# Building a tRNA thermometer to estimate microbial adaptation to temperature

Emre Cimen<sup>(D1,2,\*,†</sup>, Sarah E. Jensen<sup>(D3,\*,†</sup> and Edward S. Buckler<sup>(D1,3,4</sup>

<sup>1</sup>Institute for Genomic Diversity, Cornell University, Ithaca, NY 14853, USA, <sup>2</sup>Computational Intelligence and Optimization Laboratory, Industrial Engineering Department, Eskisehir Technical University, Eskisehir 26555, Turkey, <sup>3</sup>School of Integrative Plant Sciences, Plant Breeding and Genetics Section, Cornell University, Ithaca, NY 14853, USA and <sup>4</sup>United States Department of Agriculture, Agricultural Research Service, Ithaca, NY 14850, USA

Received July 23, 2020; Revised October 13, 2020; Editorial Decision October 15, 2020; Accepted October 20, 2020

# ABSTRACT

Because ambient temperature affects biochemical reactions, organisms living in extreme temperature conditions adapt protein composition and structure to maintain biochemical functions. While it is not feasible to experimentally determine optimal growth temperature (OGT) for every known microbial species, organisms adapted to different temperatures have measurable differences in DNA, RNA and protein composition that allow OGT prediction from genome sequence alone. In this study, we built a 'tRNA thermometer' model using tRNA sequence to predict OGT. We used sequences from 100 archaea and 683 bacteria species as input to train two Convolutional Neural Network models. The first pairs individual tRNA sequences from different species to predict which comes from a more thermophilic organism, with accuracy ranging from 0.538 to 0.992. The second uses the complete set of tRNAs in a species to predict optimal growth temperature, achieving a maximum  $r^2$  of 0.86; comparable with other prediction accuracies in the literature despite a significant reduction in the quantity of input data. This model improves on previous OGT prediction models by providing a model with minimum input data requirements, removing laborious feature extraction and data preprocessing steps and widening the scope of valid downstream analyses.

# INTRODUCTION

Environmental temperature affects every biochemical reaction within an organism, from spontaneous protein folding to complex metabolite catalysis. Tools that infer an organism's optimal growth temperature from genomic sequence have potential biological and economic implications and can improve understanding of how both individual cell components and whole organisms adapt to their environment. However, experimentally identifying the true optimal temperature of every newly discovered micro-organism is unfeasible due to the sheer number of prokaryotes that have been identified and the difficulty in isolating and culturing many prokaryotic species. Predicting an organism's optimal growth temperature based on physical characteristics of the genome is one way to determine optimal temperature without needing to successfully culture a new species.

Temperature has a significant effect on cell biochemistry. In general, cellular processes speed up as temperature increases, but extremely high temperatures can also denature proteins and negatively affect biochemical reactions. Proteins function best within a specific temperature window that maximizes enzymatic reaction rate without denaturing the protein, and most enzymes have evolved within an optimal temperature range that is closely tied to environmental temperature (1). Few enzymes show optimal activity more than 10°C above or below the optimal growth temperature of the host organism (1,2). Maintaining catalytic function at extreme temperatures requires specific changes in genome composition, and many studies have identified differences between thermophilic and mesophilic genomes (1,3–11). Thermophilic proteins tend to have more hydrophobic residues, disulfide bonds and ionic interactions to pack amino acid residues closely together and prevent protein unfolding, while psychrophilic (cold-adapted) proteins require fewer strong interactions between amino acid residues (2). Significant shifts in genome composition have also been correlated with environmental temperature con-

Sarah E. Jensen, Edward S. Buckler, 526 Campus Rd., 175 Biotechnology Building, Buckler Lab., Ithaca, NY 14853, USA.

Published by Oxford University Press on behalf of Nucleic Acids Research 2020.

This work is written by (a) US Government employee(s) and is in the public domain in the US.

<sup>\*</sup>To whom correspondence should be addressed. Tel: +90 554 237 0746; Fax: +90 222 323 9501; Email: ecimen@eskisehir.edu.tr

Correspondence may also be addressed to Sarah E. Jensen. Tel: +1 607 255 1809; Fax: +1 607 254 6379; Email: sej65@cornell.edu

<sup>&</sup>lt;sup>†</sup>The authors wish it to be known that, in their opinion, the first two authors should be regarded as joint First Authors.

Present addresses:

Emre Cimen, Industrial Engineering Department, Eskischir Technical University, Eskischir, 26555, Turkey.

ditions, affecting genome features like GC content, codon bias and amino acid frequency (12,13).

Because physical changes in DNA, RNA and protein composition have been correlated with optimal growth temperature, multiple approaches have been taken to predict OGT from a combination of these features. Aptekmann et al. found a positive correlation between GC content of tRNA regions and OGT and between information content and optimal growth temperatures in Archaea (12). Li *et al.* implemented a machine learning workflow in order to predict the OGT of micro-organisms and enzyme catalytic optima from 2-mer amino acid composition and investigated a wide range of regression models (5). Sauer and Wang used multiple linear regression to predict the OGT of prokaryotes from genome size and tRNA, rRNA, open reading frame and proteome composition (13). Ai et al. targeted a problem related to protein thermostability and classified thermophilic and mesophilic proteins using support vector machines and decision trees (6). Similarly, Capaldi et al. predicted bacterial growth temperature range based on genome sequences with a Bayesian model (1).

These previously published approaches to predicting organism OGT tend to use a combination of genome and sequence features that cover the entire Central Dogma of molecular biology. Such models can be useful when the end goal is to predict OGT for a newly identified species, but they have limited downstream applications when the end goal is understanding how cellular components adapt or evolve under different temperature conditions, because using many cell features to predict OGT then statistically confounds downstream analyses involving the same cellular components. Optimal growth temperatures predicted with a model requiring amino acid composition as input, for example, cannot be used in subsequent analyses investigating how proteins evolve under different temperatures-the protein evolution results would be confounded with the initial OGT prediction. To address this, we set out to create a model that predicts prokaryote OGT using a minimal set of input data.

