# TorsiFlex v2021.3

# A program for the conformational search in flexible molecules

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#### About TorsiFlex License

MIT/X11 License

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# 1. TorsiFlex software

This document is a guide to use TorsiFlex, a program designed to find the conformers of flexible molecules using a combination of a preconditioned variation of the torsions and a Monte Carlo search.

The example files cited along the document are listed in Chapter 8.

#### 1.1 About TorsiFlex

• Programming language: Python 3

• Operating systems: Linux and macOS

TorsiFlex is an user-friendly code written in Python 3 designed to seek the conformers of a given molecule by adopting a combined low-level/high-level (LL/HL) methodology. [1,2] TorsiFlex is part of a bigger project where other programs of interest are included: the Cathedral package. We encourage the user to visit our GitHub webpage to know about the software we are developing in our group. [3]

#### 1.2 Terms of use

TorsiFlex is free software under MIT license. We refer to the GitHub webpage for more details about the license:

https://github.com/cathedralpkg/torsiflex

#### 1.3 TorsiFlex software contents

The source code is located in the src/ folder and contains the following files/sub-directories:

- torsiflex.py: the main executable file;
- common/: modules with diverse functions of general use;
- modtorsiflex/: modules that deal with (i) TorsiFlex options, (ii) electronic structure softwares, (iii) reading/writing of files.

# 1.4 Setting up TorsiFlex

TorsiFlex can be executed using the python3 executable:

```
>> python3 path_to_torsiflex/torsiflex.py
```

or it can be directly executed provided that execution permissions are granted:

```
>> chmod u+x path_to_torsiflex/torsiflex.py
```

and as long as the path to the python3 executable is /usr/bin/python3. If this is not the case, the user can modify the *shebang* line of torsiflex.py to define the path of the Python interpreter:

#!/usr/bin/python3

#### 1.5 Electronic structure software

At this stage, TorsiFlex can only perform electronic structure calculations using the Gaussian software (versions 09 and 16). [4,5] The path to the Gaussian executable is read from an environment variable denoted as GauExe, which can be defined and exported in the .bashrc file. For example:

export GauExe="/home/programs/Gaussian/g09"

#### 1.6 How to cite

The following publications must be cited in any work presenting results obtained with TorsiFlex:

• D. Ferro-Costas and A. Fernández-Ramos, *Front. Chem.*, 2020, **8**:16 [A Combined Systematic-Stochastic Algorithm for the Conformational Search in Flexible Acyclic Molecules].

# 1.7 Version history

## 1.7.1 List of changes in version 2021.3

- New keywords are available: nprocll, nprochl, memll, memhl, skipcon, ifqrangeLL, and ifqrangeHL.
- A bug related to the projection of the rotational degrees of freedom has been fixed.
- Other minor changes.

# 1.7.2 List of changes in version 2021.2

- Option --syst is now renamed as --prec
- A bug related to the calculation of the rovibrational partition function for transition state structures has been fixed.

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# 2. The algorithm in a nutshell

This Chapter briefly describes the algorithm implemented in TorsiFlex, which is also schematized in the flowchart of Figure 2.1. The first part of the algorithm consists on a low-level search of conformers (section 2.1). The second part comprises a geometry re-optimization at a higher level of calculation (section 2.2). We refer to our previous works<sup>[1,2]</sup> for a more detailed explanation.

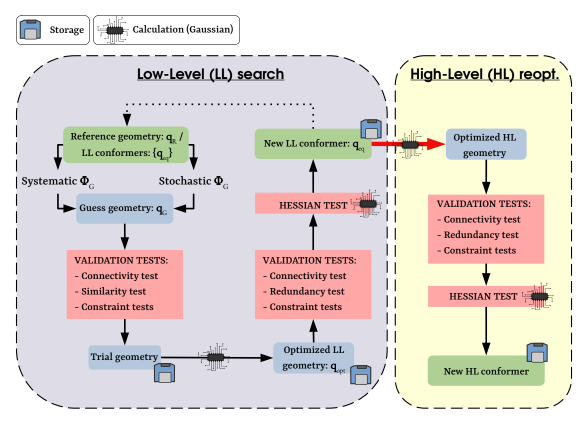


Figure 2.1: Flowchart of the search and reoptimization processes implemented in TorsiFlex.

#### 2.1 The low-level search

TorsiFlex requires a reference geometry,  $\mathbf{q}_R$ , which is given in the Z-matrix format. This geometry explicitly includes the K proper torsions of interest, which can be collected into a K-dimensional torsion vector,  $\mathbf{\Phi}^R$ . The replacement of  $\mathbf{\Phi}^R$  by a guess vector,  $\mathbf{\Phi}^G$ , renders a new configuration or guess geometry,  $\mathbf{q}_G$ .\* There are two strategies for the generation of  $\mathbf{\Phi}^G$ : preconditioned or stochastic. The former is based on chemically-intuitive dihedral angles, whereas the latter is based on the random generation of K integers between 0 and 360 (degrees).

Each guess Z-matrix,  $\mathbf{q}_G$ , is validated through a series of tests. Guess geometries that fail the tests are discarded, whereas those that pass them turn into trial structures that will be optimized geometrically. In this manner, unnecessary calculations are avoided, accelerating the search process.

The resulting optimized geometry,  $\mathbf{q}_{\text{opt}}$ , is validated through a new set of tests and, if all of them result positive, the Hessian matrix is calculated in order to confirm that we are dealing with a conformer of the system (Hessian test).

# 2.2 The high-level reoptimization

The search task described in the previous section is intended to be carried out at an inexpensive level of calculation (low-level, LL), in order to reduce the computational cost and speed up the location of conformers.

TorsiFlex can use the LL conformers as trial geometries for more accurate electronic structure calculations (high-level, HL). The HL optimized structures are validated by a new set of tests. If all of them result positive, the Hessian test is carried out by calculating the Hessian matrix. If this test is also positive, the HL optimized geometry corresponds to a new conformer.

## 2.3 The validation tests

The following tests are implemented in TorsiFlex:

• The connectivity test: positive if the connectivity of the guess/optimized structure corresponds to that of the reference Z-matrix (i.e. this test asserts that we are dealing with the same constitutional isomer).

<sup>\*</sup>The Z-matrix of the conformers located along the search process is generally more convenient than the reference one. For this reason, whenever possible, TorsiFlex uses the Z-matrix of the conformer with the closest torsion vector to  $\Phi^G$  instead of resorting to the reference Z-matrix, speeding up the geometry optimization.

 $<sup>^\</sup>dagger By$  default, the dihedral angles for the expected gauche (60° and 300°) and anti (180°) arrangements in hydrocarbons.

• The similarity test: the torsion vector of a guess structure,  $\Phi^G$ , is compared to a pool of stored torsion vectors from previous iterations,  $\{\Phi^{st}\}$ ; if  $\Phi^G$  falls outside of the domain associated to each stored vector, *i.e.*:

$$\forall p, \exists \tau : |(\mathbf{\Phi}^{G})_{\tau} - (\mathbf{\Phi}^{st}_{p})_{\tau}| > d$$
(2.1)

then the test is positive. In the previous equation, d defines the domain of each stored point and p and  $\tau$  run over all stored points and over each target torsion, respectively.

• The redundancy test: compares the optimized torsion vector,  $\Phi^{\text{opt}}$ , against the pool of stored conformers,  $\{\Phi^{\text{eq}}\}$ . If  $\Phi^{\text{opt}}$  is not stored in  $\{\Phi^{\text{eq}}\}$ , i.e.  $\Phi^{\text{opt}} \not\subset \{\Phi^{\text{eq}}\}$ , the test results positive. From a practical point of view this test is positive if:

$$\forall p, \exists \tau : |(\mathbf{\Phi}^{\text{opt}})_{\tau} - (\mathbf{\Phi}_{p}^{\text{eq}})_{\tau}| > \epsilon$$
(2.2)

where  $\epsilon$  is a threshold (not greater than 2 degrees) that accounts for numerical errors.

