# MATSDP

The materials simulation and data processing toolkit

 $\begin{array}{c} {\rm Dianwu~Wang} \\ {\rm dianwuwang@163.com} \end{array}$ 

version 0.1.4 Nov. 6, 2019

# Contents

1	Intro	oduction	5
	1.1	Functions	5
	1.2	Requirements	6
	1.3	Installation	8
	1.4	Usage	8
		1.4.1 Running with Python environment	8
		1.4.2  Running Graphical User Interface (GUI) application  .	8
	1.5	Notes	9
2	auba		11
2	-	oackage: vasp	11
	2.1	vasp_build module	11
	2.2	2.1.1 vasp_build.substitution	11
	2.2	vasp_plot module	13
		2.2.1 vasp_plot.plot_poscar	13
		2.2.2 vasp_plot.plot_poscar_for_workdir	20
		2.2.3 vasp_plot.plot_dos	23
	2.3	vasp_read module	30
		2.3.1 vasp_read.read_doscar	30
	2.4	vasp_analyze module	31
		2.4.1 vasp_analyze.nn_map	31
		2.4.2 vasp_analyze.simple_cna	32
		2.4.3 vasp_analyze.estruct	33
		2.4.4 vasp_analyze.overlap_peak_analyzer	34
	2.5	vasp_write module	35
		2.5.1 vasp write write poscar with force	35

4 CC	ONTENTS
------	---------

3	subpackage: apt	37
	3.1 apt_read module	37
	3.1.1 apt_read_read_proxigram_csv	37
	3.2 apt_plot module	38
	$3.2.1$ apt_plot.plot_proxigram_csv	38
4	subpackage: dvm	41
	4.1 dvm_build module	41
	$4.1.1$ dmv_build.sphere_vasp	
5	tests	43
	5.1 example .py test file	43
Α	Other plotting settings	57
	A.1 Named colors in the program	57

# Chapter 1

## Introduction

MATSDP is a materials simulation and data processing toolkit. The Vienna ab-initio simulation package (VASP) and the Three-dimensional atom probe tomography (APT) analyzing and data processing tools are included.

## 1.1 Functions

VASP analyzing and data processing tools:

- vasp\_build.substitution: Auto substitution of element(s) based on a POSCAR file and a \*.subst file. Required input: POSCAR, .subst file
- vasp\_read: Read DOSCAR, OUTCAR, POSCAR, and OSZICAR
- vasp\_plot.plot\_poscar: Plot POSCAR/CONTCAR model (also support color mapping of atom properties), Required files: POSCAR/CONTCAR or POSCAR with data of atom properties
- vasp\_plot.plot\_dos: Plot DOS (PDOS, LDOS, TDOS) information. Required input: DOSCAR, OUTCAR, POSCAR
- vasp\_analyze.nn\_map: Calclate the nearest ngighbor inofmation. Required input: POSCAR
- vasp\_analyze.simple\_cna: Perform simple common neighbor analysis
- vasp\_analyze.estruct: Calculate structural energy ( $E_{struct}$ ). Required files: CONTCAR, OUTCAR, POSCAR

• vasp\_write.write\_poscar\_with\_force: Write atom force information into the POSCAR

APT postprocessing tools:

- apt\_read.read\_proxigram\_csv: Read the concentration profile \*.csv file
- apt\_plot.plot\_proxigram\_csv: Plot the concentration profile

The matsdp package contains the vasp and apt modules as shown in Figure 1.1. The structures of the vasp module and the apt module are shown in Figure 1.2 and Figure 1.3.

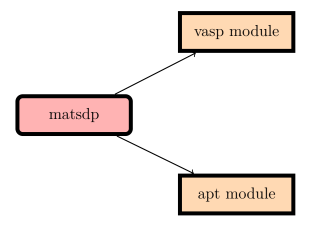


Figure 1.1: subpackages of the matsdp program.

## 1.2 Requirements

- numpy
- scipy
- scikit-learn
- matplotlib

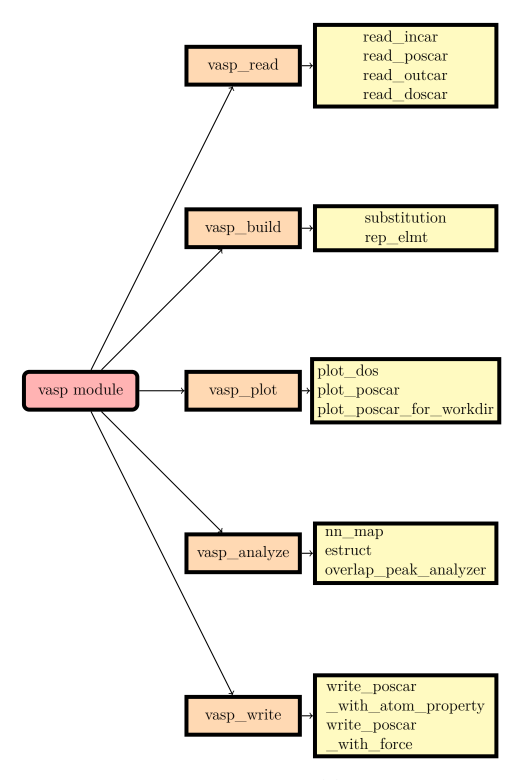


Figure 1.2: vasp module.

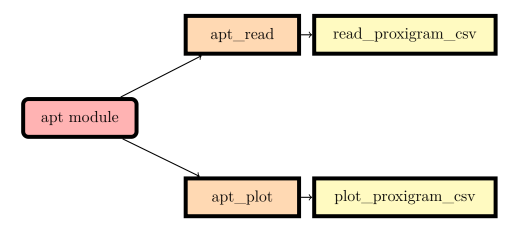


Figure 1.3: apt module.

## 1.3 Installation

For the Python users, the package can be retrieved by the following command.

```
pip install matsdp
```

For the GUI users, please run the matsdp\_gui.exe directly.

## 1.4 Usage

## 1.4.1 Running with Python environment

After installing the matsdp package, the program can be used by importing the modules and call the related functions.

## 1.4.2 Running Graphical User Interface (GUI) application

The program provides a graphical user interface (matsdp\_gui.exe). The GUI is shown in the Figure 1.4:

1.5. NOTES 9

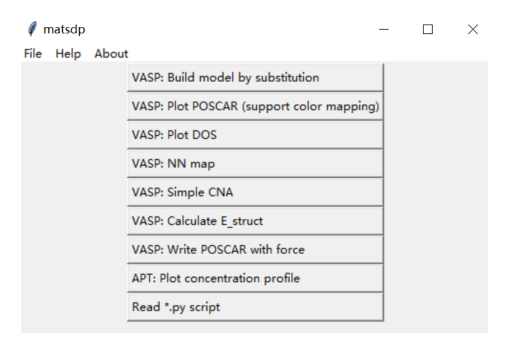


Figure 1.4: GUI for main program

## 1.5 Notes

Note that for the module that requires POSCAR/CONTCAR, OUTCAR and DOSCAR files, these files need to be in the same folder.

The following sections will introduce the settings of the parameters in the GUI.

