistributions of source code must retain the above copyright notice, this list of conditions and the following disclaimer.
- 2. Redistributions in binary form must reproduce the above copyright notice, this list of conditions and the following disclaimer in the documentation and/or other materials provided with the distribution.
- 3. Neither the name of the copyright holder nor the names of its contributors may be used to endorse or promote products derived from this software without specific prior written permission.

THIS SOFTWARE IS PROVIDED BY THE COPYRIGHT HOLDERS AND CONTRIBUTORS "AS IS" AND ANY EXPRESS OR IMPLIED WARRANTIES, INCLUDING, BUT NOT LIMITED TO, THE IMPLIED WARRANTIES OF MERCHANTABILITY AND FITNESS FOR A PARTICULAR PURPOSE ARE DISCLAIMED. IN NO EVENT SHALL THE COPYRIGHT HOLDER OR CONTRIBUTORS BE LIABLE FOR ANY DIRECT, INCIDENTAL, SPECIAL, EXEMPLARY, OR CONSEQUENTIAL DAMAGES (INCLUDING, BUT NOT LIMITED TO, PROCUREMENT OF SUBSTITUTE GOODS OR SERVICES; LOSS OF USE, DATA, OR PROFITS; OR BUSINESS INTERRUPTION) HOWEVER CAUSED AND ON ANY THEORY OF LIABILITY, WHETHER IN CONTRACT, STRICT LIABILITY, OR TORT (INCLUDING NEGLIGENCE OR OTHERWISE) ARISING IN ANY WAY OUT OF THE USE OF THIS SOFTWARE. EVEN IF ADVISED OF THE POSSIBILITY OF SUCH DAMAGE.

1.3 Requirements

In order to follow this tutorial, we will assume that the reader have a LINUX or MAC operative system with the following packages properly installed:

- The git program for cloning the codes.
- A C/C++ compiler (gcc and g++ for example)
- A Fortran compiler (gfortran for example)
- The LAPACK and BLAS libraries (GNU libblas and liblapack for example)
- The python interpreter (not essential).
- The pkgconfig and cmake programs (not essential).

On an x86_64 GNU/Linux Ubuntu 16.04 distribution the commands to be typed are the following::

```
sudo apt-get update
sudo apt-get --yes --force-yes install gfortran gcc g++
sudo apt-get --yes --force-yes install libblas-dev liblapack-dev
sudo apt-get --yes --force-yes install cmake pkg-config cmake-data
sudo apt-get --yes --force-yes install git python
```

NOTE: Through the course of this tutorial we will assume that the follower will work and install the programs in the home directory (\$HOME).

1.4 Quick installation using spack

Clone and setup the spack code:

```
cd ~
git clone git@github.com:spack/spack.git
. spack/share/spack/setup-env.sh
```

Get info on the package:

```
spack info lcc
```

Install the package, this will take a while because it'll install everything from scratch:

```
spack install lcc
```

Load the lcc module:

```
spack load lcc
```

Try lcc:

```
cd tmp ; lcc_main
spack install ovito
```

(continues on next page)

4 Chapter 1. LCC

```
spack load ovito

cd /tmp

echo "LCC{ ClusterType= Spheroid TypeOfLattice= FCC AAxis= 10.0 BAxis= 10.0 CAxis= 10.0 }

_" | tee input.in ; lcc_main input.in

ovito coords.xyz
```

1.5 Download and installation

We will need to clone the repository as follows::

```
cd; git@github.com:lanl/LCC.git
```

1.6 Compiling PROGRESS and BML libraries

The LCC code needs to be compiled with both [PROGRESS](https://github.com/lanl/qmd-progress) and [BML](https://github.com/lanl/bml) libraries. In this section we will explain how to install both of these libraries and link the code against them.

Scripts for quick installations can be found in the main folder. In principle one should be able to install everything by typing::

```
./clone_libs.sh
./build_bml.sh
./build_progress.sh
./build.sh
```

Which will also build LCC with its binary file in ./src/lcc_main.

1.7 Step-by-step installation

Clone the BML library (in your home directory) by doing¹:

```
cd
git clone git@github.com:lanl/bml.git
```

Take a loot at the ./scripts/example_build.sh file which has a set of instructions for configuring. Configure the installation by copying the script into the main folder and run it::

```
cp ./scripts/example_build.sh .
sh example_build.sh
```

¹ In order to have access to the repository you should have a github account and make sure to add your public ssh key is added in the configuration windows of github account.

The build.sh script is called and the installation is configured by creating the build directory. Go into the build directory and type:

```
cd build
make -j
make install
```

To ensure bml is installed correctly type \$ make tests or \$ make test ARGS="-V" to see details of the output. Series of tests results should follow.

After BML is installed, return to you home folder and "clone" the PROGRESS repository. To do this type:

```
cd
git clone git@github.com:lanl/qmd-progress.git
```

Once the folder is cloned, cd into that folder and use the example_build.sh file to configure the installation by following the same steps as for the bml library.:

```
sh example_build.sh
cd build
make; make install
```

You can test the installation by typing \$ make tests in the same way as it is done for BML.

Open the Makefile file in the lcc/src folder make sure the path to both bml and progress libs are set correctly. NOTE: Sometimes, depending on the architecture the libraries are installed in /lib64 instead of /lib. After the aforementioned changes are done to the Makefile file proceed compiling with the "make" command.

