#### NAME

GLStrGen.pl - Generate structures for Glycerolipids (GL)

#### **SYNOPSIS**

GLStrGen.pl GLAbbrev|GLAbbrevFileName ...

GLStrGen.pl [-c, --ChainAbbrevMode MostLikely | Arbitrary] [-h, --help] [-m, --mode Abbrev | AbbrevFileName] [-p, --ProcessMode WriteSDFile | CountOnly] [-o, --overwrite] [-r, --root rootname] [-w, --workingdir dirname] <a gray ments>...

### **DESCRIPTION**

Generate Glycerolipids (GL) structures using compound abbreviations specified on a command line or in a CSV/TSV Text file. All the command line arguments represent either compound abbreviations or file name containing abbreviations. Use *-m*, *--mode* option to control the type of command line arguments.

A SD file, containing structures for all GL abbreviations along with ontological information, is generated as an output.

### SUPPORTED ABBREVIATIONS

Current support for GL structure generation include these main classes and sub classes:

# o Monoradylglycerols

- . Monoacylglycerols
- . Monoalkylglycerols
- . Mono-(1Z-alkenyl)-glycerol

### o Diradylglycerols

- . Diacylglycerols
- . Alkyl, acylglycerols
- . Dialkylglycerols
- . 1Z-alkenyl, acylglycerols

### o Triradylglycerols

- . Triacylglycerols
- . Alkyl, diacylglycerols
- . Dialkyl, monoacylglycerols
- . 1Z-alkenyl, diacylglycerols

# **OPTIONS**

### -c, --ChainAbbrevMode MostLikely|Arbitrary

Specify what types of acyl chain abbreviations are allowed during processing of complete abbreviations: allow most likely chain abbreviations containing specific double bond geometry specifications; allow any acyl chain abbreviation with valid chain length and double bond geometry specificatios. Possible values: *MostLikely or Arbitrary*. Default value: *MostLikely*.

Arbitrary value of **-c**, **--ChainAbbrevMode** option is not allowed during processing of abbreviations containing wild cards.

During *MostLikely* value of **-c**, **--ChainAbbrevMode** option, only the most likely acyl chain abbreviations specified in ChainAbbrev.pm module are allowed. However, during *Arbitrary* value

of **-c**, **--ChainAbbrevMode** option, any acyl chain abbreviations with valid chain length and double bond geometry can be specified. The current release of lipidmapstools support chain lengths from 2 to 50 as specified in ChainAbbev.pm module.

In addition to double bond geometry specifications, valid substituents can be specified for in the acyl chain abbreviations.

#### -h, --help

Print this help message

#### -m, --mode Abbrev|AbbrevFileName

Controls interpretation of command line arguments. Two different methods are provided: specify compound abbreviations or a file name containing compound abbreviations. Possible values: *Abbrev or AbbrevFileName*. Default: *Abbrev* 

In *AbbrevFileName* mode, a single line in CSV/TSV files can contain multiple compound abbreviations. The file extension determines delimiter used to process data lines: comma for CSV and tab for TSV. For files with TXT extension, only one compound abbreviation per line is allowed.

Wild card character, \*, is also supported in compound abbreviations.

#### Examples:

```
Specific structures: MG(16:0/0:0/0:0) DG(18:1(11E)/16:0/0:0) TG(16:0/16:0/18:1(9Z)) Specific structures: MG(0-16:0/0:0/0:0) DG(P-16:0/16:0/0:0) TG(0-20:0/16:0/18:1(9Z)) Specific possibilities: DG(18:*/16:0/0:0) DG(18:1(*)/16:0/0:0) DG(*:*(9Z)/16:0/0:0) DG(*:*(9Z)/*:*(11E)/0:0) All TG possibilities: *(*:*/*:*/*:*) or *(*/*/*) All MG, DG and TG possibilities: "MG(*:*/0:0/0:0)" "DG(*:*/*:*/*:*/0:0)" "DG(*:*/0:0/*:*)" "DG(*:*/*:*/*:*/0:0)" "DG(*:*/*:*/*:*/*:*)"
```

Along with wild card character, +/- can also be used for chain lengths to indicate even and odd lengths at sn1/sn2/sn3 positions; additionally > and < qualifiers are also allowed to specify length requirements. Examples:

```
Odd and even number chains at sn1 and sn2: TG(*+:*/*-:*/*:*) Odd and even number chains at sn1 and sn2 with length longer than 10 and 20: TG(*+>10:*/*->20:*/*:*)
```

Default sn2 stereochemistry is R. However, abbreviation format also supports these additional stereochemistry specifications for sn2 position: S; U - unknown; rac - racemic mixture. Examples:

```
\label{eq:mg} \begin{array}{lll} MG(16:0/0:0/0:0)[rac] - racemic \ mixture \\ DG(18:1(11E)/16:0/0:0)[S] - sn2 \ stereochemistry \ is \ S \ instead \ of \ default \ R \\ & TG(16:0/16:0/18:1(9Z))[U] - sn2 \ stereochemistry \ is \ unknown \end{array}
```