Transfer RNAs (tRNAs) are a vital and universal part of life, with remarkably conserved structure and function. Although most tRNAs have a standard 'cloverleaf' structure with an acceptor arm, D-arm, anticodon arm and T-arm, mutants have been identified that lack the T-arm, D-arm or both (14,15). We chose to focus on tRNA sequences with this model because of the ubiquity and conservation of tR-NAs across domains, because RNA base pairing chemistry is known to be affected by temperature, and because single-base mutations in tRNAs can dramatically affect function and temperature sensitivity (16,17).

We refer to this model as a 'tRNA thermometer' because it uses tRNA sequence features as an indicator of an organism's optimal growth temperature, thus measuring the 'temperature' of the genome. Although biologists may be familiar with the concept of an 'RNA thermometer', here we use a similar term with a different context. The tRNA thermometer model uses a Convolutional Neural Network (CNN) to classify and predict prokaryote OGT using tRNA sequences as input data. Because the model uses only tRNA features, it is possible to use predicted optimal temperatures from this model in downstream analyses evaluating the effects of temperature on other cellular components, including protein and genomic features. Essentially, with only  $\sim$ 4000 bp of sequence ( $\sim$ 0.1% of the genome), this CNN model can predict OGT as well as summary data from the entire rest of the genome. Because it uses fewer features than previous OGT-prediction models and does not require feature extraction, it is also easier to use and removes researcher bias in selecting features.

### MATERIALS AND METHODS

#### Data collection and distribution

A list of species for which optimal growth temperature has been determined was obtained from Sauer *et al.* and all existing genome assemblies were downloaded for each species, resulting in an initial set of 36 529 Bacteria and 276 Archaea genomes, with optimal growth temperatures ranging from 4 to  $103^{\circ}$ C (13). tRNA sequence and positions were predicted for each genome using tRNAscan-SE (version 2.0.3; (18,19)). rRNA sequence and positions were predicted for each genome using barrnap (version 0.9; (20)).

A single genome was selected for each species, and only tRNA sequences from that genome were used in the CNN predictions. Genome assemblies were considered lowquality and removed if 16S, 23S and 5S rRNA sequences could not be predicted by barrnap. For species with multiple remaining genome assemblies, the assembly with the highest number of predicted tRNA sequences was chosen as the single 'best' assembly. Selecting genome assemblies in this way resulted in genomes for 165 unique archaea species and 2375 unique bacteria species. However, the distribution of optimal temperatures for this species set was highly skewed, with nearly 40% of archaea and nearly 70% of bacteria genomes having predicted optimal temperatures of 28, 30 or 37°C. These three temperatures are common prokaryote culture temperatures so we decided to remove observations at these temperatures to reduce the bias and help balance the dataset, since we could not be sure that these were true adaptive growth temperatures rather than culture temperatures. After removing species with OGT listed at 28, 30 or 37°C, the dataset contained 683 bacteria species with 41 853 predicted tRNAs and 100 archaea species with 5474 predicted tRNAs. In the final dataset, 70% of bacteria tR-NAs come from species with OGTs between 20 and  $40^{\circ}$ C, and 24% of archaea tRNAs come from species in this temperature range. The range of optimal temperatures included in the dataset spans nearly 100°C and as expected, tRNA structures in the dataset appear consistent despite the large variation in species' optimal growth temperatures (Figure 1). Sequences were used to train and test the model as-is and were not processed further to identify specific tRNA mutants or non-standard structures.

# **Prediction method**

We used Convolutional Neural Networks (CNNs) for OGT prediction. CNNs are neural network (NN)-based machine learning models with at least one convolutional layer. They can predict both categorical (classification) and continuous (regression) targets depending on the configuration of



Figure 1. Sample structures for Lysine tRNAs calculated using predicted MFE are consistent even in species with extreme differences in optimal growth temperature. (A) *Aequorivita sublithincola*, with OGT =  $4^{\circ}$ C, (B) *Lactobacillus frumenti*, with OGT =  $40^{\circ}$ C, (C) *Pyrolobus fumarii*, with OGT =  $103^{\circ}$ C. These tRNA structures were calculated and displayed using the ViennaRNA Web Services (41).

downstream layers. When the input is a continuous or discrete signal such as images, sensor data, or base pair sequence, CNNs are useful because they automatically extract features from the input and can capture both local and global features. Moreover, thanks to weight sharing at the convolutional layers and downsampling at the pooling layers, CNNs have fewer parameters that need to be learned than regular NNs. Thus, CNNs require less training data and have lower risk of over-fitting. Additionally, other machine learning models have no prior knowledge of how input values are organized, and are not able to take advantage of the relative positions within a sequence, (i.e. they cannot determine consecutive bases in a sequence). CNN architectures naturally have this prior neighborhood knowledge. CNNs have been shown to be successful predicting a target by using genomic sequence data: Wang et al. presented a sequence-based deep CNN model that accurately predicts the TF binding intensities to given DNA sequences (21), and Zeng *et al.*, and Zhuang *et al.* predict enhancer– promoter interactions with DNA sequence data and CNNs (22, 23).

We set two prediction problems in this study. One is a binary classification model that can take two tRNA sequences from different organisms as input. This classification model predicts which tRNA sequence in the pair comes from a micro-organism with higher optimal growth temperature, and requires only a single tRNA from each genome for the classification. The second model uses a CNN regression model that predicts the optimal growth temperatures of bacteria and archaea. In both cases, a CNN model was chosen to allow automatic feature extraction. This made it possible to use tRNA sequences as direct input and did not require manual extraction of hundreds of sequence features or assessment of the correlation of each individual feature with OGT. In both models, tRNA sequences were obtained as described above, then one-hot encoded. Because not all tRNA sequences are the same length, shorter sequences were padded with zeros to produce a final 4xL matrix of 0s and 1s, where L is the length of the longest tRNA in the input data.

