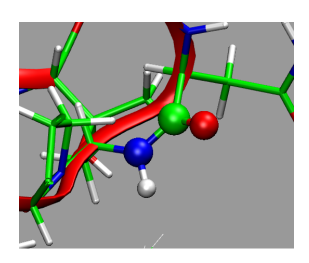
University of Illinois at Urbana-Champaign NIH Resource for Macromolecular Modeling and Bioinformatics Beckman Institute Computational Biophysics Workshop

Structure Check



Authors: Eduard Schreiner Leonardo G. Trabuco

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A current version of this tutorial is available at http://www.ks.uiuc.edu/Training/Tutorials/

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Introduction

The purpose of this tutorial is to demonstrate how to use the VMD plugins **cispeptide** and **chirality** together with **AutoIMD** to ensure proper structure of peptide bonds and correct stereochemistry of proteins and nucleic acids. The described plugins are designed to work with proteins and nucleic acids and understand atom names as used by the force fields CHARMM and AMBER as well as the standard PDB naming convention for non-hydrogen atoms.

WARNING: The current implementation of the chirality plugin does not support DNA yet.

The following points aim to motivate the necessity of the of the presented plugins:

- Structures from PDB may contain errors in stereochemistry and peptide bond configuration;
- Errors in peptide bond configuration can be introduced by modeling;
- Poor starting conditions, i.e. atoms being too close to each other, can generate huge forces during the initial optimization of the structure leading to errors in structure;
- Molecular dynamics (MD) protocols which apply forces on the molecule may generate forces large enough to introduce errors.

Since MD force fields are defined to support either type of chirality or peptide bond configuration, optimization of the structure will not correct introduced errors.

At his stage, we would like to give a brief outline of the strategy behind the described plugins. The general strategy to ensure correct peptide bond and stereochemical configuration consists of four steps.

- **1** Identify unusual configuration at chiral centers and peptide bonds All *cis* peptide bonds are regarded being unusual. In case of chiral centers, those are regarded unusual which occur less often in nature, e.g. D-form of amino acids.
- **2** Inspect the found irregularities Although it is possible to convert all peptide bonds and chirality centers into their "usual" configuration (this feature will be implemented in the future), it is safer to decide for each irregularity individually. First, different chiral forms of biomolecules and true *cis* peptide bonds are used in biology. Second, it is not always unambiguous how to correct an unusual configuration; this is especially the case for peptide bonds.
- **3** Correct the structure if necessary using structure optimization (and equilibration) The correcting happens in two steps. First one atom is moved to "preset" a proper configuration. In a second step an the structure is relaxed using an MD force field plus restraints enforcing the proper configuration.

4 If proceeding with further MD simulations, apply restraints to preserve chirality and peptide bond configuration This step may be necessary, if the molecular system is used in a simulation in which large forces may be applied, e.g., MDFF, TMD etc.

1 Chirality in proteins and nucleic acids

1.1 Introduction

All amino acids but glycine have at least one chiral center at C_{α} (see Fig. 1). Threonine and isoleucine have an additional chiral center at C_{β} . According to the D-/L- naming convention, naturally occurring amino acids are found in the L-configuration. Note, however, that D-amino acids do occur in biology, e.g., in cell walls of bacteria. Nucleic acids also have chiral centers. For example, in DNA the atoms C1', C3', and C4' are chiral, while RNA has an additional chiral center at C2'. Chirality is central to all molecular interactions in biological systems. A simple experiment demonstrates the principle: try to shake someone's left hand with your right.

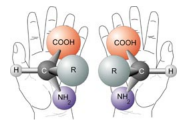


Figure 1: Chirality at the C_{α} carbon atom of amino acids. The two enantiomers are mirror images of each other, like the left and the right hand. Source: http://web99.arc.nasa.gov/ãstrochm/chirality.jpg

1.2 Checking stereochemistry of the structure

To demonstrate the usage of the plugin we will start by checking a structure consisting of the protein EF-Tu in complex with a tRNA^{Phe}. The used structure is based on the the PDB structure 10B2 in which errors have been introduced manually.

