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Conceptual Analysis

Chemistry-centric explanation of machine learning models

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Artificial intelligence (AI) is increasingly being considered across chemical disciplines, just as in many other areas of science, often with high expectations for “revolutionary” advances. Medicinal chemistry [and drug design are among the focal points of these developments [1– 4] and there is rising interest in deep neural network (DNN) architec-](#_bookmark16) tures and deep learning (DL) for the generation of novel compounds and prediction of various molecular properties [[3](#_bookmark21),[4](#_bookmark22)].

In chemoinformatics, medicinal chemistry, and drug design, the term AI is typically used synonymously with machine learning (ML), which represents only a part of the methodological AI spectrum [[5]](#_bookmark24). However, ML already has a long history in chemoinformatics and medicinal chem- istry, spanning more than two decades, and is widely applied for molecu- lar property predictions as well as the search for novel active compounds [[6]](#_bookmark26). Neural networks (NNs) were popular early on in chemoinformatics, but have been increasingly replaced over time with other approaches

[[7]](#_bookmark27) such as the support vector machine (SVM) [[8]](#_bookmark29) and random forest (RF) [[9]](#_bookmark32) algorithms or Bayesian modeling [[10]](#_bookmark33), mostly due to the in- trinsic tendency of NNs to overfit property prediction models trained on moderately sized data sets. With DL, NNs have re-emerged in this field, for the most part as DNN architectures [[11]](#_bookmark35).

In computer science, it is often observed that methodological com- plexity does not scale with predictive performance [[12]](#_bookmark37) and the same applies to medicinal chemistry and drug design [[6]](#_bookmark26). Although “big data” trends are also beginning to emerge in medicinal chemistry [[13]](#_bookmark38), ML predictions are typically based upon relatively small data regimes and well-defined molecular representations. These conditions do not play into the strengths of DNNs compared to other areas such as image anal- ysis or natural language processing where DL has made a large impact in recent years [[14](#_bookmark40),[15](#_bookmark42)]. Accordingly, standard ML methods often perform comparably well or better than DNNs in predicting biological activity and other molecular properties [[6]](#_bookmark26).

On the other hand, DL offers new opportunities in chemoinformat- ics and medicinal chemistry that were hardly possible to address in a comparable way until recently such as, for example, in chemical re- action modeling [[1]](#_bookmark16) or large-scale generative *de novo* compound de- sign [[3]](#_bookmark21). In generative design and DNN-based compound reposition- ing, successful applications charting new territory are beginning to ap-

pear [[16]](#_bookmark43) but other claimed advances remain often questionable [[17]](#_bookmark45), partly due to the lack of generally accepted evaluation criteria and stan- dards for assessing chemical novelty [[17]](#_bookmark45). Even in successful design ap- plications, newly generated or repurposed chemical entities are often viewed controversially from a medicinal chemistry perspective. Clearly, the field is in flux but ML/DL in medicinal chemistry and drug design is still largely dominated by methodological considerations (what could be done?) rather than successful practical applications (what has been done?), which are still far from being routine [[18]](#_bookmark48).

One of two particularly relevant aspects concerning ML in medicinal chemistry emerging from the discussion above is that a variety of meth- ods are being employed for property prediction and compound design, ranging from simple decision tree-based algorithms and probabilistic modeling to complex DNNs, often with comparable success. The other important aspect relates to the rationalization of predictions, as further discussed in the following.

Given the often cited “black box” character of most ML models [[12](#_bookmark37),[19](#_bookmark49)], their predictions are diﬃcult to rationalize. In the practice of medicinal chemistry, lack of transparency of predictions and model rationalization continue to hinder the acceptance of ML and limit the impact of predictive modeling on experimental programs [[6](#_bookmark26),[18](#_bookmark48)], de- spite the long tradition of ML in this field. Similar considerations ap- ply to other chemical disciplines that are primarily experimentally driven. Ideally, one would like to know how certain and reliable a given prediction is and also understand it in intuitive chemical terms. While a few studies have presented approaches for uncertainty es- timation of ML predictions [[20](#_bookmark8),[21](#_bookmark9)], robust and generally applicable methods are currently not yet available. However, in computer sci- ence and other fields, the potential of explainable or interpretable ML is a much discussed topic [[12](#_bookmark37),[22](#_bookmark10),[23](#_bookmark11)], which also is of high rel- evance for ML applications in chemistry. In computer science, inter- pretable ML refers to algorithms whose predictions can be directly reconciled (such as decision trees) and explainable ML to methods enabling the rationalization of black box models [[12]](#_bookmark37). However, in chemistry-related publications, these terms have been used more or less interchangeably concerning the rationalization of models and their decisions.

