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Trade off predictivity and explainability for ML-powered predictive toxicology: an in-depth investigation with Tox21 datasets

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Abstract

Selecting a model in predictive toxicology often involves a trade-off between prediction performance and explainability: should we sacrifice the model performance to gain explainability, or vice versa? Here we present a comprehensive study to assess algorithm and feature influences on model performance in chemical toxicity research. We conducted over 5000 models for a Tox21 bioassay dataset of 65 assays and ~7600 compounds. Seven molecular representations as features and twelve modeling approaches varying in complexity and explainability were employed to systematically investigate the impact of various factors on model performance and explainability. We demonstrated that endpoints dictated a model's performance, regardless of the chosen modeling approach including deep learning and chemical features. Overall, more complex models such as (LS-)SVM and Random Forest performed marginally better than simpler models such as linear regression and KNN in the presented Tox21 data analysis. Since a simpler model with acceptable performance often also is easy to interpret for the Tox21 dataset, it clearly was the preferred choice due to its better explainability. Given that each dataset had its own error structure both for dependent and independent variables, we strongly recommend that it is important to conduct a systematic study with a broad range of model complexity and feature explainability to identify model balancing its predictivity and explainability.

Graphical Abstract

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Keywords

Tox21 bioassay; QSAR model; machine learning; predictive toxicology; explainable artificial intelligence (AI); explainability; interpretability

Background

Artificial Intelligence (AI) has been playing an increasingly vital role in a broad range of scientific research and applications, including clinical diagnosis/prognosis, natural language processing, speech and face recognition, and machine translation. Recent development of neural networks, commonly known as Deep Learning (DL), have further speeded up development of AI by taking advantage of Big Data and increased computational power. Highlighted as one trigger event in 2012, the award-winning DL model (AlexNet) held a top-5 error rate of 15.3% in the ImageNet Large Scale Visual Recognition Challenge (ILSVRC), demonstrating a significant improvement over the second-best model's top-5 error rate of 26.2%¹. Since then, complex modeling algorithms such as DL have gained wide acceptance, leading to better model performance, especially in Big Data analysis.

In predictive toxicology, AI and Machine Learning (ML) also has been widely investigated for chemical risk assessment and drug safety evaluation. In the past decades, our group developed numerous predictive toxicology approaches and tools in this area, particularly for drug-induced liver injury (DILI)^{2–6} and toxicogenomics^{7–9}. The combination of high throughput screening and ML has also become an important direction in predictive toxicology^{10–12}. For instance, the Tox21 project has screened over 10000 chemical compounds via robotic automated high-throughput in vitro assays to measure corresponding bioactivities, an unprecedented achievement which provided millions of chemical bioactivity profiles and data points.^{10, 13}

One of the key ML applications in predictive toxicology is to predict chemical bioactivities, including toxicity with molecule structure. Traditionally known as QSARs (quantitative structure activity relationships), this field has seen significant advancements with modern machine learning approaches, such as Support Vector Machine (SVM), Random Forest, and recently DL^{9, 14, 15}. For instance, several DL approaches have been developed recently with QSAR studies^{16–19}, most of which reported improved prediction accuracy for different tasks. Along with improved performance, another advantage of some DL approaches is their innate ability to work with molecular representation as SMILES, chemical graphs or images and thus bypassing the manual feature selection process.^{20, 21} While on one side this may remove a bias of a researcher to one or another type of descriptors it may also result in DL models which are more opaque since the reasoning for the model decisions is buried amid millions of neural network weights.

The problem of model interpretations is actively pursued in chemoinfomatics²² where it is going beyond the traditional QSAR and is highly relevant in other fields of science²³. A prerequisite for a trustworthy model is that its performance can be explained. Explainability can be defined as an AI behavior that can be understood and accepted by humans, which involves many concepts and aspects such as transparency, interpretability, causality, transferability, accessibility, etc.²⁴ This is a topic of active research in explainable Artificial Intelligence and some promising development in this area were recently reported elsewhere^{23, 25}. Still some methods, such as a linear regression, k-nearest neighbors (kNN) or decision trees, which are used by researches since many years, are considered as more interpretable since, e.g., the weight of features in regression could be interpreted as its importance to the decision making. In predictive toxicology, the driving features are descriptors of a substance's biological and chemical properties. Since human experts could explain influence of physiochemical descriptors, a predictive model developed with them generally could be more easily interpretable than the one using more complex features such as hashed fingerprints, graphical/geometric depictions and/or ML-derived molecular representations. With that said, the selection of chemical features and the complexity of modeling algorithm are currently the key factors to determine the explainability in predictive toxicological research.

