Uniconf v.1.0.0

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1 System requirements

- 64bit Linux OS
- Conformer generator binary (uniconf) & input file (filename.inp)
- Input file should contain:
 - **x** Rotatable bonds (if any)
 - **✗** Initial Cartesian coordinates of the (super-)molecule
 - **x** Atomic charges
- download the code and change the permission: chmod u=x uniconf
- Running the code: "/path/to/program/uniconf filename.inp", e.g. ~/Isomers/128/uniconf m1_01_Li_VC_N3.inp

2 Quick start

The Uniconf conformer generation procedure is depicted in Scheme 1 and described in the original manuscript. Below only the technical information as well as a few hints and tips on effective using the code are provided.

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
    -3.118257632
                1.717973895 0.890951148
C
     0.543523281
                  0.285069258 -0.272235148
$END
$CHARGE
1.0
0.778632
$END
```

Uniconf 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<sub>CONF</sub>) do:
    // optimization with N<sub>ITER1</sub> steps
    \{X,Y,Z\}_i \leftarrow min(6N_{FRG} + N_{ROT}, N_{ITER1})
    // shortest non-bonded distance check
    if mindist (\{X,Y,Z\}_i) < \text{thld}:
        remove (\{X,Y,Z\}_i)
    else:
        pass
Clustering ({X,Y, Z}<sub>1</sub>, k<sub>1</sub>)
N_{CONF}=k_1
End Procedure
Procedure: final generation
for i in range(N<sub>CONF</sub>) do:
    // optimization with NITER2 steps
    \{X,Y,Z\}_i \leftarrow min(6N_{FRG} + N_{ROT}, N_{ITER2})
for i in range(N<sub>CONF</sub>) do:
        for j in range(N_{CONF}-1) do:
            // close in energies?
            if |E(\{X,Y,Z\}_i) - E(\{X,Y,Z\}_j)| \le thld:
                if RSMD(\{X,Y,Z\}_{1} - \{X,Y,Z\}_{j}) < thld:
                    N_{CONF} = N_{CONF} - 1
                    remove (\{X,Y,Z\}_i)
Clustering (\{X,Y,Z\}_1, k_2)
N_{CONF}=k_2
End Procedure
return N<sub>CONF</sub> {X,Y, Z}<sub>I</sub>
```

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

2.2 Input file main groups

2.2.1 Group \$BOND

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

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.

2.2.2 Group \$MOLECULE

Cartesian coordinates of all atoms should be provided in this section. These can be straightforwardly obtained by placing together all molecular fragments at random positions in space in any molecular builder of choice, e.g. Chemcraft available at https://www.chemcraftprog.com/

2.2.3 Group \$CHARGE

Atomic charges for each atom in \$MOLECULE group should be provided here. These can be produced by running the quantum chemical program and model chemistry of choice in the single point fashion either on each individual fragment in the system or on the whole cluster. When possible, we recommend computing the CHELPG charges and using them in the input file. For molecular or microsolvated structures we suggest calculating the charges for each molecular fragment separately and merging them afterwards. Thus generated charges are found to result in more robust conformational search than using the charges obtained with CHELPG procedure on the whole cluster.

2.2.4 Group \$ALGO

This group defines the conformer generation algorithm details with its main keywords given in Section 3. Here we restrict ourselves with only a few remarks.

- As a result of the default molecular connectivity procedure, the connectivity text file bearing the ".bond" extension is written. In the majority of cases, the internal algorithm defines the connectivity correctly. In some rare cases of failure, there is a possibility to edit the ".bond" file manually, and then re-run the code reading the connectivity. In fact, prior to the conformational search we recommend running the code in the structure optimization mode and checking the optimized structure. The resulting ".bond" file should be used for further conformational search.
- To achieve maximum diversity at the fixed CPU time, we suggest using the limited number of geometry optimization steps (ITER1) along with the maximum number of conformers to be generated.
- At the final stage, all collected after ITER1 procedure conformations are subjected to a reoptimization procedure but with a larger number of steps (ITER2). However, it has to be
 mentioned that the number of steps should be chosen thoughtfully. As the potential energy

surface (PES) employed in the Uniconf does not resemble the true one, too many steps might take the structure away from a conformation otherwise deserving consideration.

3. Group \$ALGO main keywords

```
CONNECT
              =WRITE
                            // automatic connectivity detection that is written to filename.bond
                            // filename.bond can be modified to adjust/correct the connectivity
                            // (default)
              =READ
                            // connectivity is read from filename.bond
                            // needed in cases when automatic connectivity fails
ALGO
                            // generate conformers (default)
              =CONF
                            // integer number (e.g. 10) N of initially generated conformers on-the-fly;
MK
              =int
                            // initial NCONF value in the pseudo code above
              =0
                            // systematic conformational search with respect to provided torsions
                            // see $BOND group above
COM
              =int
                            // what is considered to be center of coordinates
                            // -1 center of mass of the whole cluster 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+
SCALE
              =float
                            // any 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.
ORBSCALE =FALSE
                            // do not scale of orbital motion (default)
                            // characterizes the maximum of individual fragment com from
              =TRUE
LIBRARY
              =NLOPT
                            // optimization library for potential energy optimization (default)
                            // more at https://nlopt.readthedocs.io/en/latest/
                                   // no global optimization (recommended) (default)
       OPTGLOB
                     =none
                     =nlopt_gn_direct_l
                     =nlopt_gn_direct_l_rand
                     =nlopt_gn_direct_noscal
                     =nlopt_gn_direct_l_rand_noscal
                     =nlopt_gn_orig_direct
                     =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
```