# Chapter 2

# subpackage: vasp

Modules to import before using the vasp package

- from matsdp.vasp import vasp\_read
- from matsdp.vasp import vasp\_build
- from matsdp.vasp import vasp\_plot
- from matsdp.vasp import vasp\_analyze
- from matsdp.vasp import vasp\_write

## 2.1 vasp\_build module

## 2.1.1 vasp\_build.substitution

Descriptions

Building models by substitution of atoms

```
from matsdp.vasp import vasp_build
vasp_build.substitution(
    substitution_list_file = './example/vasp/example/
    vasp.subst',
    poscar_dir = './example/vasp/POSCAR_NoDope',
```

)

### Arguments

- substitution\_list\_file: String format. It specifies the directory of the .subst file (substitution list file)
- poscar\_dir: String format. The directory of the POSCAR file which is to be substituted. It can either be full path or relative path.

.subst file

#### Descriptions

- The .subst file (substitution list file) is required and should consists of system entries.
- A system corresponds to a specific model configuration.
- System entris specifies how atoms are substitued in different systems.
- A system entry is a block of sucessive lines without line breaks.
- Each system entris must be separated by blank lines.

File formats. A typical system entry has the following format:

```
n_subst
elment_name_to_be_substituted new_element_name
elment_name_to_be_substituted new_element_name
...
(n_subst lines of elment_name_to_be_substituted
elment_subindx new_element_name)
```

where, elment\_name\_to\_be\_substituted is he name of the element which is to be substituted. new\_element\_name is the name of the new element which take the place of the substituted atom. If new\_element\_name = Va, then a vacancy is added. As shown above, each system should start with a line which specifies a number: n\_subst. n\_subst is the number of atoms to be substituted in the system. Then each of the following n\_subst

lines specifies the element(s) to be substituted and the element(s) which take its/their place(s).

A specific example .subst file is as follow:

```
1
Ni244 W

2
Ni244 Re
Al12 Re
...
```

### GUI

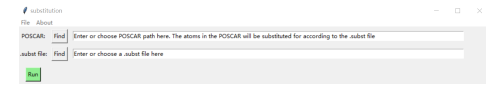


Figure 2.1: GUI for Substitution

### Outputs

Outputs: The final system name is L(line\_number)\_composition\_D(duplicate)

## 2.2 vasp\_plot module

## $2.2.1 \quad vasp\_plot.plot\_poscar$

### Descriptions

- Visualization of POSCAR model. Euler angles are used to rotate the view of the model.
- Viewer direction is in x direction. The original orientation: x direction is perpendicular to the paper, z direction is in the paper and point to upper direction

- Reference for Eulerian angles: Herbert Goldstein, Charles P. Poole Jr. and John L. Safko, Classical Mechanics (3rd Edition). Goldstein Poole & Safko, 2001.
- This module can also show the atom properties by color mapping. The POSCAR file with additional data columns used to save the data of the atom properties.

### Syntax

```
from matsdp.vasp import vasp plot
vasp_plot.plot_poscar(
    poscar_dir = './example/vasp/POSCAR',
    euler angle type = 'zyz',
    phi = -3,
    theta = 4,
    psi = 0,
    elmt_color = { 'Ni ': 'red ', 'Re': 'blue'},
    draw mirror atom = True,
    box_on = True,
    axis_indicator =True,
    plot cell basis vector label = True,
    plot_atom_label = True,
    fig_format = 'png',
    fig\_dpi = 100,
    draw colormap = False,
    colormap column indx = 1,
    colormap_vmin = None,
    colormap \quad vmax = None
    vmin color = 'blue',
    vmax\_color = 'red',
    colorbar_alignment = 'vertical'
```

### Arguments

• poscar\_dir: String format. Directory of the POSCAR file which you want to plot

• euler\_angle\_type: string of length 3. It specify the type of rotations based on Eulerian angles. Choices are 'zyz', 'zxz', 'zyx', etc.. Usually the 'zyz' type is used.

'zyz': proper Euler angle, y-convention. Performe consecutive rotations at axes counter-clockwisely. z-y-z rotation. First rotate the z axes of atoms by an angle phi, then rotate the intermidiate y axis of atoms by an angle theta, finally rotate the final z axis of atoms by an angle psi

'zxz': proper Euler angle, x-convention. Performe consecutive rotations at axes counter-clockwisely. z-x-z rotation. First rotate the z axes of atoms by an angle phi, then rotate the intermidiate x axis of atoms by an angle theta, finally rotate the final z axis of atoms by an angle psi

'zyx': Tait-Bryan angles. z-y-x rotation. Performe consecutive rotations at axes counter-clockwisely. z-y-x rotation. First rotate the z axes of atoms by an angle phi, then rotate the intermidiate y axis of atoms by an angle theta, finally rotate the final x axis of atoms by an angle psi

- phi, theta, psi: float formats. The first, second, and third rotation Eulerian angles, units in degrees.
- elmt\_color: dictionary formats. this dictionary sepcifies the color for each element. For example elmt\_color = {'Ni':'black','Al':'magenta'}
- draw\_mirror\_atom: Logical value. Whether to plot the mirror atoms at the periodic boundary
- box\_on: Logical value. Whether to plot the box or not
- axis indicator: Logic value. Whether to plot the axis indicator
- plot\_cell\_basis\_vector\_label: Logical value. Whether to plot the cell basis vector labels( i.e., to label the three basis vectors of the cell as a, b, and c)
- plot\_atom\_label: Logical value. If true, then plot the atom name of each atom.

- fig\_format: String format. Figformat is a string that defines output figure format. Supported fig\_format: 'png', 'eps', 'pdf', 'tif', 'tiff', 'jpg', 'jpeg', 'svg', 'svgz', 'pgf', 'ps', 'raw', 'rgba'
- fig\_dpi: float format. The DPI for non-vector graphics.
- draw\_colormap: Logical value. If true, the color mapping of atom properties will be performed. Default: False.
- colormap\_column\_indx: Integer value. Define which column of the atom property columns will be color mapped. Default: 1.
- colormap\_vmin: Float value. Define the minimum value of the color map. If colormap\_vmin=None, the minimum value of the original data will be used. Default: None.
- colormap\_vmax: Float value. Define the maximum value of the color map. If colormap\_vmax=None, the maximum value of the original data will be used. Default: None.
- vmin\_color = 'blue': String type. Define the color for the smallest value of the atom properties in the color map. Default: 'blue'.
- vmax\_color = 'red': String type. Define the color for the largest value of the atom properties in the color map. Default: 'red'.
- colorbar\_alignment: String type. Defines the alignment of the color bar in the figure of the color map. The value can be either 'vertical' or 'horizontal'. Default: 'vertical'.

### GUI

The GUI of the plot\_poscar module is shown in the Figure 2.2

### Outputs

Figures of POSCAR models.

### Examples

The examples are shown in the Figure 2.3, Figure 2.4, Figure 2.5 and Figure 2.6.



Figure 2.2: GUI for matsdp.vasp\_vasp\_plot.plot\_poscar

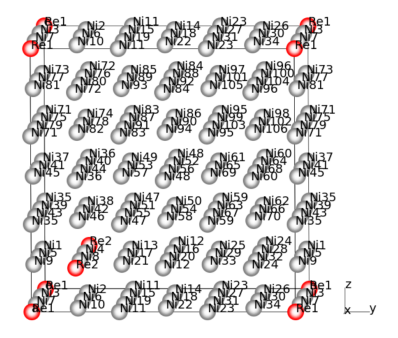


Figure 2.3: Result of the plot\_poscar module. The atom label is added.

