Machine Learning Differentiates Enzymatic and Non-Enzymatic Metals in Proteins

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Abstract
Identifying enzyme active sites is an important task in enzymology. Metalloenzymes are 40% of all enzymes and can perform all seven classes of enzyme reactions. The active sites of metalloenzymes can be difficult to distinguish from non-enzymatic metal binding sites. Because these sites have so many physicochemical similarities, finding physicochemical features that distinguish metal binding sites with enzymatic activity from metal binding sites without enzymatic activity can elucidate what is critical to enzyme function. In this work, we develop the largest structural dataset of enzymatic and non-enzymatic metalloprotein sites to date. We then use a decision-tree ensemble machine learning model to distinguish metal ions bound to a protein as enzymatic or non-enzymatic with 92.2% precision and 90.1% recall. The success of our model scores electrostatic and pocket lining features as more important than pocket volume, despite the fact that volume is the most quantitatively different feature between enzyme and non-enzymatic sites. Finally, we find our model has overall better performance in comparison against other methods that differentiate enzymatic from non-enzymatic sequences. Our model’s ability to correctly identify which metal sites are responsible for enzymatic activity could assist with identifying new enzymatic mechanisms or designing de novo enzymes.

Introduction
Enzymes are biological catalysts. They are known to increase reaction rates up to one million fold and facilitate reactions at biological conditions that would otherwise require high temperature and pressure. As of January 2021, 269,905 enzyme sequences (1) have been identified with 6,544 different reactions (2) and 108,391 protein structures (3). These enzymes are classified by the Enzyme Commission (EC) with EC numbers categorizing each reaction (4). Despite the importance and prevalence of enzymatic research, the ability to distinguish enzyme from non-enzyme sites remains an unsolved, challenging task.

Multiple physicochemical properties have been shown to be important predictors of catalytic function. Deviations in theoretical titration curves are able to identify active site residues responsible for Brønsted acid–base chemistry (5, 6). Bioinformatic studies have revealed that
catalytic residues often lie in the largest surface-accessible cleft (7) that is closest to the protein centroid (8). More recent work has shown that enzymes tend to have a network of amino acids that serve as a pathway for energy transfer (9, 10). Though these descriptive properties are helpful in enzyme site identification, they have low predictive value.

The most common way to identify enzyme sites is through homology. Native enzymes are highly homologous and so methods using homology are successful at predicting enzymatic function. Homology-based tools can detect enzyme function from the entire sequence (BLAST+) (11), sequence motif (PROSITE) (12), or a domain (Pfam) (13). These tools predict enzyme function by using previously annotated enzyme sequences or previously annotated catalytic sites and locate sequences or sites that are similar to the annotated example. Such tools allow quick mapping of previous enzymatic documentation to newly sequenced genomes. One limitation of such methods is the requirement of numerous, well-studied homologs. Another limitation is the difficulty in determining if a homolog is an enzyme or a pseudoenzyme, i.e. an enzyme homolog without catalytic activity. Though it is commonly believed that any homolog with a mutation of a catalytic residue is a pseudoenzyme, such methods over-predict pseudo enzymes by up to 1,000% (14).

An alternative strategy for predicting if a protein or site is enzymatic is to use machine learning. Machine learning generalizes important trends from training data, which can be used to make future predictions on proteins with few or no homologs. Machine learning-based methods have won the last two CASPs (15-17), a competition for de novo structure prediction, demonstrating machine learning effectiveness of predicting protein characteristics on previously unstudied proteins. There are two types of machine learning algorithms related to enzyme prediction, enzyme function predictors which mostly use sequence-data and catalytic residue predictors which mostly use structure-data.

Machine learning methods that predict enzymatic function do so by producing EC numbers (18-24). Due to the variety of enzyme functions, EC number predictors benefit from using multiple machine models, one of which predicts if a sequence is enzymatic or non-enzymatic, the next predicts the enzyme class, the next the enzyme subclass, etc. (19, 21). However, by taking a sequence level approach these algorithms miss critical active site information. A recent study demonstrated that machine learning EC number predictors and homology-based tools were rarely capable of distinguishing between native sequences and sequences where all of the residues in the enzyme site domain were mutated to alanine (20).

An alternative to enzyme function prediction is enzyme active site prediction. Unlike the layered models required for EC number prediction, methods attempting to label catalytic residues output residue-based, binary, enzymatic or non-enzymatic predictions (25-28). When identifying residues responsible for a particular class of enzymatic function, structure-based features describing the 3D neighborhood have shown tremendous success (29, 30). Methods attempting to identify catalytic residues more generally, regardless of enzymatic function, benefit from combining sequence-based features that encapsulate the large amount of available data, such
as conservation information, with structural-based features that describe the local environment (31). These methods train on datasets using all enzyme residues, labeling only the few residues responsible for catalytic activity as positives. Such imbalanced datasets result in low precision. In addition, comparing active site residues to protein surface and protein core residues may not predict catalytic activity so much as the existence of a pocket, as there are large differences in residue local environment unrelated to catalytic activity.

To address the challenges of imbalanced data and differences in local environments, we focused on the metal ions of metalloproteins. Metals – whether enzymatic or not - often lie in pockets with unusual electrostatic properties (32, 33). The residues required for coordinating metals (34) are the same charged and polar amino acids commonly used for catalytic activity (35). There are currently 53,987 crystal structures that are annotated to be metal-binding by metalPDB (36). Considering that approximately 40% of enzymes are metalloenzymes, (37), the overlap with the 108,391 structures with enzyme annotations leads to the creation enzymatic and non-enzymatic target classes with a relatively low level of imbalance compared to residue-based datasets of enzymatic and non-enzymatic residues.