- The constraint tests. For different reasons, the user may be interested in a certain set of conformers. For such cases, TorsiFlex accepts the definition of constraints based on the distance between a pair of atoms, the angle between a triad of atoms and the dihedral angle between four atoms. Specifically, TorsiFlex differenciates between two types of constraints:
  - hard constraints; the test is positive if all the specific requirements for the constraints are fulfilled.
  - *soft* constraints; the test is positive if one or more of the specific requirements for the constraints are fulfilled.
- The Hessian test: performs a test on the Hessian matrix of the optimized geometry. If all its vibrational frequencies are real, the test is positive. When dealing with the conformers of a transition state, the test results positive when all the vibrational frequencies are real but one. Notice that the calculation of the Hessian matrix is time-demanding.

In Figure 2.1 we illustrate the tests that are used to validate trial and optimized structures.

# 3. TorsiFlex options

The list of available options in TorsiFlex can be displayed by executing:

```
>> path_to_torsiflex/torsiflex.py --help
```

or just:

```
>> path_to_torsiflex/torsiflex.py -h
```

The corresponding help message is shown below.

```
Welcome to TORSIFLEX
Current version: TorsiFlex v2021.3 (2021-Nov-19)
Description:
  A program to seek the conformers of flexible molecules
  by means of a combined preconditioned-stochastic algorithm.
  The located conformers, calculated with a low-level (LL)
  electronic structure method, can be re-optimized using
  a high-level (HL) method.
Main authors:
       David Ferro-Costas
  Dr.
  Prof. Antonio Fernandez-Ramos
 Centro Singular de Investigacion en Quimica Bioloxica
  e Materiais Moleculares (CIQUS), Universidade de
  Santiago de Compostela, Galicia, Spain
Execution:
  torsiflex.py [--help /-h] [--version/-v] [--inp/--input] [--prec ] [--stoc ] [--hlopt ] [--msho ] [--mstor ] [--regen ]
 Program options
   --inp / --input
     Generates the input file and the templates for Gaussian.
   --prec [M m]
     Uses the preconditioned algorithm for the conformer location.
     In order to divide the preconditioned guesses into M groups
```

```
and deal with the m-th group, use this option as follows:
         --hlopt M m
    For example:
         torsiflex --hlopt 10 2
    divides the guesses into 10 groups and only carries out
    the calculations associated to the 2nd group.
    --> Calculations with Gaussian are carried out <--
  --stoc
    Uses the stochastic algorithm for the conformer location.
    --> Calculations with Gaussian are carried out <--
  --hlopt [nocalc]
    Re-optimizes LL conformers at HL.
    When followed by 'nocalc':
         --hlopt nocalc
    it generates the gjf files (Gaussian inputs) without
    carrying out calculations.
    Useful to send the calculations on your own.
    --> Calculations with Gaussian are carried out <--
  --msho [11/h1]
    Checks the located conformers and calculates the
    multi-structure harmonic-oscillator (MS-HO) partition
    functions.
    This option can be carried out exclusively for the
    low-level (11) or the high-level (h1) conformers if
    followed by the corresponding abbreviation:
         --msho ll
         --msho hl
  --mstor [11/h1]
    Generates the MsTor input files.
    This option can be carried out exclusively for the
    low-level (11) or the high-level (h1) conformers if
    followed by the corresponding abbreviation:
         --mstor ll
         --mstor hl
  --regen
    Regenerates the domains.txt file using the temporal files.
  --help (also -h)
    Prints this help message.
  --version (also -v)
    Prints the program version.
Extra Information
```

```
_____
(a) Assert the path to the Gaussian executable
    is defined in your .bashrc file under the name 'GauExe' and export it. Example:
    export GauExe='/home/programs/Gaussian/g09'
(b) Modify the Gaussian templates (in GauTemplates/),
    taking into account that:
    * [nproc], [mem], [level], [optmode], [charge],
      [multipl], [zmat], [modred], and [fccards]
      are TorsiFlex indications.
      They should not be removed.
    * Gaussian command line must start with '#p'
    * The command line includes 'iop(99/9=1,99/14=3)'.
      This is mandatory and should not be deleted:
         99/9=1 --> rotates to Z-matrix orientation first
         99/14=3 --> expresses final optimized structure
                       in terms of the input Z-matrix
    * 'scf=(incore)' is recommended with Hartree-Fock
      calculations. With it, Gaussian stores the full integral list in memory, speeding up the calculation.
(c) We highly recommend to firstly carry out the
preconditioned search, which should found the
    chemically-intuitive conformers.
    After it, the stochastic algorithm should be used.
```

Notice that TorsiFlex accepts several options. Of them, we highlight the following:

- --inp: generates the TorsiFlex standard input file;
- --prec: performs the LL-searching using the preconditioned algorithm;
- --stoc: performs the LL-searching using the stochastic algorithm;
- --hltop: carries out the HL-refinement of the LL-conformers;
- --msho: checks the list of conformers and calculates the corresponding multi-structural harmonic-oscillator (MS-HO) partition function.
- --mstor: generates the MsTor input files. [6,7]

# 3.1 Input generation and Gaussian templates

TorsiFlex generates a default input file by executing:

```
>> path_to_torsiflex/torsiflex.py --inp
```

This file, named torsiflex.inp (see Chapter 8), contains keywords that control the program. Their detailed description can be found in Chapter 5.

This execution also generates the following eight templates for the calculations with Gaussian inside the GauTemplates/ folder:

- min\_optLL: LL optimization of a minimum.
- min\_optHL: HL optimization of a minimum.
- min\_frqLL: LL frequency calculation of a minimum.
- min\_frqHL: HL frequency calculation of a minimum.
- ts\_optLL : LL optimization of a transition state (TS).
- ts\_optHL : HL optimization of a TS.
- ts\_frqLL : LL frequency calculation of a TS.
- ts\_frqHL : HL frequency calculation of a TS.

Although the above files are templates for the calculations with Gaussian, they contain indications for TorsiFlex (between square brackets). The [zmat] indicator is automatically replaced with a guess Z-matrix and must not be removed under any circumstance. The rest of them, listed below, can be controlled with the keywords in the TorsiFlex input file (Chapter 8):

- [nproc]
- [level]
- [charge]
- [modred]

- [mem]
- [optmode]
- [multipl]
- [fccards]

#### 3.2 Low-level search

TorsiFlex will carry out the preconditioned conformer localization when executed together with --prec:

```
>> path_to_torsiflex/torsiflex.py --prec
```

whereas the stochastic search is carried out when executed with --stoc:

```
>> path_to_torsiflex/torsiflex.py --stoc
```

The difference between the two procedures lies on the generation of the geometry guesses (see section 2.1). Optimum performance is achieved when the preconditioned generation precedes the stochastic one. There are two reasons for this choice. Firstly, the geometry optimizations of preconditioned guesses rarely fail. Secondly, two preconditioned guesses infrequently lead to the same conformer.

In order to speed up the preconditioned search, TorsiFlex can split the set of preconditioned guesses into G groups and deal with each individual group separately. Thus, TorsiFlex manages the g-th group with  $(g \le G)$ :

```
>> python3 torsiflex.py --prec G g
```

For example, to split the guess vectors into G=10 groups and take care of the second group (g=2) we use:

```
>> python3 torsiflex.py --prec 10 2
```

## 3.3 High-level re-optimization

When the LL conformers are obtained, HL refinements can be carried out by using the --hlopt option:

#### >> path\_to\_torsiflex/torsiflex.py --hlopt

This execution may computationally expensive. For this reason we recommend to run it in the background using either a queue system or the well-known nohup command.

# 3.4 Conformer analysis

Once all conformers (LL and/or HL) are obtained, the multi-structural harmonic-oscillator (MS-HO) partition function [8,9] can be calculated by:

#### >> path\_to\_torsiflex/torsiflex.py --msho

This option also classifies the located conformers by increasing energy and lists their total and Gibbs free energies. Other properties, as dipole moments and rotational constants are also listed.

