

Novel Potential Inhibitors Against SARS-CoV-2 Using Artificial Intelligence

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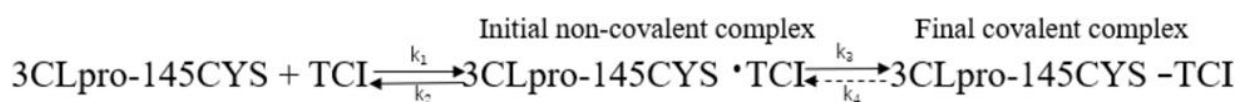
Abstract

Since known approved drugs like liponavir and ritonavir failed to cure SARS-CoV-2 infected patients, it is high time to generate new chemical entities against this virus. 3CL main protease acts as key enzyme for the growth of a virus which acts as biocatalyst and helps to break protein and ultimately in the growth of coronavirus. Based on a recently solved structure (PDB ID: 6LU7), we developed a novel advanced deep Q-learning network with the fragment-based drug design (ADQN-FBDD) along with variational autoencoder with KL annealing and circular annealing for generating potential lead compounds targeting SARS-CoV-2 3CL^{pro}. Structure-based optimization policy (SBOP) is used in reinforcement learning. The reason for using variational autoencoders is that it does not deviate much from actual inhibitors, but since VAE suffers from KL diminishing we have used KL annealing and circular annealing to address this issue. Researchers can use this compound as potential drugs against SARS-CoV-2.

1. Introduction

SARS-CoV2 coronavirus has caused pandemic worldwide which causes disease Covid-19. The number of cases and deaths is increasing day by day. This virus is similar to SARS-Covid for which there is no approved treatment. The virus becomes deadly when it causes severe acute respiratory syndrome. So, there is an urgent need to create inhibitors for this. This approach works by adding fragments instead of atoms which is not only computationally efficient but also chemically more reasonable. The rate of a reaction is defined by $r = ke^{-E_a/RT}$ where k is rate constant, E_a is the activation energy, R molar gas constant and T is the absolute temperature.

When the kinetic energy of the reactant molecules is greater than the activation energy, the reaction occurs. The reaction between inhibitor and protease is given by:



k_1 , k_2 , k_3 and k_4 are the rate constants. For inhibition to occur k_1 must be greater than k_2 for non-covalent bonding. After this step covalent bonding takes place such that $k_3 \gg k_4$ and the reverse reaction does not happen. We generated molecules using Variational Autoencoder using transfer learning on inhibitors, then applied reinforcement learning by adding fragments and calculating reward, so that molecules with the desired property are obtained which includes drug likeness, specific fragment containment and pharmacophores scores.

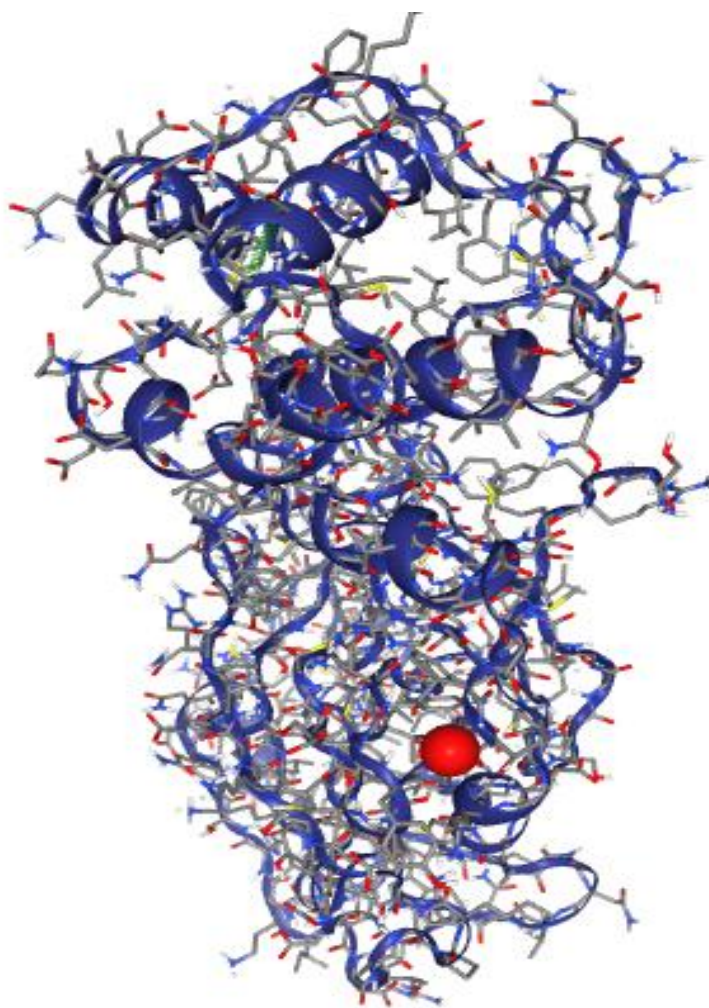


Figure1. SARS-CoV-2 main protease with unliganded active site.