In this work, we create a homology-based pipeline that identifies metalloproteins and labels the metal binding sites as enzymatic or non-enzymatic. The pipeline also removes proteins where the sequence is similar or metal binding site is similar to prevent bias and overfitting during machine learning. We then calculate structure-based features describing the environment of the metal binding sites, focusing on important catalytic properties. We use an agnostic machine learning strategy, training and optimizing several machine learning algorithms with different feature sets. The best model as determined by cross-validation was evaluated on a holdout test-set, on which it achieved a 92.2% precision and of 90.1% recall and identified enzymatic sites that were incorrectly labeled by our homology-based pipeline. We also examine the importance of the features used by our top model. Finally, we compare the performance of similar tools on our test-set, and find that our top model, using only structure-based physiochemical features, is overall superior to both enzyme function predictors and catalytic residue predictors.
Results

Data characteristics

Metalloprotein crystal structures were queried from RCSB (Research collaboratory for structural bioinformatics) and filtered for quality (Fig1A, see methods for more details). Sites were defined as the area around a metal atom. Metal atoms in the same crystal structure file within 5 Å of each other were grouped as the same site. Sites were divided into two sets. Sites from structures deposited prior to 2018 were placed in the dataset, used for algorithm optimization, model selection, and training the final model. Sites from structures deposited in 2018 or later were separated to form a holdout test-set, which was only used to evaluate the final model.

Figure 1. Workflow for dataset generation. (A) The numbers on each arrow represent the number of entries present at that step; numbers of enzymatic sites are in green, non-enzymatic sites are in blue, and not enzymatically labeled entries are in black. The final numbers are representative of the end of the computational pipeline. (B) Computational labeling of sites as enzymatic/non-enzymatic via homology expansion of M-CSA (Mechanism and Catalytic Site Atlas) (see methods for more detail).
Each site was labeled as enzymatic or non-enzymatic via a computational pipeline (Fig. 1B for further details in methods). Briefly, we identified metalloproteins that were homologous to enzymes in the manually curated Mechanism and Catalytic Site Atlas (M-CSA) database (38). For further enzymatic evidence, we required that homologs met one of two conditions: associated EC number or an “ase” suffix in structure classification or molecular name. Finally, homologs were aligned with their respective M-CSA entries to check for structural homology and to label any sites adjacent to catalytic residues as enzymatic. Any remaining sites on homologs with enzymatic sites were labeled non-enzymatic. By finding homologs to the M-CSA proteins and then removing those with structural similarity we were able to identify 12,691 metal ions located at catalytic sites, 1,089 of which were non-redundant, a nearly three-fold increase from the 316 metal ligand containing M-CSA entries (as of August 2019). Sites that were part of proteins that were not homologous to entries in the M-CSA and lacked all of the previously mentioned enzymatic evidence were classified as non-enzymatic. Because of the importance of correctly identifying enzymatic and non-enzymatic sites, any site that had some but not all of the enzymatic characteristics within our pipeline were discarded.

Biologically redundant data has shown to negatively impact machine learning models (39). Having similar data in both the training and testing set can also lead to inflated performance evaluations. To prevent these issues, similarity within and between the test-set and dataset was reduced by filtering out sites with sequence or structural similarity.

Our dataset used for ML is composed of 3,465 sites from 2,626 different PDBs; 24% of the sites are enzymatic (Supplementary file sites.csv). The test-set which is mutually exclusive from the dataset is composed of 520 sites from 400 different PDBs; 31% of the sites are enzymatic. Both sets contain sites distributed among the six major EC classes excluding the translocases a class added to the EC after the start of this project (Figure S1).

Because our test set is differentiated from our dataset by date of structure deposition, we assessed the two sets for covariate shift. Though advancements in crystallographic capabilities and changes in research funding can affect the propensities of proteins deposited in the PDB over time, no covariate shift was detectable between our final dataset and temporal test-set (see supplementary methods).

Feature analysis
In an attempt to create a machine learning model that differentiates based on physicochemical information, we used physicochemical features, including those previously mentioned to describe catalytic activity. However, we do not use features such as amino acid composition, conservation information, and secondary structure assignment (25, 26) in order to avoid biasing any algorithm towards a specific fold or function due to the wide diversity of folds utilized by enzymes and metal-binding proteins (40, 41). Moreover, metalloenzymes can be highly promiscuous with activity dependent on the identity of the metal bound (42, 43) and assigning the correct metal can be tricky in a crystallized structure (44); therefore, we also avoid using the bound metal’s identity.
In order to pass relevant, catalytic activity information to the machine learning algorithms, we developed a feature set with features from five categories—(1) Rosetta energy terms, (2) pocket geometry information, (3) terms describing the residues that line the pocket, (4) electrostatic terms, and (5) coordination geometry terms (Figure 2A). Because Rosetta energy terms are residue-based and our features are site based, different versions of the Rosetta energy term category were made by using two methods of aggregating each energy term—average or sum—and two methods of spatially defining the space around the sites—surrounding shells (0-3.5Å, 3.5-5Å, 5-7.5Å, and 7.5-9Å) or spheres (0-3.5Å, 0-5Å, 0-7.5Å, and 0-9Å) (Figure S2). In total, we used 391 features from the five categories (Table ST1), though the features were not all used simultaneously during machine learning. To efficiently search the feature space, 67 combinations of feature sets ranging in size from 4 to 181 features were evaluated during model selection (Table ST2).

To quantify the differences in feature values for enzyme and non-enzyme sites, we calculated the similarity between the enzymatic and non-enzymatic site values using a Jaccard index for discrete features and the percentage area of overlap between the kernel density estimates of the two feature curves for continuous features (see methods and Figure S3). Both methods lead to a scale where one is entirely the same between enzymatic and non-enzymatic feature-value distribution and zero is entirely different between enzyme and non-enzyme feature-value distribution. The Rosetta energy terms category that used sum sphere calculations had the most dissimilar features as demonstrated by having more points farther to the left (Figure 2B fourth row). The sum calculation magnifies differences in features by not considering effects on a per...
residue basis and the sphere calculation considers more residues. Because catalytic sites tend to be closer to the protein centroid, more residues overall contribute to van der Waals and solvation terms and sum sphere calculations are especially dissimilar. In the electrostatics category (Figure 2C first row), we find that the features farthest to the left are those that describe the deviations in the shape of theoretical titration curves for ionizable residues beyond the first shell. In the pocket category (Figure 2C second row), the volume term is substantially more dissimilar than all other terms in that category. In the lining category (Figure 2C third row), the most dissimilar features are those with describing the number and volume of residues in the pocket lining. No geometry features (Figure 2C fourth row) show particularly high dissimilarity between enzymatic and non-enzymatic sites.

