

Impressum

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Manual

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ABOUT CIMATCH™

Clmatch™ overlays molecules based on their standard isodensity surfaces and local properties projected onto them.

The simplest way to match the surfaces is a SHAPE match, which only considers the geometries of the two surfaces. Figure 1 shows the algorithm schematically:

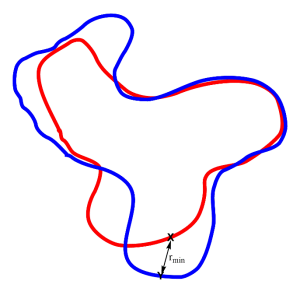


Figure 1 Schematic diagram of the algorithm used to match two surfaces

Similarity indices

The shape similarity index between surfaces A and B, S_{AB}^{R} , which is zero for identical surfaces (i.e. A =B), is given as:

$$S_{AB}^{R} = \sum_{i=1}^{N_A} r_{\min}^i \tag{1}$$

where N_A is the number of surface points for molecule A and r_{\min}^i is the minimum distance from point ion surface A to any point on surface B. Note that any point on surface B (e.g. Y in Figure 1) may be closest to more than one point on surface A, and therefore occur several times for r_{\min}^i . This means that $S_{AB} \neq S_{BA}$.

The similarity index can be weighted by any of the local properties MEP (V), EA_L (A), IE_L (I) or α_L (P) to give the following similarity indices:

$$S_{AB}^{R} = \sum_{i \neq l}^{N_{A}} r_{\min}^{i}$$

$$S_{AB}^{RV} = \sum_{i \neq l}^{N_{A}} r_{\min}^{i} \cdot \delta V_{\min}$$

$$MEP$$

$$S_{AB}^{V} = \sum_{i \neq l}^{N_{A}} \delta V_{\min}$$

$$S_{AB}^{RI} = \sum_{i \neq l}^{N_{A}} \delta I_{\min}$$

$$S_{AB}^{RI} = \sum_{i \neq l}^{N_{A}} \delta I_{\min}$$

$$S_{AB}^{RI} = \sum_{i \neq l}^{N_{A}} \delta I_{\min}$$

$$S_{AB}^{RA} = \sum_{i \neq l}^{N_{A}} \delta A_{\min}$$

In these cases, $\delta V_{\rm min}$ is the difference in the MEP projected onto the two surface points (**X** and **Y** in **Figure 1**), and analogously for the other surface properties. Note that for the similarity indices that do not include the shape, the points are still matched as shown in **Figure 1** but the distance between them does not enter the equation for the similarity index.

The closer the surface points and the smaller the difference between their local-property values, the better the match. A perfect match gives an overlay value of zero. Overlaying a molecule on itself is a good way to test program options to optimize performance. The program's default options are conservative and chosen to give reproducible results at the expense of calculational effort.

The matching algorithm

The program uses a genetic algorithm to find the best global overlay. The requested similarity index is optimized by translating and rotating molecule *B* while holding *A* constant. The optimization process consists of *repeat* iterations of complete evolution runs. Each run uses *npop* members of the population and runs for *maxgen* generations. The overall result is the lowest found in the series of evolution runs.

The size of the population (default 32), maximum number of generations (default 200) and number of repeat runs (default 20) are adjustable parameters.



CALLING CIMATCH™

<path> CImatch.exe <mol1> <mol2> <options>

Clmatch™ is designed for efficient parallel execution; it will use all available cores.

Input files

The default input file type is the ParaSurf™ output file <*mol>.psf*. However, in the absence of the *force* option, CImatch™ will check which input files are available and, if necessary, perform ne EMPIRE™ and/or ParaSurf™ calculations. In this case, a file named *CImatch.com* in the current directory is needed to define the calls to EMPIRE™ and ParaSurf™ (see below).

NOTE: Giving the *.psf* extension is optional in the program call (i.e. *Clmatch.exe mol1 mol2* and *Clmatch.exe mol1.psf mol2.psf* are identical.

NOTE: Clmatch.exe mol1 mol2 is not the same as Clmatch.exe mol2 mol1, as outlined above.