Keywords: COVID-19, SARS-COV2, 3CL Protease, Structure-based optimization policy, Deep learning, Artificial intelligence, Reinforcement learning.

2. Related Work

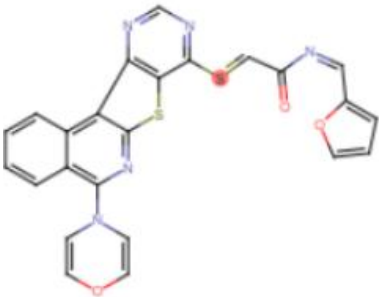
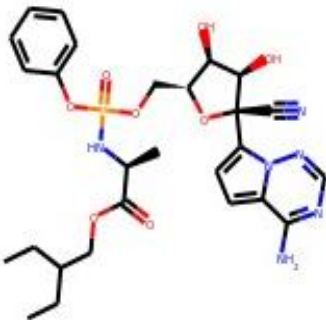
Bowen Tang, Fengming He, Dongpeng Liu, Meijuan Fang, Zhen Wu have used Reinforcement learning on SARS-CoV-2 3CLPro main protease inhibitors and defined reward function which includes other properties like fragment containment and pharmacophores score and applied efficient and novel advanced deep Q-learning network with the fragment-based drug design.

Navneet Bung, Sowmya Ramaswamy Krishnan, Gopalakrishnan Bulusu and Arijit Roy from Tata Consultancy Services Limited have trained RNN on 1.6 million data points from ChEBL dataset to learn the grammar of smile representation of molecules ,then used transfer learning on 2515 inhibitors from the same dataset and applied reinforcement learning to force generator generate molecules with desired Synthetic accessibility score, quantitative estimate of drug-likeness, partition coefficient and molecular weight.

IBM has generated 3000 novel molecules using generative models against three targets. In addition, the Hong Kong-based pharmaceutical research company, InSilico Medicine has rolled out a list of 97 potential candidate molecules designed to inhibit the 3CL protease of SARS-CoV-2.

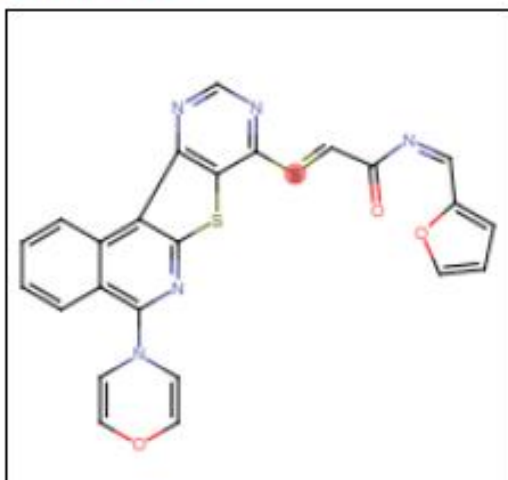
3. Results

Table 1: *Generated Molecules with Tanimoto Similarity with the already existing FDA Approved inhibitors.*

ID	MOLECULE GENERATED	FDA APPROVED INHIBITORS	Tanimoto Similarity
PM_1		 (REMDESIVIR)	0.655