Machine learning model optimization and selection
To learn and then predict enzymatic and non-enzymatic sites, we selected fourteen classification algorithms from python scikit learn that span a variety of popular machine learning methods—support vector machines (SVMs), decision-tree ensemble methods, naive Bayes, nearest neighbors, a neural network, linear and quadratic discriminant analysis (see supplement for brief explanations).

Various scoring metrics are used to evaluate binary classification models. Imbalanced data (more non-enzymatic sites than enzymatic sites) can skew some of these metrics. For example, our dataset is 76% non-enzymatic sites. Therefore, we could achieve an accuracy of 76% by predicting non-enzymatic for all of our dataset. In order to prevent the imbalance of our dataset from biasing our evaluation metrics we used the Matthews correlation coefficient (MCC, equation SE1)(45) that is less biased towards the imbalanced set. MCC values are on the [-1,1] interval, where any model predicting exclusively enzymatic or exclusively non-enzymatic would have an MCC of zero. The imbalance in our data could also produce an inflated recall, or percent of catalytic sites correctly identified, by over-predicting enzymatic sites without much of an effect on accuracy and true negative rate, or percent of non-enzymatic sites correctly identified. To avoid such over-predicting, we also prioritize precision, which is percent of catalytic site predictions that are correct.

When a model learns the details of a training set too well it can negatively impact the model performance on new data. This is called overfitting. Using our hold-out test-set and nested cross-validation (CV)(Figure S4) allowed us to use different data for model optimization, model selection, and model evaluation. During the inner CV, we optimized each algorithm for a specific feature set. We tested four different scoring metrics for optimization: accuracy, precision, MCC, or a multi-score combination of accuracy, MCC and Jaccard index. In total, 3,752 models were created (14 algorithms x 67 feature subsets x 4 optimization scoring metrics). We used the results from the outer CV to select the best of these models. However, 3,274 of the models used different “optimal” versions of the machine learning algorithm during the outer CV. To eliminate any inflated metrics that may have come from this, we re-ran the outer CV using only
the most frequently selected version of the algorithm for each model and discarded all models where large deviations persisted (Methods and Figure S5).

**Figure 3. Outer CV performance by algorithm.** (A) Each point represents the results for a specific model. Points are colored according algorithm used and grouped by classifier type; support vector machines (SVMs) are purples, decision-tree ensemble methods are blues, linear models are reds, discriminant analysis are greens, no grouping for naive Bayes, nearest neighbor, and neural network. Better performing classifiers should be close to the upper right corner. The X denotes our top model (extra trees with AllMeanSph feature set). (B) Zoomed in view of boxed region in A.

We graphed our remaining 1,668 model performances by algorithm type (Figure 3), optimization metric (Figure S6), Rosetta feature calculation method (Figure S7), feature category exclusion and feature set size (Figure S8). The only emerging trends were based on machine learning algorithm type. The neural network (neural network, Figure 3 tan) and decision-tree ensemble methods (extra trees, gradient boosting, and random forest, Figure 3 blues) perform most favorably for our prioritized metrics, MCC and precision. SVMs and linear models (logistic regression, ridge, and passive aggressive Figure 3 purples) had the highest recall (Figure S9). However, the relatively low precision of the SVM and linear models indicates that these high recall values are the result of over-predicting sites to be enzymatic.

**Top model evaluation**

The top model in our model selection was an extra-trees algorithm using all feature categories with mean, sphere calculations for Rosetta terms. We named this model Metal Activity Heuristic of Metalloprotein and Enzymatic Sites (MAHOMES) (Figure 3 ‘X’). In addition to having the best
precision relative to MCC, MAHOMES was also surprisingly stable. Three out of the four optimization strategies selected the same hyperparameter set on all seven CV folds, indicating that MAHOMES performance is not an overfit optimistic evaluation. MAHOMES was evaluated on the test-set (2018-2020 structures) where it achieved slightly higher performance metrics than its outer CV performance (Table ST3). The performance still falls within projected deviation, as observed on different test folds during outer CV, supporting the validity of the reported performance metrics.

We manually inspected the sites misclassified by MAHOMES. This included 27 of the 564 non-enzymatic sites misclassified as enzymatic (false positives) and 17 of the 185 enzymatic sites misclassified as non-enzymatic sites (false negatives) (Table ST4). Manual inspection of these sites in the literature revealed that ten of the 27 sites that had been labeled by the pipeline as non-enzymatic and as enzymatic by MAHOMES were actually correctly predicted by MAHOMES and incorrectly identified by the pipeline (Table ST4 green). All ten cases in which MAHOMES correctly predicted sites misclassified by the pipeline were sites which lacked MCS-A homologs EC numbers, and an “ase” suffix. Eight of the sites were for proteins that had not been structurally resolved before and had no homologs in M-CSA. The two sites bound to proteins that had previously been solved were both bound to sonic hedgehog proteins. Though the zinc domain of sonic hedgehog proteins was previously thought to be a pseudo-active site, more recent research indicates that it is responsible for metalloprotease activity (46, 47). In addition to the ten mislabeled sites, we were unable to definitively determine the correct label for four of the 27 the false positive sites because they are not well characterized by the literature (Table ST4 yellow). We recalculated the MAHOMES test-set performance metrics by changing the ten pipeline mislabeled sites to be true positives and excluding the four false positives we could not definitely label (Table 1).

Feature importance
Since MAHOMES is a decision-tree ensemble algorithm, it is capable of producing relative feature importance for classification (Table ST5). By graphing feature importance against our previously calculated similarity, we find that MAHOMES did not find features with high similarity to be useful in differentiating enzymatic and non-enzymatic sites (Figure 4). However, lower similarity did not always translate to higher feature importance. For example, the lowest similarity in this feature set was volume, with a similarity of 39%, meaning it is quantitatively the most useful feature, supporting previous reports of its utility for describing enzymatic active sites (7). However, volume was only the 13th most important feature for enzyme classification. So, MAHOMES finds twelve other features to be more valuable for differentiating enzymatic activity despite having more similar values for enzymatic and non-enzymatic sites. For example, the feature of the distance between the center of the active site and protein centroid, (8), is the ninth most important feature for discrimination despite a 57% feature similarity to be more important for differentiating enzymatic and non-enzymatic sites.
The most important feature for MAHOMES is an electrostatic feature derived from the shape of the theoretical titration curve of local ionizable amino acids. Specifically, it is the average of the second moment of the theoretical titration curve derivative for second shell residues (3.5Å-9Å) from the metal site. This feature was implemented due to the previous findings that residues at active site have higher deviation from the Henderson-Hasselbalch equation than other ionizable residues (5, 6, 27, 48, 49). The features of the averages of the third and fourth moments for second shell ionizable residues, while still important, were less critical, ranking tenth and nineteenth respectively. The electrostatics of the residues responsible for coordinating the metal ions (first shell) are more similar between the metalloenzyme and non-enzymatic metalloprotein sites (Table ST1) than in the second shell and this is likely preventing these descriptors from being as important. The other four of the top five features are pocket lining terms describing the number and total volume of amino acids lining the pocket.

