# Prediction of the Effect of Single Amino Acid Protein Variants Using Deep Mutational Scanning Data

University of Bologna — Master Thesis in Bioinformatics

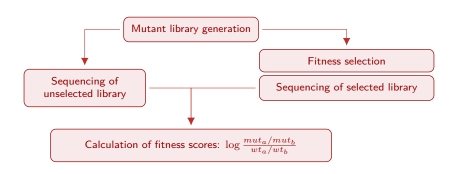
#### Pierotti Saul

Internal Advisor: Prof. Pietro Di Lena

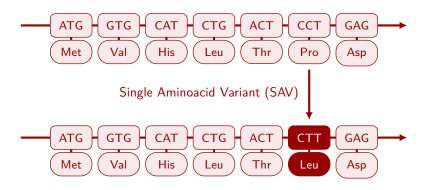
External Advisor: Prof. Arne Elofsson (Stockholm University)

#### Deep Mutational Scanning

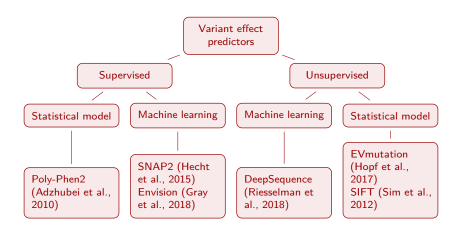
High-throughput technique for obtaining fitness information on a large number of mutations



# I Considered Only Single Amino Acid Variants

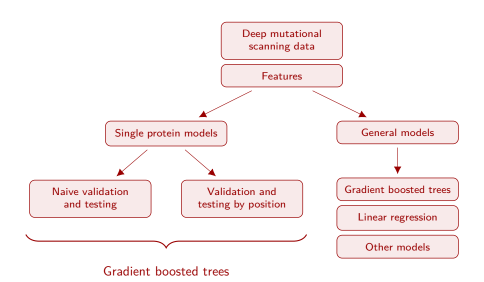


#### Some Notable Variant Effect Predictors

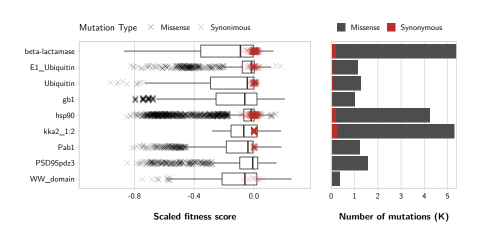


Among these predictors only Envision was trained on deep mutational scanning data

#### Structure of the Project

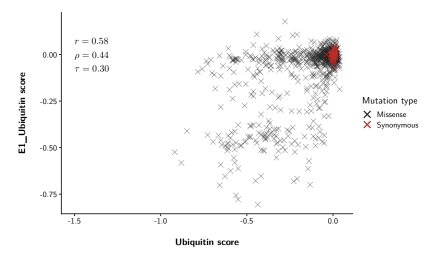


# I Used the Training Dataset of Envision (Gray et al., 2018)



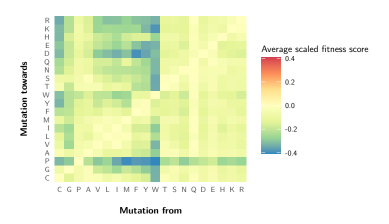
#### Poor Correlation among Experimental Results

Two independent deep mutational scanning experiments on Ubiquitin are present in the training dataset. Their correlation is low.

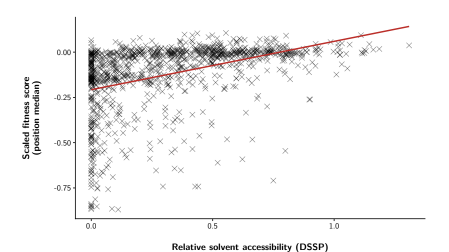


#### The Identities of the Mutated Residues Are Important

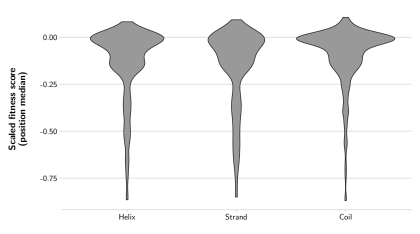
Mutations from polar residues are less detrimental than other mutations. This effect disappears when filtering by relative solvent accessibility (not shown here).