The choice of modeling algorithm frequently involves a perceived trade-off between predictivity and explainability. In other words, increasing predictivity sometimes could lead to lower explainability; the reverse also could be true. The challenge is how we can balance predictivity and explainability to achieve a trustworthy model. To achieve that, we first need to understand how much the selection of model algorithms, as well as chemical features, would impact predictivity. In this study, we report a case study using the Tox21 bioassay activity dataset.^{26, 27} We mostly focus on the transparency and interpretability versus prediction performance of analyzed methods for Tox21 endpoints, which is attributed to the types of the modeling algorithms and the descriptors. We provide a broad view of ML applications in predictive toxicology by systematically investigating the influence of assay endpoints (68 assays), modeling algorithms (twelve ML and deep learning algorithms), and features (seven chemical representations).

Methods and Materials

Tox21 Dataset

Tox21 bioassay activity data (the Tox21 Dataset¹³) were collected for 68 bioassay endpoints²⁷ and 8948 compounds, of which each was tested in at least one assay. Chemical bioactivity data was preprocessed and categorized into four major classes: active agonist, active antagonist, inactive and inconclusive²⁸. In this study, both active agonists and active antagonists were considered positive compounds, and inactives were considered negative. Only one active category was considered in each assay; i.e., if one assay contained more active agonists than active antagonists, we used the active agonists as positive in the analysis. Inconclusive compounds were excluded from all assays/endpoints. Chemicals were further deduplicated based on their InChI keys. The final number of positive and negative samples in each assay was summarized in Supplementary Table S1 and all processed data are available as Supplementary Table S2.

In addition, the chemical similarity of a Tox21 assay endpoint was calculated using a withingroup chemical similarity (S) score. S represented the chemical diversity in accordance with the endpoint, which was measured according to formula (1). In the formula, *m* and *n* represent the number of compounds in positive and negative classes, respectively. *Jaccard_{ij}* indicates the Jaccard similarity coefficient (index) between compounds *i* and *j*, which was calculated based on RDkit fingerprints. As formulated, higher S value corresponds to endpoint with compounds that are more similar within the class of actives.

$$S = Mean(\sum_{i=1}^{m} \sum_{j=1}^{m} Jaccard_{ij}) - Mean(\sum_{i=1}^{m} \sum_{k=1}^{n} Jaccard_{ik})$$
(1)

Predictive modeling algorithms

In total, 12 modeling algorithms were included in this study; These modeling algorithms can be categorized into four classes: (1) neural networks; (2) decision trees ensemble methods; (3) SVMs; and (4) simple algorithms. For neural networks, we deployed a 3-Layer neural-network (MLP-3), 7-Layer neural-network (DNN¹⁹), Associative Neural Network (ASNN-MTL)²⁹, and a Multi-task Learning version of the 7-Layer Neural-network (DNN-MTL)¹⁹. Algorithm ending with MTL were performed with multi-task learning framework. Four decision trees ensemble methods, XGBoost³⁰, AdaBoost, GradientBoosting, and Random Forest, were used. For SVMs, we applied rbf SVM (SVM)^{31, 32} and Least-Squares SVM (LS-SVM)³³ optimized for GPU-based computing. Lastly, Linear Regression and KNN were used as for simple algorithms³². These modeling algorithms provided broad coverage of modeling complexity while representing popular modeling algorithms in the field. Linear regression and KNN are simple methods that are easier to understand, while SVMs and neural networks are relatively more complicated and difficult to interpret. The use of ensembles for decision trees despite it added more complexity was also shown to significantly improve their performances^{30, 34}. Deep neural-networks are one of the recent popular modeling algorithms and often have good predictivity compared to other algorithms especially in studies dealing with large data^{14, 21, 35}.

Among the 12 studied algorithms, XGBOOST, LS-SVM, DNN, ASNN-MTL and DNN-MTL were performed in OCHEM platform (http://ochem.eu), which has LS-SVM and DNN(-MTL) implemented using GPU.³⁶ The other seven modeling algorithms were performed on local computing cluster at FDA. As designed, the training and testing compounds for two experiment sites were exactly the same, but we applied slightly different data-preprocessing strategy on each site. For example, correlation filters (>0.95) was used in OCHEM platform for feature reduction during data pre-processing. We checked the influence of the preprocessing on the performance of the Random Forest, which was calculated with the same setting of hyperparameters in OCHEM and FDA sites but did not observe any systematic bias. The parameters used in these modeling algorithms, if not specified, were set to defaults, which were optimized in multiple studies performed by their authors. The details of modeling algorithms used in this study are summarized in Table 1.