Although MAHOMES found the previously mentioned features to be the most helpful for predicting enzymatic activity, they are not solely responsible for enzymatic activity. During the outer CV, which evaluated 3,465 predictions, an extra trees model excluding the electrostatic category only made six more incorrect predictions than MAHOMES (three enzymatic and three non-enzymatic). Another extra trees model excluding the pocket lining category correctly identified two more non-enzymatic sites than MAHOMES, but it also identified fourteen fewer enzymatic sites.
Table 1: Comparison of MAHOMES performance to similar tools that make enzymatic and non-enzymatic predictions.

<table>
<thead>
<tr>
<th>Method</th>
<th>Predictions</th>
<th>Evaluation set</th>
<th>Accuracy</th>
<th>Precision</th>
<th>Recall</th>
<th>True negative rate</th>
</tr>
</thead>
<tbody>
<tr>
<td>MAHOMES (this paper)</td>
<td>Enzymatic activity by metal binding site</td>
<td>corrected test-set (this paper)</td>
<td>94.2%</td>
<td>92.2%</td>
<td>90.1%</td>
<td>96.2%</td>
</tr>
<tr>
<td>DeepEC (20)</td>
<td>Enzymatic activity by sequence</td>
<td>sequence test-set (this paper)</td>
<td>69.9%</td>
<td>59.6%</td>
<td>90.5%</td>
<td>54.8%</td>
</tr>
<tr>
<td>DEEPre (50)</td>
<td>Enzymatic activity by sequence</td>
<td>sequence test-set (this paper)</td>
<td>90.1%</td>
<td>81.3%</td>
<td>100.0%</td>
<td>82.5%</td>
</tr>
<tr>
<td>EFICAz2.5 (51)</td>
<td>Enzymatic activity by sequence</td>
<td>sequence test-set (this paper)</td>
<td>90.8%</td>
<td>88.4%</td>
<td>90.0%</td>
<td>91.3%</td>
</tr>
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Benchmarking with other methods

Since no alternative method uses metal-binding sites as input, we adjusted the MAHOMES test-set to be a sequence test-set to compare the performance of MAHOMES to other methods. The 516 metal sites in the corrected MAHOMES test-set are on only 400 proteins that were unambiguously metalloenzymes or non-enzymatic metalloproteins. So the sequence test-set consisted of those 400 unique sequences. Sequences were labeled as enzymatic or not enzymatic as described in the methods. Using this sequence test-set, we benchmarked the performance of MAHOMES against three enzyme function predictors; DeepEC (20), DEEPre (21, 50), and EFICAz2.5 (51). MCC is sensitive to the magnitude of imbalance, so it was not used as a performance metric due to the different levels of imbalance in the corrected test-set and sequence test-set.

DeepEC (20) predicts enzymatic function using three independent convolutional neural networks which only use a protein's sequence as input. The first neural network makes a binary enzymatic or non-enzymatic prediction and was trained using 153,884 enzymatic sequences and 22,158 non-enzymatic sequences which were oversampled to provide 153,884 negative training entries. We only evaluated the performance of the first neural network since the second and third neural networks make third and fourth level EC number predictions. DeepEC had a similar recall to MAHOMES, but was by far the lowest of the evaluated methods for accuracy, precision, and true negative rate.

Similar to DeepEC, DEEPre (21, 50) also uses deep learning to predict enzymatic function. DEEPre is a level-by-level predictor, meaning it has a machine learning model for every split in
the EC hierarchy for EC numbers that it is capable of predicting. We only evaluate its level zero model, which is responsible for making enzyme or non-enzyme predictions. This model was trained using 22,158 enzymatic and 22,158 non-enzymatic sequences with low homology (40% sequence similarity threshold). In addition to the sequence, DEEPre uses some features mined from the sequence data, including predicted solvent accessibility, predicted secondary structure, and evolutionary information. The evolutionary information includes the detection of any Pfam functional domains and a position-specific scoring matrix for the whole sequence produced by BLAST+. DEEPre had a remarkable 100% recall, identifying all enzyme sequences including the ones mislabeled by our pipeline. However, it over-predicted the number of enzymatic sequences by 23%, resulting in a lower precision and recall than MAHOMES.

The final sequence-based enzyme function prediction tool evaluated on our sequence test-set was EFICAz2.5 (52-54). EFICAz2.5 combined homology detection, sequence similarity, conservation information, Pfam functional domains, the PROSITE database to generate four independent predictions and two predictions made by SVMs. These six outputs are combined using a tree-based classification model which outputs an EC number. EFICAz2.5 was very consistent across all evaluation metrics. Although MAHOMES had the highest precision and true negative rate, EFICAz2.5’s came in a close second relative to DeepEC and DEEPre.

Discussion

Non enzymatic metal binding sites

Although we find metal sites to be the best most enzyme-like sites without catalytic activity, they are not simply enzyme sites denuded of activity. The non-enzymatic metal binding sites comprise a number of types of proteins some of which may be better suited for the negative dataset than others. The most frequent types of proteins are classified as transcription, transport, signaling, and binding (proteins, DNA, RNA, sugar, or membrane). Metal transport and signaling proteins are likely to only have pockets large enough to fit the metal ions. Conversely, proteins that bind other proteins, DNA, RNA, sugar, or membranes will have ligand pockets that more closely resemble the pockets of enzymes which bind substrates of various sizes. Additionally, the non-enzymatic metal ions that do not contribute to the protein function are likely to either be of structural importance or crystal artifacts. The metals bound as crystal artifacts are likely to be bound at the surface and bound less tightly than metalloenzymes. Because our dataset has three times as many non-enzymatic sites as enzymatic sites, developing a filter for the non-enzymatic sites may improve how well the model performs on predicting sites of unknown function.

