Holistic Evaluation of Biodegradation Pathway Prediction: Assessing Multi-Step Reactions and Intermediate Products

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Abstract

The prediction of metabolism and biotransformation pathways of xenobiotics is a highly desired tool in environmental and life sciences. Several systems predict single transformation steps or complete pathways as series of parallel and subsequent steps. Their performance is commonly evaluated on the level of a single transformation step. Such an approach cannot account for some specific challenges that are caused by specific properties of biotransformation experiments. That is, missing transformation products in the reference data that occur only in low concentrations, e.g. transient intermediates or higher-generation metabolites. Furthermore, some rule-based prediction systems evaluate the performance only based on the defined set of transformation rules. Therefore, the performance of these models cannot be directly compared. In this paper, we introduce a new evaluation framework that extends the evaluation of biotransformation prediction from single transformations to whole pathways, taking into account multiple generations of metabolites. We introduce a procedure to address transient intermediates and propose a weighted scoring system that acknowledges the uncertainty of higher-generation metabolites. We implemented this framework in enviPath and demonstrate its strict performance metrics on predictions of in vitro biotransformation and degradation of xenobiotics in soil. Our approach is model-agnostic and can be transferred to other prediction systems. It is also capable of revealing knowledge gaps in terms of incompletely defined sets of transformation rules.

Index Terms: speech recognition, human-computer interaction, computational paralinguistics

1. Introduction

Data requirements for environmental risk assessments of chemicals are rapidly increasing, for example in regulatory processes at the European (cf. REACH [1]) and global level, but also for the development of new chemical products with more benign profiles. This includes increasing knowledge about transformation products of these chemicals in the environment and increases the need for prediction methods of metabolism and microbial biotransformation, along with the transformation pathways.

Conceptually, biotransformation pathways represent the chemical changes a given starting compound (referred to as root compound in the remainder of the text) undergoes upon biotransformation. They are constructed from compounds (i.e. molecular structures) connected by reactions. The pathway structure can be represented as nodes and edges in a graph. Figure 1 shows the Benzyl Sulfide pathway from EAWAG-BBD as an example.

Existing methods for the prediction of biotransformation products and pathways can be categorized as either knowledge-based, machine learning-based, or hybrid. Each of the two former approaches has its strengths and weaknesses. Knowledge-based approaches use expert knowledge on the basis of sets of transformation rules, in general leading to a combinatorial explosion caused by the overly general nature of the rules. Machine learning-based approaches predict pathways solely based on existing data, the performance being limited by the lack of large data sets.

Hybrid methods, such as machine learning-based relative reasoning models [3][4][5] predict probabilities of individual transformation reactions by combining knowledge- and machine learning-based approaches. They are constructed using sets of biotransformation pathways and transformation rules as training data, such as the EAWAG-BBD [6] and EAWAG-SOIL [7] packages. These models predict which of the transformation rules that are applicable to a given compound will be correct for that compound.

Biotransformation or metabolism studies from laboratory experiments or environmental observations are the basis for both training and evaluating these models and usually report
only transformation products that were formed in high quantities. This is because low concentration metabolites are considered less relevant and are more difficult to analyze and identify. Therefore, metabolites formed in low concentrations are less likely to be reported. This becomes more relevant for higher generation metabolites, because pathways typically diverge into multiple branches and transformations occur on different time scales. Both effects lead to decreasing maximum concentrations with increasing depth in the pathway. Thus, uncertainty about the actual formation of unreported metabolites increases for higher generation metabolites.

The performance of prediction models is typically determined by comparing the predicted transformations for each individual compound against the associated transformation products in the experimental reference pathway. This approach does not take into account the position of the compound or reaction in the pathway. Problems arise when:

- Multi-step reactions are represented as a single step in the experimental data.
- Intermediate metabolites are not observed or not elucidated.
- Transformation products are incorrectly assigned to the wrong educt.
- Concentrations of downstream metabolites become too low to be observed.
- Rule-based evaluation systems fail to address observed transformations not covered by the transformation rules.

In this paper, we address these shortcomings by introducing a new Multi-Generation evaluation approach that addresses some of the problems of the state-of-the-art Single-Generation evaluation approach. Multi-Generation evaluation explicitly includes the compound positions in the graph. Instead of only comparing the reactions for each compound independently, entire predicted pathways are compared to experimentally derived validation pathways. Predictions at higher depth get reduced weights to account for the increased uncertainty due to higher likelihood of minor transformation products not being reported in the experimental reference pathway.

