“Crawling” on the self-assembly system: A molecular simulation of peptide position adjusting over self-assembly block

Tong Zhang¹, Baoying Shen¹, and Xinghua Shi²,*
¹Beihang University, Beijing, China
²The National Centre for Nanoscience and Technology, Beijing, China

Abstract. By combining non-equilibrium molecular dynamics(NEMD), umbrella sampling, and weighted histogram analysis method(WHAM), we calculated the potential of mean force of histidine peptide moving over a self-assembly structure. The reaction coordinate is along the main chain direction of the histidine peptide in the self-assembly structure. It is found that the energy needed for the histidine peptide with 3 and 5 residues while moving along the reaction coordinate is around -2.2 kCal/mol and -7.4 kCal/mol, respectively. And the histidine peptide crawls along the reaction coordinate, performing a snake-like movement. This result could illustrate how histidine peptide adjusts its position during self-assembly process.

1 Introduction

Self-assembly phenomenon is common in nature and many materials with excellent optical, electrical, catalytic properties such as nonlinear optics, chemical biosensors, information storage materials, and tissue growth scaffolding materials, are produced by self-assembly. Molecular self-assembly is the process by which molecules adopt a defined arrangement without guidance or management from an outside source. Noncovalent interactions (e.g., hydrogen bonding, metal coordination, hydrophobic forces, van der Waals forces, π-π stacking, and/or electrostatic) are crucial to the self-assembly process. [1]

Histidine is an α-amino acid that is used in the biosynthesis of proteins. It contains an α-amino group, a carboxylic acid group, and an imidazole side chain (which is partially protonated), classifying it as a positively charged amino acid at physiological pH. Histidine was first isolated by German physician Albrecht Kossel and Sven Hedin in 1896. The imidazole ring of histidine is aromatic at all pH values [2] and it can form π stacking interactions[3].

The self-assembly of histidine peptide is by hydrogen bonding between the backbone of the peptide, forming an antiparallel β-sheet second structure. What we are interested in is the self-assembly process. For the initial stage of the self-assembly process is always at random, and the final result is “neat” structure. To find out how histidine peptide adjust itself into neat structure is the target of this article. One guess is that histidine peptide is like

* Corresponding author: shixh@nanoctr.cn

© The Authors, published by EDP Sciences. This is an open access article distributed under the terms of the Creative Commons Attribution License 4.0 (http://creativecommons.org/licenses/by/4.0/).
Lego peptide[4] which can “slide” on the surface of the assembly only that instead of hydrophobic effect of the residues, there are other mechanisms for histidine peptides. Thus, we perform the following simulation, trying to find out the energy needed for the histidine peptide to adjust its position over the assembly block.

2 Models of Molecular Dynamics Simulation

2.1 Constructing histidine peptide and histidine self-assembly block

We use full atom model to do our simulation so that a clearer picture can be obtained. First, we obtain the peptides with 15 histidine residues to build an antiparallel β-sheet second structure. Then the structure is equilibrated in water at 300K and 1.0 bar of pressure for 10 ns. Two layers in the middle of the structure are taken out, one as our “reaction” surface, the other being the source of histidine peptide which will be pulled over the reaction surface, showing in Fig 1-A,B.

Then 2 types of histidine peptides are “cut” (unnecessary residues removed and only the peptide in the middle of the three chains is kept) from the layer described above, producing 2 different lengths (with 3,5 residues) of histidine peptide. The initial model showing in Fig 1-C.

![Fig. 1. A: The structure of self-assembly block: periodic only in Z direction; 3 chains of peptide interacted by π- stacking in Y direction; The blue box is the simulation box. B: The two layers chosen from the antiparallel β-sheet, one as the reaction layer, the other the source of different lengths of peptides. C: The initial model.](image)

2.2 Umbrella pulling

To calculate the PMF of the peptide along the reaction coordinate, a non-equilibrium pulling method is used. [5].

When the distance between two groups is changed continuously, work is applied to the system, which means that the system is no longer in equilibrium. However, one can use the Jarzynski relation [6] to obtain the equilibrium free-energy difference ΔG between two distances from many non-equilibrium simulations:
Lego peptide which can “slide” on the surface of the assembly only that instead of 
hydrophobic effect of the residues, there are other mechanisms for histidine peptides. Thus, 
we perform the following simulation, trying to find out the energy needed for the histidine 
peptide to adjust its position over the assembly block.

