# 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.

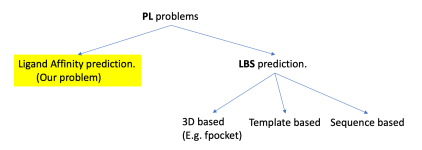


Figure: Protein-Ligand problem classifcation.



### **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.

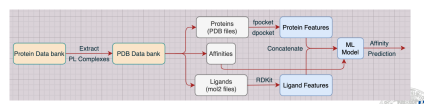


Figure: Data Input Overview.

### 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.

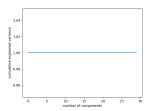
#### Concatenation:

- The (concatinated) input feature space to the model was
- R<sup>457</sup>.

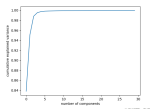
   It was less than R<sup>457</sup> if feature selection is done before model. training.

### 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.



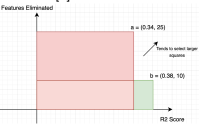
(b) With log scaled Ligand feature IPC.



#### 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]:

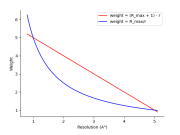




### Dealing with measurement resolution

- 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:

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

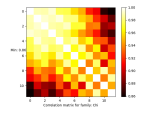






### Feature Family Correlations

- 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.



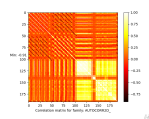
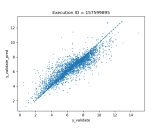


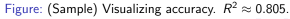
Figure: Correlation Heat Map.

### 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

#### Introduction

- It is a non-linear ensemble model of regression trees.
   (Sampling with replacement aka Bagging is used)
- For each tree, the data is split recursively till a stopping criterion. Each data subset has lesser entropy than the superset.
- There is no need for any assumption w.r.t data.
- Additional feature selection strategy: Genetic Elitism.

#### Dealing with correlated features

- The RF was forced to randomly use only 20% of the features to determine the best (feature, value) combination for each split step.
- The RF model does not depend on some features exclusively.
- The correlated features were used as an advantage.



### Random Forest Regression

#### **Feature Importances**

- Gini Importance: (Provided by the RF model)
  - The most important features are the ones that contribute the most in the reduction in entropy.
- Permutation Importance: (Model agnostic)
  - The most important features are the ones that are most relevant for prediction accuracy.

$$\operatorname{Importance}(f) = \operatorname{Accuracy}_{\operatorname{pred}}(X) - \operatorname{Accuracy}_{\operatorname{pred}}(X_{f\_shuffled})$$

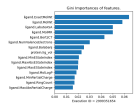
#### **Genetic feature selection**

- Fitting an RF model is expensive. However evaluation of feature importance is cheap.
- Hence the following scoring function is used,  $s = \text{Importance}(f_1, f_2...) * \text{features\_eliminated}$

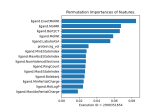




### Random Forest Regression



(a) Gini importance.



(b) Permutation importance.

Figure: Feature Importance calculation of Random Forest Regressor (With manual feature selection).

| No. Features | Feature Selection    | Weighting  | Training | Validation | OOB score | Testing |
|--------------|----------------------|------------|----------|------------|-----------|---------|
| 457          | -                    | -          | 0.961    | 0.790      | 0.791     | 0.447   |
| 457          | -                    | Hyperbolic | 0.961    | 0.773      | 0.794     | 0.448   |
| 457          | -                    | Linear     | 0.960    | 0.785      | 0.794     | 0.443   |
| 40           | Spearman Correlation | Hyperbolic | 0.930    | 0.664      | 0.671     | 0.878   |
| 40           | Pearson Correlation  | Hyperbolic | 0.925    | 0.642      | 0.645     | 0.868   |
| 229          | Genetic Elitism      | Hyperbolic | 0.958    | 0.779      | 0.793     | 0.444   |
| 394          | Genetic Normal       | Hyperbolic | 0.961    | 0.776      | 0.795     | 0.448   |
| 176          | manual               | Hyperbolic | 0.949    | 0.734      | 0.736     | 0.463   |



Table: Random Forest Regression R<sup>2</sup> Score table.