# Models

Temperature classifier model based on individual tRNA sequences. In the first model, we built a CNN classification model that can take paired sets of tRNAs and predict which tRNA belongs to a micro-organism with higher optimal growth temperature. Before presenting our classifier model, in Equations (1) and (2) we introduce the data structure. In Equation (1), dataset D has N micro-organisms. Each micro-organism m<sub>i</sub> has a corresponding OGT t<sub>i</sub>. In Equation (2), because each organism in the dataset has more than one tRNA,  $n_i$  is the number of tRNAs in micro-organism  $m_i$ . Since there are a different number of tRNAs in each micro-organism,  $n_i$  's are different for each  $m_i$ .  $tRNA_i^j$  is the j<sup>th</sup> tRNA of i<sup>th</sup> micro-organism  $m_i$  and, it is an instance in  $R^{4 \times L}$  space. The input space is  $4 \times L$  dimensional because there are 4 bp (one-hot encoding), and the length of the longest tRNA in the dataset D is L.

$$D = \{\{m_1, t_1\}, \dots, \{m_i, t_i\}, \dots, \{m_N, t_N\}\},\$$
  
$$i \in I = \{1, \dots, N\}$$
(1)

$$m_{i} = \left\{ t RNA_{i}^{1}, \dots, t RNA_{i}^{j}, \dots t RNA_{i}^{n_{i}} \right\},$$
  
$$t RNA_{i}^{j} \in R^{4 \times L}, \ j \in j_{i} = \{1, \dots, n_{i}\}, \ \forall i \in I \quad (2)$$

To predict which tRNA comes from an organism with higher OGT, we built a CNN classifier with two branches, each of which was fed an input tRNA sequence. Each branch starts with at least one pair of convolutional and pooling layers. After the last pooling layer, branches are flattened and merged. Convolutional and dense layers use a Rectified Linear Unit (ReLU) activation function. Two output nodes at the end are activated with the Softmax function which provides class probabilities. We predict the output as a binary label that indicates whether the first branch input or the second branch input has a higher OGT. The loss function of the model is the categorical cross entropy, and it is minimized with the Adam optimizer (24). Parameters of the model (e.g. layer sizes, number of layers, optimizer parameters and dropout rate etc.) were selected with hyper-parameter optimization. To train and test our classifier model, we created a dataset of tRNA pairs from the original dataset given in Equations (1) and (2). In the classification dataset each sample is in the form of {{tRNA1, tRNA2}, y}, where  $y \in \{0, 1\}$  is the class label. We used tRNA pairs only if their micro-organism OGTs were different by at least 1°C (Figure 2A).

Species OGT predictor model. The CNN regression model to predict species' OGT starts with two convolutional and subsequent maximum pooling layers. After the last pooling layer there is a single flattening layer before multiple fully connected dense layers. The activation function of the convolutional and dense layers is Rectified Linear Units (ReLU; Figure 2B). The output node is a continuous variable and is activated linearly. The loss function of the model is the mean squared error, and it is minimized with the Adam optimizer. In Figure 2, we provide the general structure of the models. Dots mean the model may have more layers of the given type. The number of layers for each model is selected with the hyper-parameter optimization.

To train and test the regression model, we created a dataset as in Equations (1) and (2), where each sample is in the form {tRNA, t}. t is OGT of the related tRNA. We trained the CNN model by considering each tRNA as independent of all other tRNAs in the organism. Once we trained the model, we had  $n_i$  tRNA-based OGT predictions for the micro-organism  $m_i$ ; one for each tRNA. We determined the OGT prediction of the species as a whole by calculating the median of all tRNA-based OGT predictions. In Equation (3),  $\hat{t}_i$  is the OGT prediction of  $i^{th}$  micro-organism  $m_i$  and,  $\hat{t}_i^j$  is the OGT prediction of  $j^{th}$  tRNA of  $i^{th}$  micro-organism.

$$\hat{t}_i = \operatorname{median}\left(\hat{t}_i^1, \dots, \hat{t}_i^{j}, \dots, \hat{t}_i^{n_i}\right),$$
  
where  $j \in j_i = \{1, \dots, n_i\}, \forall_i \in I$  (3)

#### Hyper-parameter optimization

Selecting the optimal combination of hyper-parameters is important because hyper-parameter values have a significant effect on the performance of CNN models. There are a large number of hyper-parameters and the possible values of each results in millions of potential hyper-parameter combinations. These hyper-parameters define the model structure and need to be selected by the user before training. In this study we used Bayesian optimization to determine parameter values for layer size, the number of layers, the number of filters, kernel size, pooling size, strides, dropout rate, batch size and beta1, beta2, learning rate of the Adam optimizer. The search space for each hyperparameter is listed in Supplementary Table S1 and selected hyper-parameters are provided in Supplementary Table S2.

$$x^* = \arg\min_{x \in \mathbf{X}} f(x) \tag{4}$$

In Equation (4),  $x^*$  corresponds to the best combination of hyper-parameter values. Selecting  $x^*$  from the possible selection set X is defined in Equation (4), where f(x) is the loss (e.g. mean squared error, mean absolute error and classification error) on the validation set. It is not possible to try each combination of hyper-parameters to find  $x^*$  since there are millions of potential hyper-parameter combinations. Thus, there is a need for an intelligent way to select candidate hyper-parameters. The optimization-based approach that we use maximizes the expectation of the improvement (EI) on the performance. There are several ways to estimate expected improvement, and we used a tree-structured Parzen Estimator (TPE) provided in the Hyper-opt python package to find  $x^*$  (25,26). The TPE is a sequential optimization approach: it uses historical performance to sequentially select the candidate set of hyper-parameters, and then iteratively chooses new hyperparameters to test by maximizing performance. Bergstra et. al. provide formulations and algorithms about EI and TPE in detail (25). We allow the algorithm to run 50 iterations to select the best combination of hyperparameters to use in each model.