#### Buried Residues Are More Conserved



# Secondary Structure Is Not Critical



Secondary structure (DSSP)

#### I Did Not Use Structural Information



Wild-type residue

Mutated residue

EVmutation predictions (Hopf et al., 2017)

Epistatic model

Independent model

Conservation

Mutation frequency

NetsurfP-2 predictions (Klausen et al., 2019)

Secondary structure

Solvent accessibility

Disorder

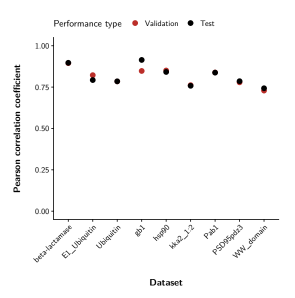
Torsion angles

HMMER (Eddy, 2011) emission probabilities

trRosetta predicted contacts (Yang et al., 2020)

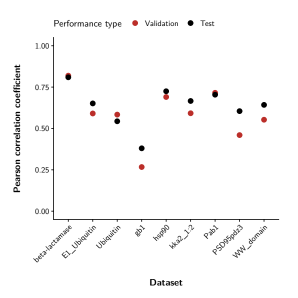
Centrality metrics

# Single Protein Models with Naive Testing



- Half of the mutations in a protein randomly set aside for testing
- ► Hyperparameters optimized in the remaining half with 5-fold cross-validation
- ► Good results but likely overfitting the testing set

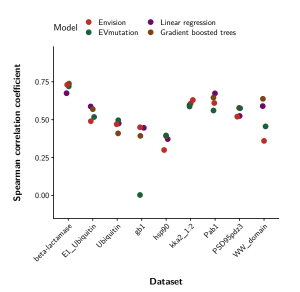
# Single Protein Models with Testing by Position



- Half of the mutations in a protein set aside for testing but avoiding mutations in the same protein position to end up in different splits
- ► Hyperparameters optimized in the remaining half with 5-fold cross-validation
- ▶ Performance more realistic

Pierotti Saul

# Leave-One-Protein-Out (LOPO) Models



- ► For the left-out protein, half of the mutations used for testing
- ➤ Spearman correlation coefficient used for evaluation
- ➤ Small difference among gradient boosted trees and linear regression
- Performances comparable to those of Envision

#### Discussion

Complex models do not improve much on linear regression

There is strong variability between datasets

Unsupervised models perform similarly to supervised models

How validation and testing are performed is crucial

Performances on par with other predictors can be reached without structural features

#### **Future Directions**

Unsupervised models seem promising and may be worth exploring more

Training on more deep mutational scanning studies

Tuning the set of features

Using residue contacts in a graph convolutional neural network

Finding a better normalization strategy for the scores from different experiments

# Bibliography I

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#### Confidence intervals in prediction performances

| Dataset name   | Model       | 95 % C.I. (Pearson) | 95 % C.I. (Spearman) |
|----------------|-------------|---------------------|----------------------|
| beta-lactamase | Naive       | 0.89 to 0.91        | 0.86 to 0.88         |
| beta-lactamase | By position | 0.79 to 0.83        | 0.75 to 0.78         |
| beta-lactamase | LOPO        | _                   | 0.68 to 0.72         |
| WW_domain      | Naive       | 0.67 to 0.82        | 0.70 to 0.84         |
| WW_domain      | By position | 0.57 to 0.73        | 0.52 to 0.72         |
| WW_domain      | LOPO        | _                   | 0.56 to 0.73         |
| PSD95pdz3      | Naive       | 0.74 to 0.83        | 0.73 to 0.80         |
| PSD95pdz3      | By position | 0.55 to 0.67        | 0.54 to 0.65         |
| PSD95pdz3      | LOPO        | _                   | 0.53 to 0.63         |
| kka2_1:2       | Naive       | 0.74 to 0.78        | 0.72 to 0.76         |
| kka2_1:2       | By position | 0.65 to 0.69        | 0.64 to 0.68         |
| kka2_1:2       | LOPO        | _                   | 0.60 to 0.64         |
| hsp90          | Naive       | 0.82 to 0.87        | 0.65 to 0.71         |
| hsp90          | By position | 0.69 to 0.76        | 0.50 to 0.57         |
| hsp90          | LOPO        | _                   | 0.38 to 0.45         |
| Ubiquitin      | Naive       | 0.75 to 0.83        | 0.78 to 0.83         |
| Ubiquitin      | By position | 0.49 to 0.60        | 0.46 to 0.57         |
| Ubiquitin      | LOPO        | _                   | 0.30 to 0.43         |
| Pab1           | Naive       | 0.80 to 0.87        | 0.75 to 0.82         |
| Pab1           | By position | 0.65 to 0.76        | 0.64 to 0.73         |
| Pab1           | LOPO        | _                   | 0.60 to 0.70         |
| E1_Ubiquitin   | Naive       | 0.75 to 0.85        | 0.65 to 0.75         |
| E1_Ubiquitin   | By position | 0.59 to 0.72        | 0.36 to 0.51         |
| E1_Ubiquitin   | LOPO        | _                   | 0.50 to 0.63         |
| gb1            | Naive       | 0.90 to 0.93        | 0.88 to 0.92         |
| gb1            | By position | 0.31 to 0.46        | 0.29 to 0.45         |
| gb1            | LÓPO        | _                   | 0.32 to 0.47         |