The new evaluation approach further introduces a way to treat intermediate metabolites in the predicted pathway. These metabolites are quickly transformed to downstream products and therefore exist only in very low concentrations. As a consequence, they are often neglected or not analyzed in experimental reference pathways. Single-Generation evaluation approaches tend to incorrectly penalize prediction of these intermediates. However, the new Multi-Generation approach can take them into consideration when the downstream products are known. Their prediction is not penalized during the scoring process, and the depths of other downstream compounds in the pathway are adjusted accordingly.

With our new evaluation approach we evaluate whole pathway predictions more realistically than before, independent of the underlying set of transformation rules, more in line with the expectations of experimentalists, and more comparable across models. Therefore, we propose to consider the pathway holistically upon evaluation of predictive performance.

Overall, our main contributions are: 1. A new scoring system that quantifies the agreement between two biotransformation pathways. 2. Consideration of compound position (pathway depth) information in the evaluation of pathway predictions via scoring weights. 3. Consideration of intermediate metabolites in the evaluation of pathway predictions. 4. Use of conditional probabilities for depth considerations in predicted biotransformation pathways. This will enable further improvements of the prediction models in future work. Our methodology is a special case of graph analysis that is particularly useful for (bio)degradation or metabolic pathways and chemical reaction networks.

1.1. Background & Related Work

In this section, we will first give an overview of state-of-the-art prediction systems for biodegradation pathways and their methods. Then we will summarize related work to our proposed evaluation of prediction systems for biodegradation pathways.

1.1.1. Biodegradation

Biochemical Network Integrated Computational Explorer (BNICE) [8] is a framework that generates all known reactions for compounds. It uses the set of enzyme reaction rules based on the enzyme commission (EC) classification system. BNICE generates metabolic pathways by first determining functional groups contained in the root compound, and then generates associated products if the reaction rules are applicable. The process is repeated on each of the products in successive generations. The iteration terminates when a threshold is reached, or when no new compounds are created.

METEOR [9] provides the option of knowledge based prediction methods as well as machine learning approaches. The knowledge based option utilizes a combination of Absolute and Relative Reasoning in their predictions of reactions. The process commences by applying biotransformation rules to the starting compound, and these generate potential metabolites. The absolute reasoning process then assigns a level of belief to each biotransformation. Biotransformations that satisfy the absolute reasoning threshold preset by the user are then ranked in the relative reasoning process. The process uses a relative reasoning threshold to calculate the resulting relative hierarchy. Static Scores and Site of Metabolism Scoring are other prediction options that make use of machine learning techniques. The first utilizes an occurrence ratio – actual occurrences over all possible occurrences. The latter further considers similarity on additional chemical properties – attributes from generated fingerprints and molecular weights. The processes in each of these options are repeated for all surviving biotransformations, until some preset stopping conditions are satisfied, such as reaching the maximum depth.

PathPred [12] executes predictions by first searching for compounds from the KEGG [13] COMPOUND database that are similar to the chosen starting compound. The results are then used as input to search through the KEGG REACTION database for matching RDM transformation patterns [14]. These patterns are defined as KEGG atom type changes at the reaction center (R), the difference region (D), and the matched region (M). Products of these matching reactant pairs are then used as input, and this process is repeated until stopping conditions are reached. The Jaccard coefficient between the query and matched compounds of each reaction is used as the reaction score to indicate its plausibility. The average of all individual reaction scores in the pathway gives the pathway score.

EAWAG-PPS (formerly UM-PPS) [6] performs pathway prediction by first determining the functional groups in the starting compound, and applies biotransformation rules to determine the transformed products. Applying these rules iteratively to the educts would lead to combinatorial explosion, and known path-
ways were used to determine biotransformation priorities \cite{15}. User input is used at the end of each transformation prediction, to determine whether prediction continues downstream of the predicted compound(s). The predicted pathway grows as this cycle is repeated.

Biotransformer \cite{16} combines a rule or knowledge based approach in conjunction with a machine learning approach, to predict metabolic reactions for compounds. It makes use of experimentally confirmed biotransformations derived from the literature, as well as precedence rules that were derived from reported observations. Many of them are from the EAWAG-PPS database. The Biotransformer Metabolism Prediction Tool (BMPT) then uses a set of random forest and ensemble prediction methods to predict reactions, for example related to Cytochrome P450 enzymes (CYP450) and Phase II metabolism. For the latter, a simple rule-based filter is applied to eliminate the most trivial non-candidates for a few chemical classes with known metabolism. Metabolic pathways are predicted progressively starting from the root compound, one reaction at a time.

OASIS TIMES \cite{17} predicts chemical toxicity by integrating metabolism simulators into models assessing toxicity of both the transformation educts and products. This has improved model performance significantly compared to traditional approaches that don’t consider metabolic transformation products. The incorporated metabolic logic accounts for enzyme interactions, channeling effects and depletion of highly reactive intermediates. The metabolism simulator aims to correctly reproduce experimentally observed metabolites, and uses xenobiotic pathway data from MetaPath \cite{18} as a reference and aims to reproduce the observed pathways. However, it simulates metabolism using a complex mathematical model rather than a rule-based approach.