PM_1

Ligand in 2D



Ligand in 3D

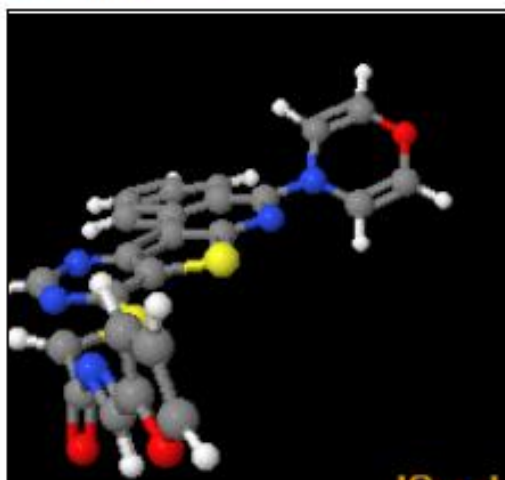


Figure2: 2d and 3d structure of PM_2



Figure3: interaction between ligand and protease

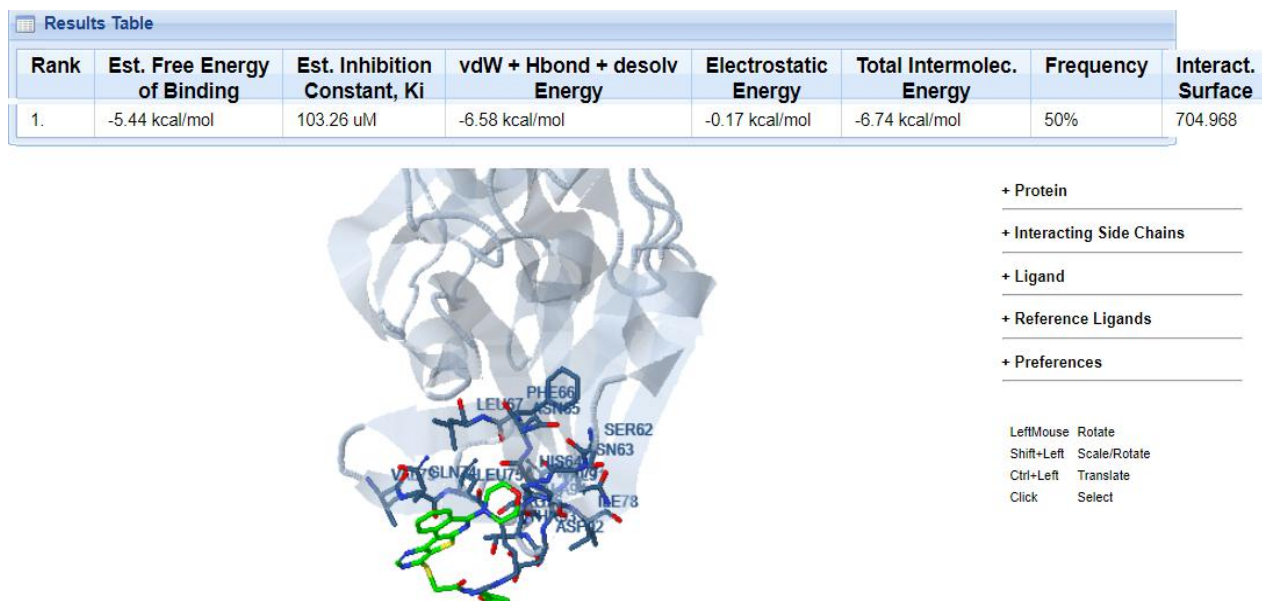


Figure4: Docking results obtained between PM_1 (ligand) and 3CL SARS-Cov2(protease) using Docking server after 2 run.

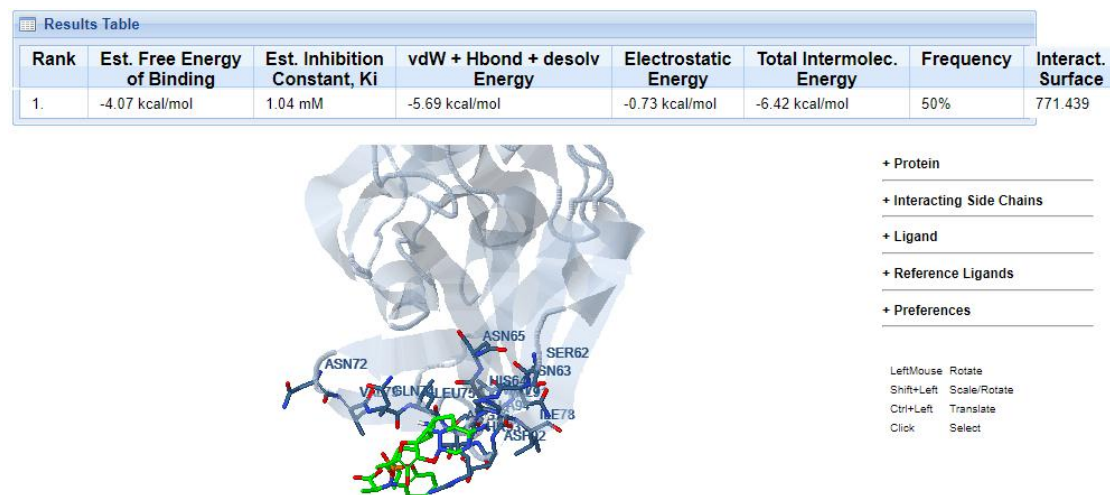


Figure5: Docking results obtained between Remdesivir and 3CL SARS-Cov2(protease) using Docking server after 2 run.

