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Machine learning applied to predicting microorganism growth temperatures and enzyme catalytic optima

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Abstract

The optimal growth temperature (OGT) of organisms is an important index to estimate the stability of enzymes encoded in their genomes. However, experimental determination of OGT for microorganisms that cannot be cultivated is difficult. Here, we report on the development of a machine learning model that can accurately predict OGT directly from proteome-wide 2-mer amino acid composition. We make use of this model to predict OGTs for 1,438 microorganisms. In a subsequent step we combine OGT data with amino acid composition of individual enzymes to develop a second machine learning model for prediction of enzyme temperature optima ($T_{\text{opt}}$). The resulting model is far superior to using OGT alone for estimating $T_{\text{opt}}$ in a dataset of 2,609 enzymes. Finally, we predict $T_{\text{opt}}$ for 6.5 million enzymes, covering 4,447 EC numbers, and make the resulting dataset available for researchers, enabling simple identification of enzymes that are potentially functional at extreme temperatures.
Introduction

The optimal growth temperature (OGT) of microorganisms is an important physiological parameter. Thus, OGT has been widely used to understand the strategies organisms use to adapt their genomes and proteomes to different environmental conditions\textsuperscript{1–3}. OGT has also been an important tool for identifying thermostable enzymes\textsuperscript{4} in the field of molecular biology (for example the Taq DNA polymerase) and in industrial applications (for example enzymes for biorefinery processes)\textsuperscript{5–7}.

Determining the OGT of a microorganism is a laborious process that requires cultivation in temperature-controlled conditions. Furthermore, the number of microorganisms that can be cultured in the lab is only a small fraction of the total diversity in nature\textsuperscript{8}. Consequently, the OGT for the vast majority of microbial organisms is currently unknown, and an easy way to computationally estimate the OGT of microbes is in demand. For such computational estimations to be feasible there must be general trends for how quantifiable biological properties change with growth temperature.

Previous studies have revealed many genomic and proteomic features that are strongly correlated with OGT. Examples include the existence of thermophile-specific enzymes\textsuperscript{9}, the presence or absence of certain dinucleotides\textsuperscript{10}, the GC content of structural RNAs\textsuperscript{11}, as well as amino acid composition of the proteome\textsuperscript{1,12}. Examples such as these indicate that estimating OGT directly from genomes or proteomes may indeed be feasible.

Statistical tools, such as regression and classification, have been used to model the correlation between OGT and biological features. For example, the OGT of 22 bacteria could be predicted using a linear combination of either dinucleotide or amino acid composition\textsuperscript{10}. Additionally, Zeldovich found that the sum fraction of the seven amino acids I, V, Y, W, R, E and L showed a correlation coefficient as high as 0.93 with OGT in a dataset consisting of 204 proteomes of archaea and bacteria\textsuperscript{12}. Jensen et al developed a Bayesian classifier to distinguish three thermophilicity classes (thermophiles, mesophiles and psychrophiles) based on 77 bacteria with known OGT\textsuperscript{13}. Training datasets containing the OGTs for a large number of organisms have been hard to obtain, something which has prevented the development of state-of-the-art machine learning models for OGT prediction.

Here we use a recently published dataset which contains OGTs for 21,498 microorganisms\textsuperscript{14}, and combine this with genomic resources to build a machine learning model. The model predicts OGT from a file containing all proteins encoded by an
organism's genome. We show that the optimal growth temperature of a microorganism can be accurately predicted solely by the amino acid sequences of its proteome. The model was then used to assign an OGT value for those microorganisms for which we had no experimental values. In a second step we make use of the resulting OGT dataset to improve the prediction of enzyme temperature optima ($T_{opt}$) by another machine learning approach. The resulting model was used to estimate $T_{opt}$ of enzymes in BRENDA. Finally, to make easy use of our OGT model and generated datasets we develop a computational method which we call Tome (Temperature optima for microorganisms and enzymes) and make it freely available for reuse (https://github.com/EngqvistLab/Tome). This method has two applications. First, it can be used to predict OGT from the amino acid composition of proteins encoded by an organism's genome. Second, it can be used to find enzyme homologs with a predicted $T_{opt}$ in a given temperature range. Both these applications will be highly valuable for the microbiology, protein engineering and biotechnology communities.