1.1.2. Pathway Evaluation and Comparison

Integrated scoring systems that attempt to quantify the quality of predictions are not always found in the systems mentioned above. PathPred \cite{12} computes the Jaccard index on compounds in each of the predicted reactions, and uses the average of all such values in a pathway as the overall score. OASIS TIMES \cite{17} takes the union of the observed and predicted pathways to tally true/false positives/negatives by comparing the metabolites. Only the first false positive in a sequence of false positives would be penalized, because the rest are conditioned from it. The system can also identify intermediates, and either reward, penalize or ignore them, based on a user-defined parameter. Prediction performances in published work for systems such as METEOR, BNICE and Biotransformer are obtained only from independent tests, without integrated options to evaluate the quality of predictions on new test sets.

A related field is the prediction of graph networks using machine learning techniques. Link prediction is a core component in many of the different approaches, such as analyzing information directly from the graph. This includes common neighbors \cite{19}, using metadata of the nodes from the application domain \cite{20}, or making use of pre-existing information on the connections between nodes in the graph \cite{21}. There are similar concepts in these approaches and the work in this paper, and we will explore them further for applicability in future work.

Research in Graph Isomorphism addresses the quantification of similarity between graphs. Many techniques focus on properties such as orientation or structural arrangements that share little relevance with biotransformation pathways. However, common metrics such as Graph Edit Distance \cite{22} can be useful in potential scoring systems or comparing predicted and observed pathways. Nevertheless, in biotransformation studies, the resulting pathways are tentative manual assignments by experts. They do not always reflect the absolute ground truth of the underlying reaction mechanism.

In summary, the work related to predicting biodegradation pathways so far does not take pathway structures into account. Our work in this paper aims to fill this gap by introducing a new approach that evaluates the predictions accordingly.

2. Methods and Experiments

In this section, we will first summarize the prediction and evaluation of models in enviPath \cite{5}, and then introduce our new Multi-Generation evaluation that overcomes the limitations of the current approach.

2.1. Single-Generation Evaluation

The standard enviPath Relative Reasoning models \cite{5,3,2,15} use a chosen set of biotransformation pathways as training data. The set of biotransformation rules consists of rules that were curated by experts. All compounds in the training pathways are independently cross-referenced with the rules for their applicability, producing effectively a quasi Boolean Matrix \cite{23} that describes their inter-relationships. The matrix connects the compounds and rules in a manner similar to:

\[
\begin{array}{cccccc}
1 & 1 & 1 & 1 & 1 \\
0 & 1 & 0 & 1 & 1 \\
0 & 0 & 1 & 0 & 0 \\
-1 & -1 & 0 & 1 & 1 \\
0 & 0 & 0 & 0 & 1 \\
\end{array}
\]

with rules \( r_n \) and compounds \( c_n \) with \( n = [1, ... 5] \) in the training. The values in the matrix elements represent Not applicable (-1), Applicable but not observed (0) and Applicable and observed (1). A machine learning model is trained on this matrix and then later used to predict probabilities for the combination of a new compound and the set of transformation rules.

In the Single-Generation evaluation, the predictions are then compared on the level of single reactions to the ground truth. That is, the known transformations are matched to the predictions for each rule and translated into true positives, true negatives, false positives, or false negatives. These counts then are translated to standard performance measures such as accuracy, recall, or precision. Multi-label approaches are used to aggregate the single transformation rule performance to one measure for the whole model. A detailed overview of the training and evaluation process is given in previous work \cite{3}.

2.2. Multi-Generation Evaluation

In contrast to the procedure outlined above, our Multi-Generation approach does not operate on the transformations of each individual compound, but first predicts a whole pathway (see next section) and then operates on the compounds as nodes in the graph. This leads to a couple of additional aspects that require consideration. Please note, however, that the underlying prediction model is identical in both approaches.

As discussed before, compounds in the first generation naturally carry higher confidence in the experimental findings, compared to transformations occurring at higher depth in the
pathway. This is due to the amount of test substance being divided into multiple reaction branches and only slow conversion over time. Thus, concentrations of higher-generation products are lower, which makes them more difficult to confirm experimentally and less likely to be reported. We therefore introduce a scoring system within our approach to account for the increasing uncertainty when comparing predicted and observed pathways.

This scoring system assigns rewards and penalties with weights according to the generation of the respective compounds. The resulting score for a pathway represents the agreement between the predicted and observed pathways. The collective scores for each of the pathways in the validation set are used to compute conventional metrics such as recall-precision curves. This new approach evaluates the pathway as a whole across multiple generations of compounds. This is in contrast to approaches in previous work where predicted reactions in each single generation are evaluated independently.