# Data splitting procedure

To evaluate the performance of each proposed model, we investigated two scenarios. First, we split the species randomly into training, validation and test sets. Second, we controlled for evolutionary relatedness and split data according to phylogenetic distance.

*Random split*. Model evaluation commonly uses k-fold cross-validation or a random split of the training, test and validation datasets. In the first part of the computational results, we held out 5% of all species as a validation set to fine-tune hyper-parameters and then used the rest of the species for 5-fold cross-validation: 76% for training and 19% for testing in each iteration of the model. Hyper-parameter op-timization was done once during the first iteration, and hyperparameters were selected according to performance on the validation set. We repeated the whole set of tests five times.

*Phylogenetic distance split.* Random train-test data splits work well for prediction models, and previous OGT-



Figure 2. OGT prediction models. Input tRNA sequences are one-hot encoded and padded with 0s to make the matrix. (A) The general structure of the temperature classifier model. The model has two channels. Both channels are fed a tRNA. Each channel starts with a convolutional layer and a maximum pooling layer. The sequence is flattened and passed to two fully connected and dropout layers. The two channels are concatenated and then passed to fully connected layers. The output layer has two neurons for binary classification. (B) The general structure of the regression model. The input layer is followed by two convolutional layers and a maximum pooling layer. Then, data are processed through fully connected dense layers, resulting in a single OGT prediction for each tRNA. This figure shows the general structure only, and the exact number of layers are selected with hyper-parameter optimization. Selected hyper-parameters are provided in the Supplementary Files.

prediction models use a random training/test split to evaluate their model (5-6,9,13). However, the random training/test split does not account for evolutionary relatedness between species. Previous machine learning studies have found that similarity between individuals can provide overly optimistic results because the training and test data sets may contain closely related species. Washburn *et al.* (27) considered this situation in the prediction of mRNA expression levels, and found that machine learning models trained without taking evolutionary history into account were able to recognize species similarity and use it to inform predictions. Additionally, Washburn *et al.* states that ignoring shared evolutionary history can exaggerate model performance and possibly lead researchers to conclude that certain model features are important, when in fact, the model test set is contaminated by similar observations that are present in both the training and test sets. In fact, if the aim of using a predictive model is only to predict accurate OGTs/class labels, splitting training and test sets randomly is valid. However, if researchers would like to obtain biological insights from the model (i.e. to answer questions like which tRNA properties are correlated with high/low OGT), a dataset split by phylogenetic distance will provide better insights. A model trained using a phylogenetic distance train/test split is also likely to be more transferable to other problems. This is because by constraining the model and removing all information from phylogenetically related species, we push the model to extract other rules from the sequence that are more transferrable.

To account for evolutionary relationships across species, we tested each model with a phylogenetically informed training/test split. A simple phylogenetic relationship between species was calculated as the relative relationship between species based on species' taxa id and the NCBI Common Tree (28, 29). The tree was converted to a simple distance matrix using the 'ape' R package (30). The phylogenetic distance matrix was used to split species into clusters. For this purpose, we applied hierarchical clustering to species by minimizing the Ward variance (31). After preserving 5% of the species for the validation set, the rest of the species were split into 10 clusters, 8 of which were used as a training set and 2 of which were used as the testing set. This procedure was repeated five times, and each cluster was included in the test set only once. We repeated the whole set of tests five times.

*Model attention.* To determine how the importance assigned to the tRNA stem structures affects model predictions, we selectively mutated each set of paired bases in the tRNA structure for 55 *Escherichia coli* tRNAs for which structure information is available (32). For each paired nucleotide we mutated the original DNA base to all three other nucleotides (e.g.  $G \rightarrow A$ , T and C in turn) which would disrupt a single Watson/Crick base pairing interaction within the transcribed tRNA. This resulted in a set of 5793 new tRNA sequences, each with a single nucleotide change that disrupted one set of pairing interactions. We then predicted OGT for these new sequences and compared predictions to OGT predictions for the original *E. coli* tRNA sequences.

*Software and computation power.* We implemented the proposed prediction method in Python 3.6 using Keras (2.2.5) to build and train the CNN model. For the distance split, species were clustered hierarchically with Sklearn (0.21.2) to find clusters related to phylogenetic distance. We have used the Keras-vis [https://github.com/raghakot/keras-vis] package to investigate model attention (33). Tests were carried out on a computer with a GeForce RTX 2080 GPU, 64 GB RAM and Intel(R) Core(TM) i7–7800X CPU running at 3.5 GHz.

# RESULTS

# Model 1: Temperature classifier based on individual tRNA sequences

We first built a model that uses a single tRNA to distinguish which of two species comes from a higher optimal growth temperature environment. Data was split into training and test sets either randomly or with phylogenetically informed distance splitting as described above. All tests were repeated five times and results represent the average and the standard deviation of these runs.

For both the random and phylogenetic data splits we calculated and presented accuracy and F1 score results when temperature differences are greater than  $0^{\circ}$ C,  $5^{\circ}$ C,  $10^{\circ}$ C,  $20^{\circ}$ C and  $30^{\circ}$ C in Tables 1 and 2. For bacteria genomes, including phylogenetic information as well as the tRNA sequence improved predictions (Figure 3A). Interestingly, the model results showed that in archaea genomes, a single tRNA pair contains enough signal to distinguish between genomes from species adapted to different optimal growth temperatures, even when phylogenetic relationships between species have been deliberately removed (Figure 3B). Unsurprisingly, the model performs better when there are larger temperature differences between the species being compared. It is possible that the model learned species OGT, rather than sequence features related to OGT, so we also verified that the model was not just predicting the same direction for a given species in the results (i.e. was not always predicting 'lower OGT' for a given species).