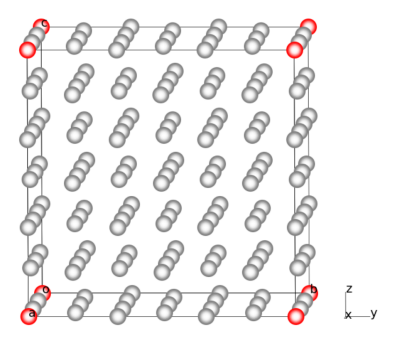


Figure 2.4: Result of the plot\_poscar module. The atom label is removed.

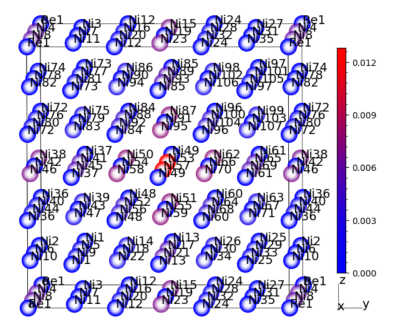


Figure 2.5: Result of the plot\_poscar module: color mapping of atom properties. The color bar is vertically aligned.

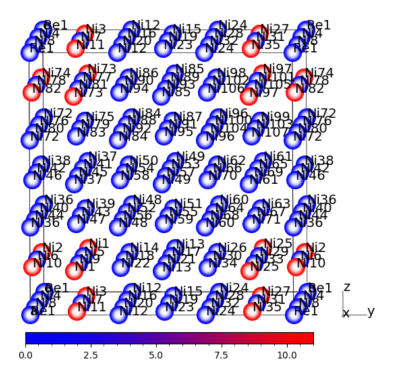


Figure 2.6: Result of the plot\_poscar module: color mapping of atom properties. The color bar is horizontally aligned.

### 2.2.2 vasp\_plot.plot\_poscar\_for\_workdir

### Descriptions

- Visualization of POSCARs.
- The mother folder needs to be specified which contains the folders with POSCARs
- Euler angles are used to rotate the view of the model
- This module can also show the atom properties by color mapping. The POSCAR file with additional data columns used to save the data of the atom properties.

```
from matsdp.vasp import vasp_plot
vasp_plot.plot_poscar_for_workdir(
    workdir = './tests/vasp/',
    euler angle type = 'zyx',
    phi = -3,
    theta = 4,
    psi = 0,
    elmt\_color = None,
    draw_mirror_atom = True,
   box_on = True,
    axis indicator =True,
    plot_cell_basis_vector_label = True,
    plot_atom_label = True,
    poscar_or_contcar = 'POSCAR',
    fig format = 'png',
    fig\_dpi = 100,
    draw_colormap = False,
    colormap\_column\_indx = 1,
    colormap vmin = None,
    colormap_vmax = None,
    vmin_color = 'blue',
    vmax\_color = 'red',
    colorbar alignment = 'vertical'
```

Arguments

# • workdir: String format. The mother folder which contains the folders with POSCARs

• euler\_angle\_type: string of length 3. It specify the type of rotations based on Eulerian angles. Choices are 'zyz', 'zxz', 'zyx', etc.. Usually the 'zyz' type is used.

'zyz': proper Euler angle, y-convention. Performe consecutive rotations at axes counter-clockwisely. z-y-z rotation. First rotate the z axes of atoms by an angle phi, then rotate the intermidiate y axis of atoms by an angle theta, finally rotate the final z axis of atoms by an angle psi

'zxz': proper Euler angle, x-convention. Performe consecutive rotations at axes counter-clockwisely. z-x-z rotation. First rotate the z axes of atoms by an angle phi, then rotate the intermidiate x axis of atoms by an angle theta, finally rotate the final z axis of atoms by an angle psi

'zyx': Tait-Bryan angles. z-y-x rotation. Performe consecutive rotations at axes counter-clockwisely. z-y-x rotation. First rotate the z axes of atoms by an angle phi, then rotate the intermidiate y axis of atoms by an angle theta, finally rotate the final x axis of atoms by an angle psi

- phi, theta, psi: float formats. The first, second, and third rotation Eulerian angles, units in degrees.
- elmt\_color: dictionary formats. this dictionary sepcifies the color for each element. For example elmt\_color = {'Ni':'black','Al':'magenta'}
- draw\_mirror\_atom: Logical value. Whether to plot the mirror atoms at the periodic boundary
- box on: Logical value. Whether to plot the box or not
- axis\_indicator: Logic value. Whether to plot the axis indicator

- plot\_cell\_basis\_vector\_label: Logical value. Whether to plot the cell basis vector labels (i.e., to label the three basis vectors of the cell as a, b, and c)
- plot\_atom\_label: Logical value. If true, then plot the atom name of each atom.
- poscar\_or\_contcar: String format. Determine whether to plot POSCAR or CONTCAR. Either 'POSCAR' or 'CONTCAR' can be used.
- fig\_format: String format. Figformat is a string that defines output figure format. Supported fig\_format: 'png', 'eps', 'pdf', 'tif', 'tiff', 'jpg', 'jpeg', 'svg', 'svgz', 'pgf', 'ps', 'raw', 'rgba'
- fig\_dpi: float format. The DPI for non-vector graphics.
- draw\_colormap: Logical value. If true, the color mapping of atom properties will be performed. Default: False.
- colormap\_column\_indx: Integer value. Define which column of the atom property columns will be color mapped. Default: 1.
- colormap\_vmin: Float value. Define the minimum value of the color map. If colormap\_vmin=None, the minimum value of the original data will be used. Default: None.
- colormap\_vmax: Float value. Define the maximum value of the color map. If colormap\_vmax=None, the maximum value of the original data will be used. Default: None.
- vmin\_color = 'blue': String type. Define the color for the smallest value of the atom properties in the color map. Default: 'blue'.
- vmax\_color = 'red': String type. Define the color for the largest value of the atom properties in the color map. Default: 'red'.
- colorbar\_alignment: String type. Defines the alignment of the color bar in the figure of the color map. The value can be either 'vertical' or 'horizontal'. Default: 'vertical'.

### Outputs

Figures of POSCAR models.

### 2.2.3 vasp plot.plot dos

Descriptions

\* Plot PDOS, LDOS, TDOS, now only available for LORBIT = 11. \* There are three types of input arguments: atom related input arguments, subplot related input arguments, and others

```
from matsdp.vasp import vasp_plot
DOS1_Dir = './ tests/vasp/DOSCAR'
vasp_plot.plot_dos(
    atom_doscar_dir_list = [DOS1_Dir],
    atom_sysname_list = ['C5'],
    atom_indx_list = ['Ni1'],
    atom_palette_list = ['black'],
    atom_subplot_arg_list = [111],
    subplot arg list = [111],
    subplot_xlo_list = [-6.5],
    subplot_xhi_list = [4.0],
    subplot_ylo_list = [None],
    subplot whi list = [None],
    subplot xtick list = [True],
    subplot_ytick_list = [True],
    subplot_xlabel_list = [False],
    subplot ylable list = [False],
    subplot_share_xy_list = [False, False],
    mainplot_axis_label_list = [True, True],
    dos\_mode = None,
    fermi shift zero = True,
    peak analyzer = False,
    fig_format = 'png',
    fig\_size = [13.0, 9.5],
    fig dpi = 600,
```

```
vasp_plot.plot_dos(
    atom_doscar_dir_list = [DOS1_Dir, DOS1_Dir],
    atom_sysname_list = ['C1', 'C1'],
    atom_indx_list = ['Ni1', 'Re1'],
    atom_palette_list = ['black', 'red'],
    atom\_subplot\_arg\_list = [111, 111],
    subplot_arg_list = [111],
    subplot_xlo_list = [-6.5],
    subplot_xhi_list = [4.0],
    subplot ylo list = [None],
    subplot_yhi_list = [None],
    subplot_xtick_list = [True],
    subplot_ytick_list = [True],
    subplot xlabel list = [False],
    subplot_ylable_list = [False],
    subplot_share_xy_list = [False, False],
    mainplot_axis_label_list = [True, True],
    dos_mode = { 'Ni ': [ 'd '], 'Re': [ 'd']},
    fermi_shift_zero = True,
    peak_analyzer = False,
    fig_format = 'png',
    fig\_size = [11.0, 9.5],
    fig\_dpi = 600,
```

### Arguments

#### Atom related Args

- atom\_doscar\_dir\_list: list format. Conatains DOSCAR files for each atom. The directory of DOSCAR files can either be full path or relative path
- atom\_sysname\_list: system name for each atom, it corresponds to the atoms in the atom\_doscar\_dir\_list. This is for the purpose of labeling the DOS curves in the legend.