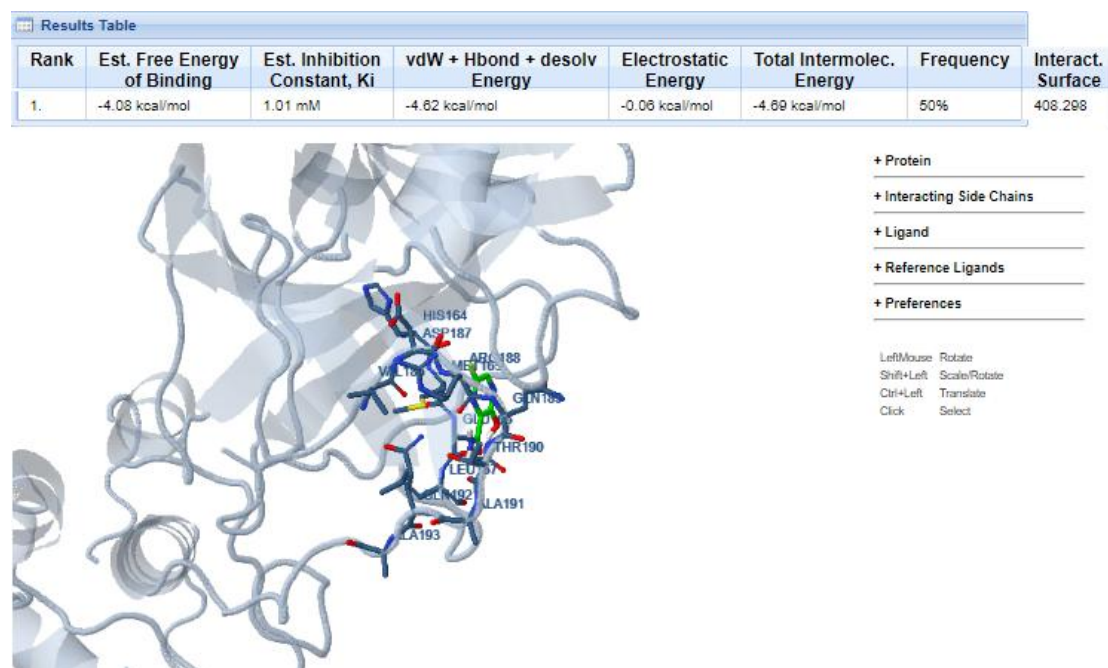
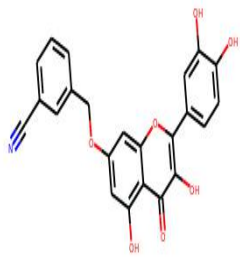
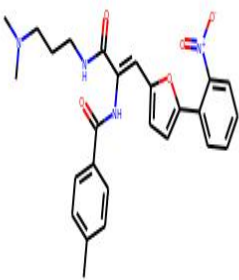
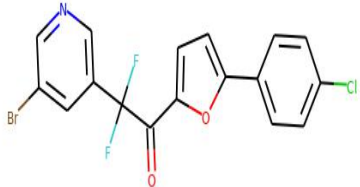
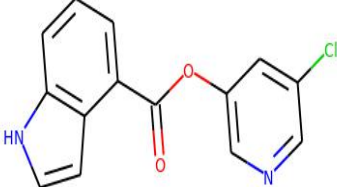
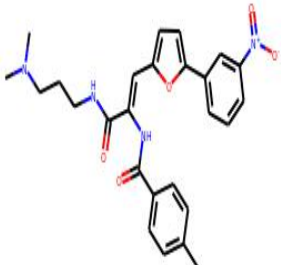
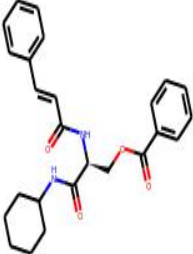
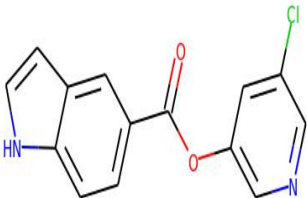
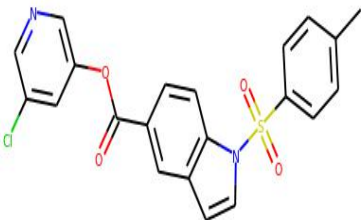
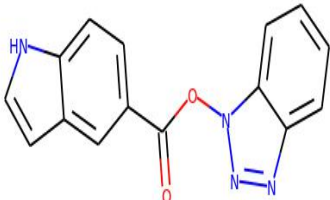
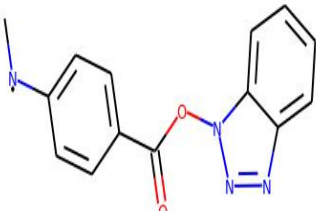
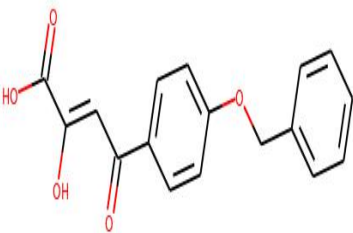
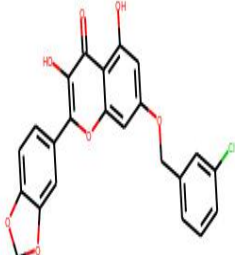


Figure6: Docking results obtained between Favipiravir and 3CL SARS-Cov2(protease) using Docking server after 2 run.