	=nlo =nlo =nlo =nlo =nlo	pt_ln_newuoa pt_ln_newuoa_b pt_ln_neldermea pt_ln_sbplx pt_ln_auglag pt_ln_auglag_eq pt_ln_bobyqa	nd	
ITER1	=int		terations for the first (prelinimary) conformer generation	
DIST	=float	// minimal no	n-bonded distance threshold for conformer exclusion	
NONHD	=FALSE	// DIST relate	inary and final conformer generation (default: 1.25) d keyword. When searching for the minimal non-bonded o not skip the pairs including hydrogens (e.g. C – H)	
	=TRUE		rs that contain hydrogens	
SORT	=e	// and final co	al distance test (see DIST) that occurs after preliminary onformer generation, all conformers are sorted based on perty. E stands for sorting by energies (default)	
	=d1	// sort by the @ // molecule ce	average distance from each fragment's com to the whole enter of mass	
	=d2		but in case of COM!=-1 (i.e. when center is not the whole	
MAXCONF	=int	// after sorting	g above, take that amount of conformer is taken	
KMEANS	=int	// first cluster // -3: K (numl	· /	
KMEANS1	=int	// only for STI // -3: K (numl		
CLUSTERPROPERTY		=IXX // the clustering property; the whole molecule sorted principal // moments of inertia (3) =distcommol // distance from each individual fragment com to the		
		=ixxfrg	// molecular COM // three sorted principal moments of the whole molecules	
		G	// and each individual fragment (default)	
		=eixx	// the whole molecule energy and three principal moments // of inertia	
		=trans =rot	// translational variables for each fragment // 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
RMSD	=float	// (default: 0) // RMSD criteria for difference between the two structures for discrarding // at the DISCARDING procedure after final conformer generation // utilized by Quatfit algorithm
MAXCONFI	PRINT =int	// number of conformers for final priting to the file
STRATEGY	=0	// strategies to 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)

```
ALGO
              =OPTIMIZE
                                   // simple optimization of a given structures
                                   // no descrete values of bond torsions are allowed
       ITER2
                     =int
                            // the number of geometry optimization steps for ALGO=OPTIMIZE
                            // is given with ITER2
PAREN
              =FALSE
                            // Parallel mode for potential energy single-point calculation (default:
                            // FALSE)
                            // DO NOT USE THIS, NO SPEED UP IS OBSERVED
              =TRUE
ITER
              =int
                            // clustering-related interation number (default: 1)
TESTSIZE
              =float
                            // NLOPT optimization-related keyword (default: 1e-3)
              =float
                            // scaling coefficient for electrostatic energy (default: 1)
KEL.
                            // scaling coefficient for the DFT-D2 Grimme's dispersion
KGD
              =float
                            // sometimes it helps for stronger binding of the fragments (default 0)
KVW
                            // scaling coefficient for the van der Waals UFF energy (default: 1)
              =float
HB1
              =float
                            // A hydrogen bonding coefficient, see main article (default: 6.71)
              =float
                            // B hydrogen bonding coefficient, see main article (default: 3.55)
HB2
                            // alpha hydrogen bonding coefficientm, see main article (default: 0.75)
HB3
              =float
ALGO
              =FIT
                            // Fitting parameters to reproduce the initial geometry specified in
                            // $MOLECULE group
       FIT
              =false
                            // no fitting (default)
                            // fitting electrostatic energy coefficient
              =KEL
                            // fitting van der Waals non-bonded UFF energy
              =KVW
              =KELKVW
                            // fitting both electrostatic and van der Waals energies
                            // fitting the hydrogen bond coefficients
              =HBOND
                                   // number of iterations for fitting (default: 1000)
       ITERFIT
                            =int
                                   // number of geometry optimization steps inside fitting step
       ITERFIRGEOM
                            =int
                                   // (default 1000)
                                   // global optmization algorithm for fitting (default:
       OPTGLOBFIT
                            =str
                                   // nlopt_gn_orig_direct)
                                   // local optimiztion algorithm for fitting (default:
       OPTLOCFIT
                            =str
                                   // nlopt ln sbplx)
                                   // local optimization algorithm for geometry optimization
       OPTLOCFITGEOM =str
                                   // inside fitting (default: nlopt ln sbplx)
                            =K-MEANS
                                                 // (default) algorithms for clustering
CLUSTERALGORITHM
                            =K-MEDOIDS
                                                 // (memory intensive, use with caution)
PRINTDATA
                                   // (default) no detailed output for clusterization
                            =()
                            =1
                                   // detailed output for clusterization
                                   // (default) do not use the centroid center, use minimum E
CENTROIDCENTER
                            =()
                                   // structure
                                   // use the cluster centroid
                            =1
PRINTLEVEL
                            =0
                                   // (default: 0). Do not print coordinates after ITER1 & ITER2
```

=-1 // print all coordinates at the intermediate steps.