1.8 Testing the code

A test script can be run as follows:

./run_test

1.9 Quick example run

Assuming the code is installed in the \$HOME directory, we will run a simple example::

```
cd /tmp
echo "LCC{ ClusterType= Spheroid TypeOfLattice= FCC AAxis= 10.0 BAxis= 10.0 CAxis= 10.0 }

→" | tee input.in ; $HOME/LCC/build/lcc_main input.in
```

This will generate a spherical structure with an FCC lattice using default parameters. One can quickly get an input file sample by running the code without giving any input file. The available keywords can be listed by running lcc_main -h

6 Chapter 1. LCC

1.10 Contributors

Christian Negre, email: cnegre@lanl.gov Andrew Alvarado, email: aalvarado@lanl.gov

1.11 Contributing

Formally request to be added as a collaborator to the project by sending an email to cnegre@lanl.gov. After being added to the project do the followig:

- Create a new branch with a proper name that can identify the new feature (git checkout -b "my_new_branch"
- Make the changes or add your contributions to the new branch (git add newFile.F90 modifiedFile.F90)
- Make sure the tests are passing (cd tests; ./run_test.sh)
- Commit the changes with proper commit messages (git commit -m "Adding a my new contribution")
- Push the new branch to the repository (git push)
- Go to repository on the github website and click on "create pull request"

SUGGESTION: Please, avoid committing a large number of changes since it is difficult to review. Instead, add the changes gradually.

1.12 Citing

If you find this code useful, we encourage you to cite us. Our project has a citable DOI (DOI:10.1088/1361-648X/acc294) with the following bibtex snipped:

```
@ARTICLE { lcc.
  title
           = "A methodology to generate crystal-based molecular structures for
            atomistic simulations",
          = "Negre, Christian F A and Alvarado, Andrew and Singh, Himanshu and
 author
            Finkelstein, Joshua and Martinez, Enrique and Perriot, Romain",
 abstract = "We propose a systematic method to construct crystal-based
            molecular structures often needed as input for computational
            chemistry studies. These structures include crystal 'slabs' with
            periodic boundary conditions (PBCs) and non-periodic solids such
            as Wulff structures. We also introduce a method to build crystal
            slabs with orthogonal PBC vectors. These methods are integrated
            into our code, Los Alamos Crystal Cut(LCC), which is open source
            and thus fully available to the community. Examples showing the
            use of these methods are given throughout the manuscript.",
 journal = "J. Phys. Condens. Matter",
 volume
          = 35,
 number
          = 22,
 month
          = mar,
          = 2023.
 keywords = "crystal structures; extended structures; miller indices; quantum
            chemistry; unit cells",
```

(continues on next page)

1.10. Contributors 7

```
language = "en"
}
```

8 Chapter 1. LCC

CHAPTER

TWO

BUILDING LCC DOCUMENTATION

The folder (src/docs:) contains all the documentation relevant to both users and developpers.

2.1 Prerequisites

- [pdflatex] Latex GNU compiler. pdfTeX is an extension of TeX which can produce PDF directly from TeX source, as well as original DVI files. pdfTeX incorporates the e-TeX extensions.
- [doxygen] Doxygen is a documentation system for C++, C, Java, Objective-C, IDL (Corba and Microsoft flavors) and to some extent PHP, C#, and D.
- [sphinx] Sphinx is a documentation generator or a tool that translates a set of plain text source files into various output formats, automatically producing cross-references, indices, etc. That is, if you have a directory containing a bunch of reStructuredText or Markdown documents, Sphinx can generate a series of HTML files, a PDF file (via LaTeX), man pages and much more.
- Any pdf viewer.
- Any web browser.

These programs can be installed as follows:

```
sudo apt-get install pdflatex
sudo apt-get install doxygen
sudo apt-get install dot2tex
sudo apt-get install python3-sphinx
pip3 install PSphinxTheme
pip3 install recommonmark
```

2.2 Build the full documentation

This will build all three types of docs (Sphinx, Doxygen, and latex):

make

The documentation that is build with Sphinx can be tested as follows:

```
firefox lcc.html
```

The file can be explored using any web browser.

One can also build any of the documentations separatly. For example, to build the Sphinx documentation, we can do:

make sphinx

2.3 Documenting

In order to add a documentation using Sphinx follow these steps:

- 1) make a file with a proper name under ./sphinx-src/source/. For example: MYPAGE.md.
- 2) Add the documentation inside the file using "markdown" syntax.
- 3) Modify the file in ./sphinx-src/source/index.txt to include the documentation.

After modyfing this file, recompile Sphinx by typing make sphinx.

CHAPTER

THREE

DEVELOPING

In order to develop within the code, the reader is encouraged to check the Modules details.

CHAPTER

FOUR

INPUT FILE CHOICES

In this section we will describe the input file keywords. Every valid keword will use "cammel" syntax and will have and = sign right next to ie. For example, the following is a valid keyword syntax JobName= MyJob. Coments need to have a # (hash) sign right nex to the phrase we want to comment. Example comment could be something like: #My comment.

4.1 JobName=

This variable will indicate the name of the job we are sunning. It is just a tag to distinguish different outputs. As we mentioned before and example use sould be: JobName= MyJob

4.2 Verbose=

Controls the verbosity level of the output. If set to 0 no output is pronted out. If set to 1, only basic meesages of the current execution point of the code will be printed. If set to 2, information about basic quantities are also printed. If set to 3, all relevant possible info is printed.

4.3 CoordsOutFile=

This will store the name of the output coordinates files. Basically if CoordsOutFile= coords two output file will be created: coords.xyz and coords.pdb.

4.4 PrintCml=

By setting PrintCml= T will also print create coords.cml which can be readed by avogadro. In order to have this option working one needs to install [openbabel] In order to read a cml file one needs to have [avogadro] installed. On gnu linux:

```
sudo apt-get avogadro
sudo apt-get obabel
```

4.5 ClusterType=

This variable will define the type of shape/cluster/slab we want to construct. There are many options including Bulk, Planes, Bravais and Spheroid. We will explain all these in the following sention.