# 3.5 Torsional anhamonicity

A more accurate method than MS-HO to evaluate the partition function can be obtained by means of the coupled torsional anharmonic approximation, MS-T(C), which incorporates couplings in the kinetic and potential energies between the torsions. [10–12] This partition function can be calculated using the MsTor software. [6,7]

TorsiFlex is able to generate the standard MsTor input files when executed with the --mstor option:

#### >> path\_to\_torsiflex/torsiflex.py --mstor

Firstly, TorsiFlex calculates the local periodicity parameter for each conformer ( $M_i$  for the j-th conformer) adopting a Monte Carlo approach:

$$M_j = \left(\frac{N_{\text{tot}}}{N_j}\right)^{1/K} \tag{3.1}$$

In this equation,  $N_{\text{tot}}$  is the total number of random points in the torsional space and  $N_j$  is the number of samples assigned to structure j. The number of torsions is represented by K.

The standard error of each  $M_i$  is given by:

$$\sigma_{M_j} = \frac{M_j}{K\sqrt{N_i}} \tag{3.2}$$

The program increases  $N_{\rm tot}$  until all  $\sigma_{M_j}$  are smaller (or equal) than the value associated with the sigmamj keyword.

Once these  $M_j$  parameters are calculated, TorsiFlex generates the two MsTor input files: mstor.dat and hess.dat.

# 4. The reference Z-matrix

The user has to provide TorsiFlex with a file containing the Z-matrix of the system. We highly recommend Molden<sup>[13,14]</sup> to build up the molecule. Molden saves the Z-matrix in the correct format by selecting the "Gaussian" option and by clicking on the "Write Z-Matrix" button (see Figure 4.1).

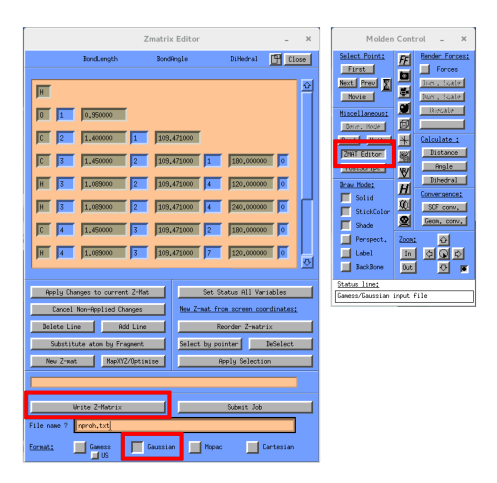


Figure 4.1: The Molden interface.

We highlight that the torsions of interest should be defined univocally: the file must contain **only one proper torsion about each torsional bond**; otherwise the TorsiFlex algorithm will fail.

# 5. The TorsiFlex input file

# 5.1 Keywords

An example of the TorsiFlex input file, torsiflex.inp, can be found in Chapter 8. This file contains different keywords, which are listed in Table 5.1. A detailed description of their meaning can be found in the following sections.

#### 5.1.1 System-specific keywords

- zmatfile: the file containing the Z-matrix of the system. This file can be created with Molden. [13,14] See Chapter 4.
- enantio: yes if torsional enantiomers are generated upon internal rotations, no otherwise.
- ts: yes if a transition state (TS) structure is being studied, no otherwise.
- cfactor: a factor that control the distance criterium needed to obtain the connectivity graph of the system. For a pair of atoms, the higher the value of cfactor, the easiest is for the two atoms to be 'connected'. We recommend a value of 1.3 for a proper connectivity.
- skipcon: accepts pair(s) of atoms, (*a*, *b*), *a* and *b* being the numerical label of each atom. The connectivity between the selected pair(s) of atoms will be omitted from the connectivity test. Several pairs of atoms can be defined in each line, for instance:

skipcon 
$$(1,2)$$
  $(4,5)$ 

Several skipcon lines can be used.

## 5.1.2 Keywords associated with the target torsions

In the keywords, X must be replaced by the corresponding integer.

• torsionX: the name of the target torsion in the Z-matrix file (zmatfile variable).

- precondX: defines the preconditioned angles of torsionX. By default, these values are 60, 180 and 300 degrees.
- tdomainX: defines the allowed domain of torsionX. See section 5.3 for more information on the definition of domains. By default the domain is [0,360).
- tsigmaX: the torsional symmetry number for torsionX. By default, it is set to 1.
- pcfile: a file containing preconditioned angles. It overrides precondX. See section 5.2.

#### 5.1.3 Keywords associated with the search procedure

• ncycles: the number of steps in the random search of conformers (active when TorsiFlex is executed with the --stoc option).

#### 5.1.4 Keywords associated with the HL re-optimization

- hlcutoff: a cutoff for HL calculations. LL conformers with relative Gibbs free energy smaller than hlcutoff (in kcal/mol) are re-optimized at HL. If this keyword is deactivated, all LL conformers are re-optimized.
- tempGibbs: temperature, in Kelvin, for the calculation of the Gibbs free energy.

## 5.1.5 Keywords associated with the validation tests

• testsG: controls which tests are carried out on a **Guess geometry**. This keyword must be followed by four integers, each one controlling one test. These tests are the connectivity, similarity, hard-constraint and soft-constraint tests (given in this order). Use 1 to activate a test and 0 to switch it off. By default, its value is:

```
testsG 1 1 1 1
```

For example, the connectivity test for the guess geometry is deactivated with the following line:

```
testsG 0 1 1 1
```

• tests0: controls which tests are carried out on a **Optimized geometry**. This keyword must be followed by four integers, each one controlling one test. These tests are the connectivity, redundancy, hard-constraint and soft-constraint tests (given in this order). Use 1 to activate a test and 0 to switch it off. By default, its value is:

```
tests0 1 1 1 1
```

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• dist1D: the domain about a torsional angle (in degrees). An angle  $\phi_i$  belongs to the domain of another angle  $\phi_j$  if their difference is within dist1D (d):

$$\phi_i \in \text{Dom}(\phi_i) \text{ if } |\phi_i - \phi_i| < d$$

This variable defines the threshold in the similarity test.

• epsdeg: a criterium to decide whether two angles are equal or not (in degrees). Two angles,  $\phi_i$  and  $\phi_j$ , are considered identical if their difference is within epsdeg  $(\epsilon)$ :

$$\phi_i = \phi_i \text{ if } |\phi_i - \phi_i| < \epsilon$$

This variable defines the threshold in the **redundancy test**.

• hconstr: defines a *hard* constraint (one per line). The argument of a hconstr line consists of an internal coordinate (a distance, an angle or a torsional angle) and a domain:

```
hconstr icoord domain
```

The internal coordinate can be either defined in the Z-matrix or can be defined by listing the involved atoms. As an example, for the latter, 1–2–3 refers to the angle defined by the atoms labelled as 1, 2 and 3. The definition of domains can be found below, in section 5.3. We notice that all hard-constraints are mandatory and a guess/optimized geometry is **discarded** if **a single** hard constraint is **not fulfilled**.

• sconstr: defines a *soft* constraint (one per line) using the same format as in hconstr:

```
sconstr icoord domain
```

In this case, a guess/optimized geometry is **discarded** if **all** the soft constraints are **not fulfilled**.

- ifqrangeLL: restricts the imaginary frequency to the defined interval (for LL conformers of saddle point structures). When the imaginary frequency falls outside this interval, the structure is discarded.
- ifqrangeHL: same as ifqrangeLL but for HL conformers.

