mm_agl V.1.0.2

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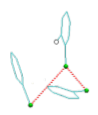
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mm_agl



1.1 Introduction

About this program:

• Program that creates PDB file with chosen agglomerate

Developer:

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```
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```

License:

• GPL

1.2 How to use

Usage:

2 mm_agl

Install

2.1 Requirements

The application mm_agl requires the following external stuff:

```
• cmake >= 2.8
```

• gcc >= 4.8

2.2 How to install

2.2.1 Linux

```
* mkdir build && cd build

* cmake -DCMAKE_INSTALL_PREFIX=/usr -DCMAKE_BUILD_TYPE=Release ../

* make

* make install
```

2.2.2 Windows

```
* create project file using 'cmake'
* compile project
```

You may also download compiled executable file for Win_x86.

Install

Changelog

V.1.0.2 (2013-07-27)

• initial release

Changelog 6

File Index

4.1 File List

Here is a list of all files with brief descriptions:

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File Documentation

5.1 src/add_main.c File Reference

```
#include <stdio.h>
#include "messages.h"
```

Functions

- int error_checking (const char *aglinp, const float *cell, const char *input, const char *output) function that checks errors in input variables
- int print_message (const int quiet, FILE *std_output, const int log, FILE *f_log, const int mode, const char *str)

function that prints message in log and stdout

• int set_defaults (char *aglinp, float *cell, char *input, int *log, char *output, int *quiet)

function that sets default values of variables

5.1.1 Function Documentation

```
5.1.1.1 int error_checking ( const char * aglinp, const float * cell, const char * input, const char * output )
```

function that checks errors in input variables

```
* error_checking (aglinp, cell, input, output);
```

Parameters

aglinp	agglomerate file name
cell	massive of cell size
input	input file name
output	output file name

Returns

- 11 error in 'cell'
- 12 error in 'input'
- 13 error in 'output'
- 14 error in 'aglinp'
- 0 exit without errors

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5.1.1.2 int print_message (const int quiet, FILE * std_output, const int log, FILE * f_log, const int mode, const char * str)

function that prints message in log and stdout

```
* print_message (quiet, stdout, log, f_log, 0, str);
```

Parameters

quiet	status of quiet-mode
std_output	stdout
log	status of log-mode
f_log	log file
mode	number of message in "messages.c"
str	additional text in message

Returns

0 - exit without errors

```
5.1.1.3 int set_defaults ( char * aglinp, float * cell, char * input, int * log, char * output, int * quiet )
```

function that sets default values of variables

```
* set_defaults (aglinp, cell, input, &log, output, &quiet);
```

Parameters

aglinp	agglomerate file name
cell	massive of cell size
input	mask of trajectory files
log	status of log-mode
output	output file name
quiet	status of quiet-mode

Returns

0 - exit without errors

5.2 src/coords.c File Reference

```
#include <stdio.h>
#include <stdlib.h>
```

Functions

• int reading_coords (const int mode, const char *filename, const int type_inter, const int *label_atom, const float *cell, int *num_mol, int *num_atoms, int *true_label_mol, int *label_mol, int *type_atoms, float *coords, char *ch_type_atoms)

function that reads coordinates from special file format

5.2.1 Function Documentation

5.2.1.1 int reading_coords (const int *mode*, const char * *filename*, const int *type_inter*, const int * *label_atom*, const float * *cell*, int * *num_mol*, int * *num_atoms*, int * *true_label_mol*, int * *label_mol*, int * *type_atoms*, float * *coords*, char * *ch_type_atoms*)

function that reads coordinates from special file format

Parameters

mode	mode of reading; '1' is statgen, '2' is envir or frad, '3' is agl
filename	input file name
type_inter	number of needed atoms (number of needed molecules)
label_atom	massive of needed atom types (massive of needed molecules)
cell	massive of cell size
num_mol	number of molecules
num_atoms	number of atoms
true_label_mol	massive of true numbers of molecule for atoms
label_mol	massive of numbers of molecule for atoms
type_atoms	massive of atom types
coords	massive of coordinates
ch_type_atoms	massive of char atom types

