Biocatalysed synthesis planning using data-driven learning

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Enzyme catalysts are an integral part of green chemistry strategies towards a more sustainable and resource-efficient chemical synthesis. However, the use of biocatalysed reactions in retrosynthetic planning clashes with the difficulties in predicting the enzymatic activity on unreported substrates and enzyme-specific stereo- and regioselectivity. As of now, only rule-based systems support retrosynthetic planning using biocatalysis, while initial data-driven approaches are limited to forward predictions. Here, we extend the data-driven forward reaction as well as retrosynthetic pathway prediction models based on the Molecular Transformer architecture to biocatalysis. The enzymatic knowledge is learned from an extensive data set of publicly available biochemical reactions with the aid of a new class token scheme based on the enzyme commission classification number, which captures catalysis patterns among different enzymes belonging to the same hierarchy. The forward reaction prediction model (top-1 accuracy of 49.6\%), the retrosynthetic pathway (top-1 single-step round-trip accuracy of 39.6\%) and the curated data set are made publicly available to facilitate the adoption of enzymatic catalysis in the design of greener chemistry processes.
Chemistry fostered the unprecedented rise of overall human wealth and well-being during the past two centuries, and today it is our trump card for averting and mitigating global crisis while reshaping our lives towards a more responsible use of natural resources. Innovation in synthetic chemistry will be critical in making chemical processes and products more sustainable, resource-efficient and CO2-neutral. While the design and development of catalysts are at the heart of greening chemistry, biocatalysis, together with chemoinformatics and artificial intelligence, can already accelerate the adoption of existing sustainable catalytic processes.

At the core of biocatalysis are enzymes, an integral part of all living organisms used in important industrial processes thanks to the multiple key advantages over conventional chemical reagents. In addition to their extremely high catalytic activity, enzymes catalyse chemo-, regio- and stereo-selective reactions and are both reusable and allow for an easy recovery of products when immobilised. A further advantage of enzyme-catalysed reactions is that they are usually performed in water under mild conditions and thus significantly reduce waste. Moreover, enzymes themselves are fully degradable in the environment, and as such, they represent an important strategy towards greener chemistry. Therefore, it is not surprising that enzymes are one of the key enablers of sustainable chemical processes, with a growing adoption of enzymatic processes fostering the unprecedented rise of overall human chemistry, biocatalysis, together with chemoinformatics and artificial intelligence, can already accelerate the adoption of existing sustainable catalytic processes.

The knowledge gap between large corpora of enzymatic chemical reactions data and the human understanding of the structure-activity relationship hinder the ability to predict successful routes when the substrates of interests are not directly associated with an enzyme. Machine learning and data-driven approaches may be useful strategies to capture the hidden patterns in large enzymatic data sets, similar to the proven ability to learn chemical reaction patterns from complex chemistry knowledge collections. The extraction of chemical reaction rules from large data sets of traditional organic chemistry reactions is one of the most successful examples of providing transparency and explainability through AI applications in chemistry to date.

Despite the impact on traditional synthetic organic chemistry, computer-aided design planning tools using biocatalytic reactions are in the early days of their development. Currently, only rule-based methods for predicting biosynthesis pathways have been examined, such as the ATLAS of Biochemistry or RetroRules. Lately, RetroBioCat became the first chemoinformatic approach for easing the adoption of biocatalytic reactions specifically for chemical synthesis. However, the implementation relies on a set of expertly encoded reaction rules coupled with a system for retrieving database records to enable the use of biocatalysis in synthetic organic chemistry. This use of reaction templates slows down the curation of newly collected data, requiring the intervention of human experts, and suffers from the limitation in capturing the effects on the reaction centre of long-range substituents. Shortly after, Kreutter et al. presented a forward reaction prediction model based on the Molecular Transformer. This approach exploits a multitask transfer learning to train a Molecular Transformer architecture, originally trained with chemical reactions from the US Patent Office (USPTO) data set, with 32,000 enzymatic transformations, each one annotated with the corresponding enzyme name. The enzymatic transformer model predicts the products formed from a given substrate and enzyme in the forward prediction task, reaching an accuracy of 54% when using the enzyme name information only and 62% when using the complete enzyme information as a full sentence (often also including the organism name). The approach addresses some of the concerns around scalability and data curation of reaction templates. However, the use of enzyme names as reaction tokens adds an additional level of challenge when trying to learn chemical reactivity patterns among enzymes with different names but belonging to closely related families. In addition, the lack of a corresponding backward model in this work does not allow for retrosynthetic planning.