The prediction quality of Relative Reasoning models depends on the compatibility between the transformation rules and the training set, as well as the test set. Rule sets with low compatibility can lead to problematic scenarios, e.g. where no applicable rules can be applied to the target compound structure. In the Single-Generation evaluation process, such scenarios would result in all (if any) observed reactions from the educt being ignored. However, if there are further reactions for the product compound in the data, they would still be evaluated. Alternatively, in the Multi-Generation evaluation approach, the prediction would terminate at the initial educt and no further scores will be rewarded besides false negatives for the observed products.

Figure 2 demonstrates this difference between the two evaluation approaches with a simple example, an experiment that begins with compound A. The observed pathway has compound A transformed to B then to C, with the reaction from B → C described by a transformation rule (r_B→C) but none for A → B (r_A→B). The Single-Generation evaluation approach would only evaluate B → C (with a reward +) and ignore A → B, since no rule can be applied.

The Multi-Generation evaluation approach assigns penalties to both compounds B and C for not being predicted in the pathway. Although the model assigns a high probability to reaction B → C, the missing transformation rule for reaction A → B prevents any progression along that path. This puts strong emphasis on the knowledge gap in the set of transformation rules and provides a more realistic evaluation metric for the prediction accuracy on the overall pathway level.

2.2.1 Multi-Generation Pathway Construction

We predict pathways in the test or validation set starting from their root compound. Each of the possible reactions is predicted using the supplied transformation rules, which can be represented as a possible branch evolving from the educt. We then calculate conditional probabilities for reactions according to their position in the pathway.

This procedure takes into account the relationships between the probabilities of upstream reactions with the current reaction. We adjust the preset threshold value dependent on the depth and use it in the pruning process with the resulting conditional probability. This conditional probability is defined by the product of the probability value assigned to the current reaction, multiplied with values from all of the upstream reactions. An example pathway beginning from compound A is shown in Figure 3.

The example shows root compound A with probabilities P_B and P_C, for reactions that transform A into compounds B and C, respectively. A probability threshold x of value P_B > x > P_C is used in the example, to demonstrate the scenario where compound C is predicted to be not observed. The algorithm then continues to determine the possible reactions for compound B, transforming to compounds D and E at the second generation of the pathway, with respective probabilities P_D and P_E.

These values are multiplied with P_A, generating conditional probabilities, to obtain the conditional probabilities P_B × P_D and P_B × P_E. They are then tested against the threshold value adjusted for reactions at second generation, at x^2. This part of the example demonstrates the scenario where
are computed as follows:

\[ W = \prod \left( P_a \times P_b \right) \]

These definitions are used with the following Weighting System: 2.2.2. Performance Calculation

We calculate the pathway prediction performance based on standard true/false positive/negative counts, with the notable difference that we apply a weighting system and account for intermediates as described below. The quantities \( TP, FP, FN \) are the true positives, false positives, and false negatives, respectively, and \( W_D \) represents the weight multiplier that is dependent on the depth of the metabolite in the pathway. For an example see Figure 4. This metric avoids the infinite number of potential true negatives. The multipliers are then used to quantify the agreement between predicted and experimental pathways. We use the Jaccard Index as metric for pathway similarity. It is defined as:

\[ Sim = \frac{\sum(TP \times W_D)}{\sum(TP \times W_D) + \sum(FP \times W_D) + \sum(FN \times W_D)} \]

where \( TP \) and \( FP \) are the True and False positives, respectively, and \( FN \) represents the False negatives. \( W_D \) represents the weight multiplier that is dependent on the depth of the metabolite in the pathway. For an example see Figure 4. This metric avoids the infinite number of potential true negative

These definitions are used with the following Weighting System and treatment of intermediate metabolites.

2.2.3. Weighting System

We propose a mathematical model to compare two pathways with multiple generations. In accordance to the natural decrease in experimental certainty along the pathways, the compounds are assigned decreasing weights as their generation or depth level increases. These weight values start at \( 1/2 \) for compounds at generation or depth level one, and decrease by 50% for each increasing level. The weights are used as multipliers to conventional classification metrics such as counts of true/false positives/negatives. The multipliers are then used to quantify the agreement between predicted and experimental pathways. We use the Jaccard Index as metric for pathway similarity. It is defined as:

\[ Sim = \frac{\sum(TP \times W_D)}{\sum(TP \times W_D) + \sum(FP \times W_D) + \sum(FN \times W_D)} \]

where \( TP \) and \( FP \) are the True and False positives, respectively, and \( FN \) represents the False negatives. \( W_D \) represents the weight multiplier that is dependent on the depth of the metabolite in the pathway. For an example see Figure 4. This metric avoids the infinite number of potential true negatives. The multipliers are then used to quantify the agreement between predicted and experimental pathways. We use the Jaccard Index as metric for pathway similarity. It is defined as:

\[ Sim = \frac{\sum(TP \times W_D)}{\sum(TP \times W_D) + \sum(FP \times W_D) + \sum(FN \times W_D)} \]

