

Modeling of Sortase-Linker Interactions for the iGEM Competition

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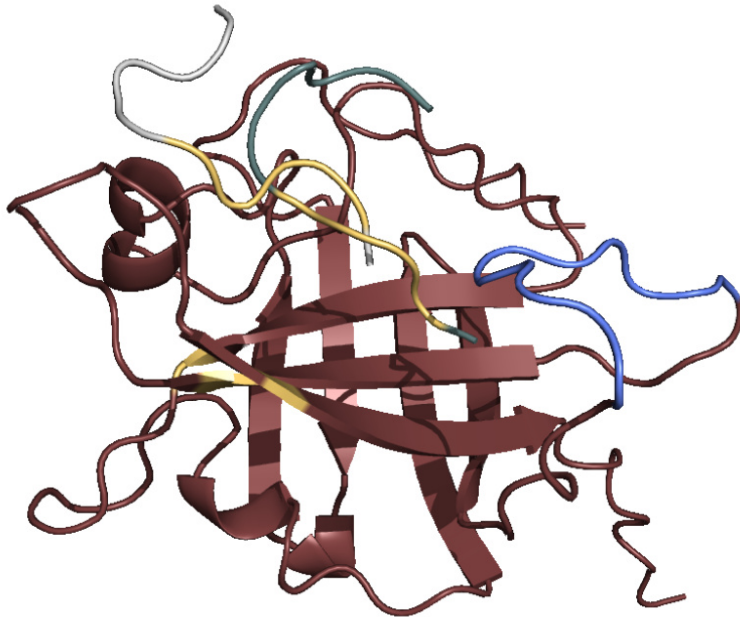
Project Term
2019 - 2019

Clusters
Lichtenberg Cluster Darmstadt

Software
GROMACS

Additional Software
RosettaCommons

University
Technische Universität Darmstadt



Introduction

Sortase A7M is an enzyme, which catalyzes the linking reaction of the N-terminal and the C-terminal region of two proteins. To improve the efficacy of this reaction we set out to model the structure of the enzyme, understand the mechanism and improve the binding to the active site of by mutating the linkers used.

Methods

Our first step was to model the 3D structure of the enzyme using *RosettaCM*. *RosettaCM* is a comparative modeling algorithm implemented in the *RosettaCommons* software package. The next step was to identify the active site. To this end, we employed molecular dynamics simulations using the *GROMACS* software package. In the last step we once again used *RosettaCommons* to perform docking simulations to find the best possible linker.

Results

After modeling the structure we used principal component

analysis to find the active site. The analysis revealed a Sortase substructure called $\beta 7/\beta 8$ loop to be crucial for the mechanism as literature confirms [1]. This substructure seems to move towards and away from the active site. We performed docking simulations and found out that the C-terminus with the specific LPETG motif favors binding to the active site 1, while the N-terminus with a poly-G motif (GGGGG) favors binding to the loop 2. Combining this finding with the insights concerning the loop movement, we postulate that the loop transports bound poly-G peptides towards the active site, where the LPETG peptides are located. This leads to the reaction and ultimately to the linkage. In terms of linker optimization we were not able to find sequences with better binding affinities as most linkers achieved a similar binding score.

Discussion

With the computations performed we were able to model a *Sortase A7M* structure and postulate a mechanism. While we were not able to find ideal linkers for the reaction the insights we gained helped us develop a better understanding of the mechanism. To fully understand the mechanism and find linkers that are better suited for our reactions, we would need to perform hybrid molecular dynamic/quantum mechanics simulations as those would not only consider binding interactions but also reactions.

Figures

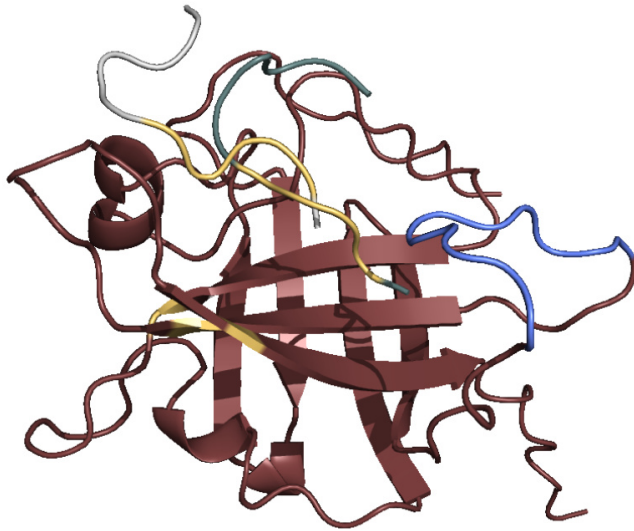


Figure 1:

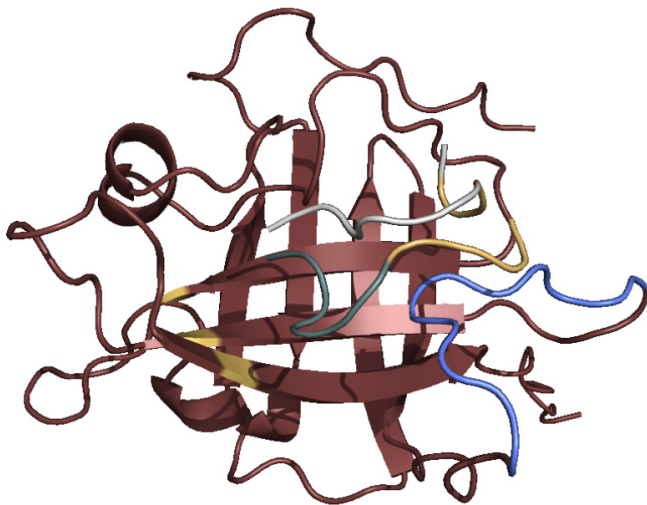


Figure 2:

Reference

- [1] Clancy, K.W.; Melvin, J.A.; McCafferty, D.G.: "Sortase transpeptidases: insights into mechanism, substrate specificity, and inhibition", *Peptide Science*, vol. 94, no. 4, pp. 385-396, 2010.
<https://doi.org/10.1002/bip.21472>

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