Currently, the **chirality** plugin can be used from the text console and through a GUI. The available commands provided by the pugin can be obtained by typing **chirality** on the Tk Console or the VMD console. The following information should be printed in the console:

Commands:

check	· identify	chirality errors
list	· list ider	ntified chirality errors
minimize	· fix chira	ality errors using energy minimization
move	· move hydr	rogen atom to fix chirality
reset	· reinitial	lize plugin state
restrain	- generate	NAMD extrabonds file to prevent chirality changes
show	· visualize	e identified chirality errors

In the same way, the usage information for any of the provided commands can be obtained. For example, typing **chirality restrain** will show the syntax of the command that generates restraints for the impropers at chiral centers which can be used in NAMD.

In the following, we will describe how to use the **chirality** plugin through its GUI, but all of the tasks can also be accomplished from the console.

- 1 Load the files chir_testcase.psf and chir_testcase.pdb into a new session of VMD.
- 2 Open the chirality GUI by selecting Extensions \rightarrow Modeling \rightarrow Fix Chirality Errors in the VMD Main menu. In the upper part of the GUI (see Fig. 2) the user can specify the molecule and the atom selection to be tested.

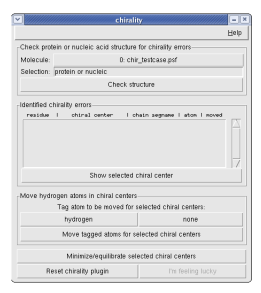


Figure 2: The chirality GUI window.

- **3** The button Check structure initializes the test. Identified unusual chiral center configurations are displayed in the Identified chirality errors form (see Fig. 3). In the present case there are 9 chiral errors identified in both the protein and RNA parts of the loaded structure.
- 4 By selecting an entry in the Identified chirality errors form and hitting the button Show selected chiral centers it is possible to visually inspect all the individual identified unusual chiral centers. In the created representation (see Fig. 4) the chiral center is highlighted by a transparent purple sphere, while the four bonded atoms are shown in CPK.

✓ ////////	chirali	ity				-×
						<u>H</u> elp
Check prof	Check protein or nucleic acid structure for chirality errors					
Molecule:	0: chir_testcase.psf					
Selection:	protein or nucleic					
	Check str	ructu	re			
Identified c residue URA 12 CYT 48 ADE 35 GLU 55 PRO 72 ILE 188 THR 115 ILE 188	hirality errors I chiral center I c H37 C37 C47 C27 C37 H47 C47 C57 C37 C47 H47 C47 C57 C37 C47 H47 C47 C58 H47 C47 C58 H47 C47 C58 H47 C47 C58 H47 C47 C47 H47 C47 C47 H47 C47 C47 H47 C47 H47 C47 C47 H47 C	2 2 2 P P P P P	N1 N1 P1 P1 P1 P1 P1 P1	e i atom i H H H H H H H H H	noved yes yes yes yes yes yes yes	
-Move hydr	ogen atoms in chiral centers	;				
	Tag atom to be moved for	seled	cted chir	al centers	:	
	hydrogen none					
	Move tagged atoms for selected chiral centers					
	Minimize/equilibrate selected chiral centers					
Re	Reset chirality plugin I'm feeling lucky					

Figure 3: Identified unusual chiral center configurations in the chirality GUI window.

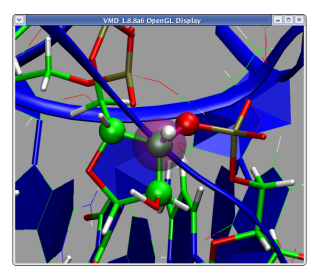


Figure 4: Representation of the chiral error in U12 of the tRNA

1.3 Correcting chirality

1 If the shown chiral center has the wrong chirality, the user can tag an atom to be moved to flip the chirality at the selected center. This is

done by hitting the button hydrogen. Note, currently only the moving of a hydrogen atom at a chiral center is supported. If the chirality at the displayed center is correct, nothing needs to be done. It is also possible to untag a tagged atom by selecting it and hitting **none**.