Table 2: Generated Molecules with high Tanimoto Similarity with the already existing 3CL Main Protease Inhibitors of SARS-CoV2.

ID	MOLECULES GENERATED	3CL MAIN PROTEASE INHIBITORS OF SARS-CoV2	Tanimoto Similarity
M_1			0.957

M_2			0.929
M_3			0.850
M_4			0.810
M_5			0.759
M_6			0.747

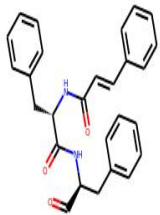
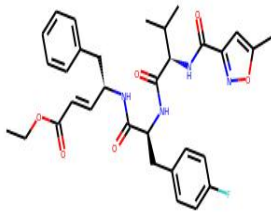
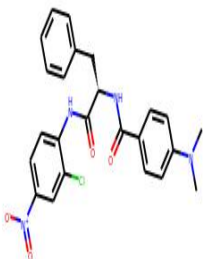

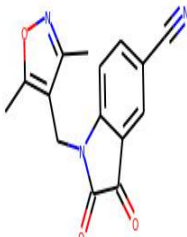
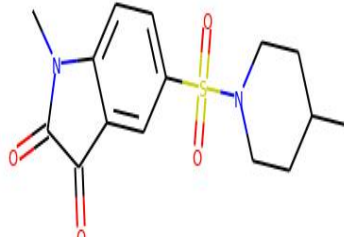
M_7			0.692
M_8			0.689
M_9			0.627

Table 3: Drug Likeness Properties of the Generated Molecule which are selected based on Lipinski's rule.

ID.	MW	Log P	HBA	HBD
M_1	417.37	3.733	8	4
M_2	412.61	5.73	3	0
M_3	476.53	4.00	6	2
M_4	272.61	3.43	3	1
M_5	278.27	2.18	4	1
M_6	298.29	2.97	5	2

M_7	426.51	3.35	3	2
M_8	466.92	4.29	5	2
M_9	281.27	1.89	5	0
PM_1	495.63	3.956	8	1

MW – Molecular Weight; Log P – Partition coefficient; HBA – Hydrogen Bond Acceptor; HBD – Hydrogen Bond Donor

4. Discussion

If the activation energy E_a , which is the difference in energy between reactants and activated complex is less, which is also called the Gibbs energy of activation represented by the equation,

$$\Delta G^\ddagger = \Delta H^\ddagger - T \Delta S^\ddagger.$$

then the rate of the reaction is more, assuming temperature is constant. So, if the rate of reaction in forward direction is more in both the steps of inhibitor reaction then inhibition will take place. So, we can reward the agent if the molecule formed after adding fragments has less activation energy and penalize if it has higher activation energy. Further, Variational Autoencoder can be improved. The conventional VAE uses KL divergence as a regularization term, instead of this, Wasserstein Distance can be used which is better than KL divergence. We can also use a generalized version of log likelihood and KL divergence similar to Coupled-VAE. Conditional GANs where QED score, fragment containment scores and pharmacophores are used as conditional input can also be used.

