Modelling Somatic Hypermutations with Reinforcement Learning

Applied Biotechnology

for PD1 and Pembrolizumab

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Somatic Hypermutations (SHM) are an important part of the

We express the biological process of AM as a Markov decision

The <u>relative configuration of PD1-Pembro complex remains the</u>

Maturation

Figure 1. Broad overview of the affinity maturation (AM) process by which

antibodies (Abs) evolve against vaccine-candidate antigens (Ags) in a germina

Tfh cell B cell

T cell-Dependent

Selection

Affinity-Based

Selection

Apoptosis

End Simulation

episode > maximur

episodes

INTRODUCTION

Model Assumptions:

positions

affinity maturation (AM) in B-cells.

mutations in loops on Pembro

GC Seeding

Recycling

Differentiation

center (GC) reaction (Faris J et al., 2022)

Secreted Abs

GC Exit

Score to Reward

process (MDP), creating a RL model of SHM.

ABSTRACT

We present a reinforcement learning (RL) model for Somatic Hypermutations (SHM), which mimics the natural selection process in a very short time scale. In this model, the agent can learn to preferentially mutate amino acids in the antibody, leading to affinity maturation. The model thus predicts a higher binding affinity antibody than the initial antibody-antigen complex. We have used the Pembrolizumab-PD1 (5b8c) complex to create the model, as Pembrolizumab (Pembro) is widely used in immunotherapy.

We use Q-Learning in RL to model SHM on a reduced state space to provide better binding affinity antibodies. We validated the structure of the antibodies predicted by the RL model using AlphaFold2 and C-alpha distance plots to check for proper folding of chains and protein-protein interactions. This study provides a proof of concept that RL can be used for modeling the biological process of SHM and can further be employed for creating novel antibodies.

OBJECTIVES

itiate Agent class with actio

list and hyper-parameters

Major Objective: To create a reinforcement learning model for somatic hypermutations using Pembro and PD1.

Minor Objective: To find states/antibodies which have better binding affinity than Pembro, and validating the good states using AlphaFold2 and C-alpha distance plots of the predicted structures.

Environment

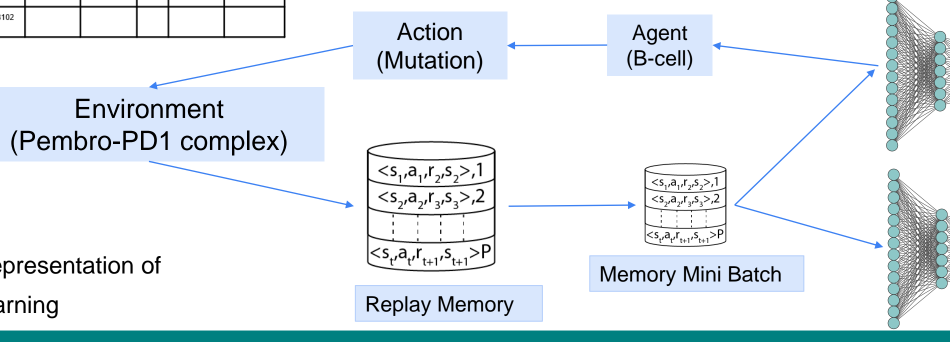


Soft-max Perform Action **Predict Actions** →Analysis by PRODIGY Initiate Q-table Initiate simulation Update Q-value Next Episode (PyMOL) **Q-Table** Epsilon greedy ACTIONS ,A53,A57,A95,A97,B30,B33,B35,B52,B54,B55,B58,B59,B99,B101,B10; S, Y, Y, Y, S, D, T, Y, Y, N, S, N, T, N, R, Y, A

Assign Score

Figure 4. A Flowchart of Basic Q-learning, B Representation of Q-Table, C a Scoring vector b Score to reward function