4.6 ClusterType= Bulk

This will just cut a "piece of bulk" by indicating how many lattice point we want. For example, the following will create a bulk/lattice with 50 points on each a,b,c direction:

```
ClusterType= Bulk
LatticePoints= 50
```

The following, instead, will create a bulk/lattice with 100 lattice points in the x direction and 50 on the rest:

```
LatticePointsX1= 1
LatticePointsX2= 100
LatticePointsY1= 1
LatticePointsY2= 50
LatticePointsZ1= 1
LatticePointsZ2= 50
```

4.7 ClusterType= Spheroid

This will produce a "spheroid" center at the origin. And example follows:

```
ClusterType= Spherid
LatticePoints= 50  #This is necesary to construct the initial bulk
AAxis= 1.0 #Radius in direction x
BAxis= 2.0 #Radius in direction y
CAxis= 2.0 #Radius in direction z
```

See section *regular* to see another example.

4.8 ClusterType= Planes

This will cut a shape using Miller indice. This is an important tool to construct a slab to study a surface. The cut does not gurantee periodicity. In order to have a periodic structure different plane boudaries need to be tried and the structures needs to be checked using a molecular sivualizer. An example is given as follows:

```
NumberOfPlanes= 6
Planes[
0 1 1 2.5
0 -1 -1 1.5
0 -1 1 4.5
0 1 -1 3.5
1 0 0 4.5
```

(continues on next page)

```
-1 0 0 3.5
]
```

Three first number on each row indicate the Miller indices. The fourth number indicates how many Miller planes from the origin will be cut out. If the number of planes is 6, then the system tries to get the slab peridicity vectors since if the Miller planes are orthogonal to each other, the shape will be a "Parallelepiped". If instead the number different than 6, then the periodicity vectors are given by the "Boundaries" of the minimal box that contains the shape.

4.9 CenterAtBox=

If set to T, the shape will be centered at the box (the periodicity vectors of the shape/cluster)

4.10 Reorient=

If set to T this, will reorient the shape, such that vector "a" will be aligned with the x dierction. This is important when making slabs needed to study a surface.

4.11 AtomType=

This will sed the atom symbol if the lattice basis is not read from file.

4.12 TypeOfLattice=

This will set the Lattice unit cell. if set to SC or FCC either a simple cubic or face centered cubinc lattice is built provided we set LatticeConstanta= to the lattice constant value. For general unit cell we can set TypeOfLattice= Triclinic, and provide the lattice parameters as in the following example:

```
      LatticeConstanta=
      6.5329400000000000

      LatticeConstantb=
      11.022100000000000

      LatticeConstantc=
      7.3568800000000000

      LatticeAngleAlpha=
      90.00000000000000

      LatticeAngleBeta=
      102.6520000000000

      LatticeAngleGamma=
      90.00000000000000
```

4.13 RandomSeed=

To generate random positions in the lattice. This will need to be used in conjunction with RCoeff= which controll the degree of deviation from the lattice positions.

4.9. CenterAtBox=

4.14 PrimitiveFormat=

This will indicate if the lattice needs to be constructed out of a,b,c and angle parameter or primitive lattice vectors. If PrimitiveFormat= Angles (default), then the lattice parameters will need to be passed as in the following example:

If instead, PrimitiveFormat= Vectors then the primitive vectors will need to be passed as in the following example:

```
LatticeVectors[
    2.0 0 0  #First lattice vector
    0.0 2.0 0
    0.0 2.0 2.0 1
```

4.15 UseLatticeBase=

This is an important tool that allows us to "dress" every lattice point with a basis of choice. The basis is defined to be the minimal set of corrdinates and atom types needed to define a crystal system lattice point. The basis here will be red from file by providing the latticebase LatticeBaseFile= wich will contain our atom types and coordinates. If ReadLatticeFromFile= is set to T, then, the lattice parameters will be read from the lattice basis file. If is set to F, the the lattice parameters will need to be passed as explaines before. Another important keyword is the BaseFormat=. If this is set to abc, then the basis coordinates stored in the file are assumed to be given in fractional coordinates of the lattice parameters. If is set to xyz, the it will be assumed to be given in catesian coordinates.

4.16 SymmetryOperations=

If the basis needs to be constricted from symetry operation, then one needs to pass all these operation to the code as follows:

```
| SymmetryOperations= T
| NumberOfOperations= 4
| Translations[
| 0 0 0 0.0
| 1 1 1 0.5
| 1 1 0 1.0
| -0.5 1.5 0.5 1.0
| Symmetries[
| 0 0 0
| -1 1 -1
| -1 -1 1 1
```

The first block indicates the "translations" within the unit cell. The first three rows indicating the directions of the translation and the fourth indicating the intensity. The second block indicates the symmetry of operations. For example, if an operation is indicated as (-x+1/2,-y,-z) then there will be a traslation 0.5 0 0 1.0 and a summetry -1 0 0:

CHAPTER

FIVE

BUILDING A LATTICE

In this section we briefly explain how to build a lattice using LCC. The finite set of points obtained in this ways has the shape that is bound by crystal faces which are parallel to the "canonical Miller planes" (100), (010), and (001) We will first execute lcc without any input file to create a sample input. Syntax follows:

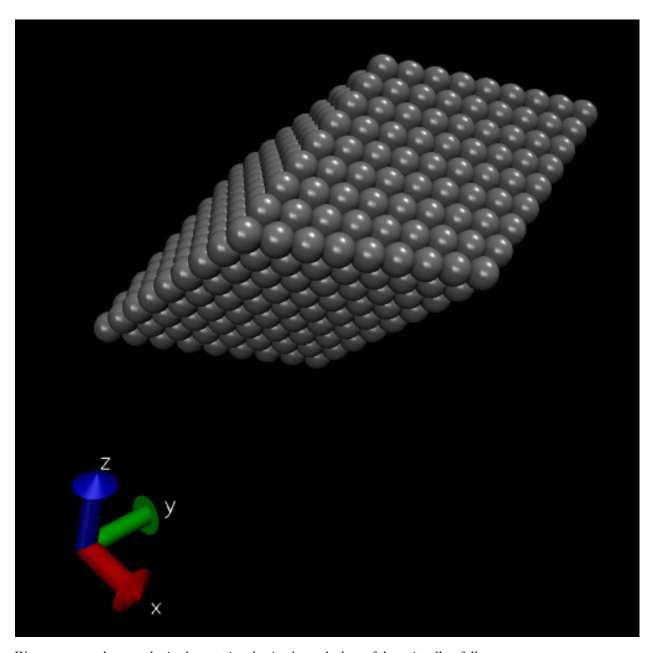
```
./lcc_main
```

This will generate a sample input file called sample_input.in. You can either edit this file or make a new one having the following:

In order to run the code, just type:

```
./lcc_main sample_input.in
```

The run will produce two coordinate files *_coords.xyz and *_coords.pdb. If we visualize this with VMD we get the following "piece of bulk" for Silver



We can recover the same lattice by entering the Angles and edges of the unit cell as follows:

#Lcc input file.		
LCC{		
JobName=	AgBulk	#Or any other name
ClusterType=	Bulk	
TypeOfLattice=	Triclinic	
LatticePoints=	8	#Number of total lattice points in each direction
AtomType=	Ag	
PrimitiveFormat=	Angles	#Will use angles and edges
LatticeConstanta=	2.885	
LatticeConstantb=	4.08	
LatticeConstantc=	2.885	
LatticeAngleAlpha=	45	

(continues on next page)

```
LatticeAngleBeta= 45
LatticeAngleGamma= 60
}
```

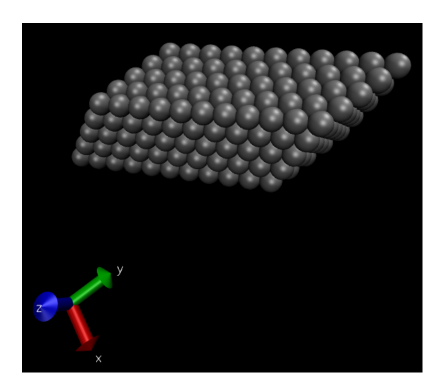
Yet another way of constructing an fcc lattice is by providing the lattice vectors directly which can be done by doing:

```
#Lcc input file.
LCC{
JobName=
                          AgBulk
                                         #Or any other name
ClusterType=
                          Bulk
TypeOfLattice=
                          Triclinic
                                         #Number of total lattice points in each direction
LatticePoints=
AtomType=
                          Ag
PrimitiveFormat=
                          Vectors
                                        #Will use primitive vectors
LatticeVectors[
        2.885 2.885 0.000
        0.000 4.080 0.000
        0.000 2.885 2.885
}
```

If we want a bulk with a particular number of lattice points on each direction we can use the following input parameters:

```
#Lcc input file.
LCC{
JobName=
                          AgBulk
                                         #Or any other name
ClusterType=
                          Bulk
TypeOfLattice=
                          Triclinic
LatticePointsX1=
                                         #Number of point in the direction of the first.
                          -2
→Lattice Vector
LatticePointsX2=
                          8
                          -2
LatticePointsY1=
                          2
LatticePointsY2=
LatticePointsZ1=
                          -2
LatticePointsZ2=
                          2
AtomType=
                          Ag
PrimitiveFormat=
                                         #Will use angles and edges
                          Angles
LatticeConstanta=
                          2.885
{\tt LatticeConstantb} =
                          4.08
LatticeConstantc=
                           2.885
LatticeAngleAlpha=
                          45
LatticeAngleBeta=
                           45
LatticeAngleGamma=
                           60
```

The latter will produce a "bulk" enlarged in the direction of the first lattice vector.



BUILDING/CUTTING A SHAPE

6.1 Growing shapes from a seed file

The Bravais theory says that a crystal face will grow faster is the atom/unit cell that is added to the face finds a higher coordination. In this way, faces that have a high reticular density will grow slower since the adatom will potentially find only a "top" position.

Here we give an example of how to grow a shape from a seed using only geometrical parameters which are: the MinCoordination and the RCut. RCut is used as a criterion to search for the coordination. If the adatom (possible atom to be included in the shape) has 3 atoms that are within RCut, the coordination of such an adatom will be 3. If MinCoordination = 2, the the adatom with coordination = 3 will be included in the shape.