Options (case insensitive)

function= <fn></fn>	function= <fn> <fn> defines the similarity index to optimized. It may be one of:</fn></fn>		o be
	R	Shape only ($S^{\scriptscriptstyle R}_{\scriptscriptstyle AB}$)	
	RV	$Shape \times MEP \ (\ S^{\mathit{RV}}_{\mathit{AB}} \)$	Default
	V	$MEP(S^{V}_{AB})$	
	RI	Shape \times IE _L (S_{AB}^{RI})	
	I	$IE_L (S^I_{AB})$	
	RA	$Shape \times EA_L (S^{\mathit{RA}}_{\mathit{AB}})$	
	Α	$EA_L\left(\left.S_{AB}^{A}\right. ight)$	
	RP	Shape $ imes$ $lpha_{L}$ (S^{Rlpha}_{AB})	
	Р	$lpha$ L (S^lpha_{AB})	
npop=< <i>n</i> >		Defines the size of the population in the genetic algorithm.	Default=32

Default=200 Defines the number of maxgen=<n> generations within a single optimization run. repeat=<n> Defines the number of times the Default=20 optimization should be repeated. Forces Clmatch™ to request force=<s> new EMPIRE™ and/or ParaSurf™ calculations Do both new EMPIRE™ and $\langle s \rangle = EMPIRE$ ParaSurf™ calculations for template and target, regardless of which files are present. $\langle s \rangle = PARASURF$ Only do new ParaSurf™ calculations for template and target, regardless of which files are present. mode=<s1>Defines the precision with which the program works. Possible values are: $\langle s \rangle = QAD$ "Quick and dirty" mode (npop=8, maxgen=100, repeat=3). This mode is intended for fast initial scans. $\langle s \rangle = MED$ "Medium" mode (npop=24, maxgen=100, repeat=10). This mode is intended as a compromise between extensive searching and computational speed. $\langle S \rangle = FIN$ "Fine" mode (npop=32, Default maxgen=200, repeat=20). This is the default mode and represents a very conservative setup for extensive sampling at the expense of cpu time. mode=<s1>,<s2> Requests that a first scan be made with multiple templates at the level <s1>, followed by selection of the best template conformations for calculations at level <s2>. The default option is given by select alone. select=<i> Defines the number of templates Default is to be selected. <*i>* = the larger of 3 or the number of targets/10

PROGRAM INPUT FILES

The minimum requirement for Clmatch™ input (provided EMPIRE™ and ParaSurf™ are available) are <mol1>.sdf and <mol2>.sdf files for template and target as input for EMPIRE™. A second alternative starts with <mol1>_e.sdf and <mol2>_e.sdf output files from EMPIRE™ and only perfoms ParaSurf™ calculations. If <mol1>.psf and <mol2>.psf ParaSurf™ files are present, Clmatch™ also requires that the two ParaSurf™ output SDF files <mol1>_p.sdf and <mol1>_p.sdf be present in the same directory as the input ParaSurf™ files. These SDF-files are required to write the Clmatch™ output SDF file, which contains the overlaid geometries of the two molecules and their bonds, which are taken from the input ParaSurf™ SDF files. Both EMPIRE™ and ParaSurf™ SDF files may contain multiple molecules, which will all be processed.

The complete process to perform a Clmatch™ overlay is therefore:

- 1. Perform EMPIRE™ single-point calculations or geometry optimizations on the two molecules to obtain EMPIRE™ <*mol>_e.sdf* files, which are used as input for
- ParaSurf™, which calculates the input <mol> .psf files and updates the input <mol> _e.sdf files to <mol>_p.sdf. The default ParaSurf™ grid size for the surface is too small for efficient Clmatch™ calculations, so that the option mesh=0.5 should be used.
- 3. Perform the Clmatch™ calculation as outlined above.

This process is performed automatically if the necessary files are not present. In this case, an addition file named *Clmatch.com* in the working directory is necessary to define the calls to EMPIRE™ and ParaSurf™ (see below).

The Clmatch.com command file

If EMPIRE™ and/or ParaSurf™ calculations are to be performed, the call for the two programs must be given in a file named *Clmatch.com* in the working directory. For Windows, a typical *Clmatch.com* might be:

```
c:\bin\empire_AM1spt.bat
c:\bin\parasurf AM1.bat
```

where *empire_AM1spt.bat* is a script to run an AM1 single-point calculation with EMPIRETM and *parasurf_AM1.bat* to run a subsequent ParaSurfTM calculation. It is important in the latter that the **ParaSurf**TM **options** psf=on and mesh=0.5 are given.



PROGRAM OUTPUT

ClmatchTM writes a text file $< mol1>_< mol2>_< FUNCTION>.log$, an SDF file of the overlaid molecules $< mol1>_< mol2>_< FUNCTION>.sdf$ and one of the target in its overlaid coordinates $< mol2> @< mol1>_< FUNCTION>.sdf$.