- 2 Once a chiral center has been inspected, and the atom to be moved has been tagged, the actual moving of the atom can be executed by hitting the button Move tagged atoms for selected chiral centers.
- 3 Since the described procedure (simple moving of the hydrogen atom) generates an unphysical geometry of the molecule, it is necessary to optimize the structure using an MD force field. The chirality plugin uses the AutoIMD plugin for this purpose. This final step is accomplished by selecting the chiral centers which should be relaxed and hitting the button Minimize/equilibrate selected chiral centers. This will open the AutoIMD Controls window (see Fig. 5).

AutoIMD Con	trols	////////			
File Settings		Help			
1. Type in the atom selections for imd (top molecule):					
imdmolten: (none or (segname N1 an	d resid 30) o	r (segname N1 a			
imdfixed: within 8 of imdmolten					
2. Choose where and how to run the simulation:					
Local 🛁 using 1 / 1 processors					
3. Submit your job:					
Submit Connect Pause	Finish	Discard			
status: Waiting for user input					

Figure 5: AutoIMD window called by chirality plugin.

- 4 Select Settings → Minimization Mode in the AutoIMD Controls window and hit then the Submit button. Once the AutoIMD session started, hit the Connect button and a minimization will start. Finish the simulation when the structure relaxed. Usually, it takes up to a few thousand steps until atoms do not move anymore. Is the structure minimized, hit the Finish in the AutoIMD Controls window. For more details about using the AutoIMD plugin, the user is referred to the AutoIMD user's guide http://www.ks.uiuc.edu/Research/vmd/plugins/autoimd/ug/.
- 5 WARNING: Depending on the quality of the initial structure, it can happen that not all errors can be fixed in one run. Thus, it is important to check the final result again and correct the remaining errors, if necessary. In order to check the structure after the performed minimization/equilibration using AutoIMD, the chirality plugin should be reset. This can be done by hitting the Reset chirality plugin button at the bottom of the chirality window. Note, in the current implementation it will

be necessary to select the molecule again in the top part of the chirality window. Note also that if the molecule to correct is solvated, it may be necessary to equilibrate the minimized parts of the structure. This can be done by selecting Settings \rightarrow Equilibration Mode and submitting a new AutoIMD run.

- 6 Finally, it is a good advice to minimize the structure in the very last step. This should be done by selecting Settings → Minimization Mode in the AutoIMD Controls window, which is still open from the last AutoIMD session, and hit then the Submit button. As mentioned in the introduction, additional restraints during the correction of chirality errors. The restraints will be used always when AutoIMD is started from the chirality window. The equilibrium values for these restraints are very approximate and bring the configuration in the correct region. Thus, in order to obtain a optimized structure, it is necessary NOT to use the restraints defined by chirality. This is achieved by reusing the AutoIMD window from the previous minimization/equilibration; the restraints will not be taken into account.
- 7 Once all minimization/equilibration steps are performed, the corrected structure should be saved. This can be done in the following way. Select the molecule in the VMD main window. Create a representation containing all atoms in the Graphical Representations window. Select File \rightarrow Save Coordinates in the VMD main window. Select all in the Selected atoms and hit Save. Finally, specify the file name in the Chose filename to save trajectory and hit OK.

1.4 Preserving stereochemistry during MD simulation

As mentioned in the introduction, structure optimizations and MD protocols applying external forces can introduce chirality errors into the simulated structure. To preserve stereochemistry, the **chirality** plugin offers the possibility to generate restraints in form of **extrabonds** which can be used in a NAMD simulations. This feature currently is available only from the console.