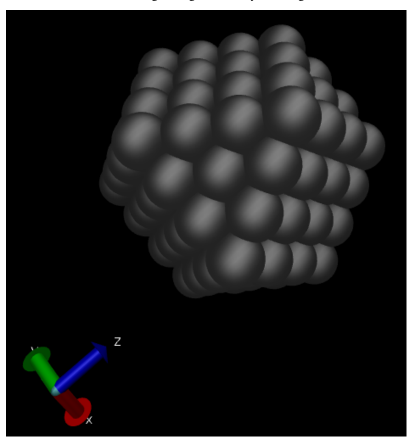
An exaple input file is given as follows:

```
#Lcc input file.
LCC{
JobName=
                          AgBulk
                                         #Or any other name
ClusterType=
                          BravaisGrowth
NumberOfIterations=
RTol=
                          0.01
MaxCoordination=
                          1
RCut=
SeedFile=
                          "seed.pdb"
TypeOfLattice=
                          FCC
                                         #Number of point in the direction of the first
LatticePointsX1=
                          -8
→Lattice Vector
LatticePointsX2=
LatticePointsY1=
                          -8
LatticePointsY2=
                          8
                          -8
LatticePointsZ1=
LatticePointsZ2=
                          8
AtomType=
                          Ag
                                         #Will use angles and edges
PrimitiveFormat=
                          Angles
LatticeConstanta=
                          4.08
```

The NumberOfIterations parameter controls the cycles of gorwing that we want. The SeedFile parameter is the name of the file containg the "seed" from where the shape will grow. For this particular exaple we will used a seed (seed.pdb) file with the following content"

```
Seed File
REMARK
TITLE coords.pdb
CRYST1 137.192 231.464
                         154.494 90.00 102.65
                                                90.00 P 1
MODEL
MOTA
                        1
                                0.000
                                        0.000
                                                0.000
                                                                           Ag
          1 Ag M
                                                       0.00
TER
END
```

This means that we will be growing from "only one" Ag atom center at the origin. The result is the following shape:



6.2 Cutting using planes

A crystal shape can also be cut using planes. This could be usefull to comput a Wulff type of crystal shape by listing the planes and the surface energies or just for creating a "slab" to study a particular surface. An example of cutting by planes is provided as follows:

<pre>#Lcc input file. LCC{ JobName= Verbose=</pre>	AgPlanes	#Or any other name
TypeOfLattice= LatticePoints= LatticeConstanta=	FCC 50 4.08	#Number of point in each direction

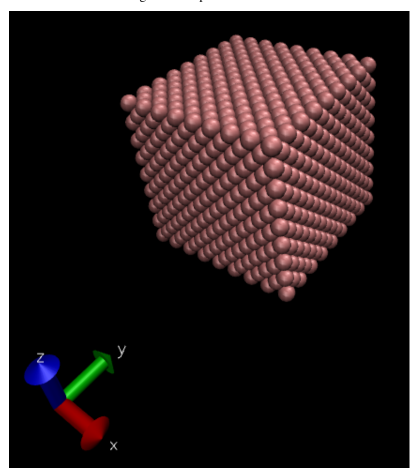
(continues on next page)

```
AtomType= Ag

ClusterType= Planes

NumberOfPlanes= 6
Planes[
    1 0 0 4.1
    -1 0 0 4.1
    0 1 0 4.1
    0 -1 0 4.1
    0 0 -1 4.1
    0 0 0 1 4.1
]
}
```

This creates the following cubic shape:



BUILDING A SLAB

In this tutorial we will explain the steps to construct a crystal "slab" that could be used to study a particular surface.

We will hereby use sucrose as an example. We will build the (0 1 1) surface and create a periodic slab. The data (CIF file) on sucrose was downloaded from svn://www.crystallography.net/cod/cif/3/50/00/3500015.cif

To this end we will use the following input file:

```
LCC{
 ClusterType=
                        Planes
 ClusterNumber=
                        2
  Verbose= 3
  LatticeBaseFile= "lattice_basis.xyz"
  WriteCml= F
  CheckPeriodicity= T
 ReadLatticeFromFile= F
                        Triclinic
  TvpeOfLattice=
  LatticePoints=
                        30
  CheckLattice=
  PrimitiveFormat=
                        Angles
  AtomType=
                        Αt
  UseLatticeBase=
                        Т
  BaseFormat=
                        abc
  CutAfterAddingBase=
  LatticeConstanta=
                        7.789
  LatticeConstantb=
                        8.743
  LatticeConstantc=
                        10.883
 LatticeAngleAlpha=
                         90
  LatticeAngleBeta=
                         102.760
  LatticeAngleGamma=
                         90
 RCoeff= 0.0
  CenterAtBox=
                     Τ
  Reorient=
                     Т
  \#+X, +Y, +Z
  \#-X, 1/2+Y, -Z
  SymmetryOperations= T
```

(continues on next page)

```
NumberOfOperations= 2
  OptimalTranslations= T
  Translations[
   0 0
           0
    0
      1
           0
               0.5
  Symmetries[
   1 1 1
    -1 1 -1
  1
NumberOfPlanes=
Planes[
 0 1
        1
             2.5
             1.5
       -1
 0 - 1
 0 -1
        1
             4.5
 0
   1
       -1
             3.5
 1
    0
         0
             2.5
 -1
    0
             1.5
]
```

The "basis" needs to be provided via the *lattice_basis.xyz* file. The content of such file is provided below:

```
45
#Sucrose basis
0 0.63189
              0.34908
                          0.62279
0 0.7136
              0.2018
                         0.41867
0 0.6440
               -0.0665
                         0.6512
0 0.2978
               -0.0008
                         0.69117
0 0.2529
              0.3114
                          0.77094
0 0.60891
              0.40061
                         0.82857
0 0.68400
              0.65323
                         0.78776
0 0.3785
              0.5127
                         0.97000
0 0.9607
               0.5091
                          0.67341
0 1.0893
              0.6500
                         1.02195
0 0.7957
              0.42950
                         1.07412
C 0.7053
              0.1955
                          0.64075
C
  0.5578
              0.0769
                          0.6265
C
  0.4362
              0.1116
                          0.71451
C 0.3651
              0.2728
                         0.6871
C 0.5149
               0.3897
                          0.70028
C
  0.8157
               0.1767
                          0.5431
  0.6306
C
               0.5556
                          0.87572
C 0.8718
              0.6862
                          0.82381
C
  0.9441
               0.5804
                          0.93500
C
  0.7861
              0.5573
                         0.99233
C 0.4569
              0.6161
                         0.8967
C 0.9532
              0.6662
                         0.7110
H 0.7813
               0.1873
                          0.7252
H 0.4894
               0.0781
                          0.5393
H 0.5018
               0.1046
                          0.8022
H 0.2953
              0.2763
                          0.6004
```