Here, we generalise the use of the Molecular Transformer by adopting a tokenisation system based on enzyme classes and introducing an extension of the retrosynthetic algorithm by Schwaller et al. to biocatalysis. Compared to the previous work by Kreutter et al., a backward model allows to predict substrates and enzyme classes given a target product, enabling retrosynthetic pathway prediction. In addition, we incorporate the EC (enzyme commission) number into the reaction SMILES, rather than encoding enzymes with their natural language name. Enzymatic reactions and the accompanying EC numbers were extracted from four databases, namely Rhea, BRENDA, PathBank and MetaNetX and merged into a new data set, named ECREACT, containing enzyme-catalysed reactions with the respective EC number, as shown in Fig. 1.

The forward prediction model achieves an accuracy of 49.6%, 63.5% and 68.8% top-1 and top-5, and top-10 respectively, while the single-step retrosynthetic model shows a round-trip accuracy of 39.6%, 42.3% and 42.6%, top-1, top-5 and top-10, respectively.

Results

Data set. The enzymatic reaction data set with related EC (enzyme commission) numbers was created by merging entries extracted from Rhea (n = 8659), BRENDA (n = 11130), PathBank (n = 31047) and MetaNetX (n = 34485). This data set was then further processed by (1) removing products that occur as reactants in the same reaction, (2) removing known co-enzymes...
and common byproducts from the products in reactions that exceed one product (Supplementary Tables 1 and 2), (3) removing molecules with a heavy atom count < 4 from the products and (4) removing reactions with >1 or <1 products or no reactants. The resulting data set contains 62,222 unique reaction–EC-number combinations. The data set is available in 5 different token schemes: With no EC number (EC0, n = 55115), only EC-level 1 (EC1, n = 55707), EC-levels 1-2 (EC2, n = 56222), EC-levels 1-3 (EC3, n = 56579) and EC-levels 1-4 (EC4, n = 62222). The different token schemes result in different set sizes as the removal of EC-levels leads to duplication and removal of extended reaction SMILES. Given the low specificity of enzyme information in EC1 and EC2 tokens, and the insufficient sampling for EC4, which is often confined to one enzyme-substrate example only, the EC3 data set remained the only one containing sufficient variability in terms of enzyme-substrate examples across individual tokens. Figure 2a shows the composition of the data set with token scheme EC3, containing 62,403 unique enzymatic reactions. At EC-level 1, which corresponds to enzyme classes, EC 2.x.xx (transferases) account for 53.5% of total entries, EC 1.x.x.x (oxidoreductases) for 24.5%, EC 3.x.x.x (hydrolases) for 10.7%, EC 4.x.x.x (lyases) for 6.3%, EC 6.x.x.x (ligases) for 2.3%, EC 5.x.x.x (isomerases) for 2.2% and EC 7.x.x.x (translocases) for 0.4%. The high fraction of transferase-catalysed reactions is a consequence of the large number of non-primary lipid pathways stored in PathBank. Among transferases, the most common subclasses at EC-level 2 are EC 2.7.x.x (transferases transferring phosphorus-containing groups) at 24.5%, EC 2.3.x.x (acyltransferases) at 16.8% and EC 2.1.x.x (transferases transferring one-carbon groups) 7.5%. The complete information on the distribution of samples across EC-levels 2 and 3 is provided in the supplementary information (Supplementary Tables 3 and 4, with a breakdown of the data set by data source shown in Supplementary Fig. 2).

