

# MCCE Program Run and Debug

## Commands and Tools

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### getpdb

*Command line tool to download pdb file from PDB*

**Author:** Yanjun Wang

**Syntax:**

```
getpdb pdbID [file]
```

This program gets a pdb file from Protein Data Bank, and saves to a file with its PDB name or a user named file.

**Example:**

```
getpdb 1akk  
Download with url https://files.rcsb.org/download/1akk.pdb.  
Download completed.
```

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### vdw\_bk.py

*Calculate conformer to backbone vdw (Lennard-Jones potential).*

**Author:** Junjun Mao

**Syntax:**

```
vdw_bk.py [-h] [-c cutoff] confID
```

Compute vdw1 breakdown at conformer level. The conformer ID can be found in head3.lst.

positional arguments:

confID      Conformer ID as in head3.lst

optional arguments:

-h, --help   show this help message and exit

-c cutoff    Cutoff value of displaying conf vdw pairwise

### Example:

```
(base) jmao@Jupiter: ~/projects/lakk$ vdw_bk.py GLU-1A0090_005 -c 0.1
LEUBKA0064_000   -0.104
METBKA0065_000   -0.499
LEUBKA0068_000   -0.825
GLUBKA0069_000   -0.758
ILEBKA0085_000   -0.571
LYSBKA0086_000   -0.990
LYSBKA0087_000   -1.005
LYSBKA0088_000   -1.011
THRBKA0089_000   -0.876
GLUBKA0090_000    1.424
ARGBKA0091_000    2.383
GLUBKA0092_000   -0.204
ASPBKA0093_000   -0.113
LEUBKA0094_000   -0.120
Total            -3.616
```

## vdw\_conf2conf.py

*Calculate conformer to conformer pairwise vdw (Lennard-Jones potential).*

**Author:** Junjun Mao

### Syntax:

```
vdw_conf2conf.py [-h] [-c cutoff] [-v] confID confID
```

Compute detailed conformer to conformer vdw.

positional arguments:

confID Conformer ID as in head3.lst, two IDs required.

optional arguments:

-h, --help show this help message and exit

-c cutoff Cutoff value of displaying atom to atom vdw

-v Turn on verbose mode, displaying more details

This program calculates Lennard-Jones potential between a conformer pair, which should be consistent with the number in opp file under energies directory. The pair could be self to self, which is vdw0 term in head3.lst. The programs reports atom to atom interaction and atom connectivity as well.

The conformer ID can be found in head3.lst.

### Example:

```
(base) jmao@Jupiter: ~/projects/lakk$ vdw_conf2conf.py GLU-1A0090_005 THR01A0089_002 -c 0.01 -v
      ATOM1          ATOM2          vdw    dist    cnct      r1      e1      r2      e2
R_sum  E_par
  CB  GLU0090A005 ->  CB  THR0089A002:   -0.024    5.446   none    1.9080  0.1094  1.9080  0.1094
3.8160  0.1094
  CB  GLU0090A005 ->  OG1THR0089A002:   -0.010    6.361   none    1.9080  0.1094  1.7210  0.2104
3.6290  0.1517
  CG  GLU0090A005 ->  CB  THR0089A002:   -0.016    5.883   none    1.9080  0.1094  1.9080  0.1094
3.8160  0.1094
GLU-1A0090_005 - THR01A0089_002: -0.150
```

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## vdw\_pw.py

*Update Lennard-Jones potential of step 3 in files head3.lst and energies/\*.opp*

**Author:** Junjun Mao

### Syntax:

```
vdw_pw.py
```

This program updates Lennard-Jones potential of step 3 in

- file head3.lst (vdw0 and vdw 1) and

- files energies/\*.opp (vdw column).

A copy of head3.lst will be made as head3.lst\_bak and the copy of energies directory will be made as energies\_bak. It corrects the some parameter issues in mcce step 3 and offers a chance to rerun vdw calculation without running PB solver again. Also it checks the possible inconsistency in parameter files and comes with two other tools: vdw\_conf2conf.py and vdw\_bk.py to inspect the vdw interaction clashes.

### Example:

```
vdw_pw.py
```

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