If sysnameList = None, then the label of system name will not shown in the legend

For example, sysnameList = ['System1', 'System1', 'System2']

• atom\_indx\_list: list format. Atom index list, it corresponding to the atoms in atom\_doscar\_dir\_list. If it is integer type then it denotes the atom index, if it is string type then it denotes the atom name

atom\_indx\_list = [1,2,45] denotes the 1st, 2nd, and the 45th atoms in the POSCAR

atom\_indx\_list = ['Ni1','Al3','Re3'] denotes Ni1, Al3, and Re3 in the POSCAR

atom\_indx\_list = ['TDOS'] and atom\_indx\_list = [0] donotes the total dos

- atom palette list: list format. Color for DOS curves of each atom.
- atom\_subplot\_arg\_list: list format. Defines the DOS curves of the atom are in which subplot. For example, atom\_subplot\_arg\_list = [221, 222] denotes that the DOS curves of the first and the second atoms are in the subplot(221) and subplot(222) subplots, respectively.

### Subplot related Args

- subplot\_arg\_list: list format. The subplot argument list, for example subplot\_arg\_list=[221,222] corresponds to subplot(221) and subplot(222)
- subplot\_xlo\_list: list format. Low boundary of the x axis for each subplots. If None value is given, the low boundary of x axis in the data set will be chosen.
- subplot\_xhi\_list: list format. High boundary of the x axis for each subplots. If None value is given, the high boundary of x axis in the data set will be chosen.
- subplot\_ylo\_list: list format. Low boundary of the y axis for each subplots. If None value is given, the low boundary of y axis in the data set will be chosen.

- subplot\_yhi\_list: list format. High boundary of the y axis for each subplots. If None value is given, the high boundary of y axis in the data set will be chosen.
- subplot\_xtick\_list: list of logical values. If the list element is True (False), then the tick of the x axis will be shown (removed).
- subplot\_ytick\_list: list of logical values. If the list element is True (False), then the tick of the x axis will be shown (removed).
- subplot\_xlabel\_list: list of logical values. Defines whether the x-label of each subplots are shown, it won't work for subplot=(111) figure.
- subplot\_xlabel\_list: list of logical values. Defines whether the y-label of each subplots are shown, it won't work for subplot=(111) figure.
- subplot\_share\_xy\_list: list of logical values of length two. Defines whether the x or y axis will be shared or not. [False, False] denotes both x and y axes will not be shared.

### Other Args

- mainplot\_axis\_label\_list: list of logical values of length two. Defines whether the x or y labels of the main figure will be shown or not. [False, False] denotes both x and y labels of the main figure will not be shown.
- dos\_mode is a dictionary that defines which partial DOS or whether LDOS is plotted for different element type. e.g.: dos\_mode = {'Ni':['s','p','d'], 'Al':['s','p']} or dos\_mode = {'Ni':['dxy','dx2']}, or dos\_mode = {'Ni':['LDOS']}.
- fermi\_shift\_zero is a logical value which determines whether to shift Fermi energy level to zero.
- peak\_analyzer:logical value. Determines whether to analyze peaks in DOS. if True, the peaks will be labeled
- Figformat: String format. Defines output figure format. Supported fig\_format: 'png', 'eps', 'pdf', 'tif', 'tiff', 'jpg', 'jpeg', 'svg', 'svgz', 'pgf', 'ps', 'raw', 'rgba'
- fig\_size: list of floats. Defines the size of the figure, e.g. fig\_size = (7.0,6.0)
- fig\_dpi: float format. The DPI for non-vector graphics.

27

#### GUI

The GUI is shown in Figure 2.7. The panel can be devided into several control regions and the several control regions are shown in Figure 2.8. The settings for the plot\_dos function is shown in Figure 2.9. The subplot layout is shown in Figure 2.10

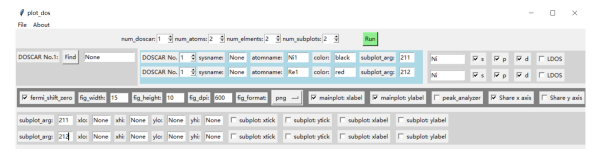


Figure 2.7: GUI for plot dos

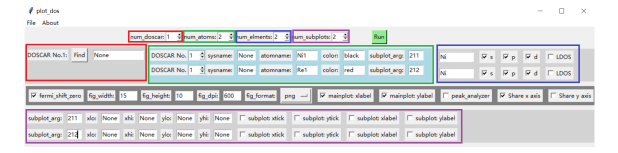
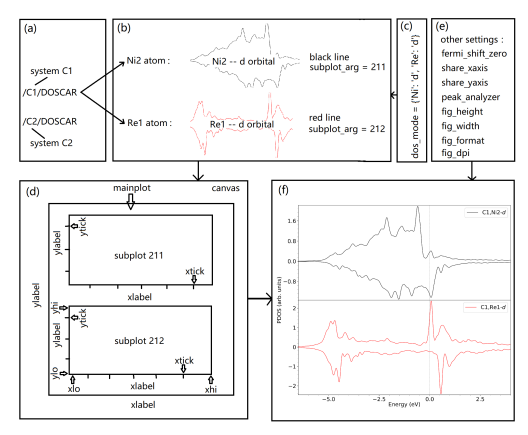


Figure 2.8: Control regions in the plot dos panel

Some of the parameters are listed below:

- num\_doscar: Number of DOSCAR files, this region can be used to import different DOSCAR
- num\_atoms: Number of atoms for plotting the DOSs
- num\_elements: Number of elements
- num\_subplots: Number of subplots
- subplot\_arg: The position of the subplot. The illustration of the subplot is shown in Fig. 2.10



- (a) DOSCAR related settings; (b) atom related settings; (c) element related settings;
- (d) subplot related settings; (e) other settings; (f) figure output

Figure 2.9: plot\_dos settings

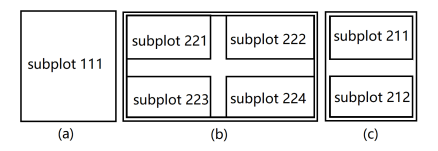


Figure 2.10: Subplot layout

If only one DOS curve will be plotted, then set num\_doscar=1 and num\_atom=1. The value of subplot\_arg then can be subplot\_arg=111. For example, if the PDOSs of "Ni1" and "A2" are to be compared, the parameter num\_atom should be taken as num\_atom=2.