Returns

- 1 file \$filename does not exist
- 2 unknown mode
- 0 exit without errors

Work blocks

```
reading file translation free memory
```

5.3 src/main.c File Reference

```
#include <math.h>
#include <stdio.h>
#include <stdlib.h>
#include <string.h>
#include "add_main.h"
#include "coords.h"
#include "messages.h"
#include "print_struct.h"
#include "read_agl.h"
#include "select_mol.h"
#include "set_center.h"
```

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Functions

• int main (int argc, char *argv[])

5.3.1 Function Documentation

```
5.3.1.1 int main ( int argc, char * argv[] )
```

Returns

- 1 error in error_checking
- 2 input file does not exist
- 3 memory error
- 4 unknown flag
- 0 exit without errors

5.4 src/messages.c File Reference

```
#include <stdio.h>
#include <time.h>
```

Functions

• int message (const int log, const int mode, const char *text, FILE *output) function that prints messages to output

5.4.1 Function Documentation

5.4.1.1 int message (const int log, const int mode, const char * text, FILE * output)

function that prints messages to output

```
* message (log, mode, text, output);
```

Parameters

log	equal to 1 if print to logfile
mode	number of message
text	additional text
output	output file (may be stdout)

Returns

- 1 unknown mode
- 0 exit without errors

5.5 src/print_struct.c File Reference

```
#include <stdio.h>
```

Functions

• int print_structure (const char *output, const int num_needed_mol, const int *needed_mol, const int num_atoms, const int *label_mol, const char *ch_type_atoms, const float *coords)

function that prints structure to pdb file

5.5.1 Function Documentation

5.5.1.1 int print_structure (const char * output, const int num_needed_mol, const int * needed_mol, const int num_atoms, const int * label_mol, const char * ch_type_atoms, const float * coords)

function that prints structure to pdb file

Parameters

output	output file name
num_needed	number of needed molecules
mol	
needed_mol	massive of number of needed molecules
num_atoms	number of atoms
label_mol	massive of numbers of molecule for atoms
ch_type_atoms	massive of char atom types
coords	massive of coordinates

Returns

0 - exit without errors

5.6 src/read_agl.c File Reference

```
#include <stdio.h>
```

Functions

• int reading_agl (const char *aglinp, int *num_needed_mol, char *agl_class, int *needed_mol) function that reads agglomerate from statgen-formated file

5.6.1 Function Documentation

```
5.6.1.1 int reading_agl ( const char * aglinp, int * num_needed_mol, char * agl_class, int * needed_mol )
```

function that reads agglomerate from statgen-formated file

```
* reading_agl (aglinput, &num_needed_mol, agl_class, needed_mol);
*
```

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Parameters

aglinp	agglomerate file name
num_needed	number of needed molecules
mol	
agl_class	agglomerate class
needed_mol	massive of numbed of needed molecules

Returns

0 - exit without errors

5.7 src/select_mol.c File Reference

```
#include <math.h>
```

Functions

int select_molecule (const float *centr_coords, const int num_needed_mol, int *needed_mol)
 function that selects molecules from array of translated molecules

5.7.1 Function Documentation

 $\textbf{5.7.1.1} \quad \text{int select_molecule (const float} * \textit{centr_coords}, \ \text{const int } \textit{num_needed_mol}, \ \text{int} * \textit{needed_mol} \)$

function that selects molecules from array of translated molecules

```
* select_molecule (centr_coords, num_needed_mol, needed_mol);
```

Parameters

centr_coords	massive of centered coordinates
num_needed	number of needed molecules
mol	
needed_mol	massive of number of needed molecules

Returns

0 - exit without errors

5.8 src/set_center.c File Reference

Functions

int set_center (const int num_atoms, const int num_mol, const int *label_mol, const float *coords, float *centr-coords)

function that searchs center mass of molecules

5.8.1 Function Documentation

5.8.1.1 int set_center (const int *num_atoms*, const int *num_mol*, const int * *label_mol*, const float * *coords*, float * *centr_coords*)

function that searchs center mass of molecules

```
* set_center (num_of_atoms, num_of_molecules, label_molecules, coords, centr_coords); *
```

Parameters

num_atoms	number of atoms
num_mol	number of molecules
label_mol	massive of numbers of molecule for atoms
coords	massive of coordinates
centr_coords	massive of centered coordinates

Returns

0 - exit without errors

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