## 5.1.6 Keywords associated with the calculations with Gaussian

- optmode: defines the arguments of the opt keyword in the Gaussian input file (it replaces the [optmode] string in the template). It can be:
  - 0: to use opt(z-matrix)
  - 1: to use opt(modreduntant).
- fccards: when set to yes, the LL Hessian matrix is used in the HL optimization. This keyword will also modify the [optmode] string in the template by adding the fccards argument:

- opt(z-matrix,fccards)
- opt(modreduntant,fccards).
- charge: the charge of the system (it replaces the [charge] string in the Gaussian templates).
- multipl: the spin multiplicity of the system (it replaces the [multipl] string in the Gaussian templates).
- lowlevel: the low-level to be inserted in the Gaussian template. This keyword replaces the [level] string in the Gaussian template.
- highlevel: same as lowlevel but to define the level of calculation in the HL templates.
- nproc: number of threads for Gaussian calculations. This keyword replaces the [nproc] string in the Gaussian template.
- procll: same as proc but it only applies to LL Gaussian templates.
- prochl: same as proc but it only applies to HL Gaussian templates.
- mem: dynamic memory for Gaussian calculations. This keyword replaces the [mem] string in the Gaussian template.
- memll: same as mem but it only applies to LL Gaussian templates.
- memhl: same as mem but it only applies to HL Gaussian templates.

## 5.1.7 Keywords associated with the partition functions

- tempsPF: temperatures, in Kelvin, for the calculation of partition functions. Each line can contain several temperatures. More than one line can be used.
- freqscalLL: a scaling factor for the harmonic frequencies of the LL conformers, if needed. By default, frequencies are not scaled (i.e. freqscalLL is set to 1).
- freqscalHL: a scaling factor for the harmonic frequencies of the HL conformers, if needed. By default, frequencies are not scaled (i.e. freqscalHL is set to 1).
- sigmamj: the standard error in the calculation of the local periodicity parameters for MsTor (see section 3.5). This value must be  $\geq 10^{-2}$ .

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### 5.1.8 Keywords associated with the storage of conformers

- dirll: the folder where the LL conformers are stored.
- dirhl: the folder where the HL conformers are stored.
- tmpll: the folder where the temporal LL Gaussian files are stored. By default this folder is set to /scratch/user/LL\_xxxxx/ where user is the name of the user and xxxxx is a random sequence of characters. The folder LL\_xxxxx/ is automatically created by the program when executed. If the creation fails, the program will let the user know that the folder must be created by themselves.
- tmphl: the folder where the temporal HL Gaussian files are stored. See also tmpll.

Table 5.1: Available keywords in the TorsiFlex input file.

Keyword	(Type of) value	More info
zmatfile	string	path to Z-matrix file
enantio	yes, no	are there torsional enantiomers?
ts	yes, no	is the system a transition state?
cfactor	float	controls system connectivity
skipcon	pair(s) of atoms (a,b)	excludes pairs of atoms in connectivity test
torsionX	string	the coordinate name of the X-th torsion
		in the Z-matrix file; replace X with integer;
precondX	a list of integers	preconditioned angles for torsionX;
•		replace X with integer;
		e.g. precond1 0 180
tdomainX	a domain	domain for torsionX; replace X with integer;
		e.g. tdomain1 (-60,60)
tsigmaX	integer	torsional symmetry number of torsionX;
· ·		replace X with integer
pcfile	string	path to file with preconditioned angles
ncycles	integer	number of cycles in stochastic search;
v		forstoc
hlcutoff	float (kcal/mol)	Gibbs free energy cutoff for HL re-optimizations
tempGibbs	float (Kelvin)	temperature for the calculation of Gibbs free energy
testsG	four integers (0's or 1's)	selected validation tests for Guess geometries
tests0	four integers (0's or 1's)	selected validation tests for Optimized geometries
dist1D	integer (in degrees)	≥ 2.0; domain about an angle
epsdeg	integer (in degrees)	≤ 2.0; max. difference between equal angles
hconstr	internal coord. and domain	one line per hard constraint
sconstr	internal coord. and domain	one line per soft constraint
ifqrangeLL	domain	restricts values for the imaginary frequency (LL)
ifqrangeHL	domain	restricts values for the imaginary frequency (HL)
optmode	0,1	defines the argument of the opt Gaussian keyword
fccards	yes,no	use LL Hessian in HL optimization
charge	integer (in a.u.)	charge of the system
multipl	integer	spin multiplicity of the system
lowlevel	string	low-level methodology
highlevel	string	high-level methodology
nproc	integer	number of threads in Gaussian template (e.g. 2)
nprocll	integer	same as nproc but only for LL
nprochl	integer	same as nproc but only for HL
mem	string	dynamic memory in Gaussian template (e.g. 2GB)
memll	string	same as mem but only for LL
memhl	string	same as mem but only for HL
tempsPF	list of floats (Kelvin)	temperatures for the partition functions
${ t freqscallL}$	float	scale factor for harmonic frequencies (LL conformers)
${ t freqscalHL}$	float	scale factor for harmonic frequencies (HL conformers)
sigmamj	float	$\geq$ 0.01; formstor
dirll	string	folder for storing LL conformers
dirhl	string	folder for storing HL conformers
tmpll	string	folder for LL Gaussian calculations
tmphl	string	folder for HL Gaussian calculations

# **5.2 The file with preconditioned angles: pcfile**

The preconditioned angles can be also introduced as plain text using the pcfile keyword. Different combinations are possible for the selected torsions.

The first line in this file consists of the target torsions, whereas the rest of the lines define their corresponding values (separated by blank spaces or by an underscore). For example, the next file:

```
1 3
100_100
200_300
```

defines two combinations:

- (torsion1, torsion3) = (100, 100)
- (torsion1, torsion3) = (200, 300)

The value of the disregarded torsions, if any, will be defined according to the default angles, *i.e.* the (60,180,300) triad or those defined through precondX. In this manner, if the system contains three torsions (torsion1,torsion2,torsion3), the previous lines would lead to the next points:

```
(100,60,100)
(200,60,300)
(100,180,100)
(200,180,300)
(200,300,300)
(200,300,300)
```

# 5.3 Specifying a domain

Keywords hconstr, sconstr and tdomainX require the definition of a domain. An interval can be defined with the format (a,b), where a and b are the limits of the interval (a < b). Several intervals can be used to define a given domain through their union (denoted by a capital U). For example, (a,b)U(c,d), where a < b < c < d.

In the case of tdomainX, the domain corresponds to a torsion angle (in degrees). However, hconstr or sconstr can be specified to restrict the domain of a distance (in Angstrom) or of an angle (in degrees).

# 6. Some tips

#### 6.1 Low-level calculations at the HF level

If Hartree-Fock (HF) is the LL method of choice for the searching algorithm, we highly recommend to modify the Gaussian template and include the argument incore in the scf Gaussian keyword:

```
scf = (verytight, incore)
```

In this manner, the full integral list will be stored in memory along the self-consistent field method (SCF). This should speed up the calculation. Obviously, if incore is used, enough memory should be available.

# **6.2** Systems with many LL conformers

The HL re-optimizations are carried out sequentially, which may not be convenient for systems with a big number of conformers. In such a cases, we recommend to execute TorsiFlex as follows:

```
>> path_to_torsiflex/torsiflex.py --hlopt nocalc
```

This will generate the Gaussian input files for the HL re-optimization of all the LL conformers. In this manner, the user can execute all Gaussian jobs at the same time if desired (i.e. the calculations are managed by the user and not by TorsiFlex). Once the optimizations are complete, the corresponding Gaussian input files for the frequency calculations can be also generated by executing TorsiFlex again:

```
>> path_to_torsiflex/torsiflex.py --hlopt nocalc
```

When all optimization and frequency calculations are finished, TorsiFlex can be executed normally and will automatically read the generated output files:

```
>> path_to_torsiflex/torsiflex.py --hlopt
```

## 6.3 Focusing on low energy conformers

The user may also be interested into re-optimize only the most stable conformers. In such cases, we recommend to use the hlcutoff keyword. Only those LL con-

formers whose Gibbs free energy is below the value associated to hlcutoff are optimized at HL.

For example, in order to optimize at HL those conformers whose relative Gibbs free energy is smaller than 3 kcal/mol at 300 K, we need the following line in the input file:

```
hlcutoff 3.0 300.0
```

The first argument of hlcutoff defines the cutoff for the Gibbs free energy, whereas the second argument defines the temperature for the calculation of such an energy. If not indicated, the default temperature is 300 K.