### Output

Figures of DOS curves

### Examples

The examples are shown in the Figure 2.11 and Figure 2.12.

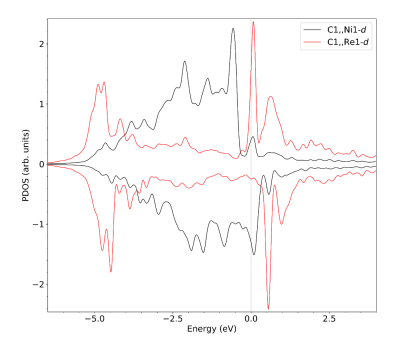


Figure 2.11: Result of the plot\_doscar module.

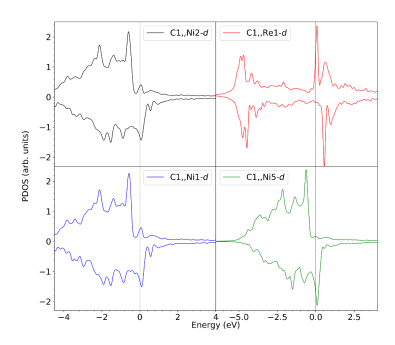


Figure 2.12: Result of the plot $\_$ doscar module.

## 2.3 vasp\_read module

## 2.3.1 vasp\_read.read\_doscar

### Descriptions

Read DOSCAR and dump density of states in oformation file into the folder where the DOSCAR lies

```
from matsdp.vasp import vasp_read
vasp_read.read_doscar(
   doscar_dir = './tests/vasp/DOSCAR',
   atom_indx = 2,
   save_dos_arr = True,
)
```

### Arguments

- doscar\_dir: String format. The directory of the DOSCAR file. It can either be full path or relative path
- atom\_indx: Integer format. The real atom index in the POSCAR. If there are N atoms then the atom indices are frim 1 to N. Note that atom indx = 0 means to extract TDOS information
- save\_dos\_arr: logical format. If save\_dos\_arr = True, the density of states inoformation will be saved to files. If save\_dos\_arr = False, the density of states inoformation will not be saved to files

### Outputs

DOS information file for the specified atom

## 2.4 vasp analyze module

## 2.4.1 vasp\_analyze.nn\_map

Calculate the nearest neighbor (NN) map.

### Descriptions

Calculate the nearest neighbor (NN) map.

```
from matsdp.vasp import vasp_analyze
vasp_analyze.nn_map(
   poscar_dir = './tests/vasp/POSCAR',
   a0 = 3.545,
   n_shell = 2,
   )
```

### Args

- poscar\_dir: String format. It specifies the directory of the POSCAR file
- a0: Float format. The lattice constant of the model. Unit in Angstrom
- n\_shell: Integer format. It determines up to which crystallographic shell the nearest neighbour map calculates

### GUI



Figure 2.13: GUI for nn\_map

## 2.4.2 vasp\_analyze.simple\_cna

Peform simple common neighbor analysis (CNA).

### Descriptions

Peform simple common neighbor analysis (CNA). Atom A is the common neighbor of element E1 and E2, this module will count the times that A appeared as the common neighbor of E1 and E2.

```
from matsdp.vasp import vasp_analyze
vasp_analyze.simple_cna(
    poscar_dir = './tests/vasp/POSCAR',
    a0 = 3.545,
    common_neighbor_elmt_list = ['Re', 'W', 'Ta', 'Ni']
    )
```

### Args

- poscar\_dir: String format. It specifies the directory of the POSCAR file
- a0: Float format. The lattice constant of the model. Unit in Angstrom
- common\_neighbor\_elmt\_list: List format. It determines what elements are taken into account in the common neighbor analysis. If common\_neighbor\_elmt\_list = ['Re', 'W', 'Ta'], then the common neighbor to Re-Re, Re-W, Re-Ta, W-W, W-Ta, Ta-Ta will be counted and printed.

### GUI



Figure 2.14: GUI for simple\_cna

### 2.4.3 vasp\_analyze.estruct

### Descriptions

- Calculates the structural energies at each atomic site
- The definition of  $E_{struct}$  can be found in the literature of the author Chongyu Wang [2, 3]

```
from matsdp.vasp import vasp_analyze
vasp_analyze.estruct(
   doscar_dir = './tests/vasp/DOSCAR',
   sysname = 'DOS1',
)
```

### Arguments

- dOSCAR\_dir: String format. It specifies the directory of the DOSCAR
- sysname: String format. User defined system name

### GUI



Figure 2.15: GUI for estruct

### Output

The first column is the atom name, the second column is  $E_{struct}$  for each atom

### 2.4.4 vasp analyze.overlap peak analyzer

### Descriptions

- Finding the overlapped orbitals of two neighboring atoms in the DOS analyis.
- DOS peak analyses for selected atoms with their neighboring atoms.
- Find the overlapped orbitals and their corresponding energy levels.

```
from matsdp.vasp import vasp_analyze
vasp_analyze.overlap_peak_analyzer(
    doscar_dir = './tests/vasp/DOSCAR',
    sysname = 'DOS1',
    atom_indx_list = ['Ni1', 'Re1'],
```

```
n_shell = 2,
a0 = 3.52,
dos_mode = {'Ni':['d'], 'Re':['d']},
fermi_shift_zero = True,
)
```

### Arguments

- doscar\_dir: String format. The directory which contains the DOSCAR file, abstract path can be accepted
- sysname: String format. A string character which specifies the name of the system, this string will be used as part of the output file name
- atom\_indx\_list: List of strings. Specifies the list of selected atoms.
- n\_shell: float format. Up to which crystallographic shell(up to which nearest neighbor) of the selected atom will be considered
- a0: float format. The approximate lattice constant of the crystal
- dos\_mode: dictionary format. Determins which orbital will be considered, f, d, p, s, dxy, dyz, ... can be used
- fermi\_shift\_zero: A logical value determining whether the energy levels of the DOS will be shifted to zero

#### Outputs

overlapped peak files

## 2.5 vasp write module

### 2.5.1 vasp write.write poscar with force

### Descriptions

write atom force data into the POSCAR file.

### Syntax

```
from matsdp.vasp import vasp_write
vasp_write.write_poscar_with_force(
   outcar_dir = './tests/vasp/OUTCAR',
   ionic_step = 'last',
   output_poscar_file_name = None
)
```

### Arguments

- outcar dir: String format. It specifies the directory of the OUTCAR
- ionic\_step: String format or interger type. If string type value is taken, either ionic\_step='last' or ionic\_step='first' can be taken. If integer type value is taken, ionic\_step defines the ionic step number. ionic\_step = 3 denotes that the force of each atom for the third ionic step will be written to the POSCAR file.
- output\_poscar\_file\_name: String type or None. If string type is taken, this parameter lets the user define the POSCAR file name which contains the atom force information. If output\_poscar\_file\_name=None, the program determines the name of the output POSCAR file.

### GUI



Figure 2.16: GUI for write\_poscar\_with\_force

### Output

The POSCAR file which contains the force on each atom.

## Chapter 3

# subpackage: apt

Modules to import before using the apt package

- $\bullet \;$  from matsdp.apt import apt\_read
- from matsdp.apt import apt\_plot
- 3.1 apt\_read module
- 3.1.1 apt\_read.read\_proxigram\_csv

### Descriptions

- Read the concentration profile file (\*.csv file)

### Syntax

```
from matsdp.apt import apt_read apt_read.read_proxigram_csv(proxigram_csv_dir)
```

#### Arguments

• proxigram\_csv\_dir: string type. The concentration profile file.