5. Methods

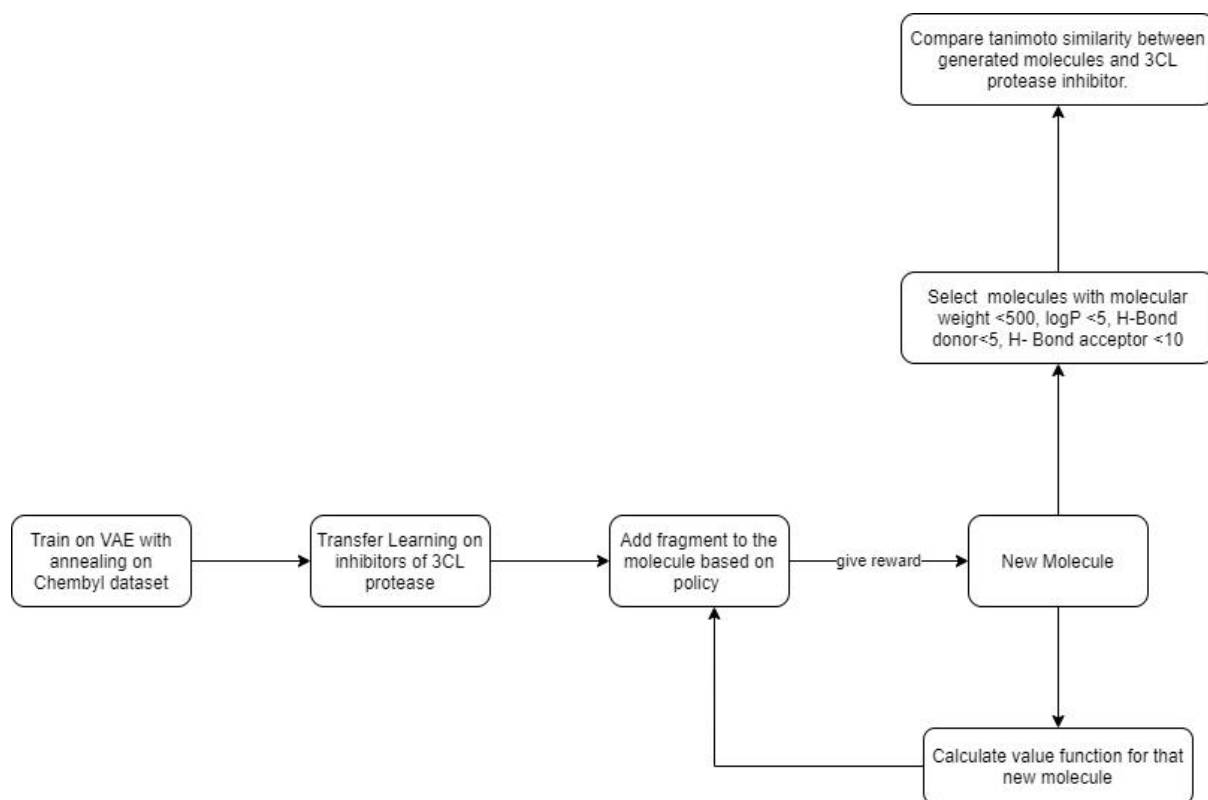


Figure2: Flow diagram for the method

5.1 Variational Autoencoder With KL Divergence And Circular Annealing

Given a datapoint x , the goal of VAE is to find at least one latent vector which is able to describe it; one vector that contains the instructions to generate x . If we formulate it using the law of total probability, we get $P(x) = \int P(x|z)P(z)dz$. The VAE training objective is to maximize $P(x)$. $P(x|z)$ is modeled using a multivariate Gaussian $N(f(z), \sigma^2 * I)$ where σ^2 is the hyperparameter that will be multiplied by I which represents the identity matrix. $f(z)$ is modeled using a neural network.

The formula for $P(x)$ is intractable, so it is approximated using Monte Carlo method which is described below:

1. Sample $\{z_i\}_{i=1}^n$ from the prior $P(z)$.
2. Approximate using $P(x) \approx \frac{1}{n} \sum_{i=1}^n p(x|z_i)$.

Since most sampled z 's won't contribute anything to $P(x)$, therefore, a new distribution $Q(z|x)$ is introduced. $Q(z|x)$ is trained to give high probability values

to z 's that are likely to have generated x . Now the Monte Carlo estimation can be calculated using much fewer samples from $Q(z|x)$.

Since

$$\log(t) \leq t - 1$$

$$\begin{aligned} -D_{\text{KL}}(q(x) \parallel p(x)) &= \int q(x) \log\left(\frac{p(x)}{q(x)}\right) dx \leq \int q(x) \left(\frac{p(x)}{q(x)} - 1\right) dx \\ &\leq \int p(x) dx - \int q(x) dx = 1 - 1 = 0 \end{aligned}$$

$$\Rightarrow D_{\text{KL}}(q(x) \parallel p(x)) \geq 0$$

This means

$$D_{\text{KL}}(p(q(z|x) \parallel p(z|x)) = -\int p(z|x) \log\left(\frac{p(z|x)}{q(z|x)}\right) dx \geq 0$$

Applying Bayes' theorem we get

$$\begin{aligned}
& -\int q(z|x) \log \left(\frac{p(x|z)p(z)}{q(z|x)p(x)} \right) dz \geq 0 \\
& \Rightarrow -\int q(z|x) \left[\log \left(\frac{p(x|z)p(z)}{q(z|x)} \right) - \log(p(x)) \right] dz \geq 0 \\
& \Rightarrow -\int q(z|x) \log \left(\frac{p(x|z)p(z)}{q(z|x)} \right) dz + \int q(z|x) \log(p(x)) dz \geq 0 \\
& \Rightarrow -\int q(z|x) \log \left(\frac{p(x|z)p(z)}{q(z|x)} \right) dz + \log(p(x)) \int q(z|x) dz \geq 0 \\
& \Rightarrow -\int q(z|x) \log \left(\frac{p(x|z)p(z)}{q(z|x)} \right) dz + \log(p(x)) \geq 0 \\
& \Rightarrow \log(p(x)) \geq \int q(z|x) \log \left(\frac{p(x|z)p(z)}{q(z|x)} \right) dz \\
& \Rightarrow \log(p(x)) \geq \int q(z|x) \log \left(\frac{p(x|z)p(z)}{q(z|x)} \right) dz \\
& \Rightarrow \log(p(x)) \geq \int q(z|x) \log \left(\frac{p(z)}{q(z|x)} \right) dz + \int q(z|x) \log(p(x|z)) dz \\
& \Rightarrow \log(p(x)) \geq -D_{KL}(q(z|x) || p(z)) + E_{z \sim q(z|x)} [\log(p(x|z))]
\end{aligned}$$

We took $\log(p(x))$ out of integral because if x is given then $\log p(x)$ will be constant, so maximizing log likelihood which is intractable is the same as minimizing its lower bound. Since it is not possible to differentiate with respect to random variable reparameterization trick is used if we sample ϵ from standard normal distribution and substitute $z = \mu + \epsilon * \sigma$.