2.2.4. Intermediate Metabolites

Intermediate metabolites are compounds with enhanced reactivity. They are quickly transformed to downstream metabolites, and therefore exist only in very low concentrations. These intermediates are sometimes included in the experimental data and sometimes not. This depends on the choice of the author of the experimental study report or the data package and the underlying experimental evidence. If they are not included, the transformation of the educt is reported to lead directly to the downstream metabolite. While prediction of such an intermediate would be mechanistically correct, they might not be present in the available data. Such a scenario would incorrectly inflate the count of false positives during the Single-Generation

\[ \text{Sim} = \frac{\sum(TP \times W_D)}{\sum(TP \times W_D) + \sum(FP \times W_D) + \sum(FN \times W_D)} \]

As the observed data is based on experiments where we can only be sure of observed compounds, we can never know for sure if a not observed compound is a truly negative or just not observed yet.
evaluation, and would be even more detrimental in the Multi-
Generation evaluation procedure. The intermediate metabolite
would be penalized, along with all metabolites downstream to
it, as they would appear at an incorrect depth in the pathway.

In order to correctly accommodate the intermediate
metabolites in the evaluation procedure, we have designed a
process that adjusts the depth level of the downstream com-
ounds accordingly. The process first determines a list of com-
ounds that are present in both the predicted and observed path-
ways. Then it checks if any of them are immediately down-
stream to one another in the observed pathway. The compound
pairs which fit this criterion are examined to test if additional
compounds are between them in the predicted pathway. These
compounds are then added to the list of intermediates. Such in-
termediate metabolites might still be correctly predicted with-
out the downstream node from the observed pathway. However,
the use of a correctly predicted downstream nodes is required
to identify them in a reliable manner and treat them properly.
In other words, we can correct the evaluation of intermediates
if and only if they have downstream products in the reference
pathway that were correctly predicted.

The list of intermediate compounds is used to adjust depth
levels in the predicted pathway accordingly. The shortest path
between each of the compounds in the pathway and the root
compound is determined using a Breadth-first search. The list
of in-between compounds is determined and the depth level
of the end compound is then decreased by the number of in-
termediate compounds that are in this list of in-between comp-
ounds. The intermediate compounds are ignored by the Multi-
Generation evaluation scoring algorithm.

Figure 5 shows an evaluation example incorporating con-
cepts from both the Weighting System and the treatment of in-
termediate metabolites.

2.3. Experimental Setup

We carried out several experiments to assess the proposed eval-
uation approach. We used a combination of pathway data se-
lection as well as several experiment designs to evaluate the
validity and difference in performance compared to the Single-
Generation evaluation approach.

2.3.1. Biotransformation Pathway Data

Several sets of biotransformation pathways were used in this
work:

EAWAG-BBD The set of biodegradation pathways con-
tained in the EAWAG Biocatalysis / Biodegradation Database
package [6] contains primarily xenobiotic chemical compounds
and microbial biocatalytic reactions. Information on such mi-
crobial enzyme-catalyzed reactions carries great importance
in the fields of biotechnology and environmental research.

EAWAG-SOIL The set of biodegradation pathways in the
EAWAG-SOIL package [7] contains pesticide degradation path-
ways compiled from laboratory soil degradation studies. These
pesticides are registered in the EU, and their degradation path-
ways are freely accessible regulatory data.

From the EAWAG-SOIL package we selected diverse sub-
sets of pathways as training and test sets that evenly cover
the chemical space. This is done to obtain a representative set with-
out over-representation of certain compound clusters. The se-
lection is based on the Tanimoto similarities from Morgan2 finge-
print algorithm [25].

\[
T(a, b) = \frac{N_{ab}}{N_a + N_b - N_{ab}}
\]

where \(N_a\) and \(N_b\) are the numbers of 1 bits present in the
fingerprints of compounds \(a\) and \(b\), and \(N_{ab}\) is the number of
1 bits occurring in both fingerprints. We used the MaxMin al-
gorithm [24] to incrementally pick compounds with the least
similarity to the most similar compound from the already se-
lected set. We selected 80% of the EAWAG-SOIL pathways to
become the TRAIN-SOIL package for model training purposes.
The remaining 20% make up the TEST-SOIL package which is
to be used as a test set. We excluded pathways that are not rep-
resentative for typical organic chemistry, i.e. when their root
compounds are inorganic salts, much larger than the rest, or
contain heavy metal elements.

