#### MicroSolv v.1.0.0

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#### 1. Brief theoretical introduction

#### Microsolv generator algorithm **Input:** Starting cluster geometry {X,Y,Z}<sub>0</sub>, atomic charges {Z<sub>i</sub>}, rotatable bonds N<sub>ROT</sub> **Output:** N<sub>CONF</sub> cluster geometries {X,Y,Z}<sub>i</sub> // determination of connectivity in the molecule Connectivity: write (automatic) or read **Initialization: for** *i* in range( $N_{CONF}$ ) **do**: // setting up random values $6N_{FRG} + N_{ROT} \leftarrow rand()$ **Procedure:** preliminary generation **for** *i* in range( $N_{CONF}$ ) **do**: // optimization with N<sub>ITER1</sub> steps $\{X,Y,Z\}_i \leftarrow min(6N_{FRG} + N_{ROT}, N_{ITER1})$ // minimal non-bonded distance check if mindist $(\{X,Y,Z\}_i)$ < thld: remove $(\{X,Y,Z\}_i)$ else: pass Clustering ( $\{X,Y,Z\}_1,k_1$ ) N<sub>CONF</sub>=k<sub>1</sub> **End Procedure Procedure:** final generation **for** i in range( $N_{CONF}$ ) **do:** // optimization with N<sub>ITER2</sub> steps $\{X,Y,Z\}_i \leftarrow min(6N_{FRG} + N_{ROT}, N_{ITER2})$ **for** *i* in range( $N_{CONF}$ ) **do**: **for** *j* in range( $N_{CONF}$ -1) **do**: // close in energies? **if** $|E(\{X,Y,Z\}_i) - E(\{X,Y,Z\}_i)| < thld:$ **if** RSMD( $\{X,Y,Z\}_i - \{X,Y,Z\}_i$ ) < thld: $N_{CONF}=N_{CONF}-1$ remove $(\{X,Y,Z\}_i)$ Clustering ( $\{X,Y,Z\}_1,k_2$ ) N<sub>CONF</sub>=k<sub>2</sub> **End Procedure** return N<sub>CONF</sub> {X,Y, Z}<sub>I</sub>

**Scheme 1.** Pseudo code of the *microsolv* spatial structure generator.

### 2. Quick start

#### 2.1 Prerequisites

- 64bit Linux OS
- Microsolv program (microsolv) & input file (filename.inp)
- Input file should contain:
  - **x** Rotatable bonds (if any)
  - **✗** Initial Cartesian coordinates of the (super-)molecule
  - **x** Atomic charges
- Running the code: "/path/to/program/microsolv filename.inp", e.g. ~/Isomers/127/microsolv m1\_01\_Li\_VC\_N3.inp

### 2. Main Keywords

### 2.1 Input file general design

Each keyword group starts with "\$" symbol and should be closed with "\$END" expression.

```
$BOND
28 29 120 240 15
$END
$ALGO
$END
$MOLECULE
Li
    -3.118257632
                 1.717973895
                               0.890951148
C
     0.543523281
                  0.285069258
                              -0.272235148
$END
$CHARGE
1.0
0.778632
$END
```

### 2.2 Group \$BOND

Here the rotational bonds should be specified in the following format:

```
Atom1 Atom2 Angle_Increment Angle_MAX_value Angle_MAX_deviation 1 2 120 240 15
```

In the example above the rotational bond is between atom 1 and atom 2. This torsion is specified to have three values: 0, 120, 240. With 0 value here the starting torsion angle value is meant. In case of no rotatable bonds, leave this group empty.

# 2.3 Group \$MOLECULE

Cartesian coordinates of all atoms should be provided in this section.

# 2.4 Group \$CHARGE

Atomic charges for each atom in \$MOLECULE group should be provided here.