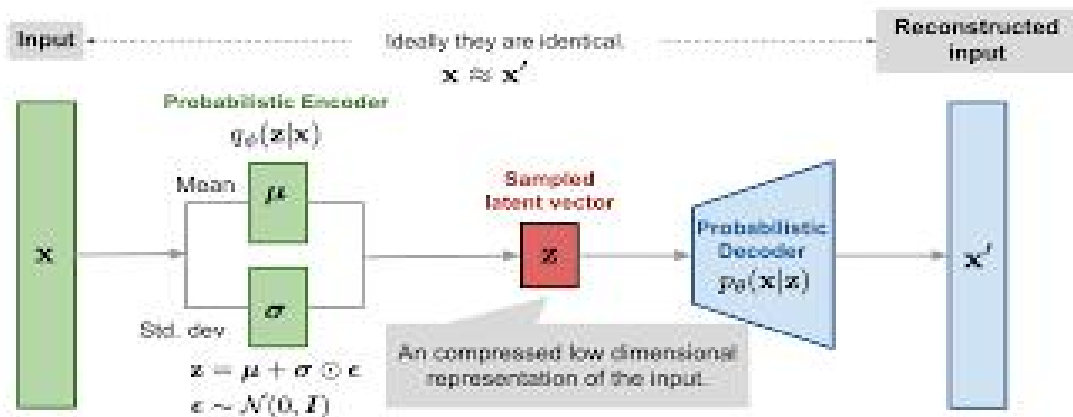


Figure 3. *Vanilla VAE*

The loss function consists of negative likelihood along with a regularizer term and a hyperparameter β which is increased slowly in case KL annealing and periodically in

case of circular annealing. This solves KL diminishing problem which makes VAE to act as simple RNN.

$$Loss(\theta, \varphi) = -E_{z \sim q_{\theta}(x)}[\log \log(p_{\varphi}(x|z))] + \beta KL(q_{\theta}(z|x)||p(z))$$

5.2 Markov Reward Process

This is an extension of Markov decision process for Reinforcement learning. This consists of states, actions, transition probability, rewards and discounting factors. In our case state will be the molecule structure at time step t will be state and adding and removing fragments will be action. Here instead of decreasing discounting factor for higher time step, it is increased by using λ^{T-t} where T is the time step at the end of the episode.

5.3 Agent Design

Agent fits the $Q^{\pi}(s_t, a_t)$ function which is the cumulative reward starting from state at time step t choosing action based on policy π such that it has maximum cumulative reward using advanced deep Q learning. The state-action and the value of the state are defined as

$$\begin{aligned} Q^{\pi}(s_t, a_t; \theta, \alpha, \beta) &= E_{a_t \sim \pi(s_t)} \left[\sum_{n=t}^T \gamma^{T-n} \cdot R(s_n, a_n) \right] \\ &= E_{a_t \sim \pi(s_t)} [R(s_t, a_t) + \gamma E_{a_{t+1} \sim \pi(s_{t+1})} (Q^{\pi}(s_{t+1}, a_{t+1}))] \end{aligned}$$

$$V^{\pi}(s_t) = E_{a_t \sim \pi(s_t)} [Q^{\pi}(s_t, a_t)]$$

In order to make RL more stable and to handle the problem of overestimation of Q-values a duel DQN network as target network is used. We keep its parameters θ', α', β' fixed and copy from dueling DQN Q^{π} every m steps.

$$TD = Q^{\pi}(s_t, a_t; \theta, \alpha, \beta) - [R(s_t', a_t) + \gamma \cdot Q_{tar}^{\pi}(s_{t+1}, \arg \max_{a_{t+1}} Q^{\pi}(s_{t+1}, a_{t+1}; \theta, \alpha, \beta); \theta', \alpha', \beta')]$$

Q function depends on parameters which come from dueling Q networks. Temporal difference is used in which reward is given at time step instead of end of episode. The parameters of Dueling Q network are updated as

$$loss(\theta, \alpha, \beta) = E[f(TD)]$$

RL agent is trained to minimize this loss function where f is the hubber function.