2.3.2. Experiment Designs

We use the set of validated biotransformation rules from the
EAWAG-BBD package to build relative reasoning models with
compound structures from pathways inside specified training
packages. We have set the probability threshold for reactions to a low value of 0.1 for all experiments, in order to efficiently capture differences between the two evaluation approaches. The following are the experiment designs used for examination, and data sets for both evaluation approaches:

**Validation Test** A procedure to strictly validate the accuracy of the proposed mathematical approach that compares biotransformation pathways. Three sub-procedures are performed:

- **Full Pathway** Evaluate each pathway against itself. The result is expected to be 1.
- **Empty Pathway** Evaluate each pathway against only its starting compound. As the comparison of the pathway starting compound is ignored in the scoring system, the result is expected to be 0.
- **Half Full Pathways** A random process is performed to remove all but the starting compound in approximately 50% of a cloned set of pathways. Each pathway in the original set is evaluated against the associated one in the cloned set. Results of some metrics such as Accuracy and Recall are expected to be close to the ratio of unmodified pathways in the cloned set.

**Evaluation with Test Sets** A procedure where the entire chosen list of compounds is used to train a relative reasoning model once. Then we carry out the evaluation on the nominated test set TEST-SOIL. This procedure is performed on these pathway set combinations: TRAIN-SOIL, EAWAG-BBD + TRAIN-SOIL.

**Evaluation with Holdout** This procedure uses a random process to select approximately 66% of the chosen molecules to train a relative reasoning model. The model is then evaluated on the remaining 34% of the data. A list of compounds extracted from all selected pathways is used for selection for the Single-Generation evaluation approach, and the list of pathways is used for the Multi-Generation approach. The process is repeated 100 times, and the results of each individual run are averaged. This approach additionally allows an opportunity to also repeatedly examine the model’s prediction ability on data that is new to the training set. This procedure is performed on these pathway set combinations: EAWAG-BBD, EAWAG-SOIL, TRAIN-SOIL, EAWAG-BBD + EAWAG-SOIL, EAWAG-BBD + TRAIN-SOIL.

3. Results and Discussion

To assess the effectiveness and validity of our Multi-Generation evaluation approach, we summarize the results from the procedures detailed in the Experiments section in the following. We calculated Accuracy, Precision, Recall and Area under the Precision-Recall Curve (AUPRC). Due to the nature of the Multi-Generation evaluation approach, where pathways potentially have an infinite number of true negatives, the false positive rate cannot be computed. In the Single-Generation evaluation approach, the number of true negatives can be calculated from the applicable transformation rules, which are neither predicted (i.e. below the threshold) nor observed experimentally. The Area under the Receiver Operating Characteristic curve (AUROC) is hence only available for the Single-Generation approach and is provided as an indicator.

3.1. Illustrative Evaluation of an Example Pathway

We demonstrate the main differences between the two evaluation approaches on an illustrative example using the 1,1,1-Trichloroethane pathway from the EAWAG-BBD package (Fig. 6). For simplicity and better readability, we removed false positive predictions in Figure 6 and the evaluation below, which would distract from the main points of this demonstration. Please note, however, that five of these false positives are mentioned in the textual description of the pathway as minor products or are reported in the literature (trichloroacetic acid, dichloroacetic acid, dichloroacetic acid, ethane, 2-chloroethanol, acetic acid), which highlights the difficulty related to the incompleteness of such minor products in the reference data. The Multi-Generation approach mitigates the problem by reduced scoring weights for metabolites at higher depth in the pathway or zero weight for transient intermediate metabolites. Our example demonstrates the impact of not recognizing such intermediate metabolites. It is indicated in the metadata of the observed pathway that the final metabolite acetaldehyde (6) is formed indirectly from chloroethane (4) via intermediates. Indeed, there is no transformation rule in EAWAG-BBD for this transformation. Therefore, the model predicts acetaldehyde only via ethanol (5) or 2-chloroethanol (not shown) as intermediate steps.