# Model 2: Species OGT prediction

We next asked whether it was possible to use information from all tRNAs within a species to predict overall species OGT using a regression CNN model. As with the classification model, prediction accuracy was compared using both random and phylogenetically informed data splits. Data were split into hyper-parameter validation, training and test sets as described in the classification model. Root mean squared error (RMSE) and the coefficient of determination  $(r^2)$  were used to evaluate model performance for both the random and phylogenetic distance split models. A good model will have both low RMSE and high  $r^2$ . Low RMSE indicates how closely the model can pinpoint a species' OGT, while high  $r^2$  indicates that most of the variance in the true OGT dataset is explained by the OGT predictors. **RMSE** and  $r^2$  were compared for each domain individually as well as the combined domains (Table 3 and Figure 4).

Results indicate that tRNA sequence alone can accurately predict both archaea and bacteria OGTs. Performance in all three datasets is highest for the randomly split dataset, achieving 0.862, 0.818 and 0.875  $r^2$  in archaea, bacteria and combined archaea & bacteria datasets, respectively (Table 3 and Figure 4).

One drawback of randomly splitting data into training and test sets is that similarity between individual observations in the training and test sets may lead to overly optimistic model performances. When the end goal of a model is prediction, species relatedness is less of an issue. However, biological insights are more difficult to draw from a model that does not control for population structure within the dataset, since causal elements (in this case, causal nucleotides in the tRNA sequence) are confounded by structure due to shared evolutionary history. The model accounting for phylogenetic distance achieved 0.772, 0.370 and 0.590  $r^2$  in archaea, bacteria and combined archaea & bacteria datasets, respectively (Table 3 and Figure 4)

#### **Regression model attention**

One criticism of convolutional neural networks is that they create a 'black box' that can be difficult to interpret, making it hard to draw meaningful biological insight from a model. To try to understand what portions of the tRNA sequence matter for OGT prediction, we correlated tRNA features to OGT predictions, then calculated attention statistics and evaluated the effects of directed mutagenesis on model predictions in bacteria. We found that both GC content and

Temperature difference	Archaea		Bac	teria	Archaea + bacteria		
	Random	Phylogenetic distance	Random	Phylogenetic distance	Random	Phylogenetic distance	
>0°C	$0.814 \pm 0.011$	$0.684 \pm 0.025$	$0.756 \pm 0.001$	$0.538 \pm 0.001$	$0.728 \pm 0.023$	$0.564 \pm 0.029$	
>5°C >10°C	$0.889 \pm 0.010$ $0.930 \pm 0.005$	$0.749 \pm 0.029$ $0.807 \pm 0.025$	$\begin{array}{c} 0.824 \pm 0.002 \\ 0.830 \pm 0.005 \end{array}$	$0.557 \pm 0.012$ $0.612 \pm 0.016$	$0.788 \pm 0.030$ $0.853 \pm 0.033$	$0.585 \pm 0.039$ $0.605 \pm 0.0512$	
>20°C >30°C	$\begin{array}{c} 0.974  \pm  0.004 \\ 0.992  \pm  0.002 \end{array}$	$\begin{array}{r} 0.902  \pm  0.011 \\ 0.954  \pm  0.008 \end{array}$	$\begin{array}{c} 0.882  \pm  0.005 \\ 0.921  \pm  0.007 \end{array}$	$\begin{array}{c} 0.660  \pm  0.024 \\ 0.694  \pm  0.035 \end{array}$	$\begin{array}{r} 0.914  \pm  0.032 \\ 0.950  \pm  0.037 \end{array}$	$\begin{array}{c} 0.622  \pm  0.063 \\ 0.636  \pm  0.071 \end{array}$	

Table 1. Classification model accuracy performance with archaea, bacteria and combined dataset



Temperature difference	Archaea		Bac	teria	Archaea + bacteria		
	Random	Phylogenetic distance	Random	Phylogenetic distance	Random	Phylogenetic distance	
>0°C >5°C >10°C >20°C >30°C	$\begin{array}{c} 0.845 \pm 0.008 \\ 0.911 \pm 0.006 \\ 0.944 \pm 0.003 \\ 0.980 \pm 0.003 \\ 0.994 \pm 0.001 \end{array}$	$\begin{array}{c} 0.722  \pm  0.013 \\ 0.786  \pm  0.018 \\ 0.840  \pm  0.015 \\ 0.925  \pm  0.006 \\ 0.966  \pm  0.005 \end{array}$	$\begin{array}{c} 0.716 \pm 0.005 \\ 0.801 \pm 0.003 \\ 0.825 \pm 0.004 \\ 0.893 \pm 0.005 \\ 0.934 \pm 0.006 \end{array}$	$\begin{array}{c} 0.512 \pm 0.025 \\ 0.529 \pm 0.027 \\ 0.614 \pm 0.033 \\ 0.688 \pm 0.036 \\ 0.731 \pm 0.039 \end{array}$	$\begin{array}{c} 0.737 \pm 0.022 \\ 0.805 \pm 0.025 \\ 0.879 \pm 0.026 \\ 0.933 \pm 0.025 \\ 0.961 \pm 0.027 \end{array}$	$\begin{array}{r} 0.579 \pm 0.053 \\ 0.612 \pm 0.062 \\ 0.644 \pm 0.072 \\ 0.669 \pm 0.079 \\ 0.681 \pm 0.087 \end{array}$	



Figure 3. CNN classification results for models built for each phylogenetic domain and with either (A) randomly split training and test datasets or (B) phylogenetically informed training and test datasets.