2 Models of Molecular Dynamics Simulation

2.1 Constructing histidine peptide and histidine self-assembly block

We use full atom model to do our simulation so that a clear picture can be obtained. First, 
we obtain the peptides with 15 histidine residues to build an antiparallel β-sheet second 
structure. Then the structure is equilibrated in water at 300K and 1.0 bar of pressure for 10 
ns. Two layers in the middle of the structure are taken out, one as our “reaction” surface, 
the other being the source of histidine peptide which will be pulled over the reaction surface, 
showing in Fig 1-A, B.

Then 2 types of histidine peptide are “cut” (unnecessary residues removed and only the 
peptide in the middle of the three chains is kept) from the layer described above, producing 
2 different lengths (with 3, 5 residues) of histidine peptide. The initial model showing in Fig 
1-C.

Fig. 1. A: The structure of self-assembly block: periodic only in Z direction; 3 chains of peptide 
interacted by π-stacking in Y direction; The blue box is the simulation box. B: The two layers chosen 
from the antiparallel β-sheet, one as the reaction layer, the other the source of different lengths of 
peptides. C: The initial model.

2.2 Umbrella pulling

To calculate the PMF of the peptide along the reaction coordinate, a non-equilibrium 
pulling method is used. When the distance between two groups is changed continuously, work is applied to the 
system, which means that the system is no longer in equilibrium. However, one can use the 
Jarzynski relation to obtain the equilibrium free-energy 

\[ \Delta G_{AB} = -k_B T \log\left(e^{-\beta W_{AB}}\right)_A \]  

where \( W_{AB} \) is the work performed to force the system along one path from state A to B, \( k_B \) is the Boltzmann constant, \( T \) is temperature and the angular bracket denotes averaging over 
a canonical ensemble of the initial state A and \( \beta = 1/k_B T \)

The pull code applies forces or constraints between the centres of mass of one or more 
pairs of groups of atoms. In this article, an umbrella pulling is used, in which case a 
harmonic potential is applied between the centres of mass of two groups. Thus, the force is 
proportional to the displacement. And the peptide, in our case, is “dragged” by the force 
along the reaction coordinate.

Fig. 2 illustrates the umbrella pulling simulation. The alpha-carbon of the first residue is 
attached to a virtual spring with the force constant being 1500. The pulling is in the 
direction of the main chain and the other two directions’ movements are restrained. The pulling simulation duration is 500 PS of MD, saving snapshots every 1 PS and the distance 
pulled is 5.0 nm. After the pulling simulation, a series of configurations are generated and 
selected so that configurations are of 0.2 nm spacing.

Fig. 2. The pulling simulation. The red arrow represents the force applied by the virtual spring.

2.3 Umbrella sampling

The binding energy (\( \Delta G_{\text{bind}} \)) is derived from the potential of mean force (PMF), extracted 
from a series of umbrella sampling simulations. A series of initial configurations is 
generated along the reaction coordinate \( \xi \), as described above, each corresponding to a 
location wherein the ligand is harmonically restrained at increasing centre-of-mass (COM) 
distance from a reference molecule using an umbrella biasing potential. This restraint 
allows the ligand to sample the configurational space in a defined region along a reaction 
coordinate between itself and its reference molecule.

In this article, we sample COM distances from 2.0 - 5.0 nm (and 1.1 - 3.4 nm for the 3-
residue model) along the x-axis using roughly 0.2-nm spacing. Start by running a brief NPT 
equilibration in each window. Then each window is sampled 10 ns.

2.4 PMF extraction using WHAM

The Weighted Histogram Analysis Method (WHAM) is one of the earliest methods that 
take into account information from all Intermediate States. The precursor to WHAM and 
first version of multiple histogram relighting techniques came from Ferrenberg and 
Swendsen;[7] WHAM was developed later for alchemical simulations[8]. WHAM is 
included in GROMACS as the wham utility, which this article uses to extract the potential 
of mean force.
3 Results of Molecular Dynamics Simulation

The potential of mean force profile is extracted by WHAM. We can thus calculate the energy needed for the histidine peptide to “crawl” on the reaction surface, which is -7.4kCal/mol for peptide with 5 residues of His and -2.2kCal/mol for peptide with 3 residues. From the PMF curve, we can also see that the process is periodic, which is reasonable as the peptide moving along the reaction coordinate, antiparallel β-sheet structure is periodically formed. The bottom of the curve means that such antiparallel β-sheet is formed and the peak means that nearly all of the hydrogen bonds between the main chain of histidine peptide and the assembly block are broken.