**Results**

**Collection of optimal growth temperature and proteomes of microorganisms**

To build a machine learning model that can predict OGT from the amino acid composition of proteins encoded by an organism's genome we first established a training dataset. To this end, we downloaded an OGT dataset (https://doi.org/10.5281/zenodo.1175608), which contains data for 21,498 microorganisms, including bacteria, archaea and eukarya. Using this dataset, all proteins from 5,761 organisms from RefSeq (https://www.ncbi.nlm.nih.gov/refseq/) and Ensembl genomes (http://ensemblgenomes.org/) could be associated with an OGT value (we refer this as the annotated dataset), while proteins from an additional 1,803 organisms could not be associated with an OGT value (we refer this as the unannotated dataset) (Fig. 1a).

For each organism in both the annotated and unannotated dataset we calculated the global amino acid monomer and dipeptide frequencies. However, some organisms in the dataset contain only a small number of protein sequences, as a consequence the amino acid composition obtained from those sequences may not represent the true amino acid composition of the complete proteome. To address this problem we applied a filtering step. As it was unclear how many protein sequences are required to obtain a stable amino acid composition we designed three different metrics (see Methods for details) to test how much protein sequence data was needed to obtain a stable amino
acid composition. (Fig. 1b). For each organism in the annotated dataset the three metrics were calculated for every protein sequence added in order to observe at which point the values stop fluctuating. Using this analysis on amino acid monomer frequencies we found that at least $10^5$ amino acids are needed to get a stable amino acid composition (Fig. 1c, d, e). Repeating this analysis for amino acid dipeptides resulted in the same threshold (Supplementary Fig. 1).

A further concern was that the order in which proteins appear in the input files may affect our cutoff analysis. For this reason, proteins from 17 organisms with different sizes of available proteomes were randomly selected. For each of these organisms the order in which protein sequences appear was shuffled and the three metrics were calculated. The shuffling, with subsequent analysis, was repeated 100 times. As expected, the analysis shows a high initial variability, where few sequences have been analyzed, but with increasing numbers of averaged proteins the values stabilized and converged (Fig. 1f, g, h). From this analysis it is clear that the arrangement of proteins in a proteome has a negligible effect when the proteome size is larger than $10^5$, and we therefore only chose organisms with at least $10^5$ amino acids in the dataset for further analysis.

This approach resulted in a dataset with 5,532 organisms annotated with OGT and 1,438 un-annotated organisms. In the annotated dataset the magnitude of protein number in each organism follows a normal distribution centered around 3,000 (Fig. 1i). The OGT distribution is, however, highly skewed with the majority of organisms having an OGT in the range 25-30°C and at 37°C (Fig. 1j). The number of organisms in the data set with an OGT higher than 40°C is 425 (340 for higher than 50°C). The majority of microorganisms in the dataset are bacteria, with around 12% being archaea and eukarya (Fig. 1k).
Figure 1 | Variability in amino acid frequencies decrease log-linearly with proteome size. (a) Schematic overview of process to build a machine learning model to predict OGT. Protein records from Ensembl genomes (bacteria and fungi) and RefSeq (bacteria, archaea and fungi) were downloaded. Sequences were annotated with the growth temperature of the organism from which they originate. Sequences from organisms that could not be annotated, i.e. for which there is no available information about the OGT for the organism, were retained in a separate un-annotated dataset. Amino acid frequencies of the annotated sequences were used to train a statistical model. This model was in turn used to predict growth temperatures for the un-annotated dataset. OGT: optimal growth temperature. (b) The frequency of each amino acid was plotted against the number of amino acids used to calculate the
frequency. The final third part was fitted to a linear model to get the absolute slope value (|a|), as well as its frequency variance (a_i) and varying range (R). The maximal |a|, a_i^2 and R of 20 amino acids of each proteome (r_{abs,max}, a_{max}^2 and R_{max}) give measures of whether frequencies were stable. The calculated (c) a_{max}^2, (d) r_{abs,max} and (e) R_{max} of all species in the dataset were plotted against the number of amino acids in the proteome. The dashed line indicates the cutoff for the selection of proteomes based on size. Effect of protein order on (f) a_{max}^2, (g) r_{abs,max} and (h) R_{max}. 17 proteomes with different size were randomly selected. Proteins in each proteome were shuffled 100 times and the three metrics for each shuffled proteome were calculated. (i) Distribution of proteome sizes in the annotated dataset after filtering. (j) Distribution of growth temperatures in the annotated dataset after filtering. (k) Proportion of species belonging to the three different taxonomic superkingdoms in the filtered dataset.