Network



Update Policy network Loss Function Network Sync

Figure 2. The interaction of agent and environment in a MDP Agent: B-cells Action: Point mutations on Pembrolizumab (using PyMOL) Environment: Affinity Maturation of Pembrolizumab-PD1 State: Pembro with mutated amino acids

same irrespective of the type of amino acids mutations at the 17 PD-L1 Pembrolizumab CDR-L1 The 3D folds of light and heavy chains are not disrupted by the CDR-L2 Lys131 ^HArg102 **=** ^HThr30 ▶ HTyr33 Lys124 Chain A: Pembro light chain variable region Arg125 Chain B: Pembro heavy chain variable region Programmed cell death protein 1 (PD1-Antigen) CDR-H2 Phe19 _{N-ter} **A** Reference crystal Pembrolizumab (PDB ID: 5B8C), B A schematic diagram of interactions between PD1 and antiPD1. Direct protein/protein hydrogen bonds are in blue, water-mediated hydrogen bonds are in green and salt

> сь Table. Score to Reward function C a Binding Affinity (kcal/mole) < -12.6 >= -12.6 < -12.6 3>score>-1 3 > score > -1> -12.6 -1>= score > -10< -10 Percentage Charged NIS Residues

> > Prodigy Parameters

bridges are in red (Horita S et al., 2016)

RESULTS

Figure 5. General Representation of

Deep Q-Learning

34.A53.A57.A95.A97.B30.B33.B35.B52.B54.B55.B58.B59.B99.B101.B10

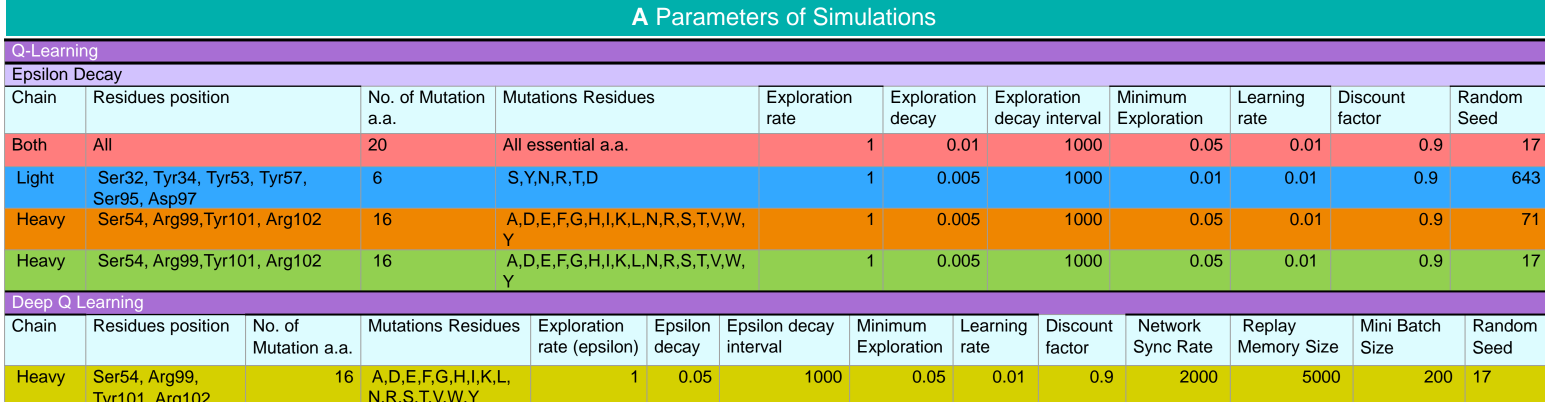


Figure 6 A Parameter Table, B Simulations and AlphaFold2 (AF) Predicted structures with Binding Affinity(BA) and RMSD. The parameters for the simulation. The simulations and the higher affinity states from them are shown in the Simulation section corresponding to the colour in the Parameter table as Red, Blue, Orange and Green. The DQL simulation (Yellow) is mentioned in the Discussion and Future work section. Structure Validation using AlphaFold2 and C-alpha distance plots