Method	Archaea		Bacteria		Archaea + bacteria	
	RMSE	r <sup>2</sup>	RMSE	r <sup>2</sup>	RMSE	r <sup>2</sup>
Random split Phylogenetic distance split	$8.06 \pm 0.967$ 11.06 \pm 1.005	$\begin{array}{c} 0.862  \pm  0.024 \\ 0.772  \pm  0.043 \end{array}$	$\begin{array}{c} 6.76  \pm  0.106 \\ 12.67  \pm  0.862 \end{array}$	$\begin{array}{c} 0.818  \pm  0.005 \\ 0.370  \pm  0.085 \end{array}$	$\begin{array}{c} 7.31  \pm  0.115 \\ 13.23  \pm  1.643 \end{array}$	$\begin{array}{c} 0.875  \pm  0.003 \\ 0.590  \pm  0.108 \end{array}$

minimum free energy of folding (MFE) were correlated with OGT predictions for individual archaea tRNAs (r = 0.79 and -0.63, respectively), and more moderately correlated with OGT predictions for individual tRNAs in bacteria (r = 0.27 and -0.26 for GC content and MFE, respectively; Supplementary Figure S1). There was also a trend in OGT predictions for different amino acid species, with certain amino acids being consistently assigned particularly high or low OGT values relative to other tRNAs (Supplementary Figure S2).

To determine which parts of the tRNA were important for model OGT predictions, we calculated the CNN activation values to determine the attention paid to each nucleotide in the tRNA, then normalized the activation values by tRNA length to get relative attention per base. While attention varied per nucleotide, results were consistent across all models and indicated that on average, the model paid most attention to nucleotides in the T arm and the anticodon arm (Figure 5A and Supplementary Figure S3). We used the predicted structure to determine the start and end of each stem-loop structure in each tRNA and summed the normalized CNN activation values over the length of the structure to determine total attention for each stem-loop structure. The anticodon arm and T arm sequences are sig-



Figure 4. CNN model performance when data are split randomly (A–C) and split with phylogenetic distance (D–F). Purple = bacteria, orange = archaea.



Figure 5. Model attention differs across the tRNA for the model trained on Archaea data with a phylogenetic data split. (A) Mean and standard error for the relative percent attention paid to each nucleotide in the tRNA, averaged across all tRNAs. Colors indicate the average positions of the stem-loop structure across all tRNAs: purple = acceptor stem, green = D arm, orange = anticodon arm, blue = T arm. (B) Average percent attention paid to each arm as a whole; Acc = acceptor, D = D arm, AC = anticodon arm, T = T arm.

nificantly more important for model predictions than the acceptor arm or D arm for both the random data split and for the phylogenetically informed data split (Figure 5B and Supplementary Figure S3, ANOVA P < 2e-16).

We selectively mutated 55 *E. coli* tRNAs to disrupt Watson/Crick base pairing and predicted OGT for each new sequence to determine how alterations in stem structure affect model OGT predictions. Results varied between the random-split and phylogenetic-split models. Mutations in the T arm of the tRNA led to a significantly different variance in OGT predictions for the random-split model, with some tRNA OGT predictions changing by 20 or 30°C (Bartlett test of homogeneity of variances, p = 1.917e-15), although the mean remained unchanged as did the mean and variance of predictions for other tRNA arms. In the phylogenetically informed model, disrupting Watson-Crick base pairing did not affect model variances.

# DISCUSSION

#### A machine learning model to predict prokaryote OGT

In this paper we discuss the application of a tRNA thermometer machine learning model that predicts prokaryote optimal growth temperatures. Unlike previous models, we aimed to produce a model concentrating on only one element of cell biochemistry - the tRNA sequence - to predict OGT. An initial classification model was able to distinguish between pairs of tRNAs to identify which came from a species with higher OGT, suggesting that even individual tRNA sequences contain signatures of thermal adaptation. A second model uses the aggregate of effects from all tRNAs in an organism - together these sequences contain enough signal to predict organism OGT using a CNN regression model.

The classification model was able to classify species OGT from a single tRNA with an accuracy greater than 0.8 for all temperature differences above  $10^{\circ}$ C in models where data

was split randomly. In all cases, models performed best when phylogenetic information was available to help with predictions, suggesting that species-specific differences in tRNA composition are indicative of tRNA thermal adaptation. This extra information about species relatedness was not available in the phylogenetic split model, resulting in lower classification accuracies. However, classification accuracies remained relatively high when classifying Archaea sequences, which may be due to the wider range of OGTs available for Archaea species. The joint model with both archaea and bacteria tRNA sequences frequently performed worse than either the archaea or bacteria model alone, likely because it needed to learn relevant sequence features for two separate domains and learn rules that applied across a much larger phylogenetic distance.

Literature results using multiple linear regression to predict OGT achieved an  $r^2$  of 0.938 for archaea, 0.767 for bacteria, and 0.835 for a combined dataset by using genomic, tRNA, rRNA, open reading frames and proteome derived features and splitting training and test sets randomly (13). However, when only genomic and tRNA derived features are used by the authors, they achieved 0.616  $r^2$  (13). On the other hand, the proposed random-split CNN model achieves 0.875  $r^2$  and shows a 42% improvement over literature  $r^2$  results for both the bacteria dataset and the combined bacteria and archaea dataset with only tRNA sequences as input. It is interesting that the CNN model outperforms in bacteria and in the combined dataset, but not in the archaea dataset. This may be due to the small number of archaea species with genome assemblies and OGT information that were available to train the model.

# Model predictions are correlated with GC content and tRNA MFE

Although the regression model merges information from all tRNA species to produce a final OGT prediction, its pre-

dictions for individual tRNA species vary by amino acid. In the archaea models, there was a strong negative correlation between tRNA MFE and predicted OGT and a strong positive correlation between tRNA GC content and predicted OGT. The correlations in bacteria were weaker, but had the same directionality. These correlations suggest that the model is learning information about secondary and tertiary tRNA structure and using it to make predictions, despite the fact that only the primary sequence was explicitly provided. Both MFE and GC content affect tRNA stability and function (16).