Chemical Features

We evaluated seven different types of chemical features to represent the chemical structures, five chemical fingerprints (i.e., RDKit³⁷, ECFP4, FCFP4, Extended Functional Groups (EFG)³⁸ and ToxPrint³⁹) and two QSAR descriptors (i.e., MordRed⁴⁰ and Mold2⁴¹). RDKit fingerprint was developed by RDKit³⁷ and was calculated with default parameters (nbits=2048). Extended-Connectivity Fingerprints (ECFP4 and FCFP4) are atom-based and feature-based chemical fingerprints, both of which were calculated by using Morgan fingerprints generated by the RDKit, with radius=2 and bit length 1024. ToxPrints are based on the publicly available ToxPrint chemotypes (v2.0_r711, https://toxprint.org/) generated within the associated ChemoTyper application (https://chemotyper.org/). EFG is an extension of a functional group set previously implemented by the CheckMol⁴² that also covers heterocyclic compound classes and periodic table groups³⁸. ToxPrint chemotypes consist of 729 uniquely defined chemical features coded in XML based Chemical Subgraphs and Reactions Markup Language (CSRML). The numbers of features generated for each type of descriptors are listed in Table 2.

Results

Study workflow

The overall study design is depicted in Figure 1a. We used all publicly available assays from the Tox21 dataset¹³ to take advantage of the diversity in assayed endpoints (i.e., 68 bioactivity endpoints). In total, 8948 compounds were profiled by at least one assay. After data pre-processing to remove chemical duplicates, 7660 compounds were retained for the analyses. These compounds were split on training and test set comprising 7000 and 660 compounds, respectively (see Supplementary Table S2). We conducted binary classification and for each assay only active and inactive compounds were considered. The active compounds could either be active agonists or active antagonists, depending on the majority group of the assay (see Materials and Methods for details). For duplicated compounds, only compound with consistent bioactivities in the same assay were considered: others were labeled as inconclusive and discarded. Finally, we removed three assays that did not have enough active compounds (<=20), therefore 65 Tox21 assays remained for

modeling analysis. The final processed-ready dataset is available for download from http://ochem.eu and is also included as Supplementary Table S2.

For each endpoint, we developed 84 models with exhaustive combination of seven types of molecular features in conjunction with 12 modeling algorithms. Specifically, we used RDkit, ECFP4, FCFP4, EFG, ToxPrint, Mold2 and MordRed to measure/represent different types of compound fingerprints or QSAR descriptors. Four major categories of 12 modeling algorithms were used to represent the varying degrees of complexity in modeling algorithms.

Figure 1b shows the general modeling pipeline with a single experiment (i.e., one feature set and one modeling algorithm applied to data from a single assay), in which a model pairing a modeling method with chemical features was evaluated by 5-fold cross-validation. During the cross-validation, we split the training dataset into 5 folds, where 4 folds were used for training and the other fold was used for validation. Next, the models were tested on the hold-out samples from the test set, which contains 660 unseen compounds. Final model performance thereafter was measured by the average AUC of the training set as well as the AUC of testing set.

Model predictive performance

Overall results on the testing data are shown in Figure 2a. As shown, each cell in this hierarchical clustering map (HCA) is the average testing AUC from the nested cross validation results in one model. The x-axis contains 65 Tox21 assays. The HCA map contains 84 rows, which represented all combinations of seven feature sets and 12 modeling algorithms.

First, we found that these 65 Tox21 assay endpoints showed very different performance patterns. Some endpoints always had a high AUC regardless of the type of feature or modeling algorithm used. Contrary to that some endpoints showed a consistently low AUC across all feature-algorithm combinations. With respect of their performance similarity models using the same chemical features tended to cluster together.

The overall predicting performance for each Tox21 assay was summarized in the box-plot representation (Figure 2b). Each bar represented a collapsed result of one particular modeling algorithm, by combining all features and endpoints. Training (i.e., Cross-validation) and testing (i.e., hold-out) results were presented in blue and orange boxes, respectively. As shown, there was no significant performance gap between Training and Testing results, indicating that the developed models were robust. Detailed model performances (AUC) for each Tox21 assay were summarized in Table S1.

Overall performance of model algorithms and features

Further analysis was applied to inspect the overall influence of chemical features and modeling algorithms, by averaging the results across all 65 Tox21 endpoints. The 5-fold CV and the hold-out testing results were summarized in Table 3.

