Binding Affinity Prediction of Protein-Ligand complexes using Machine Learning

MSc Project

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Biological Background

What are proteins and ligands?

- **Proteins:** Complex molecules that are work-horses (machines) of a living organism.
- Ligands: Molecules that bind to (receptor) proteins.
- Proteins and ligands bind together to form protein-ligand complexes.







Biological Background

Protein-Ligand complexes

- Ligands bind to proteins at "cavity" like locations called pockets.
- The pockets and the ligands are complementary in shape.

Drugs

- Drugs are ligand molecules that bind to proteins.
- They cause a therapeutic effect after binding to the proteins.







Biological Background

Protein-Binding Affinity

- Assume a dynamic system in which protein P and ligand L are binding and unbinding continuously.
- Let [P] be the concentration of the protein and [L] be the concentration of the ligand. Let [PL] be the concentration of the protein ligand complex.
- Binding affinity can be quantified by using [P], [L] and [PL] (at equilibrium).

BindingAffinity =
$$\frac{[P][L]}{[PL]}$$



Problem Definition

- Determining if a potential drug (ligand) can bind to a target protein is very expensive [3].
- The project tries to reduce the drug discovery costs by eliminating bad leads.
- Problem definition: Predict protein-ligand binding affinity using "In-Silico" methods.



Figure: Project Overview

Problem Definition

There are various problems in the protein-ligand domain. The following figure shows the classification tree.

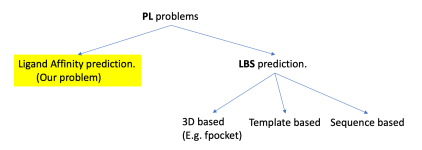


Figure: Protein-Ligand problem classifcation.



Problem Definition

- The input data to the ML model is extracted from a database called PDB Data bank.
- fpocket and RDKit were used to extract the features of proteins and ligands.
- The input features contain information about the 3D structures of the proteins and the ligands.

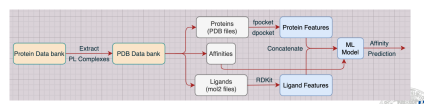


Figure: Data Input Overview.

Feature Extraction

Protein Features:

- fpocket is an LBS prediction algorithm used to predict ligand binding pockets.
- There can be multiple binding pockets for a PL complex.
- Using *dpocket*, 55 descriptors were obtained for every (potentially) binding pocket as real values.

Ligand Features:

 Using RDKit. Chem. Descriptors module, 402 features were extracted as real values.

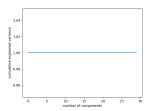
Concatenation:

- The (concatinated) input feature space to the model was
- R⁴⁵⁷.

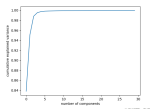
 It was less than R⁴⁵⁷ if feature selection is done before model. training.

Data Preprocessing

- Data points containing NaN (Not a number) values were removed from the data.
- PCA (Principle Component Analysis) was used to find the variance contribution of the features.
- Feature IPC was log scaled for numerical safety during training.



(a) With original Ligand feature IPC.



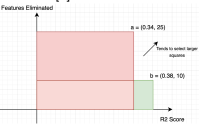
(b) With log scaled Ligand feature IPC.



Feature Selection

Feature selection strategies:

- Manual: 121 ligand descriptors + all protein descriptors.
- Output Correlation: Features with the best Pearson or Spearman correlation w.r.t the affinity score (output) were selected.
- **Genetic Algorithms**: Genetic algorithms with a population score function "**R**2 score * Features Eliminated" was used to select the best features [4]:

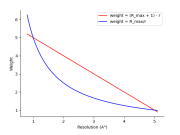




Dealing with measurement resolution

- In PDB bind databank, each complex has a corresponding measurement resolution (Å units).
- The structural detail of the 3D image is inversely proportional to the measurement resolution.
- The weighting of each data point was done according to hyperbolic formulae (or) linear formulae:

$$W_{\text{Hyperbolic}} = \frac{\max R_{1...n}}{R_i} \quad W_{\text{Linear}} = (\max R_{1...n} + 1) - R_i$$

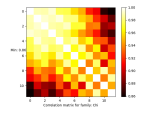






Feature Family Correlations

- Features can be divided into families.
- Important ones are AUTOCORR2d_, Chi, EState_VSA, PEOE_VSA, SMR_VSA, SlogP_VSA, VSA_EState, and fr_.
- Within Chi and AUTOCORR2d_, the features are correlated.
- ML models need to take into account this issue.



