Machine-learning models able to predict phage-bacteria interactions

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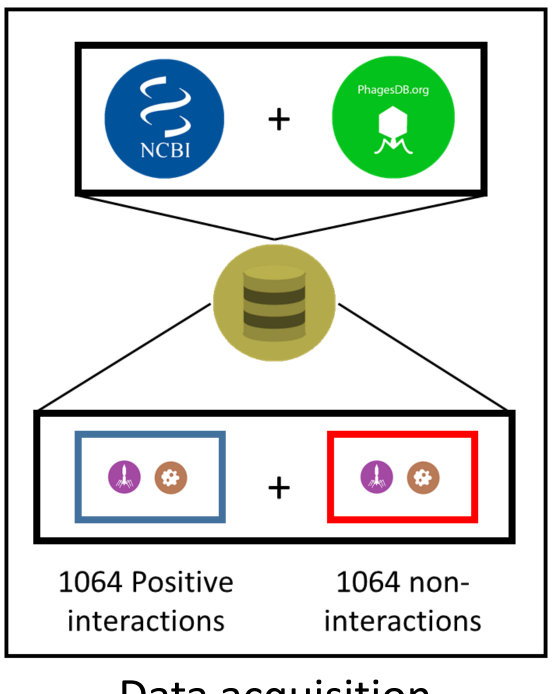
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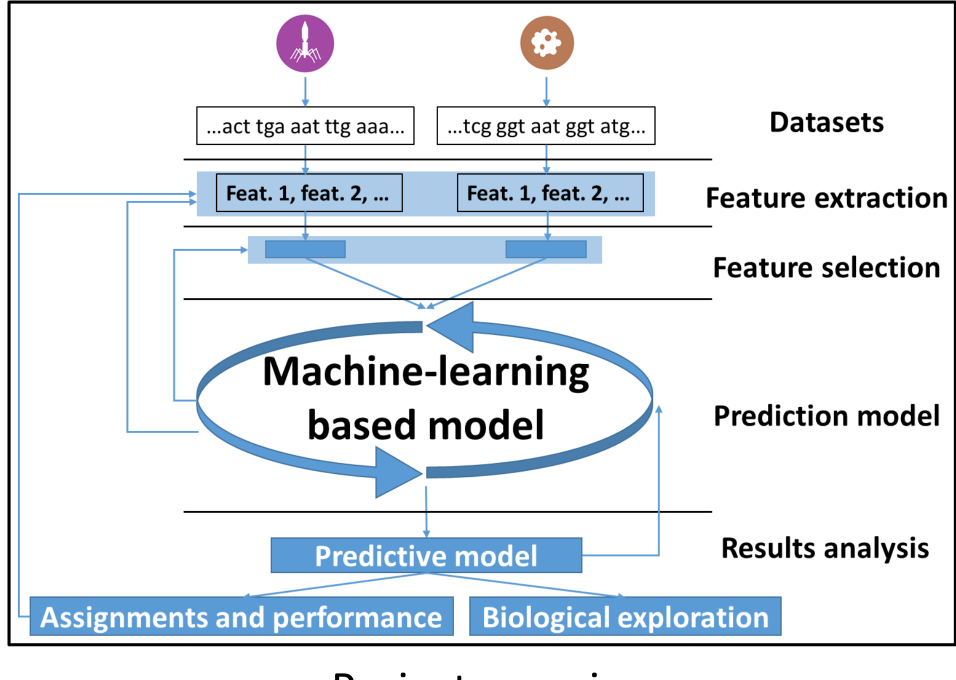


Abstract

Phage-therapy, a promising alternative to antibiotic-resistance, uses phages to infect and kill pathogenic bacteria. It requires finding perfectly matching phage-bacterium pairs, a time and money-consuming task, currently achieved empirically in laboratory. Our project aims at improving this task by predicting, *in-silico*, if a given phage-bacterium pair would interact. Predictions are performed on the base of public genomic data combined with machine-learning algorithms. With such an approach we have obtained around 90% of predictive power. In order to improve these results, we will extend our methodology and we will validate it with newly-generated clinically-relevant data.

Overview: prediction of phage-bacteria interactions





Data acquisition Project overview

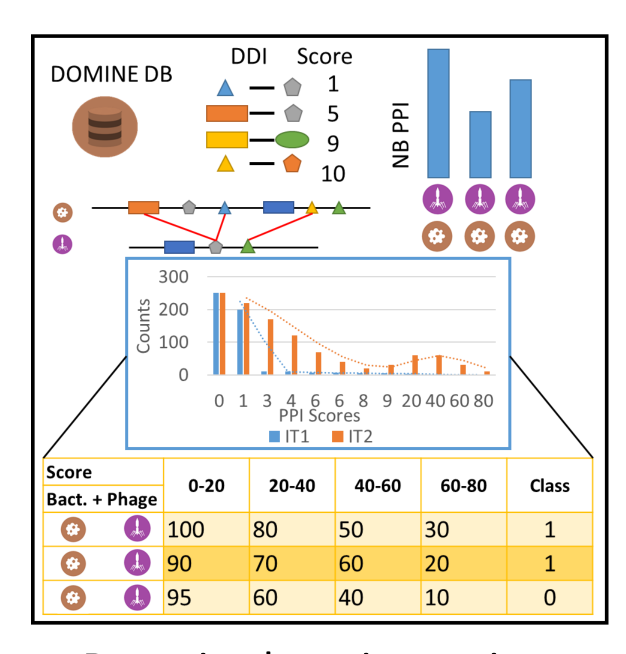
For our project we:

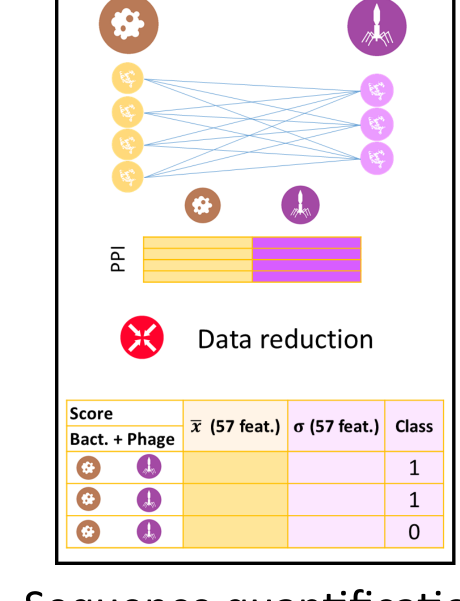
- Acquired data from public databases as NCBI [1] and phagesdb.org [2]. From GeneMarkS [3] we predict genes in bacterial and phages genomes.
- Constituted a positive dataset with 1064 phage-bacteria interactions pairs.
- Developed a model able to predict phage-bacteria interactions

Feature engineering: obtention of informative features

On our data we:

- Extracted two kinds of features based on:
 - · Protein interactions (PFAM Domain-domain interactions) 18 datasets
 - Genomic sequences (% of amino acids, chemical components and weight) 1 dataset
- Corrected the over-representation of the Mycobacterium smegmatis mc2 155 bacterium, reducing it from 86% to 14%

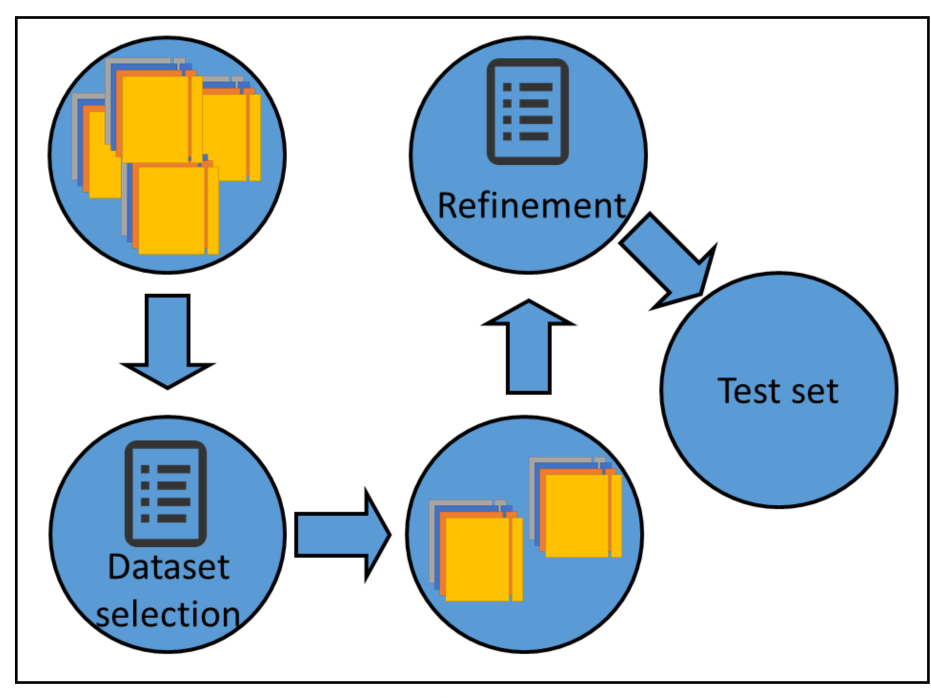




Domain-domain scoring

Sequence quantification

Machine-Learning based modeling



Process of research used

- In our machine-learning search we used:
 - 19 datasets (12'918 samples)
 - Predictive model building with 10-fold cross-validation
 - Four approaches: K-Nearest Neighbors (kNN), Random Forests (RF), Support Vector Machines (SVM), and Artificial Neural Networks (ANN)

In order to identify the best datasets and configurations of algorithms, we performed the following:

- 1. Selected the datasets exhibiting the highest scores across the four approaches
- 2. Further modelling to refine the algorithms' hyperparameters

Main results and conclusions

The best results on the test, obtained by ANN with 9 neurons and 50 epochs are presented below:

Datasets	Accuracy	F-Score	Sensitivity	Specificity
NB50	89.78%	90.13%	89.56%	90.12%
NBN50	89.79%	90.13%	89.56%	90.12%
S1e ⁻⁶	85.79%	86.24%	85.43%	86.35%

Future work:

- Explore other types of features to further increase model performance.
- Improve the predicitvity of the dataset by prunning redundant/ correlated variables that may perturb modeling
- Search for new relevant interactions allowing to predict interactions for different strains of a given bacterial species
- Improve domain interactions scores



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