Peptide Self-Assembly Design Code User Manual

Running the code:

1. Prepare input structure

To run the Peptide Self Assembly (PepSAD) code the first step is to build the initial coordinate file (PDB fortmat) of the desired structure. The structure can be constructed in a molecule design package like Discovery Studio 3.5. Once the desired structure is built, it should be relaxed by running a short molecular dynamics simulation using a moleucular dynamics package to eliminate unfavourable atomic overlaps.

Save the PDB file as comp.pdb. Add N and C to the terminal amino acids of each peptide (for example, NHIE). Delete TER lines.

2. Edit the input file

The input file looks like the following. The input file can be edited as per the requirements of the project

filename Name of the initia (e.g. : comp.pdb)	ıl input structure f	recalcu_switch Equal to zero if starting from the original pdb file (eg: comp.pdb); equal to 1 if restarting from another file			
nstep_start The starting step	number	nstep_terminal The final step number (typically 10000)			
idum The seed for the r	ekt A value determined empirically determined (typically \leq 1) for each project. It is referred to as $kT_{sequence}$.				
fpho No. of hydrophobic residues	fpol No. of hydrophilic residues	fchg No. of charge residu	ed	foth No. of charged residues	
num4category(i) Sheet number for a β sheet type structure					
peptideID(i) Peptide number on the sheet (i)	peptideID(i+1) Peptide numbe on the sheet (i	er	peptideID Peptide nu on the she	ımber	peptideID(i+3) Peptide number on the sheet (i)

3. Submitting the job

To be able to submit a PepSAD join the following files should be present in the directory:

- a. main.f90 (Main code file)
- b. comp.pdb (starting input structure file)

- c. lib folder (directory containing the amber forcefield data)
- d. pdbfiles (an empty directory where the PDB files from the design will be stored)

Once all the above files are present in the directory the main.f90 code can be compiled using a fortran compiler and the job can be submitted. An example of a command that can be used to compile the code using the ifort compiler:

ifort -O3 -r8 -o main main.f90

Details of PepSAD output files

	energydetails.txt										
	step	attempt	score	binding energy	entropy	score4hydration	$P_{aggregation}$	Type of move: Not SCMF: Residue mutation Intra SCMF: Intra chain residue exchange			
	energyprofil	e.txt									
	step	score	binding ene	ergy entropy		score4hydration	$P_{aggregation}$				
minimumenergy.txt											
	step	attempt	score								