From the simulation result we can see that the peptide moves like snake to adjust its positions along the reaction coordinate. The simulation result shows in Fig.3-A,B.

![Simulation Results](image)

**Fig. 3.** A: The peptide with 5 residues of His. “crawling” along the reaction coordinate, with labels from “a” to “e” indicating different state on the PMF curve. B: Peptide with 3 residues, from one stable state to another, and PMF curve. C: Backbone molecules’ hydrogen bonds amount change in one period, with black representing peptide with 5 His, red 3 His. The Position 1 to 5 are positions shown in A, from position “a” to position “e”, respectively.

4 Conclusions

From the PMF curve obtained above, we can find that the energy needed for the histidine peptide to adjust its position along the reaction coordinate is not much (-7.4kCal/mol and -2.2kCal/mol for peptide with 5 His residues and 3 His residues, respectively). This means the peptide could easily move on the assembly surface and find its stable position by “crawling” and form a neat β-sheet structure. And the difference between the peptides of two different lengths (3 residues and 5 residues), we guess, probably suggests that the
energy required to make position adjustment is in associate with the hydrogen bonds amount which is formed between the backbone atoms of histidine peptides. The hydrogen bonds (on backbone atoms) average amount’s change in one period of the “crawling” process proves that what we guess is true.

Firstly, the hydrogen bonds of the residue which the virtual spring is attached are broken, causing the hydrogen bonds on the residue next to it broken too. The “free” residue moves to either side of the reaction layer due to the effect of the side chain interaction. From the Fig. 3-C we can see the amount change of hydrogen bonds in one moving period, and position 1 to 5 are.

To sum up, the histidine peptide could adjust its position via interacting with other histidine peptide chains in the self-assembly surface, doing a snake-like “crawling”. This could indicate that in the self-assembly process, the histidine peptides would form the assembly block in a similar pattern.

References

1. L. a Estroff and A. D. Hamilton, “Water Gelation by Small Organic Molecules,” Chem. Rev., vol. 104, no. 3, pp. 1201–1218, Ma doi:10.1103r. (2004).
2. A. Mrozek, J. Karolak-Wojciechowska, and K. Kieć-Kononowicz, “Five-membered heterocycles. Part III. Aromaticity of 1,3-imidazole in 5 + n hetero-bicyclic molecules,” J. Mol. Struct., vol. 655, no. 3, pp. 397–403, (2003).
3. L. Wang, N. Sun, S. Terzyan, X. Zhang, and D. R. Benson, “A histidine/tryptophan π-stacking interaction stabilizes the heme-independent folding core of microsomal apocytochrome b5 relative to that of mitochondrial apocytochrome b5,” Biochemistry, vol. 45, no. 46, pp. 13750–13759, (2006).
4. H. Yokoi, T. Kinoshita, and S. Zhang, “Dynamic reassembly of peptide RADA16 nanofiber scaffold,” Proc. Natl. Acad. Sci., vol. 102, no. 24, pp. 8414–8419, (2005).
5. D. van der Spoel et al., “Gromacs user manual,” in SpringerReference, Berlin/Heidelberg: Springer-Verlag, (2010), p. 7778.
6. C. Jarzynski, “Nonequilibrium equality for free energy differences,” Phys. Rev. Lett., vol. 78, no. 14, pp. 2690–2693, (1997).
7. A. M. Ferrenberg and R. H. Swendsen, “Optimized Monte Carlo data analysis,” Phys. Rev. Lett., vol. 63, no. 12, pp. 1195–1198, (1989).
8. S. Kumar, J. M. Rosenberg, D. Bouzida, R. H. Swendsen, and P. A. Kollman, “THE weighted histogram analysis method for free-energy calculations on biomolecules. I. The method,” J. Comput. Chem., vol. 13, no. 8, pp. 1011–1021, (1992).