# 6.4 Dealing with similar molecules

When performing conformational studies, it may occur that we have to deal with two molecules with common fragments. For the sake of simplicity, let us name these two molecules as AM and BM, both containing the M moiety. In such cases, it is very likely for the conformations of the M moiety in A and in B to be similar. If we have performed a full study on AM, it would be very convenient to use the conformations of its M moiety as a guess for BM. This may be achieved by defining the pre-conditioned angles through pcfile instead of with the precondX keyword.

For instance, let us assume that molecule AM consists of three torsions, torsion1 to torsion3, the last two torsions being located in the M fragment, whereas BM consists of four torsions, torsion1 to torsion4, the last two corresponding to M. TorsiFlex can initialize torsion3 and torsion4 of BM according to the values of torsion2 and torsion3 of AM. The procedure is as follows:

- (a) Go to the working directory of AM and enter in the folder where conformers are stored.
- (b) List the angles using:

```
>> ls *log | awk -F. '{print $2}'
```

The output would look like:

```
069_284_314

071_172_060

071_182_301

072_268_179

073_080_304

075_093_192

075_280_051
```

(c) As we are only interested in the second and third columns of this list (torsion2 and torsion3), the first column should be eliminated. This can be achieved with a slight modification of the command:

```
>> ls *log | awk -F. '{print $2}' | \
awk -F"_" '{print $2, $3}'
```

leading to:

```
284 314
172 060
182 301
268 179
080 304
093 192
280 051
```

(d) Save these lines in a new file and use it for BM through the pcfile keyword. A first line, indicating the target torsions, must be added to this file:

```
3 4
284 314
172 060
182 301
268 179
080 304
093 192
280 051
```

(e) Now, the initial angles for torsions torsion3 and torsion4 of BM are set to those in AM, whereas starting values for torsion1 and torsion2 are defined by the default triad, (60,180,300).

An example of this situation may be found when studying the amino acids. For example, alanine and valine, or cysteine and methionine, et cetera.

Finally, we notice that (i) precondX defines preconditioned values for individual torsions whereas (ii) pcfile defines preconditioned (sub)vectors in the torsional space.

# 7. Worked examples

This Chapter contains the following step-by-steps examples:

• WE1: n-butanol

• WE2: L-alanine

• WE3: L-proline

• WE4: Transition state for the H abstraction from n-butanol at the  $C-\alpha$  atom by the H radical

The level of calculation for the search procedure is set to HF/3-21G (low-level, LL), whereas B3LYP/6-31G is employed as the accurate high-level (HL) for the re-optimizations.

#### **Torsions of interest**

#### Torsions for WE1:

- dih4 (H1-O2-C3-C4)
- dih5 (O2-C3-C4-C5)
- dih6 (C3-C4-C5-H6)

#### Torsions for WE3:

- dih4 (C1-C2-C3-C4)
- dih5 (C2-C3-C4-N5)
- dih17 (C3-C4-N5-H17)
- dih7 (C2-C1-C6-O7)
- dih8 (C1-C6-O7-H8)

#### Torsions for WE2:

- dih4 (H1-O2-C3-C4)
- dih5 (O2-C3-C4-C5)
- dih12 (C3-C4-N8-H12)

#### Torsions for WE4:

- dih4 (H1-O2-C3-C4)
- dih5 (O2-C3-C4-C5)
- dih6 (C3-C4-C5-C6)

#### 7.1 WE1: conformers of n-butanol

In this worked example, we find the conformers of n-butanol (Figure 7.1)

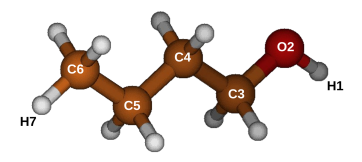


Figure 7.1: Labeling for n-butanol.

#### How to proceed

- Assert that the GauExe variable with the path to your Gaussian version is exported in the .bashrc file.
- Create the folder WE1/ and enter to it.

```
>> mkdir WE1/
>> cd WE1/
```

- Copy the corresponding Z-matrix file (we01.zmat, see Chapter 8) from the tests/ folder to the current directory (or create it with the help of Molden).
- Create the input file and the templates for the Gaussian calculations:

```
>> path_to_torsiflex/torsiflex.py --inp
```

• Modify/add the following keywords in the input file:

```
zmatfile we01.zmat torsion1 dih4
enantio yes torsion2 dih5
ncycles 20 torsion3 dih6
lowlevel hf/3-21g
```

Keyword tmpll may be also modified.

- In principle, it is not necessary to modify the templates for Gaussian. However, the incore argument could be included in the scf Gaussian option to speed-up the LL calculations (min\_optLL file inside GauTemplates/).
- Execute TorsiFlex with the --prec option:

```
>> path_to_torsiflex/torsiflex.py --pred
```

In a few minutes, the program finds the conformers from the preconditioned guesses at the HF/3-21g level. A total of 14 conformers should be found.

• Execute TorsiFlex with the --stoc option:

#### >> path\_to\_torsiflex/torsiflex.py --stoc

In this case, TorsiFlex considers random angles to find new conformers. Execute TorsiFlex as many times as needed. A new conformer should be found in this random search (086\_305\_086 or its torsional enantiomer, 274 055 274).

- A total of 15 LL conformers should be found (14+1). Notice that there exists an enantiomer for any of the conformers, but the one with Cs symmetry. Consequently, there are 29 conformers for n-butanol  $(14 \cdot 2 + 1)$ .
- List your LL conformers and calculate the corresponding MS-HO partition functions using --msho:

```
>> path_to_torsiflex/torsiflex.py --msho ll
```

• Modify again the input file to indicate the high-level of calculation:

```
highlevel b3lyp/6-31g
```

Keyword tmphl may be also modified.

• Execute TorsiFlex to carry out the HL optimizations based on the LL structures:

```
>> path_to_torsiflex/torsiflex.py --hlopt
```

This execution may take a few minutes.

• List your HL conformers and calculate the corresponding MS-HO partition function using the --msho option:

```
>> path_to_torsiflex/torsiflex.py --msho hl
```

#### List of low-level conformers

1. (295,304,176) 6. (177,302,291) 11. (295,186,291)

2. (176,300,177) 7. (285,062,179) 12. (283,058,065)

3. (295,306,290) 8. (180,180,180) 13. (181,183,292)

4. (297,296,073) 9. (178,293,078) 14. (274,055,274)

5. (296,183,180) 10. (297,181,069) 15. (292,084,299)

## List of high-level conformers

1. (297,301,177) 6. (298,303,293) 11. (297,186,293)

2. (298,183,180) 7. (177,299,294) 12. (182,183,294)

3. (291,065,179) 8. (300,181,066) 13. (179,290,075)

4. (176,297,179) 9. (300,296,073) 14. (293,081,298)

5. (180,180,180) 10. (292,064,066) 15. (283,063,274)

# 7.2 WE2: conformers of L-alanine

In this example, we find the conformers of L-alanine (see Figure 7.2).

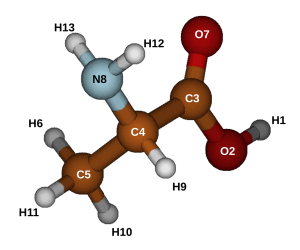


Figure 7.2: Labeling for Alanine.

### How to proceed

- Assert that the GauExe variable with the path to your Gaussian version is exported in the .bashrc file.
- Create the folder WE2/ and enter to it.

```
>> mkdir WE2/
>> cd WE2/
```

- Copy the corresponding Z-matrix file (we02.zmat, see Chapter 8) from the tests/ folder to the current directory (or create it with the help of Molden).
- Create the input file and the templates for the Gaussian calculations:

```
>> path_to_torsiflex/torsiflex.py --inp
```

• Modify/add the following keywords in the input file:

```
zmatfile we02.zmat torsion1 dih4
ncycles 20 torsion2 dih5
lowlevel hf/3-21g torsion3 dih12
precond1 0 180
```

Notice that keyword precond1 is used to set the preconditioned angles for the carboxyl group to 0 and 180 degrees. Keywords tmpll and tmphl may be also modified.