1 Assuming the structure of interest is loaded in the top molecule in VMD and one wishes to obtain restraints for all chiral centers in proteins and nucleic acids, the command reads

chirality restrain -o chirality-extrabonds.txt

The file chirality-extrabonds.txt contains now two improper restraints for each chiral center. One includes the chiral center and its non-hydrogen binding partners. In the second restraint, the chiral center is replaced by the hydrogen bound to it. The equilibrium values of the two impropers are set to the values present in the used structure.

WARNING: Note, the specified restraints modify the potential energy function of the system, and thus should not be used in equilibrium production runs.

1 CHIRALITY IN PROTEINS AND NUCLEIC ACIDS

Rather, these restraints are supposed to be used only in simulations in which large forces occur due to poor initial geometry or due to externally applied forces, e.g., MDFF or TMD. Note also that although the final structure resulting from such simulations will be correct, the behavior of the system during the simulation is artificial.

2 *cis* peptide bonds in proteins

2.1 Introduction

In naturally occurring proteins most peptide bonds are in the *trans* configuration (see Fig. 6). However, sometimes *cis* peptide bonds do occur. The vast majority of *cis* peptides is observed at a proline, Xaa-Pro, Xaa being any amino acid. But non-proline Xaa-non Pro *cis* bonds are also found in proteins, although they occur much less frequently than Xaa-Pro (see A. Jabs *et al.*, JMB, **286**, 291–304 (1999)).

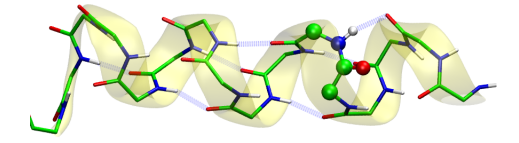


Figure 6: Structurally optimized α -helix containing a *cis*peptide bond. The *cis* peptide bond is shown in CPK, while hydrogen bonds within the helix are shown as thick dashed blue lines. The *cis* configuration of the peptide bond disrupts the hydrogen bond network stabilizing the helix. Note, the hydrogen bond network is broken not only locally.

The configuration of the peptide bond is central to the sort of secondary structure the protein backbone can adopt. It is easy to understand by imagining a normal α helix and converting a *trans* peptide bond into its *cis* form. The result is that the hydrogen bond network stabilising the helix is broken (see Fig. 6) and the helix will be unstable in long simulations.

2.2 Checking configuration of peptide bonds

To demonstrate the usage of the plugin we will start by checking a structure consisting of the protein EF-Tu in complex with a tRNA-Phe, although it is clear that in this example only the protein part will contain peptide bonds. The used structure is based on the the PDB structure 10B2 in which errors have been introduced manually.

Currently, the **cispeptide** plugin can be used from the text console and through a GUI. The available commands provided by the pugin can be obtained by typing Type **cispeptide**. The following information should be printed in the console:

```
Usage: cispeptide <command> [args...]
Commands:
  check
           -- identify cis peptide bonds
  list
           -- list identified cis peptide bonds
 minimize -- fix cis peptide bonds using energy minimization
           -- move specified atom to convert to trans
 move
           -- reinitialize plugin state
 reset
  restrain -- generate NAMD extrabonds file to restrain peptide bonds
           -- define how to modify a given cis peptide bond
  set
           -- visualize identified cis peptide bonds
  show
```

In the same way the usage information for any of the provided commands can be obtained. For example, typing **cispeptide restrain** will show the syntax of the command that generates restraints for the dihedral angle at the peptide bond which can be used in NAMD.

In the following, we will describe how to use the **cispeptide** plugin through its GUI, but all of the tasks can also be accomplished from the console.

- 1 Load the files pept_testcase.psf and pept_testcase.pdb into a new session of VMD.
- 2 Open the cispeptide GUI by selecting Extensions \rightarrow Modeling \rightarrow Fix Cis Peptide Bonds in the VMD Main menu. In the upper part of the GUI (see Fig. 7) the user can specify the molecule and the atom selection to be tested.
- **3** The button Check structure initializes the test. Peptides identified to be in *cis* configuration are displayed in the Identified cis peptide bonds form (see Fig. 8). In the present case there are 4 *cis* peptide bonds identified in EF-Tu.
- 4 By selecting an entry in the Identified cis peptide bonds form and hitting the button Show selected cis peptide bond it is possible to visually inspect all the individual identified *cis* peptides. In the created representation (see Fig. 9) the *cis* peptide bond is highlighted in CPK representation.