**OGT can be accurately predicted from amino acid composition of the proteome**

Protein amino acid composition is strongly correlated with OGT^{10,12}. For this reason we decided to train machine learning models using the amino acid composition as features. For each organism in the annotated dataset we calculated the global amino acid monomer frequencies (20 features) as well as amino acid dipeptide frequencies (400 features). To get the best feature set and statistical model for the prediction of OGT, we tested six different regression models and compared their performance on the monomer dataset and the dipeptide dataset. As shown in Fig. 2a, a 5-fold cross-validation was applied to evaluate the performance of different regression models. Using the 20 amino acid frequencies, non-linear models (SVR and Random forest) perform much better than linear models (Linear, Elastic net, Bayesian ridge regression). The superior performance of non-linear models suggests that there are important non-linear relationships between amino acid frequencies and OGT. In contrast, all models except decision tree show an almost identical performance when using dipeptide frequencies. Since models trained on each of the two datasets individually show good performance we reasoned that models trained on the combined datasets may be even better performing. However, contrary to this expectation the six models trained using both monomer dataset and dipeptide dataset together do not show improvement (Fig. 2a).

**Validation of the SVR model for growth temperature prediction**

In our comparison of different models we found that an SVR model trained on the dipeptide dataset produces the best OGT prediction results. The final SVR model was trained on the whole dipeptide dataset and stored for further use. This model can explain an astounding 95% (88% by cross-validation) of the variance in OGT (Fig. 2b). Leveraging this model, the OGT of 1,438 organisms in the unannotated dataset were predicted (Fig. 1a).

The OGT predictions were validated using two separate approaches. First, we
performed a manual literature search to find experimentally obtained OGTs for a subset of
the organisms (for which no experimental OGT was present in our original dataset).
We randomly sampled 54 of the organisms with predicted OGTs, in a manner that
ensured even spread across temperatures. For 45 of the 54 organisms, OGT values
could indeed be found in published peer-reviewed articles (Supplementary Table 1).
The agreement between the predicted OGT and the ones collected from literature is
very high, with a Pearson’s correlation coefficient of 0.96 (Fig. 2c). Second, we seized
on the fact that the average temperature optimum of catalysis ($T_{\text{opt}}$) of at least five
enzymes from an organism shows a Pearson correlation above 0.75 with growth
temperature$^{14}$. Essentially, the catalytic optimum of an enzyme tends to be close to the
organism growth temperature. The correlation between OGT and $T_{\text{opt}}$ provides a way to
validate predicted OGTs using experimental data for enzyme temperature optima
obtained from the BRENDA database (https://www.brenda-enzymes.org/). Of the 1,438
organisms with predicted OGT only 23 were found to have at least five enzymes with
$T_{\text{opt}}$ available in BRENDA. Plotting the mean enzyme optima against the predicted OGT
for these organisms reveals a strong correlation, with a Pearson’s correlation coefficient
of 0.77 (Fig. 2d). Indeed, this correlation is the same as that obtained with
experimentally determined organism OGTs$^{14}$, again showing that the predicted OGTs
are very accurate.

Figure 2 | Model development for OGT prediction. (a) $R^2$ score obtained by a 5-fold cross-validation for
six different regression models. Error bars represent the standard derivation of $R^2$ scores. (b) Performance of the final SVR (support vector regression) model trained on dipeptide data. The correlation
between predicted organism growth temperatures and those present in the original annotated dataset

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Improved estimation of enzyme temperature optima using machine learning