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In medicinal chemistry, there is an urgent need for ML model ra- tionalization. Although the reluctance of medicinal chemists to rely on black box predictions in their compound design and optimization efforts is a widespread phenomenon, only a limited number of investigations have addressed the explanation or interpretation of ML predictions in the context of structure-activity relationship (SAR) analysis [[24]](#_bookmark12). With increasing use of complex DNN architectures in the field [[3](#_bookmark21),[4](#_bookmark22)], this widens the gap between model availability and acceptance and further limits the impact of ML.

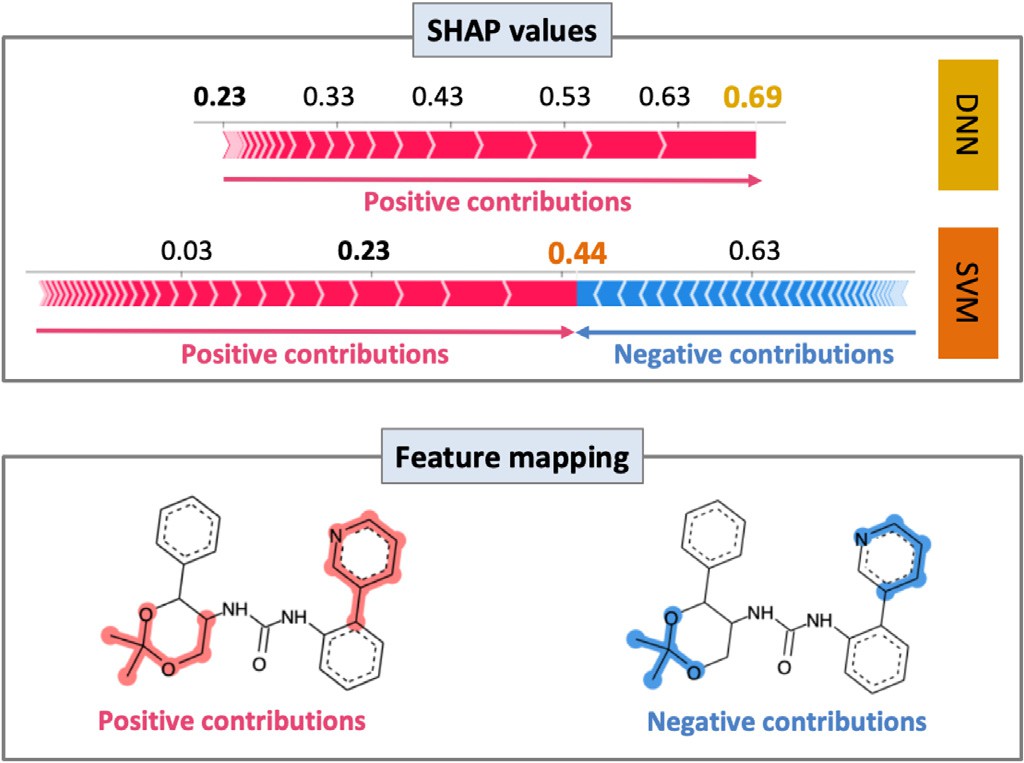
For activity prediction models, available rationalization strategies typically aim to identify molecular representation features that de- termine individual predictions, mostly in a model-dependent manner. These approaches predominantly employ feature weighting techniques to rationalize predictions of kernel-based [[25](#_bookmark13),[26](#_bookmark14)] or Bayesian [[27]](#_bookmark15) clas- sification models. In addition, weight gradients across different NN lay- ers can also be determined to trace determinants of predictions and ex- plain (D)NN models. However, these gradients tend to be unstable, fre- quently resulting in different interpretations of very similar predictions [[28]](#_bookmark17).