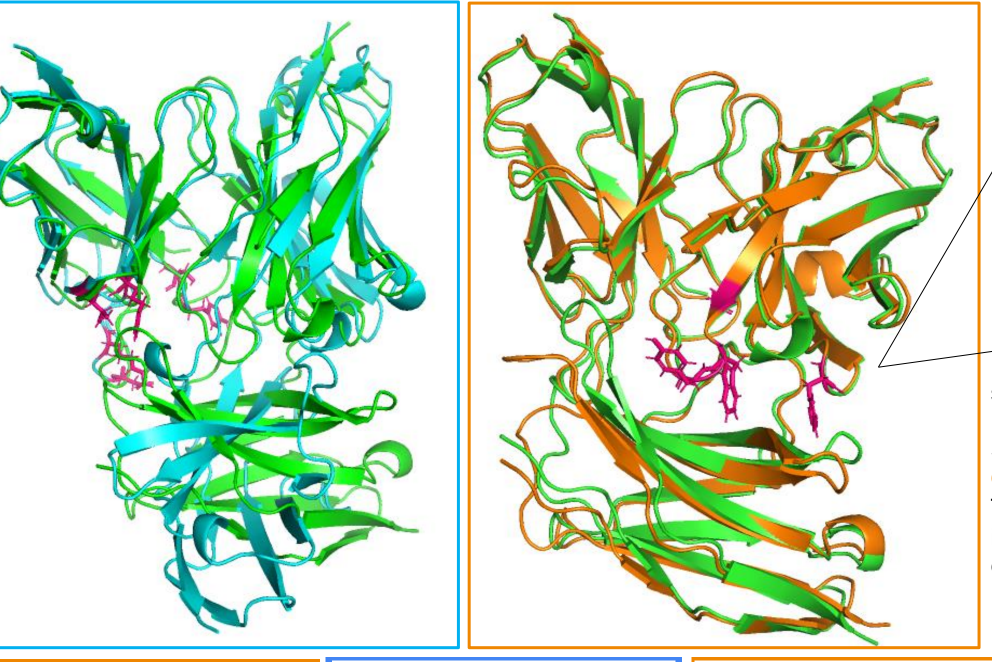


Figure 7 A State DRRTSYTYYNSNTNRYR from simulation Blue folded improperly aligned with Pembro(shown as green), **B**. State SYYYSDTYYNFNTNAWY from simulation folded properly aligned with Pembro, C The C-alpha distance plots of state A and B a intra chain B **b** inter chain A vs chain B **c** inter chain B vs chain C

Histogram of last 10000 episodes Binding affinity Histogram of last 10000 episodes

Binding affinity

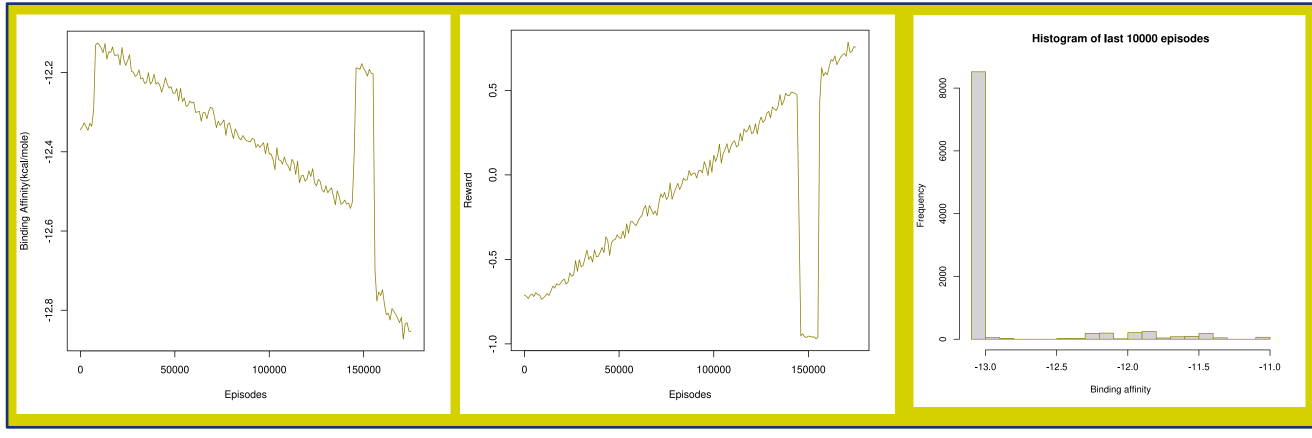
State	Freq	Reward	Crystal BA	AF BA	RMSD
SYYYSDTYYNWNTNTHL	60	1	-12.9	-13.5	2.195
SYYYSDTYYNYNTNIFN	53	1	-13.0	-13.5	2.172
SYYYSDTYYNYNTNWAK	143	1	-13.0	-13.4	2.101
SYYYSDTYYNFNTNAWY	45	10	-14.0	-13.3	2.217
SYYYSDTYYNFNTNIYY	39	10	-14.1	-13.0	2.185