In biotechnology and protein engineering OGT is often used to guide the discovery of thermostable enzymes\textsuperscript{7}. Even though this strategy has proven very useful, OGT yields an imprecise estimation of enzyme temperature optima ($T_{\text{opt}}$), as indicated by the Pearson correlation between the $T_{\text{opt}}$ of individual enzymes and OGT being as low as 0.48\textsuperscript{14}. We hypothesized that the accuracy of this estimation could be improved using machine learning. We collected 2,609 enzymes that have both $T_{\text{opt}}$ and protein sequence data in the BRENDA database\textsuperscript{15}, and that could also be mapped to organisms with experimental OGT using the annotated dataset (Fig. 3a, b). We first tested the accuracy of using OGT as an estimation of $T_{\text{opt}}$ and found that only 25% of the enzyme $T_{\text{opt}}$ variance could be explained (Fig. 3c, black bar). Then, to improve the accuracy of this estimation using machine learning, we extracted three feature sets from the enzyme sequences, namely amino acid frequencies, dipeptide frequencies and other basic protein properties like length, isoelectric point etc. (See Methods). Six regression models were trained and tested on these feature sets individually, as well as the two and three sets combined, with a 5-fold cross-validation approach. As shown in Fig. 3c, the best model (SVR) trained on amino acid frequencies achieved a slightly improved accuracy compared to OGT, as quantified by an $R^2$ score of around 30%. Using dipeptide frequencies alone in combination with amino acid frequencies did not further improve the accuracy.

Since OGT and sequence-derived features each produce estimates of similar accuracy (25% and 30%, respectively) we tested whether their combined use could boost predictive power. Surprisingly, the best model (random forest) trained on the combination of amino acid frequencies and OGT almost doubled the model predictive accuracy to over 50%. Further inclusion of other basic enzyme properties (see Methods) did not further improve the accuracy (Fig. 3c).

To generate a final model for the prediction of enzyme temperature optima the random forest model was re-trained using the full set of amino acid frequencies and OGT data.
(Fig. 3d). In this final model, OGT of the source organism is the most informative individual feature, whereas the 20 amino acid frequencies combined contribute over half of the predictive power of the model (Fig. 3e).

**Figure 3 | Model development for prediction of enzyme temperature optima.** (a) Schematic overview of process to build a $T_{\text{opt}}$ prediction model. (b) The distribution of enzyme temperature optima in training dataset. (c) 5-fold cross-validation results for five regression models on different feature sets. The $T_{\text{opt}} = \text{OGT}$ bar shows the explained variance when using OGT as the estimation of enzyme $T_{\text{opt}}$. Error bars shows the standard deviation of $R^2$ scores obtained in 5-fold cross validation. AA, amino acid frequencies; Dipeptide, dimer frequencies; OGT, optimal growth temperature of source organism; Basic, basic information of proteins, like length, isoelectric point etc., see details in Methods section. (d) Performance of the final random forest model trained on AA+OGT data. The correlation between predicted and experimental $T_{\text{opt}}$ was evaluated. RMSE: root mean square error. Colors indicate the density of the points. (e) The feature importance in the final random forest model. Error bars indicates the standard deviation of feature importances of 1,000 estimators.
Annotating enzymes in BRENDA using OGT and predicted $T_{opt}$

Currently, a main resource for $T_{opt}$ data is the BRENDA database\textsuperscript{15}. However, there are approximately 12 million native protein sequences in BRENDA while there are only about 33,000 $T_{opt}$ records, many of which are not connected to a protein sequence. Due to the very small number of features (20 amino acid frequencies plus the OGT of the source organism) required to predict $T_{opt}$ using our random forest model, it is computationally feasible to carry out that prediction for millions of enzymes in BRENDA.

First, experimentally determined OGTs\textsuperscript{14} and the 1,438 OGTs predicted with the SVR model (Fig. 2b) were combined to generate a dataset containing the OGT of 22,936 microorganisms. Using this combined OGT dataset 6,507,076 out of 12,115,011 enzymes (54\%) in BRENDA could be annotated with the OGT value of their source organism, of which 909,954 enzymes (14\%) were contributed by the predicted OGT values. In a second step, these OGT annotations were combined with the amino acid frequencies extracted from each enzyme sequence. Our random forest model (Fig. 3c) was applied to this data to estimate the $T_{opt}$ of each individual enzyme. This prediction dramatically added to the experimental $T_{opt}$ values in BRENDA, increasing them 197-fold (Fig. 4a) and covering 4,447 different EC numbers (Fig. 4b). Moreover, the temperature coverage, i.e. the minimal and maximal $T_{opt}$ for an enzyme class, of the vast majority these EC numbers (3,721 of 4,447) were expanded (Fig. 4b). The predicted enzyme $T_{opt}$ and annotated OGT values of these enzymes are freely available for download and re-use (https://zenodo.org/record/2539114, https://doi.org/10.5281/zenodo.2539114).