Regarding to the feature types, the best feature type was Mold2 (AUC-CV=0.82; AUC-test=0.84). The weakest feature type was ToxPrint (AUC-CV=0.76; AUC-test=0.78). In

general, QSAR descriptors (i.e., Mold2 and MordRed) performed better than chemical fingerprints. We also observed that EFG outperformed ToxPrint, both of which were kinds of structural alerts or functional groups.

On the other hand of modeling algorithms, we found GBoost, RF and SVMs showed better predicting performance than other categories of algorithms, and the best modeling algorithms among 12 we tested was RF (AUC-CV=0.84; AUC-test=0.84); the relatively weakest modeling algorithm was KNN (AUC-CV=0.73; AUC-test=0.75). In addition, we did not observe better performance of more complicated models such as the neural networks, compared to simple models of Linear Regression, which provided similar results on average. In addition, XGBOOST did not improve the model predictivity compared to GBoost. We also observed that Multi-task learning framework may improve the modeling predictivity since DNN-MTL outperformed DNN thus confirming results of other studies.

For the single pair of model-feature combination, LS-SVM with MordRed held the best performance in average 65 Tox21 endpoints prediction (AUC-CV=0.87; AUC-test=0.88). Note that the predictivity differences among top combinations were marginal; such as LS-SVM-Mold2, LS-SVM-MordRed, RF-Mold2 and RF-MordRed, all of which held AUC around 0.88 in hold-out testing result.

Parameter influence on predictivity of SVM and RF

Based on the overall prediction results, SVM and Random Forest (RF) showed a good predictivity among all feature types and endpoints we tested. Both RF and SVM have hyper-parameters, which may have significant impacts on the model performance. In aims to understand how much the hyper-parameter tuning could affect the predictivity of both SVM and RF, we performed a grid search analysis to fine-tune "cost parameter" (C) and gamma (g) for SVM (with default RBF kernel) and "number of trees" (n) and "minimal number of samples in leaf node" (L) for RF, respectively.

As shown in Figure 3a, the hyper-parameters showed a large impact on SVM models. As a "proper" set of hyper-parameters (e.g., C=10 and g = "*scale*")⁴³ would achieve over 0.8 AUC across all types of features and endpoints where an "improper" set of hyper-parameters (e.g., C=0.1 and g=1) would fail completely (Averaged AUC<0.5). On the other hand, we found that the influence of hyper-parameters on RF models was much smaller in comparison to SVM models (Figure 3b), as the "worst" set of hyper-parameters (e.g., n=10; L=20) still could get an averaged AUC around 0.75. Taken together, our findings demonstrated that hyper-parameter selection can have a larger impact on some modeling algorithms such as SVM, while less affects other modeling algorithms such as RF.

Influence of sample size and chemical similarity on endpoint predictability

Figure 4 shows an assessment of the other two important innate properties of a dataset that affect model performance, sample size and chemical similarity. Due to data availability, sample size varied from endpoint to endpoint, and the positive sample size (i.e., number of active agonist or antagonist in the dataset) was usually much smaller than negative sample size. A smaller positive sample leads to more imbalanced dataset, which could affect the model performance. We therefore analyzed the correlation between model performance and

positive sample ratio for the analyzed endpoint (Figure 4, light gray bars). As shown, we found that the positive sample size had little effect on the averaged model performance. However the sample size influenced the variation of the performance and we observed wider boxes and quantiles in assays with smaller positive sample sizes. In addition, lower performed assays also tended to have larger variations of performance, which indicated the modeling algorithm and feature selection could more affect these low-performed and less sample size endpoints.

With respect to the chemical similarity of endpoints, the S scores were negative for most of the Tox21 assays, indicating that for most assays, compounds within the respective positive/ active class shared low structural similarity (Figure 4, Red/Green/Black bars). Meanwhile, we found that the S scores for eight of ten endpoints with high predictability were positive, implying that higher structural similarity within the positive/active compound class may contribute to the higher predictive performance of the models for these assays. However, endpoints with negative S score did not necessarily result in poor performance; for example, although assays of tox21-ahr-p1 and tox21-pr-bla-antagonist (green bars in Figure 4) had negative S score –0.018 and –0.037 respectively, they still had high AUC (>0.85) in both training and testing results.

