BPU11 congress

11th International Conference of the Balkan Physical Union 28 August – 1 September 2022, Belgrade, Serbia



Self-Association of Antimicrobial Peptides in Mono- and Multicomponent Solutions: a Computational Study

Elena Lilkova¹, Peicho Petkov², Rositsa Marinova², Leandar Litov², Nevena Ilieva¹

¹Institute of Information and Communication Technologies, Bulgarian Academy of Sciences, ²Faculty of Physics, Sofia University "St. Kliment Ohridski" *Correspondence: rosie.marinova@gmail.com, elena.lilkova@iict.bas.bg











Antimicrobial peptides



01.09.2022

Molecular Dynamics

• Newton's equations:

$$\vec{v}(t) = \frac{d}{dt}\vec{x}(t)$$
 $\vec{F} = \frac{d^2}{dt^2}\vec{x}(t) = -\frac{\partial V(\vec{x}(t))}{\partial \vec{x}}$

• Potential energy (forcefield):

$$V = \sum V_{s} + \sum V_{a} + \sum V_{t} + \sum V_{vdw} + \sum V_{e} + \dots$$

Bond strength
Sum over all
bonds
$$Van der Waalsinteractions$$
$$Coulombinteraction$$



Indolicidin aggregation on a bacterial membrane



Elena Lilkova

01.09.2022

BPU11

Belgrade



Self-assembly of bombinin in water



Elena Lilkova

01.09.2022

BPU11

Belgrade

Self-assembly of bombinin in water



Our model

- AMPs do not exist in monomeric form after secretion in the bodily fluids as part of multicomponent mixtures
- They aggregate and form nanosized clusters in solution, prior to their attack on the target membrane
- The clusters are how AMPs get delivered to the membrane in high enough local concentration

Case study – snail mucus peptides

Recently discovered glycine-rich peptides, isolated from the mucus of the garden snail *Helix Aspersa*

Peptide	Sequence	Charge	M _r [kDa]
p1	KVKDNQWRP	+2	1.17
p2	VNVVGGGGGGVGGGGGGGM	0	1.57

Ilieva, N. et al. (2020). In Silico Study on the Structure of Novel Natural Bioactive Peptides. In: LSSC 2019. LNCS, 11958, 332–339, Springer, https://doi.org/10.1007/978-3-030-41032-2_38

Elena Lilkova

Monocomponent p1 solution

30

27 p1 peptides in a cubic box, $C_m = 48 \text{ mM}$





Maximal cluster size

Largest p1 aggregate – fairly loose, monomers are well separated.

Monocomponent p2 solution

27 p2 peptides in a cubic box, $C_m = 29 \text{ mM}$



Largest p2 aggregate – tightly packed globule



Multicomponent p1+p2 solution

25 p1 and 10 p2 peptides in a cubic box



Mixed aggregate – 7 p1 and 9 p2 peptides



Multicomponent p1+p2 solution





Mixed 16-mer aggregate – coloured by peptide type (p1 – cyan, p2 – pink)

Residue type surface representation (basic aa – blue, acidic – red, polar – green, non-polar – white

US dissociation free energies

Quantitative cluster stability assessment:

- P2 highly hydrophobic \rightarrow no dissociation expected from the cluster.

- P1 peptides might dissociate due to the electrostatic repulsion from other p1 monomers potentially overcoming the attraction due to the hydrophobic effect.

=> Calculate the dissociation free energy of the most solvent exposed p1 monomers, using umbrella sampling to estimate the potential of mean force along the collective variables ξ_1 , ξ_2 , ξ_3 , (COM distance between the respective p1 monomer and the aggregate)





Elena Lilkova

01.09.2022

BPU11

Belgrade

Conclusions

- AMPs aggregate and form stable clusters.
- Their size and properties depend on the composition of the solution. Generally like globular proteins a non-polar hydrophobic core and exposed to the solvent charged and polar residues.
- Aggregation causes inter-peptide hydrogen bonds number increase, which causes formation of new and stabilization of already formed secondary structure elements.
- The so-formed structures provide the perfect transport system locking the hydrophobic uncharged residues in the core of the cluster prevents the interaction with the eukaryotic membranes with lower surface charge density, and positioning the charged residues on the cluster surface enables for electrostatic interaction with the bacterial surface.
- In addition, the peptide folding, promoted by the amphiphilic structure in the aggregates, allows for high enough local concentration of AMPs to be delivered to the target membrane in a functionally active conformation.

Acknowledgments

- This work was supported in part by the Bulgarian National Science Fund under Grant KP-06 OPR-03-10/2018.
- Computational resources were provided by BioSim HPC cluster at the Faculty of Physics, Sofia University "St. Kliment Ohridski" and by CI TASK (Centre of Informatics – Tricity Academic Supercomputer & networK), Gdansk (Poland).





THANK YOU!