### Support Vector Regression

- It is a non linear model.
- It fits a  $\epsilon$  radius pipe to the data.
- The model was not suitable for running genetic algorithms (for both normal and permutation genetics). Training took  $\approx 99.07s$  and validation took  $\approx 27.01s$ .
- Features were selected using either output correlation or manual feature selection.
- Contrary to other models, linear weighting of data gave the best results.

| No. Features | Feature Selection    | Weighting  | Training | Validation | Testing |
|--------------|----------------------|------------|----------|------------|---------|
| 457          | -                    | -          | 0.311    | 0.290      | 0.314   |
| 457          | -                    | Hyperbolic | 0.330    | 0.323      | 0.327   |
| 457          | -                    | Linear     | 0.344    | 0.319      | 0.335   |
| 40           | Spearman Correlation | Linear     | 0.260    | 0.268      | 0.262   |
| 40           | Pearson Correlation  | Linear     | 0.198    | 0.196      | 0.197   |
| 40           | Spearman Correlation | Hyperbolic | 0.254    | 0.265      | 0.256   |
| 40           | Pearson Correlation  | Hyperbolic | 0.168    | 0.171      | 0.168   |
| 176          | Manual               | Linear     | 0.311    | 0.283      | 0.310   |

Table: Table showing  $R^2$  Scores for SVR.



### Rotation Forest Regression

- Random forest trees divide data using axis aligned hyperplanes.
- The complexity of the trees can be reduced by linearly transforming (rotating) the data. The eigen vectors of the training data become the basis vector. For example:
  - Representing y = kx would take a very deep tree.
  - If y = kx is linearly transformed to y = 0, only 1 node is enough.
- Computationally very expensive It took  $\approx$  25 min 29 seconds to train.

| No. Features | Selected             | Training | Validation | Testing |
|--------------|----------------------|----------|------------|---------|
| 457          | -                    | 0.967    | 0.767      | 0.449   |
| 40           | Spearman Correlation | 0.952    | 0.650      | 0.890   |
| 40           | Pearson Correlation  | 0.951    | 0.629      | 0.883   |
| 176          | manual               | 0.960    | 0.716      | 0.471   |

Table: Rotation Forest  $R^2$  Score overview.



### Discussion

#### Notable points:

- Best models: Linear Regression and Random Forest Regression.
- Random Forest uses correlated features to make itself more robust.
- Random Forest can deal with both discrete and real valued features.
- Rotation Forest did not improvement accuracy as they are good only for Real valued features.

#### Limitations:

- Linear regression assumes data linearity.
- Random Forest has heavy reliance on ligand features.
- Both models were black box models.
- Testing results were sometimes better than validation results.

  It is because test data < validation data. But the ""

  minimal. 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)
- Le Guilloux, Schmidtke, and Tuffery; Fpocket: An open source platform for ligand pocket detection(2009)
- 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 - Definitions

• 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%.



### Appendix - Visualizing Linear Regression results

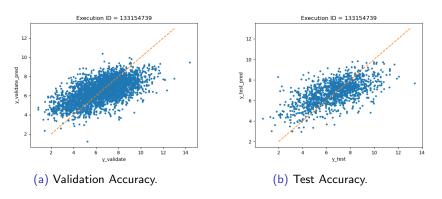


Figure: Linear Model using Hyperbolic weighting and all 49 features selected by genetic algorithm.

### Appendix - Visualizing Random Forest results

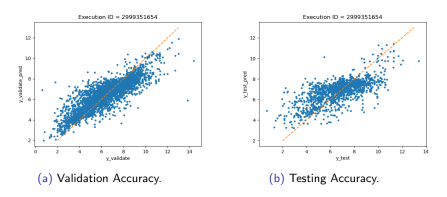


Figure: Random Forest Regressor with 176 manually selected features and hyperbolic weighting.

### Appendix - Visualizing SVR Results

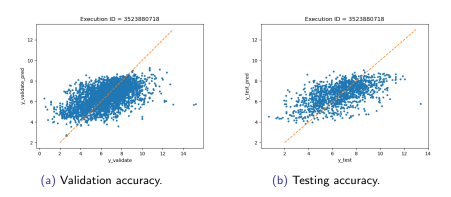


Figure: SVR accuracy visualization for all features 457 and Linear weighting.



### Appendix - Visualizing Rotation Forest Results

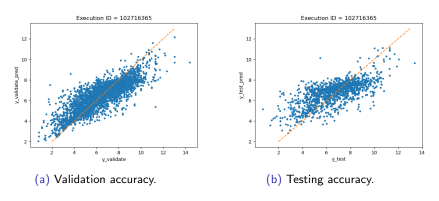


Figure: Rotation Forest Accuracy visualization for manually selected features (176). The rotation forest implementation does not support data weighting.