The distribution of the available data reveals a heavy imbalance in the distribution of the enzyme-substrate examples. Whereas transferase-catalysed reactions encompass few subclasses at EC-level 3 with large sample size, the oxireductase- and hydrolase-catalysed reactions are divided into many subclasses with a small sample size at EC-level 3. Although lyases, isomerases, ligases and translocases are split into fewer subclasses at EC-level 3, most of them contain very few samples. Therefore, the evaluation of the performance of the data-driven models will need to consider the different populations of each EC-level 3 subclass for a proper assessment.

A further property of interest regarding the reaction is the distribution of reactants and products within and across the enzyme classes at EC-level 1. The data set created with the EC3 token scheme contains 141,051 (56,017 unique) reactants and 62,403 (53,658 unique) products. Given the nature of the data sources, most reactants and products are metabolites. In Fig. 2b, c, we show the distribution of the compounds in the substrate and product chemical spaces using the 2048-dimensional binary MAP4 fingerprints and embedding them using TMAP31,32. The data points, coloured by enzyme class corresponding to EC-level 1, highlight the different distributions within the substrate set (containing cofactors) and the product set (where co-enzymes and common byproducts have been removed). Substrates and products of transferase- (class 2), lyase- (class 4) and, to a lesser extent, hydrolase-catalysed (class 3) reactions populate regions of the chemical space specific to each class (homogeneous), molecules from other classes are found in predominantly heterogeneous regions.

Model selection. A forward and backward model was trained for each of the ECREACT token schemes, with EC0 acting as a control on the influence of including enzymatic information in the reaction. The trained models were evaluated for forward, backward, round-trip and EC number prediction accuracy using 5% test splits with the condition that a product in the test split must not occur as a product in the training split (Fig. 3). The results show that the EC3 token scheme has a better forward performance than EC0 and EC2, yet performs slightly worse than EC1 and EC4 (Fig. 3a). In the backward prediction task, EC3 performs slightly worse than EC0, EC1
and EC2, but significantly better than EC4; this is most likely due to the low number of samples in each EC4 category (Fig. 3b and Supplementary Table 5). Regarding the round-trip accuracy25, EC3 performs better than both EC2 and EC4 (Fig. 3c). When solely focusing on the prediction of the correct EC number in the backward prediction, the models perform better the less detailed information they have to predict (Fig. 3d). These data show that the inclusion of enzymatic information in the form of the EC number does not affect the prediction performance negatively as long as each EC category has a sufficient number of training samples, which restricts the use of EC4 (Supplementary Fig. 16). The EC1 token, although performing well across different metrics, averages across reaction classes with different schemes and is for this reason of little interest for retrosynthetic purposes. The EC3 token scheme balances specificity of enzyme information with performance compared to the other ECREACT token schemes, resulting in a prediction performance similar to or better than EC1 and EC2, while retaining detailed information of the reaction-specific enzyme. Therefore, the relative performance among the five ECREACT token schemes EC0, EC1, EC2, EC3 and EC4, identifies EC3 as the one with the richest amount of statistically significant information.

Forward prediction. The Dataset was constructed following the details reported in the Methods, Data Sets and Model Training. We split the EC3 data set ($n = 56,579$) into a test and a training set, enforcing a zero overlap between the product distributions of the two ensembles, i.e. no product molecule present in the test set appears in the training set. Although this splitting penalises the amount of statistically significant information.

![Figure 3](https://example.com/figure3)