Feature importance

Many of the features of most importance to MAHOMES are similar to features previously described for determining active sites. Our most important electrostatic features are modeled on previous electrostatic features (5, 6, 27, 48, 49), but have subtle differences. Both use calculations of the moments for the theoretical titration curve first derivative. The previous work identifies active site residues by looking for clusters of residues with statistically significant
deviations. We find the averages of deviations for ionizable residues in the area beyond the coordinating residues (3.5Å-9Å) identifies metalloenzyme catalytic sites over sites that only bind metals. We find that the coordinating residue theoretical titration curves are more similar between enzyme an non-enzyme, likely because both our enzymatic and non-enzymatic sites are coordinating metals. The differences beyond the first shell likely indicate a networks of amino acids which serve as a pathways for energy transfer (55) which have been found for a number of types of enzymes (9). In addition, we also find the second moment to be most predictive whereas other studies find the third and fourth moment to be most predictive (6). Further investigation will be needed to determine the origin of this difference.

Our volume feature, modeled on a previous study (7), was the most dissimilar feature between enzymatic and non-enzymatic sites but was far from the most important feature for activity prediction. In contrast, a feature of distance between the center of the active site and protein centroid, also modeled on previous work (8), was 1.5 times as similar between classes as the volume feature but was a more important feature for activity prediction.

The second to fifth most important features are pocket lining features describing the number and volume of side chains and backbones lining the pocket. In all four of these features the enzyme dataset has more volume and more amino acids. There are two possible explanations for the importance of these feature. The first is that it may indicate that surface area of the pocket is superior to volume of the pocket in predicting enzymatic activity. The second is that because of the movement and flexibility known to be critical to enzyme function (9) the amino acids lining the pocket are a better proxy for true volume of the pocket than the volume calculation itself. Further research may help deconvolute these two possibilities to better explain what about pocket surface is most predictive of enzyme activity.

MAHOMES in comparison to other methods

Though catalytic residue predictors use structure-based features and therefore may be more comparable to MAHOMES, we were unable to directly benchmark MAHOMES performance against any catalytic residue predictors ((26), (56), (57) (25) (28) due to lack of availability of the model or the methods for implementation. When evaluated on a common dataset, CRPred (28), CRHunter (26), and PreVAIL (25), the precision of these catalytic residue predictors ranges between 14.7% to 28.6% (Table ST6), which is substantially lower than the 92.2% precision of MAHOMES. As precision is the percent predicted enzymatic sites that were correct it may be artificially lowered by the extreme imbalance of the common dataset used (an enzymatic: non-enzymatic ratio of 1:124). A more equitable comparison to MAHOMES would be recall. Because the common dataset contains a similar number of enzymatic sites as our test-set, recall—the percent enzymatic residues that were correctly identified—is a more equitable comparison. These methods report recalls ranging from 48.8% to 62.2% whereas MAHOMES scored 90.1% on recall (Table ST6). It may be that MAHOMES recall success is due to training using positive and negative sets that are more similar to each other instead of comparing catalytic residues to all other residues in a protein. As measured by difference in recall, we anticipate that by focusing on protein bound metal ions MAHOMES can correctly identify 1.3 times more catalytic...
sites on enzyme structures than the best of these three catalytic residue prediction methods. However, this success in recall does come at the cost of only being able to predict enzymatic activity for metalloproteins.

Benchmarking with our test-set demonstrated that MAHOMES had overall better performance at predicting enzymatic activity for metalloproteins than sequence-based enzyme function prediction tools. The one exception is the 100% recall by DEEPre which was better than MAHOMES 90.1% recall. However, DEEPre over-predicted enzymatic sites, as indicated by its 81.3% precision and its 82.5% true negative rate - the percent enzymatic residues that were correctly identified. The other neural network model, DeepEC, struggled the most with predicting metal catalysis, with a 59.6% precision and 54.8% true negative rate revealing that its 90.5 recall was due to over-predicting sites as enzymatic. Conversely, MAHOMES had a more balanced performance with a 92.2% precision and 96.2% true negative rate. EFICAz2.5, the most homology-based predictor, showed that not all EC number predictors suffer from enzymatic over-predictions with a 90.0% recall, 88.4% precision and 91.3% true negative rate, but it was still outperformed by MAHOMES for all performance metrics.

Differences in what features each tool used elucidate what can be used to successfully make enzymatic and non-enzymatic predictions. DeepEC relies heavily on deep learning, passing only the protein's sequence to its predictor with no other processing or features. DeepEC’s fourth-place performance demonstrates that relying only on sequence alone requires even more training data to allow for deep learning to extrapolate important features from the sequence. The other two sequence-based methods generate evolutionary information from the sequences which is combined with machine learning. Their large training sets, 44,316 sequences for DEEPre and 220,485 sequences for EFICAz2.5, provided ample data to allow for successful enzymatic and non-enzymatic predictions via homology and conservation. Our structure-based, physiochemical features led to MAHOMES making predictions by detecting catalytic activity, allowing MAHOMES to outperform DEEPre and EFICAz2.5 even though it was trained with only 7.8% and 1.6% the amount of data respectively.

Overall, we find that MAHOMES use of metal ion data with structure-based features proves beneficial when compared with both catalytic site predictors and enzyme activity classifiers. We find that MAHOMES has a higher precision than catalytic site predictors—where MAHOMES predicts nine out of every ten enzymatic predictions correct, catalytic residue predictors are only correct for one or two out of every six enzymatic predictions. We anticipate that MAHOMES relative success is due both to training on more similar sites and to less imbalance of the training set. By training on negative sites that were also in pockets and also coordinated metals MAHOMES was able to assign feature importance based on characteristics that were particular to enzyme activity. In addition, MAHOMES training set was up to 40 times more balanced than the catalytic residue predictor datasets.