#### Post-transcriptional modifications affect tRNA stability

Post-transcriptional modifications affect tRNA stability and can include methylation, thiolation, reduction and isomerization of nucleotide bases. Although some modifications are shared across all tRNAs, others are specific to a single tRNA species or domain of life (34,35). These modifications affect tRNA stability, maturation, degradation, and function and have been studied in organisms with a range of optimal growth temperatures, from psychrophiles to hyperthermophiles (16,36–37). Post-transcriptional modifications can have opposing effects on tRNA stability. Pseudouridine, for example, is a common and highly conserved post-transcriptional RNA modification that can stabilize tRNA stem structures, while dihydrouridine has the opposite effect and promotes stem flexibility in tRNAs (38,39). Despite the links between post-transcriptional modifications, optimal growth temperature and tRNA function, details about post-transcriptional modifications were not included in this model. Experimentally determined positions of tRNA post-transcriptional modifications exist for only a few species, and the complexity and specificity of tRNA post-transcriptional modification chemistry means that the accuracy of current prediction models varies considerably across species and is not always better than random assignment (40).

The regression model attaches more importance to the regions of the input tRNA sequence that make up the anticodon and T arm structures, and mutating the structure of the T arm significantly changed the variance of model OGT predictions for the randomly split dataset. Interestingly, the T arm and anticodon arms are the regions of a tRNA at which post-transcriptional modifications are most concentrated. Although the regression model is not given information about post-transcriptional modifications, these modifications are often specific to a certain base and are known to affect tRNA folding and stability, especially in thermophiles (16,37). The fact that both the model with randomly assigned training and test datasets and the model with phylogenetically informed training and test datasets focus on the anticodon and T arms suggests that the CNN may be identifying signals related to post-transcriptional modifications in these regions. The T arm is also important for tRNA structure because its interactions with the D arm bend the tRNA into its appropriate three-dimensional (3D) structure. In its folded state, the T arm forms the elbow region of the 3D tRNA. The elbow region is typically hydrophobic and is important for tRNA interactions with other RNAs and proteins (16).

Disrupting Watson/Crick base pairing in this region of the tRNA increases model OGT prediction variance, suggesting that the model may be recognizing the importance of this region for maintaining tRNA structure and function. Results differ slightly between the random-split model and the phylogenetic-split model. The increase in OGT prediction variance for mutations in the T arm suggests that the random-split model is looking at least partially for speciesspecific differences in this region. The increased correlation between MFE and OGT prediction suggests that the phylogenetic-split model is looking more at overall tRNA stability. However, the model is less accurate because it cannot use information about species relationships or posttranscriptional modifications that would likely influence both MFE and species OGT.

The benefits of these models are threefold. First, the number of prokaryote sequences is growing, but additional information is often not available for these species, and developing culture protocols for new species can be challenging. Understanding likely OGT for new species is useful because it provides a starting point for labs wishing to develop culture protocols and further study these species. Knowing OGT may also be useful in industrial processes requiring thermostable proteins, as this can provide insight into which species proteins are likely to be useful in such processes. Second, by using only the tRNA sequences we created a highly focused model that is independent of other cellular components. We use a minimum proportion of the overall genome sequence for predictions—only  $\sim 0.1\%$  of total DNA in prokaryotes-to predict OGT. This is beneficial for downstream comparisons of temperature effects on protein, DNA, or other RNA features of the cell, as the OGT predictions from the tRNA model are independent of other cell components. Third, by using sequence data as direct inputs to the CNN model, we made use of automatic feature extraction and allowed the model to determine which tRNA features were most relevant. This removed researcher bias and did not require initial assumptions about which components of the tRNA sequence were most important.

#### The importance of phylogenetic relationships between species

Although this model is able to accurately predict OGT within phylogenetic groups, it was unable to maintain high accuracy when predicting across phylogenetic groups, as demonstrated by the drop in prediction accuracy for the phylogenetic split in both the classification and regression models. These results indicate the importance of accounting for phylogeny when trying to extrapolate or draw biological insight from machine learning or other prediction models. In the current models, phylogeny is not part of the dataset, but the model clearly benefits from the shared evolutionary history between species in the dataset.

# **CONCLUSION AND FUTURE WORK**

The current model shows that individual tRNA sequences contain signatures of organism thermal adaptation and that a CNN can pick up on these signals to accurately predict optimal growth temperature. Certain tRNA features, including MFE and GC content seem to be particularly important for determining organism OGT. To minimize inputs and simplify data pre-processing requirements, this model uses tRNA primary DNA sequences. However, incorporating information about secondary and tertiary structures and/or post-transcriptional modifications may improve future iterations of this model, since these features are widely recognized to affect tRNA structure, function and temperature sensitivity (16). Additionally, the current model demonstrates the importance of phylogeny for model predictions, with model accuracy decreasing when phylogenetic relationships between species are hidden. Future studies may wish to determine how far insights from one group of organisms can be transferred - transfer may be limited to within species, clades or superkingdoms, depending on the traits at hand.

# DATA AVAILABILITY

Source code, trained models, predictions and data are available in the Bitbucket repository (https://bitbucket.org/bucklerlab/cnn\_trna\_ogt/). Conda environments to reproduce the computing environments can be found on CyVerse at https://doi.org/10.25739/a0g2-wb14.

# SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online.

#### FUNDING

USDA ARS; Bill and Melinda Gates Foundation [Funding ID: INV-009591/ OPP1159867]. Funding for open access charge: Agricultural Research Service; Bill and Melinda Gates Foundation.

Conflict of interest statement. None declared.