To generate all isomers for specific chains in DG and TG, use of iso designation is also supported. Stereochemistry specification support is not available with isomeric structure generation. Examples:

Additionally, all isomeric structures can also be generated by explicit specification of chains at different positions:

```
DG(18:1(11E)/16:0/0:0) DG(16:0/18:1(11E)/0:0) TG(16:0/16:0/18:1(9Z)) TG(16:0/18:1(9Z)/16:0) TG(18:1(9Z)/16:0/16:0/)
```

Wild card chain abbreviations are supported with sn2 stereochemistry but not with isomer abbreviation.

### -p, --ProcessMode WriteSDFile|CountOnly

Specify how abbreviations are processed: generate structures for specified abbreviations along with generating a SD file or just count the number of structures corresponding to specified abbreviations without generating any SD file. Possible values: *WriteSDFile or CountOnly*. Default: *WriteSDFile*.

It can take substantial amount of time for generating all the structures and writing out a SD file for abbreviations containing wild cards. *CountOnly* value of **--ProcessMode** option can be used to get a quick count of number of structures to be generated without writing out any SD file.

#### -o, --overwrite

Overwrite existing files

#### -r, --root rootname

New file name is generated using the root: <Root>.sdf. Default for new file names: GLAbbrev.sdf, <AbbrevFilenName>.sdf, or <FirstAbbrevFileName>1To<Count>.sdf.

### -w, --workingdir dirname

Location of working directory. Default: current directory

### **EXAMPLES**

On some systems, command line scripts may need to be invoked using *perl -s GLStrGen.pl*; however, all the examples assume direct invocation of command line script works.

To generate a GLStructures.sdf file containing a structure specified by a command line GL abbreviation, type:

```
% GLStrGen.pl -r GLStructures -o "MG(16:0/0:0/0:0)"
```

To generate a GLStructures.sdf file containing structures specified by a command line GL abbreviations, type:

```
% GLStrGen.pl -r GLStructures -o "MG(16:0/0:0/0:0)" "DG(18:1(11E)/16:0/0:0)" "TG(16:0/16:0/18:1(9Z))"
```

To generate a GLStructures.sdf file containing structures specified by a command line GL abbreviations with specific stereochemistry, type:

```
% GLStrGen.pl -r GLStructures -o "MG(16:0/0:0/0:0)[rac]" "DG(18:1(11E)/16:0/0:0)[S]" "TG(16:0/16:0/18:1(9Z))[U]"
```

To generate a GLStructures.sdf file containing all isomeric structures specified by a command line GL abbreviations, type:

```
% GLStrGen.pl -r GLStructures -o "DG(18:1(11E)/16:0/0:0)[iso2]"
```

```
"TG(16:0/16:0/18:1(9Z))[iso3]" "TG(16:0/17:0/18:1(9Z))[iso6]"
```

To enumerate all possible GL structures and generate a GLStructures.sdf file, type:

```
% GLStrGen.pl -r GLStructures -o "*(*/*/*)"

or
% GLStrGen.pl -r GLStructures -o "*(*:*/*:*/*:*)"

or
% GLStrGen.pl -r GLStructures -o "*(*:*(*)/*:*(*)/*:*(*))"
```

To enumerate all possible Monoradylglycerols structures and generate a MonoGLStructures.sdf file, type:

```
% GLStrGen.pl -r MonoGLStructures -o "MG(*/0:0/0:0)"
```

To enumerate all possible Diradylglycerols structures and generate a DiGLStructures.sdf file, type:

```
% GLStrGen.pl -r DiGLStructures -o "DG(*/*/0:0)"
```

To enumerate all possible Monoradylglycerols structures with one double bond on acyl chain and generate a GLStructures.sdf file, type:

```
% GLStrGen.pl -r MonoGLStructures -o "MG(*:1/0:0/0:0)"
```

To enumerate all possible Monoradylglycerols structures with even chain lengths and generate a GLStructures.sdf file, type:

```
% GLStrGen.pl -r MonoGLStructures -o "MG(*+:*/0:0/0:0)"
```

To enumerate all possible Diradylglycerols structures with odd chains longer than 10 at sn1 and even chains longer than 18 at sn2, and generate a DiGLStructures.sdf file, type:

```
% GLStrGen.pl -r DiGLStructures -o "DG(*->10:*/*+>18:*/0:0)"
```

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#### **SEE ALSO**

CLStrGen.pl, FAStrGen.pl, GPStrGen.pl, SPStrGen.pl, STStrGen.pl

#### COPYRIGHT

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