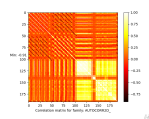
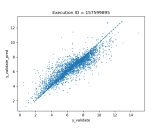


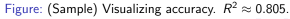
Figure: Correlation Heat Map.

Testing strategy

For every execution, we report the **random seed** used in it. This random seed can be used as a script argument (Execution ID) to reproduce the exact results. To determine the result quality we use:

- R^2 score (Coefficient of determination): $R^2 \in (-\infty, 1.0]$ where 1.0 is the best score.
- **Visualization:** The model's prediction is visualized as a 2D scatter plot. The best plot is y = x line which corresponds to the best R^2 score of 1.0.







Machine Learning models

The ML model should approximate the following function:

Binding affinity prediction : $\mathbb{R}^n \mapsto \mathbb{R}$ where $n \in \mathbb{I}^+$

The following ML models were studied

- Simple Linear Regression
- Random Forest Regression
- Support Vector Regression
- Rotation Forest Regression

Also Note:

- DNNs cold not be used due to lack of data.
- ullet The project only had pprox 16000 data points to train.
- A simple DNN a model of size [457, 20, 10, 1] has 9350 parameters.
- The DNN would overfit drastically.



Simple Linear Regression

- This model approximates the binding affinity using a linear hyperplane. (By minimizing the square of errors).
- It is the cheapest computational model.
- Assumes strong linear relationship between input features and binding affinity.
- Genetic algorithms sucessfully used for feature selection.
- Alternate weighting strategy: Data duplication.

No. features	Feature selection	Weighting	Training	Validation	Testing
457	-	-	0.461	0.415	0.320
457	-	Hyperbolic	0.454	0.427	0.337
457	-	Hyperbolic duplication	0.465	0.416	0.326
457	-	Linear	0.458	0.419	0.328
457	-	Linear Duplication	0.460	0.428	0.327
49	Genetic	Hyperbolic	≈0.377	≈0.374	≈0.364
40	Pearson Correlation	Hyperbolic	0.287	0.278	0.285
40	Spearman Correlation	Hyperbolic	0.289	0.294	0.290
176	Manual	Hyperbolic	0.362	0.346	0.331





Random Forest Regression



Support Vector Regression



Rotation Forest Regression



Discussion

Notable points:

- Best models: Linear Regression and Random Forest Regression.
- RF uses correlated features to make itself more robust.
- RF can deal with both discrete and real valued features.

Limitations:

- Linear regression assumes data linearity.
- RF has heavy reliance on ligand features.
- Both models were black box models.
- Testing results were sometimes better than validation results.
 This is because test data < validation data. But the

difference is minimal.

Discussion

Further work:

 A new weighting strategy: Weighting a pocket descriptor based on the overlap between the pocket and the ligand. For example,

$$W_{\mathrm{Total}} = W_{\mathrm{Hyperbolic}} * W_{\mathrm{Overlap}}$$

- Improvement of feature selection: Build 1 model per family of features. Use the best feature as a family surrogate.
- A more explainable model can be built.



Q & A



References

- Du, Li, Xia, Ai, Liang, Sang, Ji and Liu; Insights into Protein–Ligand Interactions: Mechanisms, Models, and Methods (2016)
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- DiMasi, Grabowski and Hansen; nnovation in the pharmaceutical industry: New estimates of R & D costs (2016)
- John H. Holland. Genetic Algorithms. (1960)
- Is rotation forest the best classifier for problems with continuous features? A. Bagnall, M. Flynn, J. Large, J. Line, A. Bostrom, and G. Cawley (2020)

Appendix

• Binding affinity between a protein and a ligand is quantified by the K_d , K_i and IC_{50} . Here K_d refers to the dissociation constant, K_i to inhibition constant, and IC_{50} to inhibitory concentration 50%.