(continues on next page)

				(commuted from previous page)
Н	0.4639	0.4900	0.6734	
Н	0.9127	0.2488	0.5604	
Н	0.8647	0.0743	0.5487	
Н	0.733	0.298	0.402	
Н	0.2287	0.0165	0.7364	
Н	0.2152	0.3986	0.7560	
Н	0.8878	0.7925	0.8526	
Н	0.9806	0.4827	0.9048	
Н	0.7738	0.6491	1.0414	
Н	0.4764	0.7140	0.9395	
Н	0.3769	0.6323	0.8158	
Н	0.3772	0.4263	0.9405	
Н	0.8853	0.7242	0.6409	
Н	1.0716	0.7077	0.7308	
Н	0.860	0.480	0.654	
Н	1.185	0.604	1.009	
Н	0.8058	0.3509	1.0352	
Н	0.553	-0.128	0.642	

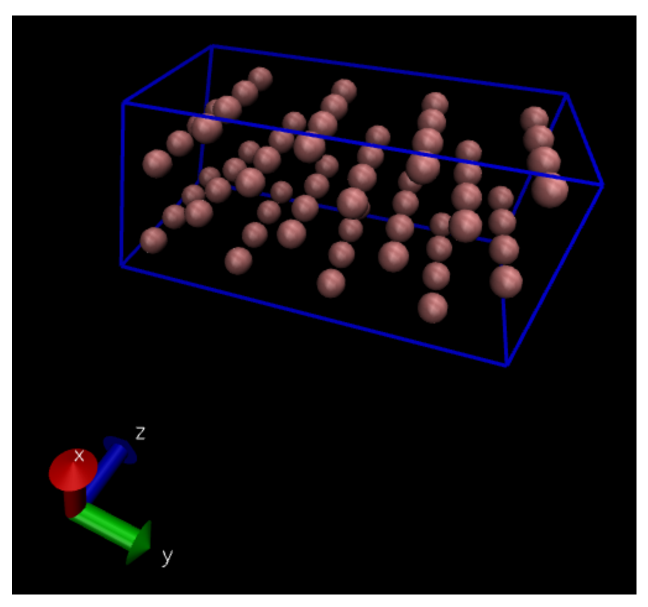
The first run we will do needs to have UseLatticeBase = F. In this way we will be able to inspect the lattice points and make sure that we get a periodic slab. The code automatically checks the periodicity. If the Miler planes are not ensuring periodicity, the code will raise an error. The lattice could be visualized with avogadro or vmd.:

```
avogadro coords.cml
```

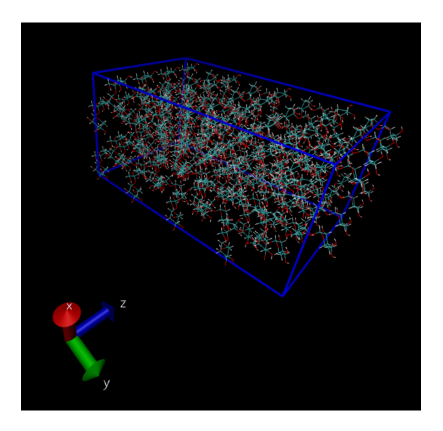
or:

```
vmd -f coords.pdb
```

This will show the following structure:



The next step is to run the code with UseLatticeBase = T to generate the final structure. Note that the input file contains the symetry operation to "complete" the unit cell. After running the code we will get the following structure:



7.1 Building a slab from three PBC vectors

Another method we have to build a crystal slab is to give the program the PBC vectors. For this, we will set *ClusterType*= to *ClusterType*= *Slab*. We will also need to give the PBC vectors and their lengths as follows:

```
Slab[
    1.0 0.0 0.0 10.0
    0.0 1.0 1.0 10.0
    0.0 0.0 1.0 10.0
]
```

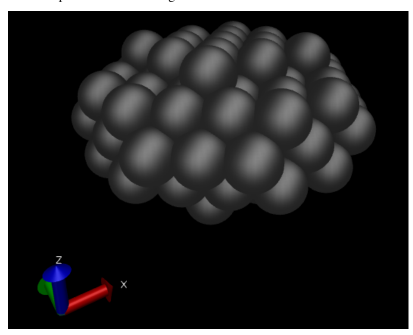
The latter input block means that the first vector will be the (1,0,0) with length 10.0. Note that three general vectors cannot guarantee that the slab will be congruent with the lattice. If we give three random vectors and have *CheckPeriodicity=T*, the code will most likely give an error. An example of construction of this type of slab can be find in *examples/build_from_vectors/*.

BUILDING REGULAR SHAPES

Congruence transformation. One can also build regular shapes, such as for example a "spheroid." The parameters to do this can be entered as follows::

```
#Lcc input file.
LCC{
JobName=
                          AgSpheroid
                                            #Or any other name
TypeOfLattice=
                          FCC
LatticePoints=
                          50
                                         #Number of point in each direction
LatticeConstanta=
                          4.08
AtomType=
                          Ag
ClusterType=
                          Spheroid
                                   #Radius in Ang for x direction
AAxis=
                          10
BAxis=
                                  #Radius in Ang for y direction
                          10
CAxis=
                          5
                                  #Radius in Ang for z direction
```

This will produce the following "oblate":



COMPUTING ROUGHNESS

In this tutorial we will explain the steps to compute a crystal surface Roughness defined as the ration between the "effective" surface and the "flat" surface.