Discussion

We conducted a systematic investigation on the choice of modeling approaches (12 ML/DL algorithms used) and chemical features (seven molecular representations) in predictive toxicology, with a specific focus on comparative analysis of model performance and explainability and the trade-off between them. Results demonstrated that the assay or endpoint itself was the largest determining factor for model performance; a good performance was reached for a predictable endpoint (high predictability) regardless of the choice of modeling algorithm or feature type used⁴⁴. As implied in Figure 4, assays with lower performance tended to have larger variations. The influence of modeling algorithm and features was more pronounced for such endpoints as well as for those with more imbalanced dataset (i.e., smaller positive sample sizes). For such endpoints it could be important to conduct a systematic analysis by using a broad range of approaches of various model complexity and feature explainability. For the Tox21 datasets studied here, we found that using simple modeling approaches such as Linear Regression in a number of cases provided models with similar performance to those of more complex approaches, such as neural networks. Such models could be more preferable in the context of the computational toxicology due to better balancing of their predictivity and interpretability. Of course, the reported in this study results could be influenced by type of the data and used descriptors, but the use of simple baseline models should not be ignored.

Not all datasets have the same complexity, which further emphasizes the need for evaluating a broad range of modeling approaches and molecular representations. Three innate data properties are of special importance: endpoint predictability, data imbalance, and size. With respect to endpoint predictability, we only examined the chemical structure similarity within and across class labels without considering the quality of the endpoints measurements

themselves. The results implied that a high chemical structure-driven predictability likely resulted in a good performance, but the reverse was not entirely apparent.

An algorithm with a simple architecture, such as linear regression may not be the most powerful, but it will be easily explainable when using interpretable descriptors. Algorithms with more complicated architectures, such as (LS-)SVM or (deep) neural networks, may have better statistical performance but can be more difficult to interpret. In this study, we did observe that some more complex modeling algorithms had overall better prediction accuracy as compared to simple algorithms such as linear regression and KNN. However, the influence of a modeling algorithm was not as significant as the nature of the endpoint itself and in a number of cases simpler and easier to interpret models with similar performances were obtained. Selecting a complex but less explainable model could hinder its use due to possible concerns with interpretation of its predictions for new data.

Therefore, when considering both predictivity and explainability in the context of chemical feature based Tox21 data analysis, we recommend do not overlook using simple models: they provide higher explainability while still can have similar performance as some more complicated approaches. Another strategy is to increase the explainability of complex models via model-agnostic approaches⁴⁵. With more such approaches being developed^{46–48} and with their wider availability, complex models such as DL networks may hold great potential for improved explainability. In their absence an interpreting a complex model is much more difficult than interpreting a simple model such as linear regression. Given that each dataset has its own error structure both for dependent and independent variables, we strongly recommend that it is important to conduct a systematic study with a broad range of method complexity and feature explainability to select a model, which balance its predictivity and explainability

Note that in this study we only considered chemical feature-based models, while recently we observed many deep learning models now directly analyse chemical structure such as SMILE string, InChI key, 3D image as the model input^{49, 50}. Directly using chemical structure instead of features may be a game changer to the predictive toxicology, just like the current image analysis nowadays will directly use the raw image rather than human-engineered features. The investigation and comparison between feature-based and feature-free models also need to be comprehensively performed. In addition, consensus modeling could also be an effective way to improve model predictivity⁵¹. Evaluation the explainability of consensus model is also a challenge and the objective of future studies. While we plan to perform such studies ourselves we also encourage the other researchers to analyze the data of this study, which contain nearly 440k measurements for 65 properties (Supplementary table S2) in order to propose and benchmark approaches balancing predictivity and explainability of models.

Future directions can also include evaluating metrics to qualitatively or quantitatively measure the interpretability on demand, in order to investigate how much interpretability could be gained by using different approaches and whether they contribute the same interpretations. In addition, multi-task learning was proved to be an efficient way to improve the model performance by sharing the modeling architectures among different

predictive endpoints^{14, 17, 19}. We also observed the same tendency in this study but a more comprehensive investigation on the effects of multi-task learning compared to single-task learning would be critical to better elucidate impact of these methods on the predictive toxicology.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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Figure 1.

The overview of the workflow used to analyze the data. (a) Overall study design. (b) Construct and evaluate predictive model with selected predictor, modeling algorithm and endpoint.

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Figure 2.

(a) Heatmap of all Tox21 assays across different modeling algorithms and features. (b) Overall, collapsed performance for all Tox21 assays. The validation and testing results are consistent across all endpoints.

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Figure 4.

Influences of positive sample ratio (light-gray bars) and sample within-group similarity (green/red/black bars) on endpoint performance (blue box plot). Green and red bars are the top ten highest performed assays with positive and negative S score, respectively.