# 2.2 Group \$ALGO

CONNECT	=WRITE =READ	// automatic connectivity detection that is written to filename.bond // filename.bond can be modified to adjust/correct the connectivity // (default) // connectivity is read from filename.bond // needed in cases when automatic connectivity fails			
ALGO	=CONF	// generate conformers (default)			
MK	=int =0	// integer number (e.g. 10) N of initially generated conformers on-the-fly; // initial NCONF value in the pseudo code above // systematic conformational search with respect to provided torsions // see \$BOND group above			
COM SCALE	=int =float	// what is considered to be center of coordinates // -1 center of mass of the geometry specified in \$MOLECULE (default) // 0 N — center of mass of individual fragment in the cluster // numeration starts from 0. It is convenient when an ion needs to // be surrounded by molecules, e.g. Li+ // any float value from 0.0 to 0.99 (default: 0.75)			
SCALE =float		// dny float value from 0.0 to 0.99 (default, 0.75)  // characterizes the maximum of individual fragment com from  // the initial center. The lesser the value the closer to the inital  // structure conformer is generated.			
LIBRARY	=NLOPT	// optimization library for potential energy optimization (default)			
// more at https://nlopt.readthedocs.io/en/latest/  OPTGLOB =none // no global optimization (recommended) (default) =nlopt_gn_direct_l =nlopt_gn_direct_l_rand =nlopt_gn_direct_l_rand_noscal =nlopt_gn_orig_direct =nlopt_gn_orig_direct_l =nlopt_gn_orig_direct_l =nlopt_gn_crs2_lm =nlopt_gn_mlsl =nlopt_gn_mlsl_lds =nlopt_gn_isres =nlopt_auglag =nlopt_auglag_eq					

```
=nlopt_gn_esch
       OPTLOC
                     =nlopt ln sbplx
                                           // local optimization method, must be specified! (default)
                     =nlopt_ln_praxis
                     =nlopt_ln_cobyla
                     =nlopt_ln_newuoa
                     =nlopt_ln_newuoa_bound
                     =nlopt ln neldermead
                     =nlopt_ln_sbplx
                     =nlopt_ln_auglag
                     =nlopt_ln_auglag_eq
                     =nlopt_ln_bobyqa
ITER1
                            // number of iterations for the first (prelinimary) conformer generation
              =int
                            // (default: 10)
DIST
              =float
                            // minimal non-bonded distance threshold for conformer exclusion
                            // after preliminary and final conformer generation (default: 1.25)
              =FALSE
                            // DIST related keyword. When searching for the minimal non-bonded
NONHD
                            // distance, do not skip the pairs including hydrogens (e.g. C - H)
                            // (default)
              =TRUE
                            // skip the pairs that contain hydrogens
SORT
                            // after minimal distance test (see DIST) that occurs after preliminary
              =e
                            // and final conformer generation, all conformers are sorted based on
                            // certain property. E stands for sorting by energies (default)
                            // sort by the average distance from each fragment's com to the whole
              =d1
                            // molecule center of mass
              =d2
                            // same as d1 but in case of COM!=-1 (i.e. when center is not the whole
                            // molecule center of mass)
                            // after sorting above, take that amount of conformer is taken
MAXCONF =int
                            // for further treatment (default: 100)
                            // first clustering option (in the preliminary conformer generation)
KMEANS
              =int
                            // -3: K (number of clusters) is estimated K=Nconf/2
                            // -1: (default) no clustering
                            // 0: K = (Nconf/2)**0.5
                            // Any int number, e.g. 10
KMEANS1
              =int
                            // second clustering option (after the final conformer generation)
                            // only for STRATEGY=1, see below
                            // -3: K (number of clusters) is estimated K=Nconf/2
                            // -1: (default) no clustering
                            // 0: K = (Nconf/2)**0.5
                            // Any int number, e.g. 10
CLUSTERPROPERTY1
                             =IXX // the clustering property; the whole molecule sorted principal
                                    // moments of inertia (3)
                             =distcommol // distance from each individual fragment com to the
                                           // molecular COM
                                           // three sorted principal moments of the whole molecules
                             =ixxfrg
                                           // and each individual fragment (default)
```

			// the whole molecule energy and three principal moments // of inertia		
			// translational variables for each fragment		
		=rot	// rotational variables for each fragment		
		=transrotbond // translational, rotational variables and torsion bonds			
			// for each fragments		
ITER2	=int	// number of iterations for the second (final) conformer generation // (default: 10 000)			
EDIFF	=float	// energy difference floating point value below which the			
		// structure geometries are compared with quatit RMSD			
		// (default: 0)	1 1		
RMSD	=float		ria for difference between the two structures for discrarding		
			ARDING procedure after final conformer generation		
MAXCONFPRINT =int		// number of conformers for final priting to the file			
STRATEGY	=0	0	treat the conformers after the final generation (default: 0)		
		// just print MAXCONFPRINT conformers to the file			
	=1	// clustering of all conformes after the final generation, followed by // sorting. After that MAXCONFPRINT conformers are printed to file.			

# 2. Detailed documentation (advanced keywords)

```
// simple optimization of a given structures
ALGO
             =OPTIMIZE
                                  // no descrete values of bond torsions are allowed
                           // the number of geometry optimization steps for ALGO=OPTIMIZE
       ITER2
                    =int
                           // is given with ITER2
                           // Parallel mode for potential energy single-point calculation (default:
PAREN
             =FALSE
                           // DO NOT USE THIS, NO SPEED UP IS OBSERVED
             =TRUE
                           // NLOPT optimization-related keyword (default: 1e-3)
TESTSIZE
              =float
              =float
                           // scaling coefficient for electrostatic energy (default: 1)
KEL
              =float
                           // scaling coefficient for the DFT-D2 Grimme's dispersion
KGD
```