- In principle, it is not necessary to modify the templates for Gaussian. However, the incore argument could be included in the scf Gaussian option to speed-up the LL calculations (min\_optLL file inside GauTemplates/).
- Execute TorsiFlex with the --prec option:

```
>> path_to_torsiflex/torsiflex.py --prec
```

In a few minutes, the program finds the conformers at the HF/3-21g level from the preconditioned guesses. A total of 11 conformers should be found.

• Execute TorsiFlex with the --stoc option:

```
>> path_to_torsiflex/torsiflex.py --stoc
```

In this case, TorsiFlex considers random angles to find new conformers. This execution may take some time, depending on the value of the ncycles keyword. No new conformers are found for this system.

• List your LL conformers and calculate the corresponding MS-HO partition function using --msho:

```
>> path_to_torsiflex/torsiflex.py --msho ll
```

• Modify again the input file to perform the HL calculations:

```
highlevel b3lyp/6-31g
```

• Execute TorsiFlex to carry out the HL optimizations based on the LL structures:

```
>> path_to_torsiflex/torsiflex.py --hltopt
```

This execution may take a few minutes.

• List your HL conformers and calculate the corresponding MS-HO partition function using the --msho option:

```
>> path_to_torsiflex/torsiflex.py --msho hl
```

The number of conformers should have been reduced to 10.

### List of low-level conformers

1. (179,298,294)

5. (178,291,033)

9. (355,294,290)

2. (354,147,088)

6. (181,307,197)

10. (351,289,023)

3. (178,162,304)

7. (182,171,175)

11. (013,183,174)

4. (182,075,288)

8. (179,069,059)

# List of high-level conformers

1. (356,139,097)

5. (178,289,027)

9. (358,293,294)

2. (179,294,295)

6. (181,309,196)

10. (355,291,016)

3. (184,197,295)

7. (180,166,175)

4. (181,077,290)

8. (180,076,058)

# 7.3 WE3: conformers of proline

In this example, we find the conformers of L-proline (see Figure 7.4).

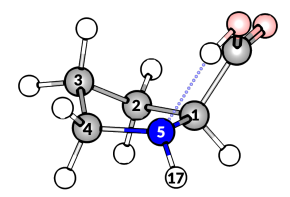


Figure 7.3: Cycle numeration in Pro. C in grey, O in red, N in blue and H in white.

We notice that L-proline is the only proteinogenic amino acid containing a flexible ring in its structure. TorsiFlex is not specially optimized for dealing with ring puckering, but it can still be used to find the corresponding conformers. If we consider the numbering shown in Figure 7.3, the ring configurations arise from the following torsions: (i)  $C_1$ - $C_2$ - $C_3$ - $C_4$ , (ii)  $C_2$ - $C_3$ - $C_4$ - $N_5$  and (iii)  $C_3$ - $C_4$ - $N_5$ - $H_{17}$ . In order to avoid huge distortions in the five-member ring, the domains of these torsions are limited to a small interval. Moreover, since the connectivity of the ring may be compromised in the guess structure, the connectivity test for guess structures should be deactivated.

# How to proceed

- Assert that the GauExe variable with the path to your Gaussian version is exported in the .bashrc file.
- Create the folder WE2/ and enter to it.

```
>> mkdir WE3/
>> cd WE3/
```

- Copy the corresponding Z-matrix file (we03.zmat, see Chapter 8) from the tests/ folder to the current directory (or create it with the help of Molden).
- Create the input file and the templates for the Gaussian calculations:

```
>> path_to_torsiflex/torsiflex.py --inp
```

• Modify/add the following keywords in the input file:

```
zmatfile
           we03.zmat
                                           (-50,50)
                               tdomain1
testsG
           0 1 1 1
                               tdomain2
                                           (-50,50)
           hf/3-21g
                                           (-160, -80) U(80, 160)
lowlevel
                               tdomain3
torsion1
           dih4
                                            -30 +30
                               precond1
                                            -30
           dih5
                                                +30
torsion2
                               precond2
                               precond3
torsion3
           dih17
                                           -120 + 120
torsion4
           dih7
                               precond5
                                              0
                                                 180
           dih8
torsion5
```

Notice that we have reduced the interval for dih4 and dih5 to (-50,50), in order to generate structures where the five-member ring is "conserved". Similarly, the interval for dih17 should account for the two arrangements in the umbrella inversion (in this case, this conformational effect leads to different conformers). The preconditioned angles for these torsions were also modified, in order to fit the previous defined intervals. As in WE2, precond5 account for the two expected conformations of the carboxylic acid group. The first argument of testsG is set to 0 to deactivate the connectivity test on the guess structures.

• Execute TorsiFlex with the --prec option:

```
>> path_to_torsiflex/torsiflex.py --prec
```

In a few minutes, the program finds the conformers at the HF/3-21g level from the preconditioned guesses. A total of 17 conformers should be found.

• Execute TorsiFlex with the --stoc option:

```
>> path_to_torsiflex/torsiflex.py --stoc
```

In this case, TorsiFlex considers random angles to find new conformers. This execution may take some time, depending on the value of the ncycles keyword. A total of 3 new conformers should be found.

• List your LL conformers and calculate the corresponding MS-HO partition function using --msho:

```
>> path_to_torsiflex/torsiflex.py --msho ll
```

• Modify again the input file to perform the HL calculations:

```
highlevel b3lyp/6-31g
```

• Execute TorsiFlex to carry out the HL optimizations based on the LL structures:

```
>> path_to_torsiflex/torsiflex.py --hltopt
```

This execution may take a few minutes.

• List your HL conformers and calculate the corresponding MS-HO partition function using the --msho option:

```
>> path_to_torsiflex/torsiflex.py --msho hl
```

The number of conformers should have been reduced to 14.

# List of low-level conformers

- 1. (320,037,211,121,358)
- 2. (321,029,123,294,180)
- 3. (016,325,171,298,181)
- 4. (012,330,274,139,354)
- 5. (323,022,136,159,179)
- 6. (013,327,171,157,179)
- 7. (322,039,195,186,186)
- 8. (040,330,239,098,007)
- 9. (323,038,198,043,176)
- 10. (037,335,225,181,184)

- 11. (031,319,171,074,182)
- 12. (025,322,272,204,186)
- 13. (027,322,271,052,175)
- 14. (037,337,206,299,180)
- 15. (007,017,081,158,355)
- 16. (320,031,117,291,354)
- 17. (022,321,170,298,357)
- 18. (029,319,169,182,012)
- 19. (331,010,246,283,352)
- 20. (356,342,172,038,352)

## List of high-level conformers

- 1. (323,036,209,117,359)
- 2. (027,324,266,127,357)
- 3. (324,028,123,294,181)
- 4. (033,322,163,296,181)
- 5. (325,023,136,145,177)
- 6. (017,326,173,151,178)
- 7. (326,036,190,188,185)

- 8. (035,330,234,195,185)
- 9. (326,034,195,040,176)
- 10. (031,325,255,049,175)
- 11. (007,017,082,156,356)
- 12. (033,322,162,297,001)
- 13. (324,030,118,292,359)
- 14. (029,321,171,196,009)

# 7.4 WE4: conformers of a transition state

In this example, we find the conformers of the transition state associated to the H abstraction by H in the  $\alpha$  position of n-BuOH (see Figure 7.4).

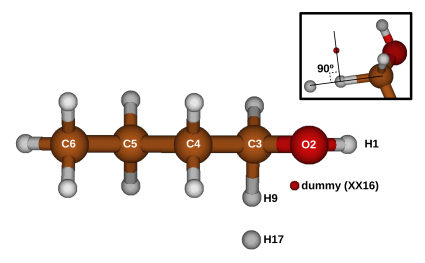


Figure 7.4: Labeling for the H-abstraction transition state. The definition of the dummy atom is also shown from another point of view.