In this scenario, the Single-Generation evaluation approach will return three true positives: two reactions from the root node leading to products 2 and 3, and one subsequent reaction from 3 to 4. The predicted reaction from chloroethane (4) to ethanol (5) is counted as false positive. The reported transformation from chloroethane (4) to acetaldehyde (6) is not evaluated at all, because there is no corresponding transformation rule in the underlying set of rules. This example demonstrates that the Single-Generation evaluation does not adequately address the likely intermediate and the final product acetaldehyde. The resulting accuracy for this example is 0.75, although the predicted
### 3.2. Validation Tests

Results of the validation tests performed on the EAWAG-BBD compounds are presented in Table 2. As expected, the evaluated full pathways from both packages achieve 1.0 for Accuracy, Precision and Recall, as there are only true positives and no false positives or negatives. The expected values for evaluated empty pathways from both packages are also 0 for all three metrics, as there are only false positives without any true positives. The “Half Full” pathways from both packages achieve 1.0 for Precision, and a value that is proportional to the amount of empty pathways (see Table 3) for Accuracy and Recall. The empty pathways will contribute with false negatives while the full pathways will contribute to the true positive score.

### 3.3. Evaluation with Test Sets

Relative reasoning models were trained with the TRAIN-SOIL package and the combination of EAWAG-BBD + TRAIN-SOIL packages. In both cases we evaluated the models on the TEST SOIL package. Tables 4 and 5 show the results, and Figure 7 gives the associated Precision-Recall curves.

The numerical values of each metric are noticeably lower for the Multi-Generation evaluation approach compared to the Single-Generation approach. This has two main reasons: First, the Single-Generation evaluation is based only on defined transformation rules, whereas Multi-Generation evaluates all nodes in the reference pathway and thus penalizes the incompleteness of the transformation rules. Second, a wrong prediction in the Multi-Generation approach is more detrimental, because all the downstream nodes from this branch will be wrong as well. In other words, for a true positive to be tallied, all upstream nodes also have to be predicted correctly. Additionally, a false positive will lead to even more false positives downstream. These two reasons make the Multi-Generation approach a much harder evaluation criterion, and thus the lower numerical values do not simply imply a worse result.

Another point worth noting from the Multi-Generation evaluation results is that the values for recall do not reach one (see Precision-Recall curve, Figure 7). The gap between the maximum recall and the value of 1.0 is caused by transformations in the reference pathways which are not covered by transformation rules and their downstream nodes. The products of such reactions can therefore never be predicted correctly, no matter how low the probability threshold is and will always count as false negatives. Moreover, as discussed above, any downstream nodes won’t be predicted either. In contrast, missing transformation rules have no effect on the Single-Generation performance, since Single-Generation is only evaluated on the existing rules. Thus, the maximum recall value at probability threshold zero can be used as an indicator for the completeness of the rules for the test set.

Furthermore, the results show that the precision values for both approaches do not reach 1, which means the number of false positives does not go down to 0, no matter how high the probability threshold is set. This is expected and implies that there are always products predicted with a high probability that are not correct. Given that the transformation rules are extremely general and can easily be triggered, the role of the machine learning models is to limit this. These results simply show that they do not predict perfectly, which will be hard to achieve with the available data.

The data in the TRAIN-SOIL package is naturally more representative for the evaluated TEST-SOIL package in terms of chemical and biological properties compared to the EAWAG-BBD package. While the EAWAG-BBD+TRAIN-SOIL configuration will provide better predictions for a broader chemical configuration.
and biological space, the model trained with only the TRAIN-SOIL package will perform better on data that share greater chemical similarity. Therefore, the relative reasoning model trained without the EAWAG-BBD package is more compatible with the evaluation data set. We can see that the Multi-Generation evaluation approach better reflects the compatibility between compound structures and the transformation rules used to train the model.

3.4. Evaluation with Holdout

We trained Relative Reasoning models with the EAWAG-BBD package, EAWAG-SOIL package, TRAIN-SOIL package, EAWAG-BBD + EAWAG-SOIL, and EAWAG-BBD + TRAIN-SOIL packages. For all cases, we repeated a holdout evaluation 100 times. Tables 6 and 7 give the results, and Figure 8 shows the associated Precision-Recall curves. A zoom-in to the Multi-Generation Precision-Recall curves from the Evaluation with Holdout experiments is presented in Figure 9.

Observations from the Evaluation with Test Sets results are
| Packages | AUPRC | AUROC |
|----------|------|------|
| BB D    | 0.64 | 0.87 |
| SOIL    | 0.47 | 0.83 |
| TRAIN_SOIL | 0.43 | 0.82 |
| BB D+SOIL | 0.56 | 0.85 |
| BB D+TRAIN_SOIL | 0.57 | 0.85 |

Table 7: Statistics of the Evaluation with Holdout experiments.

also notably present in these results from a more repetitive averaging process. The Precision-Recall curves are also notably smoother from the averaging process. The curves from the Multi-Generations approach also distinguish more clearly between results from only using the EAWAG-BBD package and other configurations. This is partially due to the fact that transformation rules from the EAWAG-BBD package are used to train all of these relative reasoning models. These rules were optimized for EAWAG-BBD but not for the other packages for which they are less suitable. Also note that soil is a much more complex system compared to the typical in vitro culture studies, which EAWAG-BBD is mainly comprised of. The outcome of a degradation experiment in soil is more difficult to predict.[7]

Figure 7 and 8 indicate that the Precision-Recall curves from each of the evaluation approaches occupy a different region in this space. Numerical values of Precision and Recall from the Multi-Generation approach seem less ideal. However, it must be emphasized that this is not due to worse predictions (same prediction model), but from a more holistic evaluation, taking into account additional aspects on the pathway level.