**Figure 3** Overall accuracies of models based on different ECREACT token schemes EC0, EC1, EC2, EC3 and EC4. Accuracies are reported for a forward prediction, b backward prediction, c round-trip prediction (a forward prediction followed by a backward prediction) and d backward EC number only prediction. Top-n indicates the accuracy when checking the top n predictions for the correct one.
of donors. The predicted reaction (1) contains an excessive number of carbon atoms. The inferred product of the reaction (2) is equivalent to the ground truth, as the linear and cyclic forms are in equilibrium. (3) shows an example of the model correcting an error in the data set and predicting the correct stereochemistry. The discrepancies also highlight the possibility of false negatives due to the prediction of zwitterions. Concerning the diester hydrolase-catalysed reaction (4) and the intramolecular lyase reaction (5), it is worth noting that the data set contains few chemical reaction records with identical substrates and EC numbers that result in different products. With sufficient data, the model would be able to recommend, given an EC-number, the various transformations of a single substrate into different products with corresponding confidence levels. However, due to the limited data volume and the random nature of the split into training, validation, and test sets, it is unlikely that all possible outcomes of a single substrate and EC number will be used for training. As a result, the two reactions are predicted incorrectly. (6) is an example of the model failing to predict the correct stereochemistry of a product. Prediction of correct
stereochemistry has been reported by Schwaller et al.\textsuperscript{25} as a major challenge for the molecular transformer and is linked to the lack of coherent stereochemical information in the USPTO data set\textsuperscript{13}. Similarly, the correct stereochemical prediction, affected by the limited data coverage on stereochemical examples, remains the major challenge for the model, especially when predicting reactions catalysed by isomerases (class 5, Fig. 4a). The removal of all stereochemical information from the predicted products increases the accuracy of isomerase-catalysed reaction prediction by a factor of two (Supplementary Fig. 5).

Inspired by the work of Schwaller et al.\textsuperscript{18} we unboxed the forward prediction model to better understand how it exploits enzyme information. Figure 6 shows an example of an enzymatic reaction and the attention relationship between the EC and the product tokens. The EC tokens relate to the centre of the nucleophilic addition as well as the reactive nucleophile. A more extended analysis of the attention patterns and additional examples can be found in the supplementary information. The analysis of the attention weights confirms the capacity of the forward Molecular Transformer to use the EC token for discerning the enzymatic reaction centre while capturing enzymatic reaction rules.

**Backward prediction.** Here, we will use both the round-trip\textsuperscript{26} and the top-\(k\) accuracy for the evaluation of the backward model. Both metrics will provide evidence of how good the model is at proposing different enzymatic reactions that can lead to the desired target (round-trip accuracy) or the one specifically reported in the ground truth (top-\(k\) accuracy). With a top-1 accuracy of 60\%, the backward model has a behaviour similar to the forward model in the performance between and within classes (Fig. 7), as well as in the correlation between the size of the training samples and accuracy (Supplementary Fig. 16). In addition to the substrates, the model also predicts the enzyme EC-level 3 token; the accuracy of predicting EC numbers only is shown in Supplementary Fig. 10. The analysis of the model shows an exceptional performance on transferase-catalysed reactions (class 2), traceable to the two large EC-level 3 subclasses EC 2.3.1.x, and EC 2.7.8.x., which contain 17\% and 20\% of all available samples, respectively (Fig. 7b). This analysis further explains the comparatively low prediction accuracy in the class of oxidoreductases (class 1) as it contains a large number of EC-level 3 subclasses, each small in size (Fig. 7 and Supplementary Table 4). Translocases are involved in catalysing the movement of molecules or ions across membranes. Together with the limited set of reaction records (191), this specific function causes the substrates and products to have lower diversity than those in other classes. The limited data reduces the statistical significance for this class, and we have thus opted to exclude Translocases from a detailed analysis.

The confusion matrix (Supplementary Fig. 12) provides further insight into the backward prediction performance. The model’s ability to assign a product to the correct enzymatic class differs significantly between classes and is again influenced by the cohort of each class. Despite the larger population of the oxidoreductases, the split into many EC-level 3 subclasses causes the backward model to perform worse in predicting substrates for oxidoreductase-catalysed reactions than for hydrolase-, lyase- and ligase-catalysed reactions (Fig. 7a). However, the prediction of the enzyme class (EC number) only shows high accuracy (71.97\%) for the class of oxidoreductase. This result shows that given a diverse reaction data set, the model can distinguish between classes but does not have enough data to predict the correct substrates. A challenge in terms of predicting the correct enzyme class are the isomerases (class 5), as they encompass intramolecular oxidoreductases, transferases and lyases. This is reflected in the relatively high misassignment of isomerases to oxidoreductases, transfrases and lyases (classes 1, 2 and 4).