#### Outputs

- data\_set: numpy array type. The original data of the concentration profile file
- elmtname\_list: List type. The elements contained in the concentration profile file
- 3.2 apt\_plot module
- 3.2.1 apt\_plot.plot\_proxigram\_csv

#### Descriptions

• Plot the concentration profile based on the proxigram \*.csv file

#### Syntax

```
from matsdp.apt import apt_plot
apt_plot.plot_proxigram_csv(
    proxigram_csv_dir = './tests/apt/profile-
    interface0.csv',
    sysname = 'M2',
    visible_elmt_list = ['Ni', 'Al'],
    interplation_on = False,
    fig_width = 6,
    fig_height = 5,
    fig_dpi = 600,
    fig_format = 'png',
)
```

#### Arguments

- proxigram\_csv\_dir: string type. The concentration profile file.
- sysname: string type. The system name.
- visible\_elmt\_list: List type. The elements which are to be plotted. For example, ['Ni','Al'].

- interpolation\_on: logical type. whether to interpolate the concentration profile or not.
- fig width: float type. Figure width.
- fig\_height: float type. Figure height.
- fig\_dpi: float format. The DPI for non-vector graphics.
- fig\_format: String format. Figformat is a string that defines output figure format. Supported fig\_format: 'png', 'eps', 'pdf', 'tif', 'tiff', 'jpg', 'jpeg', 'svg', 'svgz', 'pgf', 'ps', 'raw', 'rgba'

GUI



Figure 3.1: GUI for plot\_concentration\_profile

### Examples

The example is shown in the Figure 2.11.

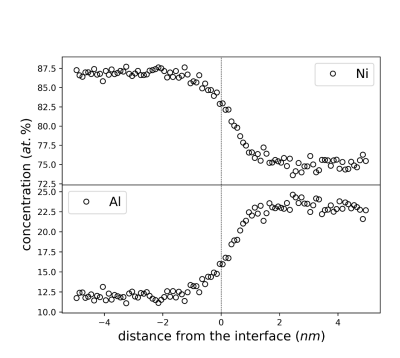


Figure 3.2: Result of the plot\_proxigram\_csv module.

## Chapter 4

## subpackage: dvm

Modules to import before using the dvm package

• from matsdp.dvm import dvm\_build

## 4.1 dvm build module

### 4.1.1 dmv\_build.sphere\_vasp

### Descriptions

• generate model (\*.incar file) for the DVM program, select atoms in a sphere that is centered around an arbitrary atom from the POSCAR file

### Syntax

```
from matsdp.dvm import dvm_build dvm_build.sphere_vasp(poscar_dir, origin_atom_name, radius, include_mirror_atoms, dvm_incar_file_name)
```

### Arguments

• radius: float type. The atoms within a distance "radius" from the original atom are selected (units in Angstroms)

- include\_mirror\_atoms: Logical value. Whether to include the mirror atoms or not
- dvm\_incar\_file\_name: user-defined dvm job name

### Outputs

- $\bullet \ \ dvm\_incar\_file\_name.incar$
- $\bullet \ \ dvm\_incar\_file\_name.vasp$
- $\bullet$  dvm\_incar\_file\_name.xyz