In addition to model-dependent techniques, model-independent methods can also be considered, which are principally preferred be- cause they are applicable to any ML model, regardless of the com- plexity of the underlying algorithm. Moreover, model-independent ap- proaches typically do not require balancing performance and inter- pretability across different models [[29]](#_bookmark18). However, with sensitivity anal- ysis [[30]](#_bookmark19), only one model-independent approach has until recently been applied to property predictions in chemoinformatics. This methodol- ogy was adapted about a decade ago to study the influence of sys- tematic feature value changes on activity predictions using ML mod- els [[31](#_bookmark20),[32](#_bookmark23)]. Sensitivity analysis generally relies on feature perturbation to study ensuing effects. In particular, partial derivatives were used to assess the impact of local perturbations of fragment descriptors on model predictions [[32]](#_bookmark23). Sensitivity analysis becomes rapidly infeasi- ble with increasing model dimensionality, which limits its applicability. Hence, this methodology has been more or less abandoned in the field or substituted by approximations that better summarize perturbations effects.

Our group has considered alternative approaches with potential for ML model-independent rationalization from different viewpoints includ- ing broad applicability, quantitative assessment of predictions, and ease of visual interpretation. In light of these criteria, the concept of *Shap- ley values* [[33](#_bookmark25),[34](#_bookmark28)] from cooperative game theory [[34](#_bookmark28),[35](#_bookmark30)] has become our methodology of choice. Shapley values were originally introduced in 1953 [[33]](#_bookmark25) to quantitatively account for contributions of individual players forming a team. These values provide a quantitative assessment of cooperative contributions to a team’s ultimate success (total gain). Accordingly, they specify a partition of merit among individual players by calculating the average of all contributions made by a given player in different team constellations.

The Shapley value concept is readily transferable to ML by applying the following analogies: The game a team engages in can perceived as a prediction task for a single instance (e.g., a compound). The merit for this task is given by the difference between its prediction and the aver- age prediction of all instances. The players participating in the game are features values of the instance that cooperate (act in concert) to obtain the merit for a given prediction. The resulting Shapley value of a given feature is then obtained as the average contribution of a feature over all possible feature combinations. Accordingly, Shapley values account for the partition of contributions over individual features comprising a feature vector or set (such as a molecular representation). A key aspect of the Shapley value concept that sets it apart from model-dependent feature weighting methods is that not only the contribution of feature presence to a given prediction can be quantified, but also the contribu- tion of feature absence.

For feature sets of increasing size, systematic calculations of Shapley values on the basis of all possible feature combinations become com-



**Fig. 1.** Interpretation of predictions. SHAP feature importance values are rep- resented as sequential arrows recording positive (red) and negative (blue) con- tributions to the prediction of activity, resulting in a cumulative output proba- bility. For ML, the given compound was represented using a topological finger- print comprising overlapping atom environment features. Top-ranked features making positive or negative contributions are mapped onto the structure. The activity of the compound was only correctly predicted using a DNN but not an SVM model. The figure was adapted from Rodríguez-Pérez and Bajorath [[39]](#_bookmark39). (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article).

putationally increasingly expensive. Therefore, to render the approach practical for ML, a *locally interpretable explanatory model* is derived that limits required Shapely value calculations. This methodology is termed *Shapley Additive exPlanations* (*SHAP*) [[36]](#_bookmark31) and can be perceived as an extension of the *Local Interpretable Model-agnostic Explanations* (LIME) approach [[37]](#_bookmark34). A brief account of the underlying theory is presented in the **Appendix** below.

We have adapted the SHAP methodology for explaining compound activity predictions and SARs [[38]](#_bookmark36). To these ends, SHAP was combined with molecular feature mapping and feature classification according to quantitative contributions to correct or incorrect ML model predictions

[[38]](#_bookmark36) and successfully applied to different compound activity prediction tasks including classification, regression, or multi-task learning [[38](#_bookmark36),[39](#_bookmark39)]. [Fig. 1](#_bookmark3) shows exemplary results of molecular SHAP analysis.

Shown is an active compound whose biological activity that was correctly predicted by a DNN model whereas an SVM classifier pre- dicted this compound to be inactive. The SHAP contribution plots of these predictions reveal that multiple features were assigned compa- rable importance for predictions with both models. However, the SVM model yielded negative contributions for a number of features that were not prioritized by the DNN. Mapping of highly weighted positive (red) and negative (blue) features from the SVM prediction on the compound structure revealed that these features formed overlapping substructures, thus detecting and explaining a model error.