As can be seen in Fig. 4c, many of the predicted enzyme $T_{opt}$ values differ significantly from the OGT of the source organism. For enzymes from organisms with OGT below 40\(^{\circ}\)C many have $T_{opt}$ higher than the OGT. In contrast, enzymes from thermophiles generally have a lower $T_{opt}$ than the OGT. These results are in good agreement with previous findings comparing experimental OGT of organisms with average enzyme $T_{opt}$\textsuperscript{14}. For three representative organisms we show that the distribution of predicted $T_{opt}$ values are indeed consistent with experimental values (Supplementary Fig. 4).

Tome: a command line tool for OGT prediction and identification of enzyme homologues with different $T_{opt}$

To ensure easy access to the OGT predictive model for the scientific community, as well as the enzyme data with estimated $T_{opt}$, we developed the command line tool Tome (Temperature optima for microorganisms and enzymes). This tool is simple to use and has two fundamental applications: (1) prediction of OGT from a file containing protein sequences encoded by an organism’s genome; (2) identification of functional
homologues within a specified temperature range for an enzyme of interest. For the prediction of OGT, a list of proteomes in fasta format is provided as input and the temperature predictions are returned as an output. While this tool will perform predictions on any input given, we stress that the tool has been trained on bacteria, archaea and a only small set of eukarya - mostly fungi and protists. Predictions on organisms which do not fall into these categories may result in inaccurate results. For the identification of enzyme functional homologs with different estimated $T_{\text{opt}}$, one can either simply specify an EC number and temperature range of interest to get all enzyme sequences from BRENDA matching the criteria. Alternatively, the sequence of an enzyme of interest can be provided in fasta format. The algorithm will then perform a protein BLAST and an additional output file will be generated containing only homologous enzymes (default e-value cutoff is $10^{-10}$) within the specified temperature range. Full instructions regarding installation and usage of the Tome tool is available online (https://github.com/EngqvistLab/Tome).

Figure 4 | Prediction of enzyme temperature optima. (a) Visual representation of the number of the enzymes with experimental $T_{\text{opt}}$ in BRENDA and the number of enzymes for which $T_{\text{opt}}$ was predicted leveraging experimental and predicted OGTs. Each box represents ~33,050 enzymes. There are 12,115,011 enzymes in total. Pred. is an abbreviation of predicted. (b) A visual representation of the $T_{\text{opt}}$ temperature coverage for each EC number after annotation. The span between the highest and lowest $T_{\text{opt}}$ for each enzyme is indicated. Experimental (BRENDA) and predicted $T_{\text{opt}}$ values are shown in different colors. (c) Comparison between OGT of source organism and predicted and experimental $T_{\text{opt}}$ values of enzymes. Colors indicate the density of the points.
Discussion

A main finding of this study is that the OGT of microorganisms from the three domains of life can, using non-linear regression models, be accurately predicted from the sequence information of proteins encoded by their genomes. The fact that the same predictive model can be used for widely different organisms implies that thermoadaptation in proteins follow generalizable evolutionary trends. What those trends are is the topic of a forthcoming study by these authors. The OGT prediction model generated in this study should prove useful in assigning growth temperatures for un-culturable organisms for which sequence data is available, and even for those microorganisms where OGT has been determined but where the information is absent from major databases.

Our OGT prediction model is the most accurate model to date when compared to other published models (Supplementary Fig. 2 and Supplementary Fig. 3). We propose that the high predictive accuracy seen in our OGT prediction model results from two features of our approach; the size and quality of the training data used, and the use of non-linear regression models. Our training dataset consisted of 5,533 microorganisms, including 4,974 bacteria, 222 archaea and 337 eukarya. This training set is much larger than those used in other approaches, such as 22 bacteria, 77 bacteria, 204 prokaryotes. As a direct consequence of the increased size of the dataset, we could train models that are more general applicable. We find that in general non-linear models outperform linear models when using amino acid frequencies (Fig. 2a). This suggests that the linear models used previously such as that from Nakashima et al. might be further improved by non-linear regression to correlate the amino acid frequencies to OGT.