Similar to the forward reaction prediction model, we report a set of successful (Supplementary Fig. 14) and unsuccessfully predicted backward reactions (Fig. 8), together with their ground truth. The successful examples reflect the models’ capability to predict substrates across enzyme classes. Among the unsuccessful ones, Example (1) highlights the prediction of a different EC-level 3 token to catalyse the reaction. Whereas the ground truth reaction is catalysed by an oxidoreductase acting on the CH-OH group of donors with oxygen as an acceptor (1.1.3.x), the prediction suggests the reaction to be catalysed by an oxidoreductase acting on the CH-OH group of donors with NAD\(^+\) as an acceptor (1.1.1.x). This choice may reflect the respective number of training samples for the two classes (3,220 and 176 for 1.1.3.x and 1.1.1.x, respectively) and could be considered a viable alternative to the ground truth. Example (2) shows a correct EC-level 3 prediction (hexokinase). However, the substrate did not match the ground truth because the model predicted the acyclic
This approach represents an alternative to gaseous hydrogen or other solid hydride sources typically employed in the reduction of carbonyls, which often represent a safety concern when employed already on gram-scale. Homoaasparte (2) can be accessed by a series of chemoselective enzymatic transformations of \( \alpha \)-erythrose to the corresponding carboxylic acid, followed by regioselective dehydration to the \( \alpha \)-ketoacid. Finally, the model infers that a transamination with glutamate on the newly-introduced keto functionality ensures the delivery of the target amino acid. Given the similar reactivity of the -OH groups within the substrate, such a series of transformations would require considerable effort using non-enzymatic approaches. For the third example, the model predicts that 4-hydroxy-L-glutamic acid (3) can be obtained from oxidation of inexpensive L-hydroxyproline in the presence of \( \text{NAD}^+ \) (catalysed by EC 1.5.5.5), followed by a further oxidation of the aldehyde intermediate with EC 1.2.1.x and \( \text{NADP}^+ \). Enzymatic reactions enable oxidations to be also carried out in the presence of \( \text{O}_2 \), as exemplified by the prediction of the synthesis of \( \alpha \)-ketoacid (4). The chemoselective oxidation of the amino group of L-tyrosine leaves the sensitive and electron-rich aromatic moiety unaltered and obviates the use of stronger oxidising agents. Lastly, the model predicts that an enzymatic Pictet-Spengler reaction catalysed by EC 4.2.1.x, can convert dopamine and the corresponding aldehyde to the alkaloid (S)-norlaudanosoline (5) enantioselectively, which typically requires the presence of organocatalysts or transition metals. It is interesting to compare the routes suggested by our model with the ones from RetroBioCat. In reaction (1) the starting substrate is styrene, which undergoes epoxidation in the presence of an epoxidase, followed by epoxide opening, partial oxidation of the primary alcohol to aldehyde and transamination. Homoaasparte (2) is instead shown to be obtained via aldol addition of sodium pyruvate on formaldehyde, followed by transamination with alanine. Similarly, RetroBioCat shows that 4-hydroxy-L-glutamic acid (3) can be prepared by treatment of pyruvic acid with glyoxylactic acid in the presence of an aldolase, delivering the target compound after transamination with an amino donor. Pyruvic acid is also the substrate suggested for the synthesis of \( \alpha \)-ketoacid (4), which is delivered upon reaction with phenol in the presence of a lyase. Alkaloid (S)-norlaudanosoline (5) is synthesised in a similar fashion as suggested by our model with a norcoclaunine synthase, which acts on the same primary amine and aldehyde substrates shown by the Molecular Transformer. With the
exception of route (5), which is highly substrate-specific, one can appreciate the dissimilarity of the synthetic pathways suggested by our model when compared with RetroBioCat. While RetroBioCat algorithmically constructs a set of potential substrates using expert-curated rules, our model incorporates both the implicit reaction rules and its knowledge about the substrates contained within the training set into the generation of potential substrates. Based on the ECREACT training set, which contains mainly biosynthetic reactions, the suggested substrates resemble natural products with potentially high affinity towards the wild-type
Fig. 9 Distribution of rxnfp fingerprints for the reactions in the combined space of ECREACT (grey) and RetroBioCat test set reactions (blue), embedded with TMAP. a The reactions from the RetroBioCat test set are forming distinct clusters in the combined reaction space. b For RetroBioCat test set (blue) reactions, the fraction of nearest neighbours \((k = 10)\) from the set itself is consistently higher compared to reactions from ECREACT (grey).