# How to proceed

- Assert that the GauExe variable with the path to your Gaussian version is exported in the .bashrc file.
- Create the folder WE4/ and enter to it.

```
>> mkdir WE4/
>> cd WE4/
```

- Copy the corresponding Z-matrix file (we04.zmat, see Chapter 8) from the tests/ folder to the current directory (or create it with the help of Molden).
- Create the input file and the templates for the Gaussian calculations:

```
>> path_to_torsiflex/torsiflex.py --inp
```

• Modify/add the following keywords in the input file:

zmatfile	we04.zmat	ifqrangeLL	(1500,2500)
ts	yes	torsion1	dih4
multipl	2	torsion2	dih5
lowlevel	hf/3-21g	torsion3	dih6
skipcon	(3,9) (9,17)		

Notice that setting ts to yes indicates that we are dealing with a transition state (different Gaussian templates are selected), whereas multipl defines the spin multiplicity of the system. Two extra keywords (skipcon and ifqrangeLL) are included to consider two specific problems that may arise.

On the one side, it may occur that the connectivity between H9 and H17 differs from one conformer to another due to a small variation in the distance between them. The skipcon keyword can be used to omit the two distances in the  $C\cdots H\cdots H$  moiety when carrying out the connectivity test.\*

On the other side, the system could end up in an internal rotation transition state.<sup>†</sup> The ifqrangeLL keyword can be used to define an expected range of values for the imaginary frequency, which should be betweem 1500*i* and 2500*i* cm<sup>-1</sup> for a H-abstraction transition-state at the HF/3-21G level.

We notice that, for this system, these two last keywords are not really required, but we recommend to include them to avoid the unique situations previously described.

• Execute TorsiFlex with the --prec option:

### >> path\_to\_torsiflex/torsiflex.py --prec

In a few minutes, the program finds the conformers at the HF/3-21g level from the preconditioned guesses. A total of 22 conformers should be found.

• Execute TorsiFlex with the --stoc option:

### >> path\_to\_torsiflex/torsiflex.py --stoc

In this case, TorsiFlex considers random angles to find new conformers. This execution may take some time, depending on the value of the ncycles keyword. A total of 3 new conformers should be found.

• List your LL conformers and calculate the corresponding MS-HO partition function using --msho:

```
>> path_to_torsiflex/torsiflex.py --msho ll
```

This also lists the imaginary frequency of each transition state conformer. A total of 25 LL conformers should be listed.

• Modify again the input file to perform the HL calculations:

```
highlevel b3lyp/6-31g
```

• Execute TorsiFlex to carry out the HL optimizations based on the LL structures:

```
>> path_to_torsiflex/torsiflex.py --hltopt
```

This execution may take a few minutes.

<sup>\*</sup>This could also be solved by modifying the cfactor keyword.

<sup>&</sup>lt;sup>†</sup>For example, if the H radical goes away from butanol during the optimization.

• List your HL conformers and calculate the corresponding MS-HO partition function using the --msho option:

>> path\_to\_torsiflex/torsiflex.py --msho hl

The number of conformers should be reduced to 19.

# List of low-level conformers

1. (056,058,184)	10. (061,304,182)	19. (286,311,176)
2. (176,306,177)	11. (177,304,081)	20. (051,278,059)
3. (182,062,183)	12. (179,321,299)	21. (290,184,180)
4. (055,068,290)	13. (053,177,290)	22. (287,304,074)
5. (056,055,069)	14. (179,181,290)	23. (285,326,300)
6. (054,176,180)	15. (063,315,296)	24. (287,185,289)
7. (181,059,068)	16. (176,168,067)	25. (293,174,067)
8. (178,179,180)	17. (056,166,067)	
9. (180,071,285)	18. (072,314,084)	

# List of high-level conformers

1. (051,063,183)	8. (049,069,290)	15. (185,298,077)
2. (187,066,181)	9. (185,064,066)	16. (051,170,070)
3. (050,177,180)	10. (049,180,293)	17. (181,171,069)
4. (183,302,179)	11. (185,074,287)	18. (048,283,061)
5. (186,180,181)	12. (184,183,293)	19. (056,306,088)
6. (052,300,182)	13. (182,311,295)	
7. (049,060,066)	14. (047,308,296)	

# 8. Example files

#### Example of file with preconditioned angles (pcfile keyword).

### Gaussian template for LL optimization of a minimum.

```
%nproc=[nproc]
%mem=[mem]
#p [level]
scf=(tight)
iop(99/9=1,99/14=3)
opt=([optmode],tight,MaxCycles=200)
--Optimization of minimum--
[charge],[multipl]
[zmat]
[modred]
[fccards]
```

### Gaussian template for HL optimization of a minimum.

```
%nproc=[nproc]
%mem=[mem]
#p [level]
scf=(verytight)
iop(99/9=1,99/14=3)
opt=([optmode],verytight,MaxCycles=200)
--Optimization of minimum--
[charge],[multipl]
[zmat]
[modred]
[fccards]
```

### Gaussian template for LL optimization of a transition state.

```
%nproc=[nproc]
%mem=[mem]
#p [level]
scf=(tight)
iop(99/9=1,99/14=3)
opt=([optmode],tight,calcfc,ts,noeigentest,MaxCycles=200)
--Optimization of transition state--
[charge],[multipl]
[zmat]
[modred]
[fccards]
```

### Gaussian template for HL optimization of a transition state.

```
%nproc=[nproc]
%mem=[mem]
#p [level]
scf=(verytight)
iop(99/9=1,99/14=3)
opt=([optmode],verytight,calcfc,ts,noeigentest,MaxCycles=200)
--Optimization of transition state--
[charge],[multipl]
[zmat]
[modred]
[fccards]
```

### Gaussian template for LL frequency calculation.

```
%nproc=[nproc]
%mem=[mem]
#p [level]
scf=(tight)
iop(99/9=1,99/14=3)
freq=noraman
--Frequency calculation--
[charge],[multip1]
[zmat]
```

### Gaussian template for HL frequency calculation.

```
%nproc=[nproc]
%mem=[mem]
#p [level]
scf=(verytight)
iop(99/9=1,99/14=3)
freq=noraman
--Frequency calculation--
[charge],[multip1]
[zmat]
```

### The default TorsiFlex input file

```
# This is a torsiflex input file
 # System
 #----#
 # Target torsions
  #----#
  torsion1
                                                                                  # name of 1st target torsion in the Z-matrix file
 # name of 1st target torsion in the precond of 1st target torsion in the precond angles for torsion in the precond angle in the precond angles for torsion in the precond angle in t
 #pcfile
                                precond.txt # file with precond. angles
 # Search Procedure #
 ncycles
                                200
                                                              # number of steps of stochastic algorithm
 # HL reoptimization #
 #-----#
#hlcutoff 5.0 # Gibbs energy cutoff (kcal/mol)
tempGibbs 298.15 # Temperature (K) for Gibbs free energy
 # Validation tests # #-----#
 testsG 1 1 1 1 # for Guess geom (Conn, Simil, Hard, Soft)
testsO 1 1 1 1 # for Opt geom (Conn, Redun, Hard, Soft)
dist1D 15 # domain size about each point (degrees)
epsdeg 2 # max diff between two identical angles (degrees)
tests0 1 1 1 1 1 # for Opt geom (Conn, Redun, Hard, Soft)
dist1D 15 # domain size about each point (degrees)
epsdeg 2 # max diff between two identical angles (de
#hconstr ic domain # hard constraint (see manual)
#sconstr ic domain # soft constraint (see manual)
#ifqrangeLL domain # restricts LL imaginary-frequency interval
#ifqrangeHL domain # restricts HL imaginary-frequency interval
 # _ _ _ _ _ #
 # Gaussian calculations #
 #----#
 optmode 1 # 0:opt(z-matrix), 1:opt(modredundant)
fccards no # Use LL Hessian in HL opt (yes/no)
charge 0 # charge of the system
multipl 1 # multiplicity of the system
lowlevel hf 3-21g # low-level of calculation
highlevel b3lvn 6-31G # high level of calculation
 highlevel
                                          b3lyp 6-31G # high-level of calculation
                                                                  # Number of threads (low-level)
# dynamic memory (low-level)
# Number of threads (high-level)
# dynamic memory (high-level)
 nprocll
                                          1
                                    1GB
 memll
 nprochl
                                          1
                                       1 GB
 memhl
 # Partition functions #
 # - - - - - - #
 # tempsPF 100 200 # temperatures (K) for part. functions tempsPF 298.15 # temperatures (K) for part. functions tempsPF 300 500 # temperatures (K) for part. functions tempsPF 750 1000 # temperatures (K) for part. functions tempsPF 2000 2500 # temperatures (K) for part. functions freqscalLL 1.0 # freq. scaling factor (LL) freqscalHL 1.0 # freq. scaling factor (HL) sigmamj 0.02 # max value for sigma(Mj); >= 0.01
  # Storage
                                      files_LL/  # folder to store LL conformers
files_HL/  # folder to store HL conformers
/scratch/david/LL_uRRXp/  # folder for LL temporal files
  dirhl
  tmpll
                                          /scratch/david/HL_uRRXp/ # folder for HL temporal files
```