5.4 Reward Design

$$R(s) = w_{prof}f(s)_{pro} + w_{csf}f(s)_{csf} + w_{pha}f(s)_{pha}$$

Pro represents QED which is drug likeness property, csf represents whether a particular fragment is present and pha stands for pharmacophores score which depends on ligand-protein interaction mode from the crystal structure.

References

- 1) Bowen Tang, Fengming He, Dongpeng Liu, Meijuan Fang, Zhen Wu. AI-aided design of novel targeted covalent inhibitors against SARS-CoV-2. *BioRxiv*(2020)
- 2) Hao Fu^{1†}, Chunyuan Li^{2†*}, Xiaodong Liu² Jianfeng Gao², Asli Celikyilmaz², Lawrence Carin¹. Cyclical Annealing Schedule: A Simple Approach to Mitigating KL Vanishing
- 3) Gorbalenya AE. Severe acute respiratory syndrome-related coronavirus—The species and its viruses, a statement of the Coronavirus Study Group. *BioRxiv*, (2020). Google Scholar
- 4) Coronavirus COVID-19 Global Cases by Johns Hopkins CSSE, available from: <https://forum.truckersmp.com/index.php?/topic/93003-coronavirus-covid-19-global-cases-by-johns-hopkins-csse/&tab=comments>. Google Scholar
- 5) Zhang L, et al. Alpha-ketoamides as broad-spectrum inhibitors of coronavirus and enterovirus replication Structure-based design, synthesis, and activity assessment. *Journal of Medicinal Chemistry*, (2020). Google Scholar
- 6) Liu X, Wang X-J. Potential inhibitors for 2019-nCoV coronavirus M protease from clinically approved medicines. *bioRxiv*, (2020). Google Scholar
- 7) Yang H, et al. Design of wide-spectrum inhibitors targeting coronavirus main proteases. *PLoS biology* 3, (2005). Google Scholar
- 8) Singh J, Petter RC, Baillie TA, Whitty A. The resurgence of covalent drugs. *Nature reviews Drug discovery* 10, 307–317 (2011). CrossRefPubMedGoogle Scholar
- 9) Tuley A, Fast W. The taxonomy of covalent inhibitors. *Biochemistry* 57, 3326–3337 (2018).Google Scholar
- 10) Wu C-Y, et al. Small molecules targeting severe acute respiratory syndrome human coronavirus. *Proceedings of the National Academy of Sciences* 101, 10012–10017 (2004).Abstract/FREE Full TextGoogle Scholar
- 11) Ghosh AK, et al. Design and synthesis of peptidomimetic severe acute respiratory syndrome chymotrypsin-like protease inhibitors. *Journal of medicinal chemistry* 48, 6767–6771 (2005). CrossRefPubMedWeb of ScienceGoogle Scholar
- 12) Shie J-J, et al. Discovery of potent anilide inhibitors against the severe acute respiratory syndrome 3CL protease. *Journal of medicinal chemistry* 48, 4469–4473 (2005).CrossRefPubMedGoogle Scholar

- 13) Shie J-J, et al. Inhibition of the severe acute respiratory syndrome 3CL protease by peptidomimetic α , β -unsaturated esters. *Bioorganic & medicinal chemistry* 13, 5240–5252 (2005).Google Scholar
- 14) Al-Gharabli SI, et al. An efficient method for the synthesis of peptide aldehyde libraries employed in the discovery of reversible SARS coronavirus main protease (SARS-CoV Mpro) inhibitors. *ChemBioChem* 7, 1048–1055 (2006). PubMedGoogle Scholar
- 15) Lu I-L, et al. Structure-based drug design and structural biology study of novel nonpeptide inhibitors of severe acute respiratory syndrome coronavirus main protease. *Journal of medicinal chemistry* 49, 5154–5161 (2006). CrossRefPubMedGoogle Scholar
- 16) Tsai K-C, et al. Discovery of a novel family of SARS-CoV protease inhibitors by virtual screening and 3D-QSAR studies. *Journal of medicinal chemistry* 49, 3485–3495 (2006). CrossRefPubMedGoogle Scholar
- 17) Wu C-Y, et al. Stable benzotriazole esters as mechanism-based inactivators of the severe acute respiratory syndrome 3CL protease. *Chemistry & biology* 13, 261–268 (2006). CrossRefPubMedGoogle Scholar
- 18) Akaji K, Konno H, Onozuka M, Makino A, Saito H, Nosaka K. Evaluation of peptide-aldehyde inhibitors using R188I mutant of SARS 3CL protease as a proteolysis-resistant mutant. *Bioorganic & medicinal chemistry* 16, 9400–9408 (2008).Google Scholar
- 19) Ghosh AK, et al. Design, synthesis and antiviral efficacy of a series of potent chloropyridyl ester-derived SARS-CoV 3CLpro inhibitors. *Bioorganic & medicinal chemistry letters* 18, 5684–5688 (2008).Google Scholar.