MAHOMES also outperformed enzyme function predictors. These predictors were able to achieve a similar or better recall to MAHOMES by combining homology with large amounts of
data, but they struggled to correctly identify non-enzymes which were the majority class of our test-set. We anticipate that MAHOMES success over enzyme function classifiers was due to the use of structural features which are more sensitive to small differences of the active site. Though structural data is less available, it is more predictive of enzyme function than homology.

MAHOMES ability to correctly detect enzymatic activity should make it especially useful to problems where methods reliant on homology are not applicable. For example, MAHOMES may be used for eliminating poor de novo metalloenzyme designs prior to experimental testing. Another use case could be detecting when a mutation eliminates enzymatic activity. As more datasets become available, we anticipate that the MAHOMES approach will be able to be refined and deployed in future protein design and catalytic-site prediction pipelines.

Materials and Methods

**Metalloprotein ion site data**

For the purposes of our study, we focused on protein structures that contain residues codes with one or two of the following metals: iron, copper, zinc, manganese, magnesium, molybdenum, nickel and cobalt (residue codes of interest are FE, FE2, FES, FEO, CU, CU1, CUA, MG, ZN, MN, MO, MOO, MOS, NI, 3CO, CO).

RCSB (3) was queried to find the list of crystal structures containing a residue code of interest. Structures with nucleotide macromolecular types were removed. Crystal structures with a resolution greater than 3.5Å were removed to ensure high-quality feature calculations. Structures with more than 40 sites were removed as those indicated large complex structures such as virus particle and ribosomes. Protein chains with less than 20 residues were removed to avoid polypeptides. For cross-referencing and to eliminate as many de novo proteins or immunoglobulins as possible, we also removed protein chains that did not have a UniProtKB accession number (58).

For uniform handling of mononuclear and polynuclear metal sites, we defined a site as the area around metal atoms of interest attached to the same protein chain within 5 Å of each other. Because homology detection was determined by chain, sites that were within 4 Å of a residue from another chain were indicative of a possible multi-chain metal-binding site and were removed. Additionally, the few sites with more than four metal atoms, such as metal storage proteins, were removed to allow for more uniform feature calculation methodology. Sites within 10 Å of mutated residues were removed to avoid studies that identified catalytic residues through loss of activity mutations. We split the sites into a dataset and a test-set – the dataset consisted of protein structures resolved prior to 2018; the holdout test-set consisted of protein structures resolved in 2018 or later.

**Computational labeling of sites as enzymatic and non-enzymatic**

The Mechanism and Catalytic Site Atlas (M-CSA) contains manually annotated reaction mechanisms with a representative PDB structure for each entry (59). A few M-CSA entries are
undesirable for our study, such as those requiring heme cofactors or dimerization. Additionally, a few M-CSA entries’ catalytic residues were missing metal coordinating residues or included unwanted residues, such as those with steric roles. We performed our own manual annotation to adjust the data we used in these cases (Table ST7). To expand the coverage of the M-CSA data, potential sequence homologs were detected for each M-CSA entry using PHMMER (60) with a cutoff E-value of $10^{-6}$ and a database of sequences in the PDB. E-values take into account database prevalence, leading to the addition and removal of detected homologues when using updated versions of the PDB. Hence, two versions of the PDB are used to allow for updating homolog detection for the test-set (currently PDB as of May 21, 2020) without affecting the homologs detected in the dataset (PDB as of May 17, 2019). Homologues of the undesirable M-CSA entries were removed from the dataset and test-set. To avoid labeling non-enzymatic homologs, like pseudo-enzymes, we threw out homologues that did not meet at least one of the following requirements; an associated EC number, a PDB classification containing “ase”, or a PDB Macromolecule Name containing “ase”.

For further support of enzymatic evidence and identification of the homolog’s active site location, each remaining homolog was aligned to its respective M-CSA representative, using TM-align (61). We chose to use a TM-score of 0.40 or greater to represent structural homology, and discarded all aligned homologs below that cut-off. All sites that met all previous requirements and aligned within 5 Å of a catalytic residue listed in the M-CSA entry were labeled as enzymatic. For chains containing enzymatic sites, unlabeled sites were labeled as non-enzymatic. To create our non-enzymatic set, sites that were not M-CSA sequence homologues, had no associated EC number, no “ase” in the PDB classification, and no “ase” in the PDB Macromolecule Name were labeled non-enzymatic. Remaining unlabeled sites in the dataset and test-set were removed at this point.

Removal of redundant sites
There are many redundant protein structures—proteins that are highly homologous or identical proteins with different ligands. To prevent redundancy from biasing our training toward one particular protein we implemented the following method of intra-set redundancy removal. First, we removed proteins that were sequence redundant. We removed all but one instance of identical sequences. Then sequence homologs were assembled and clustered using PHMMER taking from the PDB on the dates mentioned and using an E-Value of $10^{-20}$. This homolog collection and clustering were executed independently for the dataset and test-set. Next, we removed site redundancy. Chains within a cluster were aligned using TMalign and sites that aligned within 5 Å with a ≥ 0.50 TMscore were checked for local similarity. We defined local site similarity as the Jaccard index of residue identities within a 3.75 Å radius for two sites. Sites were removed if they had a local similarity greater than 0.80. Due to high preprocessing computational costs, sites that had already undergone relaxation were selected when possible. Otherwise, we used the following priority to keep the more favorable site: 1) catalytic site, 2) no other ligands present, 3) no other sites within 5 Å to 15 Å, 4) no mutations in PDB, 5) crystal structure resolution.
Evaluating machine learning models on the same data used to train and optimize them can lead to overfit, inflated results. The temporal separation of the test-set and dataset prevents the same structure from being in both sets. However, the aforementioned redundancy of protein structures, different structures of the same protein, or closely related homologs can still lead to reporting an inaccurately high machine learning performance. We used an all against all method to compare the residue identities within a 6.0Å radius of all remaining sites, removing sites with greater than 0.80 similarity.

Further data processing
The pipeline described above displayed catalytic identification errors with electron transport proteins, kinesins and NTPases, in part because they were inconstantly labeled in the training dataset. In order to prevent further errors, we removed electron transport proteins and kinesins as identified by PDB classification or molecular name. To remove all ATPase and GTPase sites, we removed all Mg sites within 10 Å of a nucleic-phosphate ligand that were labeled non-enzymatic.