In analogy to language processing, the Single-Generation approach is analogous to evaluating each predicted word written by a columnist individually. The Multi-Generation approach on the other hand, is analogous to extending this to sentences and paragraphs. That is, correct predictions in the former may be penalized in the latter for being in the wrong place. Such a relationship between the two approaches indicate that it is natural to expect this difference in resulting numerical values between the two approaches.

4. Conclusion

In this paper, we present a new Multi-Generation approach for evaluating relative reasoning prediction models, that are used to predict biodegradation pathways. It includes methodology as well as performance in specifically designed experiments. The new approach evaluates predicted pathways with multiple generations of compounds holistically, in contrast to considering each reaction independently. Our approach additionally takes into consideration the increased uncertainty of observing compounds at higher depths in the pathways. We also propose an algorithm to account for intermediate metabolites, which would otherwise be incorrectly penalized during evaluation.

Our experiments show that the Multi-Generation evaluation metrics are much harder criteria. They provide a more realistic view on the prediction quality of whole pathways and reveal the incompleteness of the underlying transformation rules. With our new approach we can now start to compare the predictivity of different models on an objective basis in a model-agnostic way, i.e. independent from the model architecture and the set of transformation rules. Furthermore, we have demonstrated that our approach is more suitable to address two important characteristics of biotransformation pathway data: missing minor products in the reference data and intermediate metabolites. Single-Generation evaluation on the other hand might still be useful for determining the predictivity for individual (defined) transformation rules or for computationally demanding steps like hyper-parameter optimization.

Overall, our experiments demonstrate that it is still a long way until biotransformation prediction models can achieve top accuracy for complete pathways. However, with the Multi-Generation approach we improved our toolbox for the evaluation and comparison of pathway prediction models, which will facilitate the development of better models. In future work, we will use this approach to improve the compatibility of the biotransformation rules. Additionally, we will integrate the new knowledge about likely intermediates into model training.

5. List of Abbreviations

AUPRC - Area Under Precision-Recall Curve
AUROC - Area Under Receiver Operating Characteristic Curve
BBD - Biocatalysis/Biodegradation Database
BMPT - Biotransformer Metabolism Prediction Tool
BNICE - Biochemical Network Integrated Computational Explorer
CYP450 - Cytochrome P450
EAWAG - Swiss Federal Institute of Aquatic Science and Technology (Eidgenössische Anstalt für Wasserversorgung, Abwasserreinigung und Gewässerschutz)
EC - Enzyme Commission
FN - False Negative
FP - False Positive
KEGG - Kyoto Encyclopedia of Genes and Genomes
METEOR - Name of a knowledge-based expert system for toxicity and metabolism prediction
OASIS - Name of the company that manages the TIMES software
PPS - Pathway Prediction System
RDM - Reaction center, the Difference region and the Matched region of the KEGG atom type changes
REACH - Registration, Evaluation, Authorization and Restriction of Chemicals (European Union Regulation)
TIMES - Name of the software from the Laboratory of Mathematical Chemistry Burgas for simulation of chemical metabolism
TN - True Negative
TP - True Positive
UM - University of Minnesota

6. Declarations

6.1. Availability of data and materials

The EAWAG-BBD and EAWAG-SOIL packages used in this study are publicly available on enviPath. A python code that demonstrates the Multi-Generation evaluation process can be accessed at the enviPath Github repository, along with two files containing the names of pathways included in the TRAIN-SOIL and TEST-SOIL packages.

6.2. Competing Interests

Jörg Wicker is one of the founders of enviPath UG & Co. KG, a scientific software development company that develops and maintains enviPath. Jörg Wicker, Tim Lorsbach, and Jason Tam
are employees of enviPath UG & Co. KG. Sebastian Schmidt is an employee of Bayer AG, a manufacturer of pharmaceutical, agricultural, and consumer health chemicals. The authors declare no competing interests.

6.3. Funding
Not applicable.

6.4. Authors’ contributions
JT proposed the initial methodology for the Multi-Generation approach, carried out the experiments and wrote the initial manuscript draft. TL, SS and JW continually provided expertise for refining the details during the entire analysis process, as well as editing and improving the manuscript. All authors read and approved the final manuscript.

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