Statistical significance of performance differences. Bonferroni-corrected  $\alpha=\frac{0.05}{27}=0.00185185$ . Starred values are significant.

| beta-lactamase Linear regression Gradient boosted trees 1 · 10 <sup>-4</sup> * beta-lactamase Linear regression EVmutation 1 · 10 <sup>-4</sup> * beta-lactamase EVmutation Gradient boosted trees 1 · 10 <sup>-4</sup> * ww_domain Linear regression Gradient boosted trees 1 · 10 <sup>-4</sup> * ww_domain Linear regression EVmutation 1 · 10 <sup>-4</sup> * ww_domain EVmutation Gradient boosted trees 1 · 10 <sup>-4</sup> * ww_domain EVmutation Gradient boosted trees 1 · 10 <sup>-4</sup> * PSD95pd23 Linear regression Gradient boosted trees 1 · 10 <sup>-4</sup> * PSD95pd23 Linear regression EVmutation 1 · 10 <sup>-4</sup> * PSD95pd23 EVmutation Gradient boosted trees 0 · 51 Linear regression EVmutation 0 · 0.51 Linear regression Gradient boosted trees 0 · 0.51 Linear regression Gradient boosted trees 0 · 0.00 Linear regression EVmutation 0 · 0.01 Linear regression EVmutation 0 · 0.01 Linear regression EVmutation 0 · 0.01 Linear regression EVmutation Gradient boosted trees 1 · 10 <sup>-4</sup> * https://doi.org/10.10.01/10.01 | Dataset name   | Model 1           | Model 2                | p value    |
|--|----------------|-------------------|------------------------|------------|
| beta-lactamase     EVmutation     Gradient boosted trees     1 · 10 <sup>-4</sup> *       WW_domain     Linear regression     Gradient boosted trees     1 · 10 <sup>-4</sup> *       WW_domain     EVmutation     EVmutation     1 · 10 <sup>-4</sup> *       WB_domain     EVmutation     Gradient boosted trees     1 · 10 <sup>-4</sup> *       PSD95pdz3     Linear regression     EVmutation     1 · 10 <sup>-4</sup> *       PSD95pdz3     EVmutation     EVmutation     1 · 10 <sup>-4</sup> *       PSD95pdz3     EVmutation     Gradient boosted trees     0.51       kka2_1:2     Linear regression     Gradient boosted trees     0.00       kka2_1:2     Linear regression     EVmutation     0.01       kka2_1:2     EVmutation     Gradient boosted trees     1 · 10 <sup>-4</sup> *       kka2_1:2     EVmutation     Gradient boosted trees     1 · 10 <sup>-4</sup> *       hsp90     Linear regression     Gradient boosted trees     1 · 10 <sup>-4</sup> *  | beta-lactamase | Linear regression | Gradient boosted trees |            |
| WW_domain     Linear regression     Gradient boosted trees $1 \cdot 10^{-4}$ *       WW_domain     Linear regression     EVmutation $1 \cdot 10^{-4}$ *       WB_domain     EVmutation     Gradient boosted trees $1 \cdot 10^{-4}$ *       PSD95pd23     Linear regression     Gradient boosted trees $1 \cdot 10^{-4}$ *       PSD95pd23     EVmutation $1 \cdot 10^{-4}$ *       PSD95pd23     EVmutation     Gradient boosted trees $0.51$ kka2_1:2     Linear regression     Gradient boosted trees $0.00$ kka2_1:2     Linear regression     EVmutation $0.00$ kka2_1:2     EVmutation     Gradient boosted trees $1 \cdot 10^{-4}$ *       hsp90     Linear regression     Gradient boosted trees $1 \cdot 10^{-4}$ *   | beta-lactamase | Linear regression | EVmutation             | 1 · 10-4 * |
| WW_domain     Linear regression     EVmutation $1 \cdot 10^{-4}$ *       WW_domain     EVmutation     Gradient boosted trees $1 \cdot 10^{-4}$ *       PSD95pd23     Linear regression     Gradient boosted trees $1 \cdot 10^{-4}$ *       PSD95pd23     Elmear regression     EVmutation $1 \cdot 10^{-4}$ *       PSD95pd23     EVmutation     Gradient boosted trees $0.51$ kka2_1:2     Linear regression     Gradient boosted trees $0.00$ kka2_1:2     Linear regression     EVmutation $0.01$ kka2_1:2     EVmutation     Gradient boosted trees $1 \cdot 10^{-4}$ *       hsp90     Linear regression     Gradient boosted trees $1 \cdot 10^{-4}$ *  | beta-lactamase | EVmutation        | Gradient boosted trees |            |
| Wb_domain     EVmutation     Gradient boosted trees $1 \cdot 10^{-4}$ *       PSD95pdz3     Linear regression     Gradient boosted trees $1 \cdot 10^{-4}$ *       PSD95pdz3     Linear regression     EVmutation $1 \cdot 10^{-4}$ *       PSD95pdz3     EVmutation     Gradient boosted trees $0.51$ kka2_1:2     Linear regression     Gradient boosted trees $0.00$ kka2_1:2     Linear regression     EVmutation $0.01$ kka2_1:2     EVmutation     Gradient boosted trees $1 \cdot 10^{-4}$ *       hsp90     Linear regression     Gradient boosted trees $1 \cdot 10^{-4}$ *   | WW_domain      | Linear regression | Gradient boosted trees |            |
| $ \begin{array}{cccccccccccccccccccccccccccccccccccc$  | WW_domain      | Linear regression | EVmutation             |            |
| $ \begin{array}{cccccccccccccccccccccccccccccccccccc$  | WW_domain      | EVmutation        | Gradient boosted trees | 1 · 10-4 * |
| $ \begin{array}{cccccccccccccccccccccccccccccccccccc$  | PSD95pdz3      | Linear regression | Gradient boosted trees | 1 · 10-4 * |
| $kka2\_1:2$ Linear regression     Gradient boosted trees $0.00$ $kka2\_1:2$ Linear regression     EVmutation $0.01$ $kka2\_1:2$ EVmutation     Gradient boosted trees $1 \cdot 10^{-4}$ * $hsp90$ Linear regression     Gradient boosted trees $1 \cdot 10^{-4}$ *   | PSD95pdz3      | Linear regression | EVmutation             | 1 · 10-4 * |
| kka2_1:2Linear regressionEVmutation0.01kka2_1:2EVmutationGradient boosted trees $1 \cdot 10^{-4}$ *hsp90Linear regressionGradient boosted trees $1 \cdot 10^{-4}$ *  | PSD95pdz3      | EVmutation        | Gradient boosted trees |            |
| kka2_1:2 EVmutation Gradient boosted trees $1 \cdot 10^{-4}$ * hsp90 Linear regression Gradient boosted trees $1 \cdot 10^{-4}$ *  | kka2_1:2       | Linear regression | Gradient boosted trees | 0.00       |
| hsp90 Linear regression Gradient boosted trees $1 \cdot 10^{-4}$ *   | kka2_1:2       | Linear regression | EVmutation             | 0.01       |
|  | kka2_1:2       | EVmutation        | Gradient boosted trees | 1 · 10-4 * |
|  | hsp90          | Linear regression | Gradient boosted trees | 1 · 10-4 * |
| hsp90 Linear regression EVmutation 1 · 10 <sup>-4</sup> *  | hsp90          | Linear regression | EVmutation             | 1 · 10-4 * |
| hsp90 EVmutation Gradient boosted trees 0.24   | hsp90          | EVmutation        | Gradient boosted trees | 0.24       |