Finally, when the structures were relaxed using Rosetta (see Supplement for RosettaScripts inputs), 728 sites with loosely bound metals, often the result of crystal artifacts, that moved more than 3Å during relaxation were removed from the dataset and test-set. Also, 179 sites were removed due to failure to complete feature calculations.

Features
Five feature categories were calculated on the relaxed structures – Rosetta energy terms, pocket void, pocket lining, electrostatics, and coordination geometry (Figure 2A; Table ST1). To prevent outlier values from affecting out models, all features were scaled (normalized) using sci-kit’s RobustScaler prior to machine learning. The scaler was fit to the 20th and 80th quantile of the dataset and used to transform the dataset and test-set.

1. Rosetta energy terms
Rosetta features values were assigned to all sites using all the terms in the energy function beta_nov16. Rosetta assigns a value for each term to each residue. We used the sum or the mean of the per-residue Rosetta energy terms as features, calculating terms in spheres and in shells using the cutoffs 3.5Å, 5Å, 7.5Å and 9Å. This results in 21 features per cutoff and 84 features per calculation method. The four groups of Rosetta terms were never included together in any model.

2. Pocket void terms
Rosetta’s pocket_measure application (62) was executed on all individual sites, using the residue closest to the site center to anchor the grid. This maps the pocket to a 0.5Å-interval grid. Using these pocket grid points, we determined the city block distance from site center to the center of the pocket opening. Volumes were calculated by Rosetta and depth was taken as the distance between pocket opening and site center.

To quantify the shape of the pocket, we took three slices of pocket points (bottom, middle, and top) and projected them into 2D space after rotating the pocket so that the z-axis runs from site
center to center of the pocket opening. For each slice, we calculated the farthest distance between two points, the area formed by a 2D slice that encompassed all points, the distance from the origin to the center of the ellipse that encompasses the convex hull, and the two radii of the ellipse. These calculations result in 20 features.

3. Pocket lining terms

The grid produced by Rosetta’s pocket_measure also allows us to identify and describe the residues that line the pocket. Residues were split into backbone-only (where exclusively backbone atoms are adjacent to a pocket grid point) and sidechain (where any sidechain atom of a residue is adjacent to a pocket grid point; adjacent sidechain and backbone atoms of the same residue are included in this group). We then calculated the average, minimum, maximum, skew, and standard deviation of the hydrophobicity for the side chain residues using two different scales—Eisenberg (63) and Kyte-Doolittle (64). We also calculated the occluding volume of the pocket as the sum of the van der Waals volume of the sidechains in the pocket which allowed us to determine the total volume of the pocket without sidechains and the Leonard Jones volume of that pocket occupied by sidechains. Finally, we calculated the surface area of the pocket walls by summing the solvent-accessible surface area for any sidechain-adjacent residue. This resulted in 16 features.

4. Electrostatics

Our electrostatics features are based on the use of theoretical titration curves by THEMATICS (65, 66) which showed that the theoretical pKa curves of ionizable side chains deviate from expected Henderson-Hasselbach behavior (48). We used blues (67) to calculate our theoretical titration curves for all ionizable residues (see supplement for command line options), which is a generalized Born solution to the Poisson-Boltzmann equation rather than a finite difference solution and therefore quickly calculates the electrostatics for our dataset.

We calculated the mean and max of the second, third and fourth central moments of the theoretical titration curves for all residues in two shells. The first shell included residues with a C$_\alpha$ atom within 3 Å of the site center; the second shell included residues with a C$_\alpha$ atom between 3-9 Å of the site center; the shells are labeled “Inside” and “Outside”, respectively. We also calculated three environmental features for the site center as defined in Somarowthu, et al. (2011):

$$x_{env}^n = \frac{\sum_r w(r)x(r)}{\sum_r w(r)}$$

where $r$ is an ionizable residue with a distance < 9Å from the site center, $x(r)$ is the second, third or fourth central moment of $r$, and $w(r)$ is the distance weighted to be inversely proportional to the squared distance. We calculated $x_{env}^n$ of the second, third, and fourth moments. The THEMATICS features are based on single-residue calculations for the third and fourth moments; we calculate per-residue terms for the second, third, and fourth moments using the average of the residues in the first shell (0-3.5Å) and second shell (3.5-9Å).
Blues (67) also provides information about the pKa shift from an ideal amino acid to the amino acid pKa in the context of neighboring ionizable residues. Because pKa shifts are observed in catalytic residues (68-70), we include the minimum and maximum of these values for the same two shells as the central moments described above. We also calculated the residuals for the deviation of the theoretical titration curve from a sigmoidal curve; we similarly calculated the mean and max of these values in the two shells as described above. Residues adjacent to active sites often rank among the most destabilizing (positive) $\Delta G_{elec}$ values of a protein (71); we use the solvation energies calculated by blues as a proxy and rank all residues from highest to lowest solvation energies. Residue ranks are then split into five bins to avoid length-dependent ranks; destabilizing ranks run from highest to lowest solvation energies while stabilizing ranks were assigned from lowest to highest solvation energies. We then calculated the mean and max in the two shells as well as the environmental rank as described above. Overall, there are 37 electrostatic features.

5. Coordination geometry

FindGeo (72) and CheckMyMetal (73, 74) are both webservers that provide information about the coordination geometry of bound metals in crystal structures. We added functionality to FindGeo’s python portion to calculate features from CheckMyMetal. FindGeo compares the coordination of a metal atom in a PDB to a library of 36 idealized geometries and, in addition to the name of the coordination geometry, determines whether the geometry is irregular, regular, or distorted. For each site, we record which of the 36 geometries are present. However, these features are not used in training because assignments were diverse enough that it lead to problems of sparse data. Instead we recorded coordination number based on the 36 ideal geometries and determined whether the geometry included a vacancy.

We also recorded the numbers of nitrogen, oxygen, sulfur, and other atoms coordinating the metals in each site. Both the total number of each coordinating atom and the average number per metal are included as features. We calculated the overall charge on the site from the FORMUL lines of the PDB. Four terms from CheckMyMetal are also calculated – the sum of the angular deviation from the ideal geometry, the maximum single angle deviation, the bond valence (which is the sum of the individual valences), and the normalized bond valence. Perfect geometries are defined to have a normalized bond valence of 1. These features all attempt to describe how far from physically ideal the metal binding site is. In total there are 24 coordination geometry features.