The Clmatch™ log file

The program log file provides details of program execution and the calculated similarity indices. **Box 1** shows a log file obtained using default parameters.

```
h
h hh
                                              СС
                    mm m m
                      m
m
                                                     hh
                               aaaa
                   m m m
              III m m m aaa a
                                                                  (c) Cepos InSilico GmbH, Obermichelbach 2017, 2019
Started at: Fri Apr 26 15:56:51 2019 on PC_TIM
<> Template molecule
<> Target molecule
<> Target function
                                 : Shape * MEP
<> Population
<> Maximum Nr. generations
<> Nr. evolutionary cycles
                                       2.0
<> Numbers of surface points : Template:
<> Maximum shift
                                      2.617 Angstrom
<> Translational resolution :
                   1.997 : Best yet :
2.035 : Best yet :
Cycle 2:
Cycle 3:
Cycle 4:
                   2.019 : Best yet :
                                                1.997
                   2.040
                                                1.997
                                               1.932
1.932
1.932
Cycle
                   1.932 : Best vet :
Cycle
Cycle
                   2.090
                            Best vet
Cycle
Cycle
                                               1.932
                   2.127
                            Best yet
                            Best yet
Cycle 10:
                   2.016
                            Best vet
                                                1.932
                   1.956
Cycle
Cycle 12:
                            Best vet
                                                1.932
Cycle 13:
Cycle 14:
                                               1.932
                   2.012
                          : Best yet
                   2.051
                            Best yet
Cycle 15:
Cycle 16:
                            Best yet
Best yet
                   1.999:
                                                1.932
Cvcle 17:
                   1.954
                            Best vet :
                                                1.932
                   1.946
1.982
Cycle 18:
                          : Best yet
                                                1.932
Cycle 19:
                            Best yet
                                                1.932
Cycle 20:
                   2.132 : Best yet
                                                1.932
<> Scores (Function optimized indicated by asterisks):
                                                             Pol(L)
                                                                         Shape*MEP Shape*IE(L) Shape*EA(L) Shape*Pol(L)
                                                                           1.932
```

Box 1 Example of a Clmatch™ log file using default parameters and a single template and target

After printing the header and the details of the calculation, Clmatch™ calculates the maximum translation allowed for the second molecular center relative to the first and gives the resolution of the translational moves. These two parameters are determined from the maximum dimensions of the molecules involved. The translational range is designed to cover all possible significant overlays and the resolution results from adapting this range to the overlay genes.

The program then prints the results of each evolutionary run (Cycle); in this case 20. The best result in all runs is stored and becomes the final output. Note that the results of the individual runs vary between 1.932 and 3.132, a typical range for multiple overlays of the same molecules

The final scores (i.e. those for the best solution) are given as a one-row table. The target parameter (in this case *Shape*MEP*) is marked by the row of nine asterisks. The scores for the other variables <u>at the geometry of the optimized overlay for the target *Shape*MEP* are given in the other columns.</u>

The Clmatch™ target .sdf file

The orientation and position of the overlaid target molecule are written in a file named <mol1>@<mol2>_e.sdf.

The Clmatch™ overlay .sdf file

ClmatchTM writes an output .sdf file with the two molecules in the calculated overlay. The bonds of the two individual molecules are taken from the input < mol >_e.sdf files.

Figure 2 shows a visualization of such an overlay:

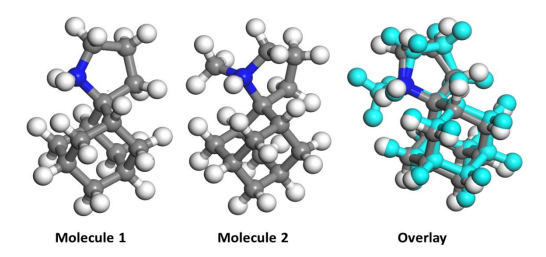


Figure 2 The two input molecules and the resulting overlay from the Clmatch™ calculation shown in Box 1

The Clmatch™ table .csv file (Template_<input1>_<function>.csv)

Clmatch™ writes an output .csv file that lists the results for all templates and targets the separator is a comma). Separate tables of all the targets are written for multiple templates and for two-stage runs an additional table for the selected targets at the second calculation mode. This file can be imported directly into spreadsheet programs.



SUPPORT

Contact

Questions regarding Clmatch™ should be sent directly to:

support@ceposinsilico.com

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LIST OF FIGURES AND BOXES

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