Fig. 10 Enzyme-catalysed synthesis of synthetically useful compounds under mild conditions. (1) Aminoalcohol, (2) Homoaspartate, (3) 4-hydroxy-L-glutamic acid, (4) \(\beta\)-ketoacid and (5) (S)-norlaudanosoline.
enzymes. The use of commercial biocatalytic reaction-specialised data sets, not accessible to the authors and the general public, would likely lead to quite different and possibly more similar predictions to the ones suggested by RetroBioCat. However, even with the biosynthetic-specific nature of our data set, the model remains flexible when tasked to predict never-seen substrates by associating enzyme classes with the centre of the enzymatic reaction (Fig. 6 and Supplementary Fig. 18). As the molecular transformer-based model allows for retraining or fine-tuning using different data sets, these examples show that the substrate-scope can be narrowed down or expanded, depending on the use-case and available data. These properties open a way to synthetically useful compounds from a variety of different inexpensive substrates once experimentally validated.

Discussion
We presented forward and backward prediction models based on Molecular Transformer trained on enzyme-catalysed reactions extended with EC (enzyme commission) numbers. Our results show that the Molecular Transformer performs well in predicting products given EC number and substrates, predicting substrates and EC number or EC number alone given a product. The enzymatic models reach an overall top-1 accuracy of 49.6%, 60% and 39.6% in forward, backward and round-trip accuracy, respectively. The accuracies correlate heavily with the amount of training data in each token class, presenting a major challenge given the limited data availability. In addition to the trained models for biocatalysed synthesis planning, we introduced an aggregated data set, ECREACT, containing preprocessed enzyme-catalysed reactions sourced from different publicly available databases. The primary limitation of the data set, and thus of the model, is the scarcity of samples for some classes, such as isomerases, which results in an imbalanced data set. However, as interest in biocatalysis grows and the research community embraces open data, we anticipate an increase in the quantity and quality of available training data, as well as experimental validation of proposed synthetic routes. In addition, the data has been sourced from databases containing mainly biosynthetic reactions due to the lack of publicly available data on the biocatalysed synthesis of non-natural products, leading the model preferably suggest natural products as substrates. Despite this bias, we showed that the model generalises by learning the association between enzyme classes and the centres of the enzymatic reactions, which is a valuable property given the limited availability of biocatalytic reaction data. The research community will be able to build on the legacy of the present work to retrain models with higher accuracy and broader scope without the limitation of humanly curating reaction rules. Finally, we presented several use-cases based on well-understood pathways that showed how template-free machine learning models trained on enzymatic reactions can play an essential role in promoting the adoption of greener chemistry strategies in daily laboratory work.

Methods
Transfer learning. We used the USPTO data set, which contains 1 million organic chemical reactions, together with the more specific enzymatic reaction data set to train the molecular transformer using multi-task transfer learning. The USPTO data set is used to learn general chemistry knowledge and the SMILES grammar, as the comparatively small enzymatic reaction data set does not provide sufficient information for these tasks. This approach was previously successfully applied to carbohydrate reactions by Pesciullesi et al.25. The reactions in the USPTO data set are encoded as so-called reaction SMILES, using the same convention of Schwaller et al.25. An example is the reaction SMILES $\text{CC} (=\text{O})\text{OCC} \rightarrow \text{OS} \rightarrow \text{(O)} \rightarrow \text{O} \rightarrow \text{OCC} \rightarrow \text{OCC} \rightarrow \text{O}$ encoding a Fischer esterification. The main conceptual difference lies in extending the reaction SMILES tokeniser to handle enzymatic reactions represented by the EC number as detailed in the section Preprocessing.