2.3 Converting *cis* peptides into *trans* configuration

WARNING: The cispeptide plugin identifies all *cis* peptide bonds. As noted in the introduction, there are cases in which *cis* peptides do occur in proteins.

2 CIS PEPTIDE BONDS IN PROTEINS

	cispeptide	•	/// =×			
			<u>H</u> elp			
Check protein struct	ure for cis peptide	bonds				
Molecule:	0: pep	ot_testcase.psf				
Selection: pro	tein					
Split chains by: 🛛 🔅	chain	 segname 				
	Check struc	ture				
-Identified cis peptide	bonds					
1st residue 2nd	residue chain :	segname atom mov	/ed			
			7			
Sh	Show selected cis peptide bond					
Select and move ato	ms in cis peptide	bonds				
Tag atom to b	e moved for selec	ted cis peptide bond	ls:			
hydrogen	hydrogen oxygen none		e			
Move tagge	Move tagged atoms for selected cis peptide bonds					
Minimize/e	Minimize/equilibrate selected cis peptide bonds					
Reset cispeptide plugin I'm feeling lucky						

Figure 7: The cispeptide GUI window.

 ✓ ✓		ispeptid	e			-×
						<u>H</u> elp
Check protein st	Check protein structure for cis peptide bonds					
Molecule:		0: pe	pt_testc	ase.psf		
Selection:	protein					
Split chains by:	C chain		۲	segname		
	Ch	eck stru	cture			
-Identified cis per	ntide honde					
1st residue		l chain	segname	l atom I	moved	
ALA 29 A ASN 51 A ASP 161 F	LA 30 LA 52 HE 162 RO 200	P P P	P1 P1 P1 P1 P1	none none none none	no no no no	
	Show sele	cted cis j	peptide	bond		
_C Select and move	e atoms in ci	s peptide	bonds-			
Tag atom	to be moved	for sele	cted cis	peptide b	onds:	
hydroger	hydrogen oxygen none					
Move tagged atoms for selected cis peptide bonds						
Minimize/equilibrate selected cis peptide bonds						
Reset cisp	oeptide plugi	n		I'm feeling	lucky	

Figure 8: Identified *cis* peptide bonds in the cispeptide GUI window.

Thus, it should be checked if a given *cis* peptide is real or is a mistake. This can be done by literature research, by comparing high resolution crystallographic

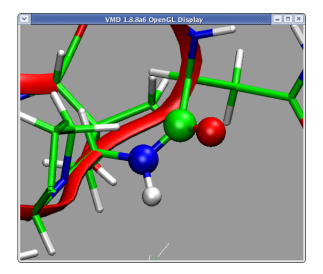


Figure 9: Representation of the cis peptide bond between Asn 51 and Ala 52 within EF-Tu

models, or by comparing crystallographic models of similar proteins.

1 If the shown peptide bond is supposed to have a *trans* configuration, the user can tag the atom to be moved to flip the configuration from *cis* to *trans*. This is done by hitting the button hydrogen or oxygen.

WARNING: The choice which atom to move is crucial and has to be made based on the local structure around the peptide under consideration. The decision is particularly critical helices and β -sheets: the relaxed backbone should integrate in the existing hydrogen bond network characteristic for the specific secondary structure motif (see Fig. 6).

If the identified *cis* peptide bond is supposed to be there, nothing needs to be done. It is also possible to untag a tagged atom by selecting the corresponding *cis* peptide and hitting none.