B Simulations and Predicted Antibodies Histogram of last 10000 episodes Binding affinity Reward Crystal BA AF BA RMSD State Freq DRRTSYTYYNSNTNRYR -13.4 10 -23.3STRNYYTYYNSNTNRYR -13.6 2.061 -13.5

-13.4 -13.4 2.058 YRNNSRTYYNSNTNRYR 141 DSRNYYTYYNSNTNRYR -13.7 -13.3 2.072 41 DTRNYYTYYNSNTNRYR -13.7 -13.3 -13.3 2.072 101 -13.3 YRNNNRTYYNSNTNRYR Histogram of last 10000 episodes 144414144414441444441444441444

0e+00 1e+05 2e+05 3e+05 4e+05 5e	e+05 6e+05		-14.0 -13.5	-13.0 -12.	5 -12.0			
Episodes		Binding affinity						
*Looping from starting state upon reaching -10 reward, simulating apoptosis								
State	Freq	Reward	Crystal BA	AF BA	RMSD			
SYYYSDTYYNFNTNRYR	5	1	-13.4	-13.1	2.179			
SYYYSDTYYNLNTNRYR	6	1	-13.5	-13	2.153			
SYYYSDTYYNWNTNRYR	6	1	-13.4	-13	2.286			
SYYYSDTYYNYNTNRYR	9	1	-13 6	-12 8	2 193			

DISCUSSION AND FUTURE WORK



Q-Learning simulated SHM and provided a better binding Pembro-PD1 complex using a reduced state space.

Deep Q-Learning can be employed to predict alternate complexes to Pembro-PD1 -Currently underway.

Protein Language Models (PLM) can be employed in conjunction with our SHM-RL model for drug discovery.

Further validation of predicted antibodies using molecular dynamics and different binding affinity tools is required.

REFERENCES

- Faris, J. G., Orbidan, D., Wells, C., Petersen, B. K., & Sprenger, K. G. (2022). Moving the needle: Employing deep reinforcement learning to push the boundaries of coarse-grained vaccine models. Frontiers *Immunology*, https://doi.org/10.3389/FIMMU.2022.1029167
- Horita, Shoichiro, et al. "High-resolution crystal structure of the therapeutic antibody pembrolizumab bound to the human PD-1." Scientific reports 6.1 (2016): 35297
- Sutton, R. S., & Barto, A. G. (2018). Reinforcement learning: An introduction, 2nd ed. In Reinforcement learning: An introduction, 2nd ed. The MIT Press.
- Vangone, A., & Bonvin, A. M. J. J. (2017). PRODIGY: A Contact-based Predictor of Binding Affinity in Protein-protein Complexes. Bio-Protocol, https://doi.org/10.21769/BIOPROTOC.2124

ACKNOWLEDGEMENT

This study is supported by Dept. of electronics, IT, BT and S&T, Government of Karnataka. We would also like to extend our gratitude to the faculty of IBAB and Mr. Balakrishna Prabhu B N, Ms. Namita Menon, Mr. Yash Chindarkar and Ms. Apoorva Ganesh for their contribution and guidance.