ADMET Property Prediction With Oloren ChemEngine

Andrew Li, David Huang

Oloren AI

Abstract

We present a set of model architectures built using the Oloren ChemEngine library for the Therapeutics Data Commons (TDC) benchmarks. We evaluate the models on ADMET benchmarks, representing various small molecule drug properties encompassing absorption, distribution, metabolism, excretion, and toxicity. The models predict values for these properties given the chemical structure of the molecule. The presented models draw from a wide set of state-of-art strategies, such as custom model gradient boosting, graph neural networks, molecular fingerprinting, and OlorenVec, a supervised learned molecular representation. We show that these models are superior to current leaderboard model approaches.

Introduction to OCE

Oloren ChemEngine¹ (OCE) is an open-source Python software package developed by Oloren AI for developing and using molecular property predictors. OCE implements current state-of-art methods such as message passing neural networks, graph neural networks, and molecular fingerprints and representations. It additionally allows for the creation of custom gradient boosting models consisting of user-defined submodels for each learner stage.

The TDC² ADMET benchmarks consist of 22 datasets composed of drug chemical structures and the property value for their respective ADMET property. Dataset sizes range from hundreds to \sim 15,000 molecules. We present the models which achieve a rank of first-place in 14 of these tasks, and top 3 in 5 tasks.

Model Architectures

We construct and test 8 unique models. This set of models consists of 1 random forest stacker, 1 Extreme Gradient Boosting model (XGBoost), and 6 custom gradient boosting models composed of various submodels.

The models are represented in the table below:

| Model Name | Model Description | Number Parameters | |
|------------------------------|---|-------------------|--|
| BaseBoosting ADkCCrwJ | Random Forest (RDKit2DNormalized), Random Forest (Morgan Fingerprint), Random Forest (OlorenVec) | 23 | |
| BaseBoosting OQRAYLPP | SPGNN, Random Forest (OlorenVec), BaseTorchGeometricModel (GINModel) | 2,003,108 | |
| BaseBoosting xXOn1QFI | Random Forest (Morgan Fingerprint), Random Forest(Morgan Chiral Counts), Random Forest (Morgan Feature Counts), Random Forest (RDKit2DNormalized), Random Forest (OlorenVec) | 27 | |
| BaseBoosting 1zpI0dIb | Random Forest (Morgan Fingerprint), Random Forest (Morgan Chiral Counts), Random Forest (Morgan Feature Counts), Random Forest (RDKit2DNormalized), Random Forest (OlorenVec) | 27 | |
| BaseBoosting ZSCWPcLT | Random Forest (Morgan Fingerprint) Random Forest (Morgan Chiral Counts) Random Forest (Morgan Feature Counts) Random Forest (RDKit2DNormalized) Random Forest (OlorenVec) | 27 | |
| BaseBoosting ktIIq91G | Random Forest (Morgan Fingerprint), Random Forest (Morgan Feature Counts), Random Forest (Morgan Chiral Counts), Random Forest (RDKit2DNormalized), Random Forest (Mordred Descriptor), Random Forest (OlorenVec) | 40 | |
| ZWK_XGBoostModel u3zq9AAV | XGBoost (RDKit2DNormalized) | 12 | |
| RFStacker jTNhN7U7 | Random Forest (RDKit2DNormalized) | 1,858,225 | |

| Random Forest (OlorenVec) Random Forest (Lipinski Descriptor) Random Forest (Morgan Feature Counts) SPGNN | |
|---|--|
|---|--|

Model names are defined by OCE's uniquely generated name for a given model. BaseBoosting models are custom-made gradient boosting models. SPGNN represents the graph neural network implemented in Hu et al. (2020)³. Certain gradient boosting models have identical model and feature architectures but differ in terms of the parameterization of the constituent submodels, which can be observed in the source code. Random forest submodels are defined at a high-level by the molecular fingerprint or representation that they use. BaseTorchGeometricModel is an implementation of the graph neural network in Brossard et al. (2021)⁴. OlorenVec is a custom supervised molecular representation developed and trained by Oloren AI, available to use in OCE.

Results

| Task | Model Name | Metric | Score | Stdev. | Leaderboard Rank |
|----------------------|----------------------------------|------------------------|-------|--------|------------------|
| НІА | RFStacker jTNhN7U7 | ROC-AUC | 0.988 | 0.002 | 1 |
| Pgp | BaseBoosting xXOn1QFI | ROC-AUC | 0.946 | 0.001 | 1 |
| BBB | BaseBoosting ADkCCrwJ | ROC-AUC | 0.923 | 0.002 | 1 |
| AMES | BaseBoosting 1zpI0dIb | ROC-AUC | 0.865 | 0.002 | 1 |
| DILI | BaseBoosting ADkCCrwJ | ROC-AUC | 0.926 | 0.005 | 1 |
| CYP3A4 Substrate | BaseBoosting ADkCCrwJ | ROC-AUC | 0.679 | 0.033 | 1 |
| Caco2 | ZWK_XGBoost Model u3zq9AAV | Mean Absolute Error | 0.289 | 0.011 | 1 |
| CYP2C9 Inhibition | BaseBoosting 1zpI0dIb | Average Precision | 0.791 | 0.005 | 1 |
| CYP2D6 Inhibition | BaseBoosting ktIIq91G | Average Precision | 0.721 | 0.001 | 1 |

| CYP3A4 Inhibition | BaseBoosting ktIIq91G | Average Precision | 0.882 | 0.001 | 1 |
|-------------------------|--------------------------|------------------------|-------|-------|---|
| CYP2D6 Substrate | BaseBoosting OQRAYLPP | Average Precision | 0.711 | 0.006 | 1 |
| Clearance Microsome | RFStacker jTNhN7U7 | Spearman | 0.625 | 0.002 | 1 |
| Half Life | BaseBoosting ktIIq91G | Spearman | 0.416 | 0.009 | 1 |
| Clearance Hepatocyte | BaseBoosting xXOn1QFI | Spearman | 0.491 | 0.006 | 1 |
| Bioavailability | BaseBoosting 1zpI0dIb | ROC-AUC | 0.736 | 0.004 | 2 |
| hERG | BaseBoosting ADkCCrwJ | ROC-AUC | 0.871 | 0.003 | 2 |
| CYP2C9 Substrate | BaseBoosting ZSCWPcLT | Average Precision | 0.419 | 0.007 | 2 |
| PPBR | BaseBoosting ktIIq91G | Mean Absolute Error | 8.503 | 0.023 | 2 |
| LD50 | BaseBoosting ktIIq91G | Mean Absolute Error | 0.613 | 0.001 | 3 |

Source Code

OCE is open-source and accessible at https://github.com/Oloren-AI/olorenchemengine. Code to define, train, and use these models is available publicly at https://github.com/Oloren-AI/OCE-TDC.

References

[1] Oloren AI, Oloren ChemEngine, (2022), Github repository,

https://github.com/Oloren-AI/olorenchemengine

[2] Huang, Kexin, et al. Therapeutics Data Commons: Machine Learning Datasets and Tasks for

Drug Discovery and Development. arXiv, 28 Aug. 2021. arXiv.org,

http://arxiv.org/abs/2102.09548.

- [3] Hu, Weihua, et al. Strategies for Pre-Training Graph Neural Networks. arXiv, 18 Feb. 2020. arXiv.org, https://doi.org/10.48550/arXiv.1905.12265.
- [4] Brossard, Rémy, et al. Graph Convolutions That Can Finally Model Local Structure. arXiv, 3 June 2021. arXiv.org, https://doi.org/10.48550/arXiv.2011.15069.