The Molecular Transformer models were implemented following the protocol introduced by Schwaller et al.25. Multi-task transfer learning was implemented, as described by Pesciullesi et al.25, using a convex weighting scheme for USPTO and ECREACT, 9 and 1, respectively. Both encoder and decoder were of type transformer with 6 layers, word vectors and RNN of size 512, the gradient was accumulated 8 times with a maximum vector norm of 0.0, and adam was used as an optimiser ($\beta_1 = 0.9, \beta_2 = 0.998$). Batch size was set to 4096, and the batch type as the gradient normalisation method to tokens. The learning rate was set to 2.0 with noam as decay method. Dropout and label smoothing ($\epsilon$) were set to 0.1. Optimiser initialisation was disabled and position encoding enabled. All models were trained using a version of OpenNMT40, adapted for the Molecular Transformer41.

Preprocessing. The standard definition of a reaction SMILES was extended to include EC numbers (e.g. the reaction catalysed by the maltose alpha-D-glucosidase is written as $\text{A5.4.99}-18$, whereas the SMILES for $\alpha$-maltose and $\alpha$-trehalose have been replaced by A and B for brevity). We denote this extension to reaction SMILES enzymatic reaction SMILES.

We adapted the tokenisation operation used by Schwaller et al.25 for the molecular transformer to handle enzymatic reaction SMILES, EC-levels 1-3 are treated as unique tokens to enable the transformer to learn the hierarchical structure of the EC numbering scheme. Because digits are already used to represent ring closures in SMILES, a number prefix is added to each level (v for EC-level 1, u for EC-level 2 and 1 for EC-level 3) during tokenisation. In addition, each EC token is encapsulated in brackets to simplify the tokenisation and detokenisation process. An example tokenisation of an enzymatic reaction SMILES is shown in Supplementary Fig. 22.

Finally, the resulting tokenised data set was split into a training, validation and test set (90%, 5% and 5%, respectively). The training set was sampled so that none of the products contained within it are present in the training and the validation data sets.

Retrosynthesis routes prediction. We adapted the methodology proposed by Schwaller et al.25, extending the retrosynthetic routes’ prediction to handle enzyme inhibition using the EC number format. The hyper-graph exploration algorithm, at each step, proposes disconnections using the backward model and computes a score for each prediction, in a Bayesian sense, based on the confidence of the forward model reweighted by the SCScore25 measured on the precursors. The pathways are then prioritised, exploiting the score by using beam search until a terminating condition is satisfied, i.e., commercial availability of the precursors (see Supplementary Fig. 23).

For the analysis of the targets from25 we used an interactive version of the approach, where the backward Molecular Transformer allowed us to explore the synthetic routes iteratively until reaching commercially available precursors and inhibitors, at the same time, enzymes (represented up to EC level 3) that catalyse the corresponding reaction.

We based the selection of the targets on a comparative analysis of the coverage of the chemistry embedded in the reactions from the RetroBioCat25 test set and the ECREACT data set. We annotated reaction SMILES for each step of the biocatalytic cascades considered in the test set from Finnegan et al.25, excluding solvents information. For each reaction SMILES we extracted fingerprints using rnxp35 and we computed among the k-nearest neighbours (k = 10), the fraction of neighbours belonging to RetroBioCat test set. The visualisation of the embedded reactions was generated using TMAP31.

Data availability
The ECREACT data set is publicly available at the URL https://github.com/ranchemistry/biocatalysis-model. The trained models are made publicly available as part of IBM RXN for Chemistry (https://ran.rex.ibm.com/).

Code availability
Code is available at the URL https://github.com/rxn4chemistry/biocatalysis-model.

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Author contributions

The project was conceptualised by D.P., M.M., F.P. and T.L.; D.P. and M.M. curated the data, developed the methodology, and carried out the formal analysis, investigation and validation. D.P., M.M. and Y.T. created the visualisations and software. D.P., M.M., Y.T. and A.C. wrote the original draft. A.C. validated the test set and curated the data for the use cases. T.L. supervised and administered the project, reviewed and edited the writing and acquired funding and resources.

Competing interests

The authors declare no competing interests.

Additional information

Supplementary information

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