## Chapter 5

## tests

## 5.1 example .py test file

The example.py test file is provided as runtests.py. Its content is listed below. The parameter "package\_path" is used to define the matsdp package directory and can be modified by the users.

```
# -*- coding: utf-8 -*-
import os
import sys
package_path = './'
sys.path.insert(0, os.path.abspath(package_path))
from matsdp.vasp import vasp_read
from matsdp.vasp import vasp_plot
from matsdp.vasp import vasp_analyze
from matsdp.vasp import vasp_build
from matsdp.vasp import vasp write
from matsdp.apt import apt_read
from matsdp.apt import apt_plot
from matsdp.dvm import dvm_build
run_nn_map = True
run_simple_cna = True
run_substitute = True
run replace elmt = True
```

```
run_get_doscar = True
run_plot_dos = True
run plot poscar = True
run_plot_poscar_for_workdir = True
run_overlap_peak_analyzer = True
run estruct = True
run_write_poscar_with_force = True
run_plot_concentration_profile = True
run_dvm_build = True
# nn_map_Calc
if run_nn_map == True:
    vasp_analyze.nn_map(
       poscar_dir = './tests/vasp/POSCAR',
       a0 = 3.545,
       n \text{ shell} = 2
# simple_common_neighbor
if run_simple_cna == True:
    vasp_analyze.simple_cna(
       poscar_dir = './tests/vasp/POSCAR',
       a0 = 3.545,
       common_neighbor_elmt_list = ['Re', 'W', 'Ta','
       Ni']
       )
    vasp plot.plot poscar (
       poscar_dir = './outputs/
       POSCAR simple common neighbor pair count ReNi'
       euler angle type = 'zyz',
       phi = -3,
       theta = 5,
       psi = 0,
       elmt_color = { 'Ni ': 'red ', 'Re': 'blue'},
```

```
draw_mirror_atom = True,
    box_on = True,
    axis indicator =True,
    plot_cell_basis_vector_label = True,
    plot_atom_label = True,
    fig_format = 'png',
    fig dpi = 100,
    draw colormap = True,
    colormap\_column\_indx = 2,
    colormap_vmin = None,
    colormap \quad vmax = None
    vmin_color = 'blue',
    vmax_color = 'red',
    colorbar_alignment = 'vertical'
vasp_plot.plot_poscar(
    poscar_dir = './outputs/
    POSCAR_simple_common_neighbor_pair_count_ReNi '
    euler_angle_type = 'zyz',
    phi = -3,
    theta = 5,
    psi = 0,
    elmt_color = { 'Ni ': 'red ', 'Re': 'blue'},
    draw_mirror_atom = True,
    box on = True,
    axis indicator =True,
    plot_cell_basis_vector_label = True,
    plot_atom_label = True,
    fig format = 'png',
    fig\_dpi = 100,
    draw colormap = True,
    colormap\_column\_indx = 2,
    colormap vmin = None,
    colormap_vmax = None,
    vmin_color = 'blue',
    vmax\_color = 'red',
    colorbar alignment = 'horizontal'
```

```
)
# run_substitute
if run substitute == True:
   vasp build.substitution (
       substitution_list_file = './tests/vasp/example
       poscar_dir = './tests/vasp/POSCAR_NoDope',
# run_replace_elmt
if run_replace_elmt == True:
   vasp_build.rep_elmt(
       substitution_list_file = './tests/vasp/example
       .subst',
       poscar dir = './tests/vasp/POSCAR NoDope',
       old_elmt= 'Re',
       elmt\_group = ['W', 'Cr'],
#plot_dos
if run_plot_dos == True:
   DOS1_Dir = './ tests/vasp/DOSCAR'
   vasp_plot.plot_dos(
       atom_doscar_dir_list = [DOS1_Dir],
       atom sysname list = ['C5'],
       atom_indx_list = ['Ni1'],
       atom_palette_list = ['black'],
       atom_subplot_arg_list = [111],
       subplot arg list = [111],
       subplot_xlo_list = [-6.5],
       subplot_xhi_list = [4.0],
       subplot_ylo_list = [None],
       subplot whi list = [None],
```

```
subplot_xtick_list = [True],
    subplot_ytick_list = [True],
    subplot xlabel list = [False],
    subplot_ylabel_list = [False],
    subplot_share_xy_list = [False, False],
    mainplot_axis_label_list = [True, True],
    dos mode = None,
    fermi shift zero = True,
    peak_analyzer = False,
    fig_format = 'png',
    fig size = [13.0, 9.5],
    fig\_dpi = 600,
vasp_plot.plot_dos(
    atom_doscar_dir_list = [DOS1_Dir, DOS1_Dir],
    atom_sysname_list = ['C1', 'C1'],
    atom_indx_list = ['Ni1', 'Re1'],
    atom_palette_list = ['black', 'red'],
    atom subplot arg list = [111, 111],
    subplot_arg_list = [111],
    subplot_xlo_list = [-6.5],
    subplot xhi list = [4.0],
    subplot_ylo_list = [None],
    subplot_yhi_list = [None],
    subplot_xtick_list = [True],
    subplot ytick list = [True],
    subplot_xlabel_list = [False],
    subplot_ylabel_list = [False],
    subplot_share_xy_list = [False, False],
    mainplot axis label_list = [True, True],
    dos_mode = { 'Ni ': [ 'd '], 'Re ': [ 'd ']},
    fermi_shift_zero = True,
    peak_analyzer = False,
    fig format = 'png',
    fig\_size = [11.0, 9.5],
    fig\_dpi = 600,
```

```
vasp_plot.plot_dos(
    atom_doscar_dir_list = [DOS1_Dir, DOS1_Dir],
    atom sysname_list = ['C1', 'C1'],
    atom_indx_list = ['Ni2', 'Re1'],
    atom_palette_list = ['black', 'red'],
    atom\_subplot\_arg\_list = [211, 212],
    subplot arg list = [211, 212],
    subplot xlo list = [-6.5, -6.5],
    subplot_xhi_list = [4.0, 4.0],
    subplot_ylo_list = [None, None],
    subplot whi list = [None, None],
    subplot_xtick_list = [True, True],
    subplot_ytick_list = [True, True],
    subplot_xlabel_list = [False, False],
    subplot ylabel list = [False, False],
    subplot_share_xy_list = [False, False],
    mainplot_axis_label_list = [True, True],
    dos_mode = { 'Ni ':[ 'd '], 'Re ':[ 'd ']},
    fermi shift zero = True,
    peak_analyzer = False,
    fig_format = 'png',
    fig size = [11.0, 9.5],
    fig\_dpi = 600,
vasp plot.plot dos(
    atom_doscar_dir_list = [DOS1_Dir, DOS1_Dir],
    atom_sysname_list = ['C1', 'C1'],
    atom_indx_list = ['Ni2', 'Re1'],
    atom palette list = ['black', 'red'],
    atom\_subplot\_arg\_list = [211, 212],
    subplot_arg_list = [211, 212],
    subplot_xlo_list = [-6.5, -6.5],
    subplot xhi list = [4.0, 4.0],
    subplot_ylo_list = [None, None],
    subplot_yhi_list = [None, None],
    subplot_xtick_list = [False, True],
    subplot ytick list = [True, True],
```

```
subplot_xlabel_list = [False, False],
    subplot_ylabel_list = [False, False],
    subplot share xy list = [True, False],
    mainplot_axis_label_list = [True, True],
    dos_mode = { 'Ni ':[ 'd '], 'Re ':[ 'd ']},
    fermi_shift_zero = True,
    peak_analyzer = False,
    fig_format = 'png',
    fig\_size = [11.0, 9.5],
    fig\_dpi = 600,
    )
vasp_plot.plot_dos(
    atom_doscar_dir_list = [DOS1_Dir, DOS1_Dir,
   DOS1_Dir, DOS1_Dir,
    atom_sysname_list = [ 'C1', 'C1', 'C1', 'C1'],
    atom_indx_list = ['Ni2', 'Re1', 'Ni1',
    atom_palette_list = ['black', 'red', 'blue',
    green'],
    atom\_subplot\_arg\_list = [221, 222, 223, 224],
    subplot arg list = [221, 222, 223, 224],
    subplot_xlo_list = [-4.5, -6.5, -4.5, -6.5],
    subplot_xhi_list = [4.0, 4.0, 4.0, 4.0],
    subplot_ylo_list = [None, None, None, None],
    subplot whi list = [None, None, None, None],
    subplot_xtick_list = [True, True, True, True],
    subplot_ytick_list = [True, True, True, True],
    subplot_xlabel_list = [False, False, False,
    False].
    subplot_ylabel_list = [False, False, False,
    False,
    subplot_share_xy_list = [False, False],
    mainplot axis label list = [True, True],
    dos\_mode = \{ 'Ni' : [ 'd'], 'Re' : [ 'd'] \},
    fermi_shift_zero = True,
    peak_analyzer = False,
    fig format = 'png',
```

```
fig\_size = [11.0, 9.5],
        fig\_dpi = 600,
    vasp_plot.plot_dos(
        atom_doscar_dir_list = [DOS1_Dir, DOS1_Dir,
        DOS1_Dir, DOS1_Dir],
        atom_sysname_list = [ 'C1', 'C1', 'C1', 'C1'],
        atom_indx_list = ['Ni2', 'Re1', 'Ni1',
                                             'Ni5'].
        atom_palette_list = ['black', 'red', 'blue',
        green'],
        atom subplot arg list = [221, 222, 223, 224],
        subplot_arg_list = [221, 222, 223, 224],
        subplot_xlo_list = [-4.5, -6.5, -4.5, -6.5],
        subplot_xhi_list = [4.0, 4.0, 4.0, 4.0],
        subplot_ylo_list = [-2.3, -2.3, -2.3, -2.3],
        subplot_yhi_list = [2.5, 2.5, 2.5, 2.5],
        subplot_xtick_list = [False, False, True, True
        subplot ytick list = [True, False, True, False
        subplot_xlabel_list = [False, False, False,
        False],
        subplot_ylabel_list = [False, False, False,
        False,
        subplot_share_xy_list = [True, True],
        mainplot_axis_label_list = [True, True],
        dos_mode = { 'Ni ': [ 'd '], 'Re ': [ 'd ']},
        fermi_shift_zero = True,
        peak_analyzer = False,
        fig format = 'png',
        fig\_size = [11.0, 9.5],
        fig\_dpi = 600,
        )
# overlap_peak_analyzer
if run_overlap_peak_analyzer == True:
    vasp analyze.overlap peak analyzer (
```

```
doscar_dir = './tests/vasp/DOSCAR',
       sysname = 'DOS1',
       atom indx list = ['Ni1', 'Re1'],
       n_shell = 2,
       a0 = 3.52,
       dos_mode = { 'Ni ': [ 'd'], 'Re': [ 'd']},
       fermi shift zero = True,
#Get DOS files with DOS info
if run_get_doscar == True:
   poscar_dir = './tests/vasp/POSCAR'
   poscar_dict = vasp_read.read_poscar(poscar_dir)
   for atom_indx in range(0, len(poscar_dict['
   atom_species_arr']) + 1):
       vasp_read.read_doscar(
           doscar_dir = './tests/vasp/DOSCAR',
           atom indx = atom indx,
           save_dos_arr = True,
#Visualization of POSCAR
if run_plot_poscar == True:
   vasp_plot.plot_poscar(
       poscar_dir = './tests/vasp/POSCAR',
       euler_angle_type = 'zyz',
       phi = -3,
       theta = 5,
       psi = 0,
       elmt_color = { 'Ni ': 'red ', 'Re': 'blue '},
       draw mirror atom = True,
       box_on = True,
       axis_indicator =True,
       plot_cell_basis_vector_label = True,
       plot atom label = True,
```

```
fig_format = 'png',
        fig\_dpi = 100,
        draw colormap = False,
        colormap\_column\_indx = 1,
        colormap_vmin = None,
        colormap_vmax = None,
        vmin_color = 'blue',
        vmax color = 'red',
        colorbar_alignment = 'vertical'
# write_poscar_with_force
if run_write_poscar_with_force == True:
   # write_poscar_with_force
   vasp_write.write_poscar_with_force(
        outcar_dir = './tests/vasp/OUTCAR',
        ionic step = 'last',
        output_poscar_file_name = None
    vasp_plot.plot_poscar(
        poscar_dir = './outputs/
        POSCAR_with_force_step_1,
        euler_angle_type = 'zyz',
        phi = -3,
        theta = 5,
        psi = 0,
        elmt_color = { 'Ni ': 'red ', 'Re': 'blue '},
        draw mirror atom = True,
       box_on = True,
        axis indicator =True,
        plot_cell_basis_vector_label = True,
        plot_atom_label = True,
        fig_format = 'png',
        fig\_dpi = 100,
        draw\_colormap = True,
        colormap column indx = 1,
```

```
colormap_vmin = None,
       colormap_vmax = None,
       vmin color = 'blue',
       vmax\_color = 'red',
       colorbar_alignment = 'vertical'
    vasp_plot.plot_poscar(
       poscar_dir = './outputs/
       POSCAR_with_force_step_1_absforce',
       euler_angle_type = 'zyz',
       phi = -3
       theta = 5,
       psi = 0,
       elmt_color = { 'Ni ': 'red ', 'Re': 'blue'},
       draw mirror atom = True,
       box_on = True,
       axis_indicator =True,
       plot_cell_basis_vector_label = True,
       plot atom label = True,
       fig_format = 'png',
       fig\_dpi = 100,
       draw colormap = True,
       colormap\_column\_indx = 1,
       colormap_vmin = None,
       colormap_vmax = None,
       vmin color = 'blue',
       vmax color = 'red',
       colorbar_alignment = 'vertical'
╫╫╫╫╫╫╫╫╫╫╫╫╫╫╫╫╫╫╫╫
#run_plot_poscar for the POSCARs in a directory
if run plot poscar for workdir = True:
    vasp_plot.plot_poscar_for_workdir(
       workdir = './outputs/example/',
       euler_angle_type = 'zyx',
       phi = -3,
```

```
theta = 5,
        psi = 0,
        elmt color = None,
        draw_mirror_atom = True,
        box_on = True,
        axis_indicator =True,
        plot_cell_basis_vector_label = True,
        plot_atom_label = True,
        poscar_or_contcar = 'POSCAR',
        fig_format = 'png',
        fig dpi = 100,
        draw\_colormap = False,
        colormap\_column\_indx = 1,
        colormap_vmin = None,
        colormap_vmax = None,
        vmin_color = 'blue',
        vmax_color = 'red',
        colorbar_alignment = 'vertical'
# run_estruct
if run_estruct == True:
    vasp_analyze.estruct(
        doscar_dir = './tests/vasp/DOSCAR',
        sysname = 'DOS1',
        )
    vasp_plot.plot_poscar(
        poscar_dir = './outputs/POSCAR_estruct_DOS1_Ef
        -7.0888,
        euler_angle_type = 'zyz',
        phi = -3,
        theta = 5,
        psi = 0,
        elmt_color = { 'Ni ': 'red ', 'Re': 'blue'},
        draw_mirror_atom = True,
        box_on = True,
        axis indicator =True,
```

```
plot_cell_basis_vector_label = True,
    plot_atom_label = True,
    fig format = 'png',
    fig\_dpi = 100,
    draw_colormap = True,
    colormap\_column\_indx = 1,
    colormap_vmin = None,
    colormap_vmax = None,
    vmin_color = 'blue',
    vmax\_color = 'red',
    colorbar_alignment = 'vertical'
vasp_plot.plot_poscar(
    poscar_dir = './outputs/POSCAR_estruct_DOS1_Ef
    -7.0888,
    euler_angle_type = 'zyz',
    phi = -3,
    theta = 5,
    psi = 0,
    elmt_color = { 'Ni ': 'red ', 'Re': 'blue '},
    draw mirror atom = True,
    box_on = True,
    axis_indicator =True,
    plot_cell_basis_vector_label = True,
    plot atom label = True,
    fig_format = 'png',
    fig\_dpi = 100,
    draw\_colormap = True,
    colormap column indx = 1,
    colormap_vmin = -80,
    colormap vmax = -40,
    vmin_color = 'blue',
    vmax color = 'red',
    colorbar_alignment = 'vertical'
```

```
#apt- plot concentration profile
if run_plot_concentration_profile == True:
   apt_plot.plot_proxigram_csv(
       proxigram_csv_dir = './tests/apt/profile-
       interface0.csv',
       sysname = 'M2',
       visible_elmt_list = ['Ni', 'Al'],
       interplation_on = False,
       fig_width = 6,
       fig height = 5,
       fig\_dpi = 600,
       fig_format = 'png',
# dvm build
if run dvm build = True:
   dvm_build.sphere_poscar(
       poscar_dir = './tests/vasp/CONTCAR',
       origin atom name = 'Re1',
       radius = 7,
       include_mirror_atoms = True,
       dvm_incar_file_name = 'dvm_example'
```