Recently, an algorithm was introduced for the exact calculation of local SHAP values specifically with decision tree-based methods, taking advantage of the tree distribution [[40]](#_bookmark41). By comparing predictions using SHAP and this algorithm directly, high correlation of greater than 80% was observed between approximated and exactly determined local fea- ture importance values, both for decision tree-based classification and regression models [[39]](#_bookmark39). These findings further supported the reliability of SHAP’s local kernel approximation.

We also note that there are conceptually distinct yet complementary approaches to SHAP for model interpretation. For example, the *Simi- larity Maps* approach depends on the removal of individual atoms from test compounds and estimation of their importance by quantifying the

resulting change of a prediction [[41]](#_bookmark44). This is in contrast to SHAP that does not rely on atom removal (i.e., introduction of a perturbation at the structural level), but assesses collaborative (i.e., combination-based) contributions of representation features. The Similarity Maps approach does not require feature mapping but cannot quantify contributions of feature absence. In addition, single-atom perturbations might not signif- icantly modify a model’s prediction but affect structural integrity.

Another conceptually distinct approach involves feature weighting of multiple models generated with the same ML algorithm, providing the basis for the determination of *feature importance correlation* [[42]](#_bookmark46). For target-based compound activity classes, this measure yielded model- internal data set signatures and revealed similar compound binding characteristics of proteins as well as functional relationships. Strong feature importance correlation between compound activity prediction models indicated functional similarities between different targets that were not related to compound binding [[42]](#_bookmark46). For RF models derived for more than 200 targets, the Gini impurity (GI) criterion [[43]](#_bookmark47) served as a measure of node-based recursive partitioning quality. GI is a metric from information theory defined in [Eq. (1)](#_bookmark5):

GI = ∑ *𝑝𝑖* (1 − *𝑝𝑖* ) (1)

*𝑛*

*𝑖*=1

Here, *𝑝𝑖* is the frequency for class *i* at a given node, and *n* is 2 for

binary classification. Accordingly, GI for a given feature is equivalent

to the mean decrease in GI, i.e., the normalized sum of all impurity decrease values for nodes in the RF where splitting was based on that feature. Thus, increasing values indicate increasing feature importance for the RF model [[43]](#_bookmark47). Correlation or statistical association across fea- ture importance values from different models is then quantified using correlation coeﬃcients.

Feature importance correlation analysis only requires model-internal

information, but does not depend on model explanations. Feature

LIME generates the explanation *𝜉* of an instance *x* according to

[Eq. (3)](#_bookmark4):

*𝜉*(*𝑥*) = argmin G(*𝑓, 𝑔, 𝜋𝑥* ) + Ω(*𝑔*) (3)

*𝑔*∈*𝐺*

where *𝐺* is a class of interpretable (linear) models, G is the loss function to minimize, *𝜋𝑥* the proximity measure between an instance *z* and *x* (kernel defining locality), and Ω(*𝑔*) an optional regularization term to

limit model complexity.

For the explanation of a given test instance *x*, the following proce- dure is applied:

1. Artificial samples are obtained by permuting features of the test instance *x*.
2. These samples are weighted by the value of a kernel calculated for them and *x*.
3. A model *g* is trained to predict *f(x)* with coeﬃcients estimating feature importance.

Hence, LIME builds a linear model *g* in a feature region proximal to the test instance, with ML model *f* typically being non-linear. The LIME approach provides the basis for the development of the kernel SHAP methodology, as explained in the following.

Shapley values account for the distribution of feature contributions to a model’s prediction for a given test instance. To determine the contri- bution of a feature *i*, all operations by which a feature might be added to

the set (*𝑁* !) and a summation over all possible sets (*𝑆*) must be carried

ture *i* is given by [*𝑓* (*𝑆* ∪ {*𝑖*}) − *𝑓* (*𝑆*)]. The resulting quantity is weighted out. For any feature sequence, the marginal contribution by adding fea-

tion of feature *I*, i.e., ( *𝑆* !), and the order in which remaining features by the number of combinations available to form the set prior to addi- might be added, i.e., (( *𝑁* − *𝑆* − 1)!). Hence, the importance of a given