While there are only a few published models predicting organism OGT, there are many methods for the estimation of protein stability. These methods fall into two main categories; predicting the stability of whole proteins, and predicting the stability change in a protein upon amino acid substitutions. Machine learning has been used extensively for the prediction of stability change upon amino acid substitutions, while only a few methods have been developed for the prediction of stability of whole protein empirically. The desire to accurately predict protein stability and stability changes largely stems from a real-world need to engineer proteins with increased thermostability, as well as identifying natural ones that are already stable, for use in industrial applications. However, computational prediction of protein stability is challenging since it usually needs an accurate calculation of Gibbs-free energy change of protein unfolding process, which relies mainly on high-quality protein structures, which are
limited in number.

Another important finding of this study is that $T_{opt}$ values of enzymes can be predicted from a combination of enzyme sequence information and physiological parameters of the source organisms (OGT in the present study). OGT has been used to estimate protein stability$^4$, which is based around the fact that a typical protein should be functional at the growth temperature optimum of its parent organism, something which is supported by a strong correlation between OGT and average enzyme $T_{opt}$ from the same organism$^{14,27}$. However, we found that OGT by itself can only explain 25% of individual enzyme $T_{opt}$ variance when tested on a dataset containing 2,609 enzymes from BRENDA (Fig. 3b, c). To improve the accuracy of this estimation, a second machine learning approach was applied. Machine learning models trained on sequence features are not significantly better than using OGT as estimation (Fig. 3c). However, the combined use of OGT and sequence features almost doubled the explained variance to 51% (Fig. 3c). This is a remarkable result that demonstrates a clear importance of physiological parameters in the estimation of protein properties. We speculate that inclusion of more samples and extraction of more descriptive features (both from sequence and physiological parameters) in conjunction with advanced machine learning models, like deep learning$^{28}$ may further improve the prediction of enzyme $T_{opt}$. The $R^2$ score of 51% obtained for $T_{opt}$ predictions in this study could be used as a benchmark accuracy for future model development.

In addition to our scientific findings we used experimental as well as predicted OGT to predict $T_{opt}$ for enzymes in the BRENDA database. We could assign $T_{opt}$ for 54% (6,507,076) of the all enzyme sequences and up to 59% of all EC numbers in the database. The number of sequences annotated with $T_{opt}$ computed using predicted OGT is 909,954. This is remarkable given that only around 33,000 enzymes in BRENDA has an experimentally determined $T_{opt}$. The annotation expanded both lower and upper bound of $T_{opt}$ for almost every EC number (Fig. 4b). This $T_{opt}$ annotation should prove very useful for identification of both thermostable enzymes$^7$ as well as cold-active enzymes$^{29}$ for applications in protein engineering and biotechnology. We provide a data file with all these annotations (https://zenodo.org/record/2539114, https://doi.org/10.5281/zenodo.2539114) as well as a script with which to query the data for unrestricted re-use as part of our Tome package (https://github.com/EngqvistLab/Tome).
Methods

Proteome dataset
The bulk of protein sequence data used in this work was obtained from Ensembl Genomes release 37, obtained in September 2017 (http://ensemblgenomes.org/). For all archaea and bacteria listed at ftp://ftp.ensemblgenomes.org/pub/bacteria/release-37/fasta/ fasta files containing protein sequences were downloaded. Similarly, fasta files containing protein sequences for all fungi listed at ftp://ftp.ensemblgenomes.org/pub/fungi/release-37/fasta/ were downloaded. As a complement to the Ensembl Genome data we made use of protein data from RefSeq release 87, obtained in March 2017 (https://www.ncbi.nlm.nih.gov/refseq/). Fasta files containing a nonredundant set of protein sequences for each organism were downloaded from ftp://ftp.ncbi.nlm.nih.gov/refseq/release/ for archaea, bacteria, fungi and protozoa.

In many cases the Ensembl Genomes and RefSeq datasets both contained information for the same organism, or for several strains of the same organism. Therefore, to combine the two datasets, the following steps were followed: First, where multiple strains from the same organism were present in the Ensembl Genomes dataset, the strain with the largest file size, indicating the greatest number of amino acids in the downloaded fasta file, was selected for analysis. Other strains for that organism were discarded. Second, where the same organism was present in both the Ensembl Genomes and RefSeq datasets the one from Ensembl Genomes was retained and the one from RefSeq was discarded. In this way a protein dataset comprising protein sequence data for 7,565 microorganisms was obtained. Of these 5,325 originated from Ensembl Genomes and 2,240 originated from RefSeq.