- **2** Once a *cis* peptide bond has been inspected, and the atom to be moved has been tagged, the actual moving of the atom can be executed by hitting the button Move tagged atoms for selected cis peptide bonds.
- **3** Since the described procedure (simple moving of one atom) generates an unphysical geometry of the molecule, it is necessary to optimize the structure using an MD force field. The cispeptide plugin uses the AutoIMD plugin for this purpose. This final step is accomplished by selecting the *cis* peptides which should be relaxed and hitting the button Minimize/equilibrate selected cis peptide bonds. This will open the AutoIMD Controls window (see Fig. 10).

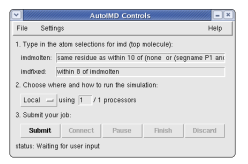


Figure 10: AutoIMD window called by cispeptide plugin.

- 4 Select Settings → Minimization Mode in the AutoIMD Controls window and hit then the Submit button. Once the AutoIMD session started, hit the Connect button and a minimization will start. Finish the simulation when the structure relaxed. Usually, it takes up to a few thousand steps until atoms do not move anymore. Is the structure minimized, hit the Finish in the AutoIMD Controls window. For more details about using the AutoIMD plugin, the user is referred to the AutoIMD user's guide http://www.ks.uiuc.edu/Research/vmd/plugins/autoimd/ug/.
- **5** WARNING: Depending on the quality of the initial structure, it can happen that not all errors can be fixed in one run. Thus, it is important to check the final result again and correct the remaining errors, if necessary. In order to check the structure after the performed minimization/equilibration using IMD, the cispeptide plugin should be reset. This can be done by hitting the Reset cispeptide plugin button at the bottom of the cispeptide window. Note, in the current implementation it will be necessary to select the molecule again in the top part of the cispeptide window. Note also that if the molecule to correct is solvated, it may be necessary to equilibrate the minimized parts of the structure. This can be done by selecting Settings \rightarrow Equilibration Mode and submitting a new AutoIMD run.
- 6 Finally, it is a good advice to minimize the structure in the very last step. This should be done by selecting Settings → Minimization Mode in the AutoIMD Controls window, which is still open from the last AutoIMD session, and hit then the Submit button. As mentioned in the introduction, additional restraints during the correction of *cis* peptides. The restraints will be used always when AutoIMD is started from the cispeptide window. The equilibrium values for these restraints are very approximate and bring the configuration in the correct region. Thus, in order to obtain a optimized structure, it is necessary NOT to use the restraints defined by cispeptide. This is achieved by reusing the AutoIMD window from

the previous minimization/equilibration; the restraints will not be taken into account.

7 Once all minimization/equilibration steps are performed, the corrected structure should be saved. This can be done in the following way. Select the molecule in the VMD main window. Create a representation containing all atoms in the Graphical Representations window. Select File \rightarrow Save Coordinates in the VMD main window. Select all in the Selected atoms and hit Save. Finally, specify the file name in the Chose filename to save trajectory and hit OK.

2.4 Preserving configuration of peptide bonds during MD simulation

As mentioned in the introduction, structure optimizations and MD protocols applying external forces can flip the configuration of a peptide bond in the the simulated structure. To preserve peptide configuration, the **cispeptide** plugin offers the possibility to generate restraints in form of **extrabonds** which can be used in a NAMD simulations. This feature currently is available only from the console.

1 Assuming the structure of interest is loaded in the top molecule in VMD and one wishes to obtain restraints for all peptide bonds in the system, the command reads

cispeptide restrain -o peptide-extrabonds.txt

The file **peptide-extrabonds.txt** contains now one dihedral restraint for each peptide bond. The equilibrium value of this dihedral is set to the value present in the used structure.

WARNING: Note, the specified restraints modify the potential energy function of the system, and thus should not be used in equilibrium production runs. Rather, these restraints are supposed to be used only in simulations in which large forces occur due to poor initial geometry or due to externally applied forces, e.g., MDFF or TMD. Note also that although the final structure resulting from such simulations will be correct, the behavior of the system during the simulation is artificial.

Acknowledgements

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