We will hereby use sucrose as an example. We will analyse (100) and (10-1) crystal faces. Lattice information on sucrose was downloaded from svn://www.crystallography.net/cod/cif/3/50/00/3500015.cif

To this end we will use the following input file which can be also found under /examples/Roughness:

```
LCC{
 ClusterType=
                        Planes
  ClusterNumber=
  Verbose= 3
  LatticeBaseFile= "lattice_basis.xyz"
  WriteCml= F
  CheckPeriodicity= T
  ReadLatticeFromFile= F
  TypeOfLattice=
                        Triclinic
  LatticePoints=
  CheckLattice=
  PrimitiveFormat=
                        Angles
  AtomType=
                        Αt
  UseLatticeBase=
  BaseFormat=
                        abc
  CutAfterAddingBase=
  LatticeConstanta=
                        7.789
  LatticeConstantb=
                        8.743
  LatticeConstantc=
                        10.883
 LatticeAngleAlpha=
                         90
  LatticeAngleBeta=
                         102.760
  LatticeAngleGamma=
                         90
  RCoeff= 0.0
  CenterAtBox=
                     Т
  Reorient=
                     Т
  \#+X, +Y, +Z
```

(continues on next page)

```
\#-X, 1/2+Y, -Z
  SymmetryOperations= T
  NumberOfOperations= 2
 OptimalTranslations= F
  Translations[
  1 0
         1
             -1
             0.5
  0 1
         2
  Symmetries[
    1 1 1
    -1 1 -1
 ]
  NumberOfPlanes=
  ComputeRoughness= T
 RoughnessParameters[
   50.0 1.0 40 80 80
 ]
 Planes[
          0
    1
      0
               2.5
    -1
       0
          0
              1.5
   0
      1
          0
              4.5
   0
               3.5
      -1
          0
   0
          1
              2.5
      0
    0
           -1
              1.5
 ]
  #Planes[
   1 0
           -1
               2.5
    -1
       0 1
              1.5
   0 -1
          0
              4.5
            1.5
      1 0
   1
      0
          1
              4.5
          -1 1.5
    -1 0
 ]
}
```

The "basis" needs to be provided via the lattice_basis.xyz file. The content of such file is provided below and can also be found under /examples/Roughness:

```
45
#Sucrose basis
0 0.63189
              0.34908
                         0.62279
              0.2018
                         0.41867
0 0.7136
0 0.6440
              -0.0665
                         0.6512
0 0.2978
              -0.0008
                         0.69117
0 0.2529
              0.3114
                         0.77094
0 0.60891
              0.40061
                         0.82857
0 0.68400
                         0.78776
              0.65323
```

(continues on next page)

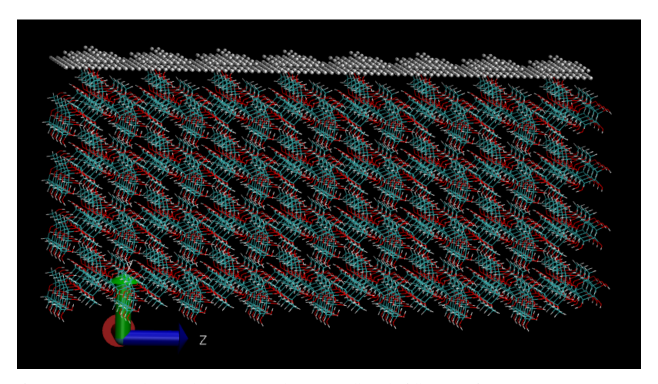
				(continued from previous page)
0	0.3785	0.5127	0.97000	
0	0.9607	0.5091	0.67341	
0	1.0893	0.6500	1.02195	
0	0.7957	0.42950	1.07412	
C	0.7053	0.1955	0.64075	
C	0.5578	0.0769	0.6265	
C	0.4362	0.1116	0.71451	
C	0.3651	0.2728	0.6871	
C	0.5149	0.3897	0.70028	
C	0.8157	0.1767	0.5431	
C	0.6306	0.5556	0.87572	
C	0.8718	0.6862	0.82381	
C	0.9441	0.5804	0.93500	
C	0.7861	0.5573	0.99233	
C	0.4569	0.6161	0.8967	
C	0.9532	0.6662	0.7110	
Η	0.7813	0.1873	0.7252	
Η	0.4894	0.0781	0.5393	
Н	0.5018	0.1046	0.8022	
Н	0.2953	0.2763	0.6004	
Н	0.4639	0.4900	0.6734	
Н	0.9127	0.2488	0.5604	
Н	0.8647	0.0743	0.5487	
Н	0.733	0.298	0.402	
Н	0.2287	0.0165	0.7364	
Η	0.2152	0.3986	0.7560	
Н	0.8878	0.7925	0.8526	
Η	0.9806	0.4827	0.9048	
Н	0.7738	0.6491	1.0414	
Н	0.4764	0.7140	0.9395	
Н	0.3769	0.6323	0.8158	
Н	0.3772	0.4263	0.9405	
Н	0.8853	0.7242	0.6409	
Н	1.0716	0.7077	0.7308	
Н	0.860	0.480	0.654	
Н	1.185	0.604	1.009	
Н	0.8058	0.3509	1.0352	
Н	0.553	-0.128	0.642	

The first run we will do needs to have the first plane listed uncommented. Note that the first plane listed will be the one used to compute the roughness. After executing lcc as follows:

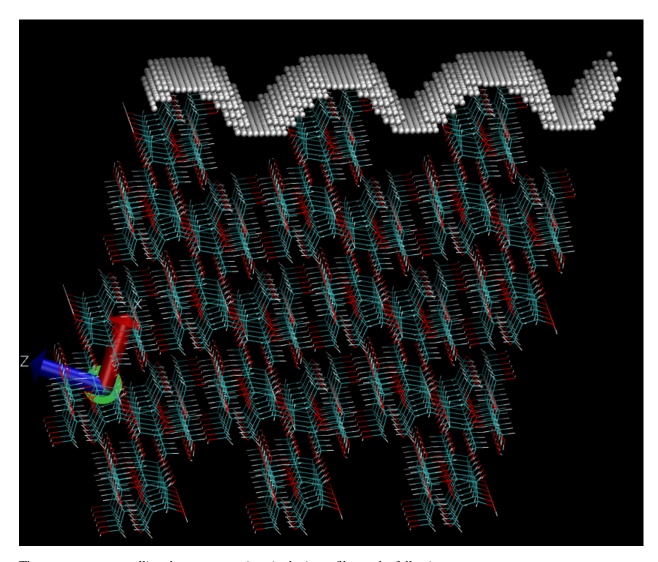
```
lcc_main sucrose.in
```

We will get information about the surfa areas S1 (effective) and S0 (flat) surface areas, together with their ratios.

The run will also produce a file called mask.xyz which contains a set of coordinates showing the surface pattern of the crystal face (vmd -e mystate.vmd).



If we now uncomment the second plane (10-1) and rerun we will see the following surface:



The parameters controlling these computations in the input file are the following:

```
ComputeRoughness= T
RoughnessParameters[
50.0 1.0 40 80 80
]
```

The first value controls the isovalue to compute the surface, the second value controls the radius of the "probe sphere" used to construct the surface, the third, fourth and fifth values control the discretization along the a1, a2, and a3 axis respectivelly.

CHAPTER

TEN

TOOLS

In this section we briefly describe some of the auxiliary tools for the code.

10.1 Kinetic Monte Carlo Crystal Growth Tool

10.1.1 About

This folder contains a Python script growth_kmc.py which performs a Kinetic Monte Carlo simulation of crystal growth from solution, starting with a spherical seed. Currently, the code supports primative tetragonal lattices, with the intent to generalize this. The core developer of this tool is Jacob Jeffries (jwjeffr@clemson.edu or jwjeffr@lanl.gov).

10.1.2 Requirements

- The python interpreter.
- The external python packages numpy and code: numba.

The example input (example_input.json) provided works for Python 3.9.12, Numpy 1.21.6, and Numba 0.55.1. Other versions are not guaranteed to be functional.

10.1.3 Testing and running the code

The code can be tested with:

python growth_kmc.py example_input.json

or:

./growth_kmc.py example_input.json

Two runs will be performed:

- A short, small run which first compiles functions. This run data will be stored in small.dump in the LAMMPS-style dump format.
- A longer run with parameters provided in example_input.json. The parameters are:

Box dimensions = (30, 30, 70) (in lattice units) Number of steps = 100,000 Dump every = 500 steps Dump file name = petn_growth.dump Initial seed radius = 75.0 angstroms Temperature = 300.0 kelvin First-neighbor cutoff distance = 7.0 angstroms Second-neighbor cutoff distance = 7.5 angstroms First-neighbor interaction energy = -0.291 electron volts Second-neighbor interaction energy = -0.186 electron volts

For new parameters, simply change the dictionary written in example_input.json to match your desired parameters.

10.1.4 Notes

It's assumed that $|\mathbf{a}| = |\mathbf{b}| = 9.088$ angstroms and $|\mathbf{c}| = 6.737$ angstroms, and that the offset for the second molecule in the unit cell is (4.54348, 4.54346, 3.36908) angstroms. These are specified in the initialize_simulation() function starting at line 134. These parameters are optimized for a PETN crystal, but are easy to change. (TODO allow the user to specify the lattice in the input file. Create lattice classes).

This code is highly parallelized, and will use all available cores. If cores are currently being used, your system might crash. (TODO allow the user to specify the number of cores to use in the input file. Default to all cores if not specified)

Adsorption and evaporation rates are both assumed to be Arrhenius with a prefactor of 1e+10 Hz. Adsorption rates are constant everywhere, with an activation energy of 0.9 electron volts, while evaporation rates are assumed to be governed by the energy change when bonds are broken. (TODO allow the user to modify the prefactors and the adsorption activation energy, or come up with a better adsorption rate expression)

10.2 Encapsulating orthogonal vectors

Code for finding encapsulating orthogonal vectors for a given lattice. Enter a given direction (surface normal) to be given two orthogonal vectors. Scalars will be generated that correspond to an error value. The error value is related to the periodicity of the orthogonal vectors.

Components in directory:

- README.md details
- orth_calc.cxx, func_v.h source code
- run.sh example run script
- latticefile.dat read in lattice vector file from command lattice=read

Compile the code with:

```
g++-std=c++11 orth_calc.cxx
```

see run. sh for example on commandline inputs Inputs:

- nx x value for direction of interest (numerical)
- ny y value for direction of interest (numerical)
- nz z value for direction of interest (numerical)
- random Selects random direction of interest (yes, no)
- lattice lattice vector input (fcc, bcc, cubic, read)
- t lattice vector scalar (numerical)
- tmax Max scalar length for direction of interest (numerical > 0)
- tstart Starting scalar for direction of interest (numerical > 0)
- tol Tolerance for Error function (numerical)
- tstep Loops through tsart to tmax with this step (numberical > 0)

42 Chapter 10. Tools

CHAPTER

ELEVEN

INDICES AND TABLES

- genindex
- modindex
- search