hcc15

dih15

109.471

240.000

#### Reference Z-matrix file for n-BuOH (WE1). h 1 oh2 0 1 coh3 С 2 co3 3 cc4 2 cco4 1 dih4 С С 4 cc5 3 ccc5 2 dih5 3 dih6 С 5 cc6 4 ccc6 6 hc7 5 hcc7 4 dih7 h h 3 hc8 2 hco8 4 dih8 h 3 hc9 2 hco9 4 dih9 h 4 hc10 3 hcc10 5 dih10 4 hc11 3 hcc11 5 dih11 h 5 hc12 4 hcc12 h 6 dih12 h 5 hc13 4 hcc13 6 dih13 7 dih14 6 hc14 5 hcc14 h h 6 hc15 5 hcc15 7 dih15 oh2 0.950000 co3 1.380000 coh3 109.471 1.500000 cc4 cco4 109.471 180.000 dih4 1.500000 сс5 ccc5 109.471 180.000 dih5 cc6 1.500000 ccc6 109.471 180.000 dih6 hc71.070000 109.471 hcc7dih7 180.000 hc8 1.070000 hco8 109.471 dih8 120.000 hc9 1.070000 hco9 109.471 dih9 240.000 hc10 1.070000 hcc10 109.471 dih10 120.000 hc11 1.070000 hcc11 109.471 dih11 240.000 hc12 1.070000 hcc12 109.471 dih12 120.000 hc13 1.070000 hcc13 109.471 dih13 240.000 hc14 1.070000 hcc14 109.471 dih14 120.000 hc15 1.070000

## Reference Z-matrix file for L-alanine (WE2).

h						
	1	oh2				
0	_					
С	2	co3	1	coh3		
С	3	cc4	2	cco4	1	dih4
С	4	cc5	3	ccc5	2	dih5
h	5	hc6	4	hcc6	3	dih6
0	3	oc7	2	oco7	4	dih7
n	4	nc8	3	ncc8	5	dih8
h	4	hc9	3	hcc9	5	dih9
h	5	hc10	4	hcc10	6	dih10
h	5	hc11	4	hcc11	6	dih11
h	8	hn12	4	hnc12	3	dih12
h	8	hn13	4	hnc13	12	dih13

oh2 0.950000 1.380000 co3 coh3 109.471 cc4 1.500000 cco4 109.471 dih4 180.000 сс5 1.500000 109.471 ccc5 dih5 180.000 hc6 1.070000 hcc6 109.471 dih6 180.000 1.380000 oc7 oco7 120.471 dih7 180.000 nc8 1.070000 ncc8 109.471 dih8 240.000 1.070000 hc9 hcc9 109.471 120.000 dih9 hc10 1.070000 hcc10109.471 120.000 dih10 hc11 1.070000 hcc11 109.471 dih11 240.000 hn12 1.030000 109.471 hnc12 dih12 180.000 hn13 1.030000 hnc13 109.471

120.000

dih13

dih17

240.000

#### Reference Z-matrix file for proline (WE3). С С 1 cc2 1 ccc3 С 2 cc3 3 cc4 2 ccc4 1 dih4 С n 4 nc53 ncc5 2 dih5 $2 \ \text{ccc6}$ 5 dih6 С 1 cc6 1 occ7 2 dih7 6 oc7 0 h 7 ho8 6 hoc8 1 dih8 6 oc9 7 dih9 0 1 occ9 h 1 hc10 2 hcc10 5 dih10 1 hcc11 2 hc11 3 dih11 h 1 hcc12 3 dih12 h 2 hc12 h 3 hc13 2 hcc13 4 dih13 4 dih14 3 hc14 2 hcc14 h h 4 hc15 3 hcc15 5 dih15 4 hc16 3 hcc16 5 dih16 h 5 hn17 4 hnc17 1 dih17 h cc2 1.500 1.500 cc3 ccc3 109.471 1.500 cc4 109.471 ccc4 dih4 0.000 1.500 nc5ncc5109.471 0.000 dih5 1.500 cc6 ccc6 109.471 -120.000 dih6 1.500 oc7 occ7 109.471 dih7 180.000 ho8 1.070 hoc8 109.471 dih8 180.000 oc9 1.380 occ9 109.471 dih9 180.000 hc10 1.070 hcc10 109.471 dih10 120.000 hc111.070 hcc11109.471 120.000 dih11 hc12 1.070 hcc12109.471 dih12 240.000 hc13 1.070 hcc13 109.471 dih13 120.000 hc141.070 109.471 hcc14 dih14 240.000 hc15 1.070 109.471 hcc15 120.000 dih15 hc16 1.070 109.471 hcc16 dih16 240.000 1.030 hn17hnc17 109.471

### Reference Z-matrix file for the H<sub>a</sub>-abstraction by H in n-BuOH (WE4).

```
h
      1 oh2
 0
 С
      2 co3
                     1 coh3
                                      1 dih4
      3 cc4
                     2 cco4
 С
 С
      4 cc5
                     3 ccc5
                                      2 dih5
                                      3 dih6
 С
      5 cc6
                     4 ccc6
                                      4 dih7
 h
      6 hc7
                     5 hcc7
 h
      3 hc8
                     2 hco8
                                      4 dih8
                                      4 dih9
 h
      3 hc9
                     2 hco9
 h
      4 hc10
                     3 hcc10
                                      5 dih10
      4 hc11
                     3 hcc11
                                      5 dih11
h
h
      5 hc12
                     4 hcc12
                                      6 dih12
 h
      5 hc13
                     4 hcc13
                                      6 dih13
      6 hc14
                     5 hcc14
                                      7 dih14
h
h
      6 hc15
                     5 hcc15
                                      7 dih15
                                      2 dih16
хx
      9 xxh16
                     3 xxhc16
      9 hh17
                    16 hhxx17
                                      3 dih17
h
oh2
            0.950000
co3
            1.380000
coh3
             109.471
             1.500000
cc4
            109.471
cco4
dih4
            180.000
             1.500000
cc5
ccc5
             109.471
dih5
            180.000
            1.500000
cc6
ccc6
            109.471
dih6
            180.000
             1.070000
hc7
hcc7
            109.471
dih7
            180.000
hc8
             1.070000
hco8
            109.471
dih8
             120.000
hc9
             1.288000
hco9
            109.471
dih9
             240.000
hc10
             1.070000
hcc10
             109.471
dih10
             120.000
hc11
            1.070000
            109.471
hcc11
dih11
             240.000
hc12
             1.070000
hcc12
             109.471
dih12
             120.000
             1.070000
hc13
hcc13
             109.471
dih13
             240.000
             1.070000
hc14
hcc14
             109.471
            120.000
dih14
hc15
             1.070000
hcc15
             109.471
dih15
             240.000
xxh16
             1.000000
             90.000
xxhc16
dih16
               0.000
             0.993000
hh17
hhxx17
             90.000
```

dih17

180.000

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