For each organism in the protein dataset we attempted to annotate it with its optimal growth temperature. In this annotation procedure organism names were stemmed to the species level (ignoring strain designations) and cross-referenced with a published dataset containing growth temperatures for 21,498 microorganisms (https://doi.org/10.5281/zenodo.1175608). Growth temperatures could be associated with the protein sequence data from 5,762 organisms, whereas 1,803 were left unannotated.

Estimation of threshold
For each proteome, the total length of each protein was calculated. Then the amino acid frequencies and the total number of residues of the first $n$ proteins ($n = 1, 2, \ldots N$, were $N$
is the total number of proteins) were calculated sequentially. The data points in the last one-third of all residues added were used to measure the stability of the calculated amino acid frequencies. Three different metrics were designed: (1) the absolute slope value \(|a|\) in the linear regression between the number of residues and amino acid frequency; (2) frequency variance of these selected frequencies \(\sigma^2\) and (3) varying range \(R\), the difference between maximal frequency and minimal frequency. Ideally, 0 was expected for all these three metrics if there is an absolutely stable amino acid frequency in a given proteome. Finally, for each proteome, the maximal \(|a|\), \(\sigma^2\) and \(R\) of 20 amino acids of each proteome \((r_{\text{abs,max}}, \sigma_{\text{max}}^2 \text{ and } R_{\text{max}})\) were used to measure whether frequencies were stable.

To test the effect of the protein order in a proteome in the above analysis, a shuffling strategy was applied. Firstly, equal coverage over the log\(_{10}\)-transformed proteome size range 3-7.5 was ensured by performing the random sampling in 20 bins. One proteome was randomly selected for each bin and this resulted in 17 selected proteomes as there is no proteome in 3 of these bins. The order of the proteins in each proteome was randomly shuffled and then \(r_{\text{abs,max}}, \sigma_{\text{max}}^2 \text{ and } R_{\text{max}}\) were calculated. Each proteome was shuffled for 100 times.

**Machine learning workflow for OGT model**

20 amino acid frequencies and 400 dipeptide frequencies were extracted for each proteome. Then, each of these features were normalized by \(x_{N,i} = \frac{x_i - u_i}{\delta_i}\), where \(x_i\) is the values of feature \(i\), \(u_i\) and \(\delta_i\) are mean and standard derivation of \(x_i\), respectively. The following six models were selected and their performance were tested on the annotated and filtered proteome dataset using single amino acid frequencies (AA), dipeptide frequencies (Dipeptide) or the two together (AA+Dipeptide): Linear regression (Linear), bayesian ridge, elastic net, decision tree, support vector regression (SVR) and random forest. 5-fold cross-validation was used for the calculation of \(R^2\) scores. For SVR, elastic net, decision tree and random forest models, an additional 3-fold internal cross-validation were used to optimize the hyperparameters. The model with the highest \(R^2\) score was selected and trained, without cross-validation, on the whole dataset. For the prediction of OGT for those un-annotated organisms, dipeptide frequencies were normalized by \(x_{N,i} = \frac{x_i - u_i}{\delta_i}\), where \(x_i\) is the values of feature \(i\), \(u_i\) and \(\delta_i\) are mean and standard derivation of feature \(i\) in the training dataset, respectively.

**OGT Model validation**

For validating the OGT prediction model we sampled 54 species with predicted growth temperatures (for which no growth temperatures were available in the original dataset)
at random. Equal coverage over the temperature range 0-100°C was ensured by performing the random sampling in 10 bins, each spanning a 10°C temperature range. The primary scientific literature was then manually searched to obtain documented experimental growth temperatures for the sampled organisms. For 45 organisms a documented growth temperature could be found, for 9 organisms it could not. The accuracy of predicted OGT was assessed by computing the Pearson correlation with experimental OGT.

In a second approach to validating the OGT prediction model we used Python scripts and the Zolera SOAP package (https://pypi.python.org/pypi/ZSI/) to extract all available experimentally determined enzyme temperature optima from the BRENDA enzyme database release 2018.2 (July 2018). Data coming from the same enzyme was de-duplicated by averaging temperature optima from records with the same EC number and originating from the same organism. For each organism with catalytic optima for more than five enzymes the arithmetic mean of those optima were calculated. Those organisms present in both the BRENDA enzyme data as well as the dataset with predicted OGT were identified through cross-referencing species names. The accuracy of predicted OGT was assessed by computing the Pearson correlation between predicted OGT and mean catalytic optima of enzymes.