# Appendix A

# Other plotting settings

## A.1 Named colors in the program

The named colors which can be used by the program is listed in Figure  $A.1^1$ .

<sup>&</sup>lt;sup>1</sup>https://matplotlib.org/3.1.0/gallery/color/named\_colors.html

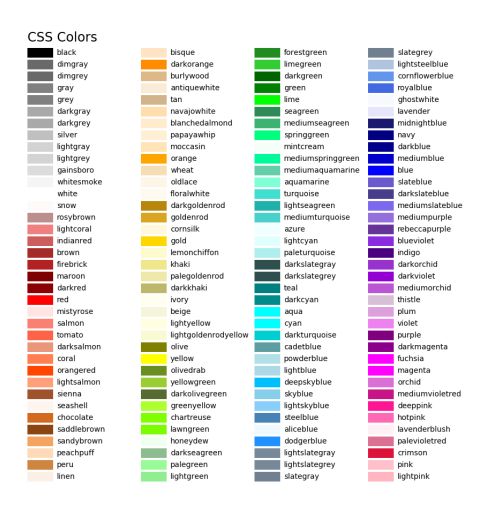


Figure A.1: The named colors supported by the current program

## Bibliography

- [1] Herbert Goldstein, Charles P. Poole Jr. and John L. Safko, Classical Mechanics (3rd Edition). Goldstein Poole & Safko, 2001.
- [2] Chongyu Wang and Feng An and Binglin Gu and Liu Fusui and Ying Chen, Electronic structure of the light-impurity (boron)—vacancy complex in iron. Physical Review B, 1988: 3905–3912.
- [3] Chong-yu Wang, Sen-ying Liu and Lin-guang Han, Electronic structure of impurity (oxygen)—stacking-fault complex in nickel. Physical Review B, 1990: 1359–1367.