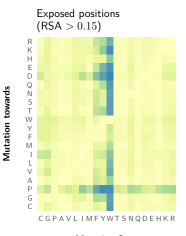
The table continues on the next slide.

Statistical significance of performance differences. Bonferroni-corrected  $\alpha=\frac{0.05}{27}=0.00185185.$  Starred values are significant.

The table continues from the previous slide.

| Dataset name | Model 1           | Model 2                | p value      |
|--------------|-------------------|------------------------|--------------|
| Ubiquitin    | Linear regression | Gradient boosted trees | 1 · 10 - 4 * |
| Ubiquitin    | Linear regression | EVmutation             | 1 · 10 - 4 * |
| Ubiquitin    | EVmutation        | Gradient boosted trees | 1 · 10 - 4 * |
| Pab1         | Linear regression | Gradient boosted trees | 1 · 10 - 4 * |
| Pab1         | Linear regression | EVmutation             | 1 · 10 - 4 * |
| Pab1         | EVmutation        | Gradient boosted trees | 1 · 10 - 4 * |
| E1 Ubiquitin | Linear regression | Gradient boosted trees | 1 · 10 - 4 * |
| E1 Ubiquitin | Linear regression | EVmutation             | 1 · 10 - 4 * |
| E1 Ubiquitin | EVmutation        | Gradient boosted trees | 1 · 10 - 4 * |
| gb1          | Linear regression | Gradient boosted trees | 1 · 10 - 4 * |
| gb1          | Linear regression | EVmutation             | 1 · 10 - 4 * |
| gb1          | EVmutation        | Gradient boosted trees | 1 · 10 - 4 * |

When filtering by relative solvent accessibility apolar residues are not more sensitive to mutations than polar residues.



Mutation from