**Machine learning workflow for $T_{opt}$ model**

UniProt identifiers for proteins with an experimentally determined catalytic optimum were obtained from the “TEMPERATURE OPTIMUM” table in the web pages of the BRENDA database, release 2018.2 (July 2018). These identifiers were filtered to retain only those associated with an organism with experimentally determined OGT. After further filtering to remove sequences containing “X” (unknown amino acid), a dataset with 2,609 enzymes was generated. The protein sequences for each of these identifiers were downloaded from the UniProt database in fasta format.

The following features were extracted for each enzyme: (1) 20 amino acid frequencies (AA); (2) 400 dipeptide frequencies (Dipeptide); (3) OGT of its source organism; (4) Basic features including protein length, isoelectric point, molecular weight, aromaticity, instability index, gravy and fraction of three secondary structure units: helix, turn and sheet. These features were extracted with the module Bio.SeqUtils.ProtParam.ProteinAnalysis in Biopython (version 1.70). Additionally, six binary features were extracted: EC=1, 2, 3, 4, 5, 6. These numbers represent the first digit in a EC number. All features except binary features were normalized as described in section “Machine learning workflow for OGT model”. The following five models were tested on the resulting dataset: bayesian ridge, elastic net, decision tree, support vector
regression (SVR) and random forest. The linear model was not used due to its poor performance on any datasets containing dipeptide frequencies (negative $R^2$ scores by cross-validation). The performance of the five regression models was tested using the same cross-validation strategy as for OGT. In addition, to test the accuracy of using OGT of the organism as an estimation of enzyme $T_{\text{opt}}$, the $R^2$ score between each enzymes $T_{\text{opt}}$ and associated OGT was calculated. The model with the highest $R^2$ score was chosen and trained on the full training dataset.

**BRENDA annotation**
Protein sequence data for each EC class was obtained by downloading comma-separated flatfiles from the BRENDA database version 2018.2 (July 2018). Each sequence in these files contain information regarding source organism as well as unique UniProt identifiers. Where possible, each protein sequence was associated with an OGT value by mapping the source organism name to the OGT dataset from (https://doi.org/10.5281/zenodo.1175608). Those sequences were firstly mapped to the existing $T_{\text{opt}}$ values in BRENDA by matching EC-UniProt id pair. For those enzymes without any experimental $T_{\text{opt}}$ values, the amino acid frequencies were calculated (ignore all ‘X’ in the sequence). All 20 amino acid frequencies as well as the OGT variable were normalized by $x_{N,i} = \frac{x_i - u_i}{\delta_i}$, where $x_i$ is the values of feature $i$. $u_i$ and $\delta_i$ are mean and standard derivation of feature $i$ in the original training dataset, respectively. Finally, the normalized values were used for the prediction of $T_{\text{opt}}$ by the previously generated random forest regressor trained on the AA+OGT datasets. The predicted enzyme $T_{\text{opt}}$ and annotated OGT values of these enzymes are freely available for download and re-use (https://zenodo.org/record/2539114, https://doi.org/10.5281/zenodo.2539114).

**Software**
All machine learning analysis were conducted with scikit-learn package (version 0.19.1)\textsuperscript{34} using Python version 2.7.14. The module and model hyperparameters used are listed in Supplementary Table S2.

**Code availability**
Python code for proteome analysis, machine learning and data visualization are available from the authors upon request. The source code for the Tome package is available under a permissive GPLv3 license at GitHub (https://github.com/EngqvistLab/Tome).
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Author contributions

GL, JN and MKME conceptualized the research. JN acquired funding to support the project. MKME generated the proteome and BRENDA datasets and performed data curation. GL performed the computational and statistical data analysis. GL, KSR and MKME interpreted results. GL wrote the computer code for the Tome package. GL and MKME created the publication figures. GL and MKME wrote the initial draft of the paper. GL, KSR, JN and MKME carried out revisions on the initial draft and wrote the final version.

Competing interests

The authors declare no competing interests.

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References


11. Galtier, N. & Lobry, J. R. Relationships between genomic G+C content, RNA secondary


its primary sequence. Protein Eng. 4, 155–161 (1990).