Feature similarity

We calculated feature similarity for discrete features – those that can take fewer than 21 unique values - using the Jaccard similarity between the proportions observed in the enzymatic and non-enzymatic sites. In this case, the Jaccard similarity is equal to the sum of the minimum values divided by the sum of the maximum values. For example, if 50% of the non-enzymatic sites have regular geometry and 50% do not, and 75% of the enzymatic sites have regular geometry and 25% do not, the Jaccard similarity is $(0.5+0.25)/(0.5 + 0.75) = 0.6$. 


We calculated feature similarity for our continuous features, as the overlap coefficient which is the area under the minimum of two curves or the shared area of two curves. First, we fit a kernel density estimator (KDE) to the enzymatic values and to the non-enzymatic values. The KDEs were evaluated at $2^{10}+1$ data points equally spaced from the minimum and maximum values of the feature across the whole training set. Then, Romberg integration was used to calculate the area under the minimum of the two evaluated KDEs.

**Machine learning**

We selected fourteen classification algorithms readily available in Python that cover a variety of popular methods for machine learning – linear regression, decision-tree ensemble methods, support vector machines (SVMs), nearest neighbors, Bayesian classification, and simple neural networks. A hyper-parameter search space of eight to twelve hyperparameter sets was selected for each algorithm (see supplemental for algorithms and search space). The algorithms require or greatly benefit from normalizing the features values. We used sci-kit’s RobustScaler with the 20th and 80th quantiles to limit the effect of outliers during scaling. The scaler was fit to the dataset and used to transform both the dataset and test-set. Due to the imbalance of target classes (more non-enzymes than enzymes), the dataset was randomly under-sampled at a ratio of three non-enzymatic sites to one enzymatic site.

A nested cross-validation strategy was used for model optimization to avoid overfitting and bias. Each inner loop used GridSearch with StratifiedShuffleSplit (in the python scikit-learn package (Pedregosa, et al. 2011)) and was optimized four times for each of four scoring terms – accuracy, precision, MCC, or a multi-score combination of accuracy, MCC and Jaccard index. An outer loop used a stratified k-fold cross validation. The most frequently selected hyperparameter set during the outer cross validation was considered optimal for the model. The dataset was under-sampled once prior to model optimization.

In total, we examined 3,752 machine learning models (14 algorithms x 67 feature sets x 4 optimization terms). For model selection, we re-ran the outer cross validation using only the optimal hyper-parameter set. During stratified k-fold cross validation, the data is divided into k groups (k = 7), each with the same number of positive and negative entries. All except for one of the groups are used to train a model and left out group is used to evaluate that model. This is repeated k times, leaving out a different group each time, and the performance is averaged. Our random-sampling, and some of the machine learning algorithms that require random sampling, are susceptible to differences in the machines on which they are executed. In order to produce more reliable performance evaluations for model selection, we repeated each iteration of the outer cross validation ten times when we re-ran it. During each repetition, a new random seed was passed to the machine learning algorithm and used to under sample the training folds. Since we used k=7, the reported outer cross validation results are the average of 70 different models (7 folds, each with 10 different random seeds).

The second run of the outer cross validation displayed a much higher performance deviation for the different folds (figure S5), supporting that a large number of models had overfit evaluations.
due to changing hyperparameter sets during the initial outer cross validation. To avoid selecting a model that only performed well for specific data, we filtered the results to keep models that met several conditions: Accuracy standard deviation <= 6.5%, True Negative Rate standard deviation <= 9%, and MCC standard deviation <= 0.11. Only 1,668 of 3,752 models considered met these requirements (Supplemental Figure S4). Our top model was an extra-trees algorithm using all feature categories with mean, sphere calculations for Rosetta terms because it had high MCC, high precision, and converged on the same optimal extra trees algorithm for three of the four optimization metrics.

A holdout test-set was used for our final performance evaluation of the selected top model. Similar to the second outer cross validation run, we repeated the predictions ten times. Each repetition used a new random seed for the extra tree’s algorithm and a different random under-sampling the non-enzymatic dataset sites. If the average of the ten predictions was < 0.5 the site was classified as non-enzymatic if the average was ≥ 0.5 the site was considered enzymatic.

Finally, we used the scikit-learn ExtraTreesClassifier feature importance to output the impurity-based feature importance. This feature importance is calculated based on how much each feature contributed to the reduction of the Gini criterion for the extra trees model. During each repetition, the output impurity-based feature importance was normalized relative to the largest value, giving feature importance between 0 and 100. The results from the ten repeated runs with different random sampling were averaged to give the final feature importance (Table ST5).

**DeepEC, DEEPRe, and ENZYMz2.5 method evaluations**

After removing the inconclusive false positives from the test-set, the set contained 516 sites on 400 unique sequences. The sequences were used for testing sequence-based classifiers for metrics comparison. Because sequences could contain both positive and negative metal binding sites, sequences were labeled as enzymatic if they met at least one of the following criteria: an associated EC number, an “ase” suffix in structure classification or molecular name, or previously identified to be enzymatic by manual inspection. Sequences that did not meet any of the previous criteria were labeled as non-enzymatic. The sequence test-set contained 170 enzymatic sequences and 230 non-enzymatic sequences.

EFICAz2.5 was download from https://sites.gatech.edu/cssb/eficaz2-5/. We used it to make a prediction for all 400 sequences in the sequence test-set. Predictions containing "No EFICAz EC assignment" were considered non-enzymatic and all other predictions were considered enzymatic. DeepEC was downloaded from https://bitbucket.org/kaistsystemsbiology/deepec/src/master/. DeepEC was only able to make predictions for the 396 sequences that were less than 1000 residues in length. The output “enzyme_prediction.txt” file was used for the performance evaluation. Finally, the 393 sequences that had between 50 and 5000 residues were uploaded to webserver link provided in DEEPre’s publication, http://www.cbrc.kaust.edu.sa/DEEPre/index.html. The prediction was considered non-enzymatic if the first digit was ‘0’ and enzymatic for all other first digits.
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**References**