[| | |](#_bookmark6) |

| |

feature *i* is defined by [Eq. (4)](#_bookmark6):

weights or importance values can be extracted using multiple strategies,

*𝜙* = 1

∑ |*𝑆* |!(|*𝑁* | − |*𝑆* | − 1)![*𝑓* (*𝑆* ∪ {*𝑖*}) − *𝑓* (*𝑆* )] (4)

depending on the underlying ML algorithm. Moreover, in contrast to SHAP-based analysis of individual predictions, feature importance cor- relation analysis is based upon global model assessment, taking many compounds into account. While SHAP and feature importance correla- tion analysis are conceptually distinct, these methods are complemen- tary and can be combined for ML model assessment. It is anticipated that increasing efforts will be expanded in chemoinformatics to develop methods for model explanation in order to further increase the accep- tance of ML in the practice of medicinal chemistry.

*𝑖*

# Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

# Appendix A

*SHAP theory*

The principal goal of an explanation model *g* is to locally approx- imate and thereby simplify a complex model *f* that is diﬃcult to un- derstand. Additive feature attribution methods generate an explanation model via a linear function of binary variables, given by [Eq. (2)](#_bookmark7):

*𝑁* ! *𝑆⊆𝑁* ∖{*𝑖*}

It follows that Shapley values represent a unique way of dividing a model’s output among feature contributions satisfying three axioms: *lo- cal accuracy* (or additivity), *consistency* (or symmetry), and *nonexistence* (or null effect).

Additive feature attribution methods typically do not consider two properties that are of high relevance for assessing feature importance, i.e., *local accuracy* and *consistency*. The SHAP formalism was specifically devised to take these axiomatic properties into account [[36]](#_bookmark31). The prop- erty *local accuracy* ensures that the sum of individual feature attributions is equal to the original prediction because SHAP allocates the model pre- diction across contributing features. Furthermore, *consistency* ensures that feature importance correctly accounts for different models on a rel- ative scale. Hence, if a change in a feature value has larger impact on model *A* than model *B*, feature importance should be larger in *A*. These properties can be accounted for by representing feature importance as Shapley values [[36]](#_bookmark31).

A weighting procedure for artificial samples is a key aspect for con- necting Shapley values to the LIME approach. In LIME, heuristic choices

are made to select G, Ω(*𝑔*), and *𝜋𝑥* . By contrast, the SHAP method in-

troduces a special kernel function that is related to the Shapley value

definition, assuming that feature weights follow the two axioms of in- terpretability. Specifically, SHAP uses the following procedure for inter- preting an instance *x*:

*𝑔*(*𝑥*′)

= *𝜙*0 +

*𝑀*

*𝑖*=1

∑

*𝜙𝑖 𝑥*′ (2)

1. Training data are organized by *k*-means clustering and the *k* sam- ples are weighted by the number of training instances they rep- resent. These samples constitute a background data set of given

where *𝑥*′ ∈ {0*,* 1}*𝑀* , *M* is the number of input features, and *𝜙𝑖* ∈ ℝ. The

*𝑖*

sents a feature contribution (*𝜙𝑖* ). Accordingly, a weight must be assigned presence or absence of a feature value impacting the prediction repre-

to each variable for which the LIME methodology [[37]](#_bookmark34) can be applied and further extended.

feature values.

1. Artificial samples are obtained by replacing features of the test instance *x* with the values from the background data set.
2. These artificial samples are weighted by the value of the SHAP kernel calculated for each sample and *x*.
3. A weighted linear regression model *g* is trained to predict *f(x)*. The model coeﬃcients are Shapley values corresponding to fea- ture importance estimates.

Sampling all possible feature subsets is avoided through permuta- tion of the feature vector by setting features on and off. A feature is assigned a large weight if its replacement with an artificial value leads to a significant change in model output. Weights of artificial samples are determined as the number of feature addition sequences of a given subset by the SHAP kernel. Coeﬃcients from local linear regression pro- vide feature weights as Shapley values, which indicate how important a feature is for a given prediction including the direction (sign) of fea- ture influence. The expected explanatory value is calculated as the mean of the model output probability (numerical value) over training set in- stances. For a given instance, the model output is then calculated as the sum of the expected (base) value and all SHAP feature values.

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