Table 1.

Brief summary of modeling algorithms used in this study.

SYNONYM	MODELING ALGORITHM	CATEGORY	PARAMETERS						
LINEAR	Linear regression	Simple Model							
KNN	K-Nearest Neighbors	Simple Model	K=5 (default)						
RF	Random Forest	Ensemble Model	n=100 (default)						
ABOOST	Adaptive Boosting	Ensemble Model	n=50 (default)						
GBOOST	Gradient Boosting	Ensemble Model	n=100 (default)						
XGBOOST	Extreme Gradient Boosting	Ensemble Model							
SVM	Support Vector Machine	SVMs	Kernel=rbf; C=100; gamma='scale'						
LS-SVM	Least-Squares SVM	SVMs							
MLP3	3 layers Multi-layer Perceptron	Neural Networks	nodes=[32, 64, 32];						
DNN	7 layers Neuro Networks	Neural Networks	nodes=[512:256:128:64:32:16]						
DNN-MTL	Multi-Task Learning of DNN	Neural Networks	Same as above						
ASNN-MTL	Associative neural network	Neural Networks	Ensemble of 64 models with one hidden layer containing 3 neurons						

Table 2.

Brief summary of feature generation tools used in this study.

SYNONYM	#FEATURES	CATEGORY	REFERENCE				
RDKIT	2048	Chemical fingerprints	https://www.rdkit.org				
ECFP4	1024	Chemical fingerprints	https://www.rdkit.org				
FCFP4	1024	Chemical fingerprints	https://www.rdkit.org				
EFG	583	Chemical fingerprints	Citation ³⁸				
TOXPRINT	729	Chemical fingerprints	https://toxprint.org, 39				
MORDRED	1825	QSAR descriptors	Citation ⁴⁰				
MOLD2	777	QSAR descriptors	Citation ⁴¹				

Table 3.

Averaged performance of seven feature types and twelve modeling algorithms.

Model/ Feature	RDKIT		ECFP4		FCFP4		EFG		TOXPRINT		MOLD2		MORDRED		Average	
	5- fold CV	Test	5- fold CV	Test	5- fold CV	Test										
DNN-MTL	0.79	0.83	0.79	0.80	0.79	0.76	0.86	0.81	0.72	0.77	0.82	0.84	0.81	0.84	0.8	0.81
DNN	0.78	0.81	0.77	0.79	0.78	0.79	0.82	0.77	0.71	0.70	0.78	0.81	0.77	0.81	0.77	0.78
ASNN- MTL	0.77	0.76	0.80	0.84	0.79	0.84	0.84	0.81	0.73	0.79	0.80	0.81	0.66	0.67	0.77	0.79
MLP3	0.76	0.80	0.75	0.78	0.75	0.77	0.77	0.75	0.76	0.80	0.83	0.83	0.83	0.85	0.78	0.8
XGBOOST	0.79	0.81	0.76	0.77	0.75	0.77	0.80	0.78	0.67	0.69	0.81	0.85	0.81	0.85	0.77	0.79
GBOOST	0.83	0.85	0.81	0.83	0.80	0.81	0.83	0.83	0.81	0.83	0.85	0.87	0.84	0.87	0.83	0.84
ABOOST	0.77	0.79	0.76	0.78	0.77	0.78	0.80	0.79	0.78	0.81	0.82	0.83	0.89	0.84	0.79	0.8
RF	0.84	0.84	0.83	0.83	0.82	0.81	0.86	0.83	0.82	0.81	0.86	0.88	0.86	0.88	0.84*	0.84*
LS-SVM	0.84	0.86	0.82	0.83	0.81	0.84	0.84	0.82	0.73	0.74	0.86	0.88	0.87*	0.88*	0.83	0.83
SVM	0.84	0.85	0.83	0.84	0.81	0.82	0.81	0.80	0.81	0.81	0.85	0.87	0.85	0.88	0.83	0.84
KNN	0.73	0.77	0.73	0.74	0.72	0.73	0.71	0.70	0.72	0.75	0.76	0.79	0.77	0.80	0.73	0.75
Linear	0.79	0.81	0.81	0.82	0.81	0.82	0.81	0.78	0.81	0.83	0.82	0.83	0.82	0.85	0.81	0.82
Average	0.79	0.81	0.79	0.80	0.78	0.80	0.81	0.79	0.76	0.78	0.82*	0.84*	0.81	0.84		

 * Stars indicate the best performing descriptor sets and algorithms.