#### REFERENCES

- Jensen, D.B., Vesth, T.C., Hallin, P.F., Pedersen, A.G. and Ussery, D.W. (2012) Bayesian prediction of bacterial growth temperature range based on genome sequences. *BMC Genomics*, 13(Suppl. 7), S3.
- Vieille, C. and Zeikus, G.J. (2001) Hyperthermophilic enzymes: sources, uses, and molecular mechanisms for thermostability. *Microbiol. Mol. Biol. Rev.*, 65, 1–43.
- van Dijk, E., Hoogeveen, A. and Abeln, S. (2015) The hydrophobic temperature dependence of amino acids directly calculated from protein structures. *PLoS Comput. Biol.*, 11, e1004277.
- 4. Rampelotto, P.H. (2013) Extremophiles and extreme environments. *Life*, **3**, 482–485.
- Li,G., Rabe,K.S., Nielsen,J. and Engqvist,M.K.M. (2019) Machine learning applied to predicting microorganism growth temperatures and enzyme catalytic optima. ACS Synth. Biol., 8, 1411–1420.
- Ai, H., Zhang, L., Zhang, J., Cui, T., Chang, A.K. and Liu, H. (2018) Discrimination of thermophilic and mesophilic proteins using support vector machine and decision tree. *Curr. Proteomics*, 15, 374–383.
- Zhang,G. and Fang,B. (2006) Application of amino acid distribution along the sequence for discriminating mesophilic and thermophilic proteins. *Process Biochem.*, 41, 1792–1798.
- Saelensminde, G., Halskau, Ø. Jr, Helland, R., Willassen, N.-P. and Jonassen, I. (2007) Structure-dependent relationships between growth temperature of prokaryotes and the amino acid frequency in their proteins. *Extremophiles*, 11, 585–596.
- 9. Zeldovich, K.B., Berezovsky, I.N. and Shakhnovich, E.I. (2007) Protein and DNA sequence determinants of thermophilic adaptation. *PLoS Comput. Biol.*, **3**, e5.

- Meruelo,A.D., Han,S.K., Kim,S. and Bowie,J.U. (2012) Structural differences between thermophilic and mesophilic membrane proteins. *Protein Sci.*, 21, 1746–1753.
- Wang,G.-Z. and Lercher,M.J. (2010) Amino acid composition in endothermic vertebrates is biased in the same direction as in thermophilic prokaryotes. *BMC Evol. Biol.*, 10, 263.
- Aptekmann, A.A. and Nadra, A.D. (2018) Core promoter information content correlates with optimal growth temperature. *Sci. Rep.*, 8, 1313.
- 13. Sauer, D.B. and Wang, D.-N. (2019) Predicting the optimal growth temperatures of prokaryotes using only genome derived features. *Bioinformatics*, **35**, 3224–3231.
- Holley, R.W., Apgar, J., Everett, G.A., Madison, J.T., Marquisee, M., Merrill, S.H., Penswick, J.R. and Zamir, A. (1965) Structure of a ribonucleic acid. *Science*, 147, 1462–1465.
- 15. Watanabe,Y.-I., Suematsu,T. and Ohtsuki,T. (2014) Losing the stem-loop structure from metazoan mitochondrial tRNAs and co-evolution of interacting factors. *Front. Genet.*, **5**, 109.
- Lorenz, C., Lünse, C.E. and Mörl, M. (2017) tRNA modifications: impact on structure and thermal adaptation. *Biomolecules*, 7, 35.
- Payea,M.J., Sloma,M.F., Kon,Y., Young,D.L., Guy,M.P., Zhang,X., De Zoysa,T., Fields,S., Mathews,D.H. and Phizicky,E.M. (2018) Widespread temperature sensitivity and tRNA decay due to mutations in a yeast tRNA. *RNA*, 24, 410–422.
- Chan, P.P. and Lowe, T.M. (2019) tRNAscan-SE: Searching for tRNA Genes in Genomic Sequences. In: Kollmar, M. (ed). *Gene Prediction. Methods in Molecular Biology*. Vol. **1962**, Humana, NY, pp. 1–14.
- Lowe, T.M. and Eddy, S.R. (1997) tRNAscan-SE: a program for improved detection of transfer RNA genes in genomic sequence. *Nucleic Acids Res.*, 25, 955–964.
- 20. Seemann,T. (2020) barrnap 0.9: rapid ribosomal RNA prediction. https://github.com/tseemann/barrnap.
- Wang, M., Tai, C., Weinan, E. and Wei, L. (2018) DeFine: deep convolutional neural networks accurately quantify intensities of transcription factor-DNA binding and facilitate evaluation of functional non-coding variants. *Nucleic Acids Res.*, 46, e69.
- Zhuang, Z., Shen, X. and Pan, W. (2019) A simple convolutional neural network for prediction of enhancer-promoter interactions with DNA sequence data. *Bioinformatics*, 35, 2899–2906.
- Zeng,H., Edwards,M.D., Liu,G. and Gifford,D.K. (2016) Convolutional neural network architectures for predicting DNA-protein binding. *Bioinformatics*, 32, i121–i127.
- 24. Kingma,D.P. and Ba,J. (2015) Adam: a method for stochastic optimization. In: *International Conference on Learning Representations (ICLR)*, pp. 1–15.
- Bergstra,J.S., Bardenet,R., Bengio,Y. and Kégl,B. (2011) Algorithms for hyper-parameter optimization. In: Shawe-Taylor,J., Zemel,R.S., Bartlett,P.L., Pereira,F. and Weinberger,K.Q. (eds). *Advances in Neural Information Processing Systems*. Curran Associates, Inc., Red Hook, NY, pp. 2546–2554.
- Bergstra, J., Yamins, D. and Cox, D. (2013) Making a science of model search: hyperparameter optimization in hundreds of dimensions for vision architectures. In: *30th International Conference on Machine Learning*. Vol. 28, pp.115–123.
- Washburn, J.D., Mejia-Guerra, M.K., Ramstein, G., Kremling, K.A., Valluru, R., Buckler, E.S. and Wang, H. (2019) Evolutionarily informed deep learning methods for predicting relative transcript abundance from DNA sequence. *Proc. Natl. Acad. Sci. U.S.A.*, **116**, 5542–5549.
- Sayers, E. W., Beck, J., Brister, J.R., Bolton, E.E., Canese, K., Comeau, D.C., Funk, K., Ketter, A., Kim, S., Kimchi, A. *et al.* (2020) Database resources of the National Center for Biotechnology Information. *Nucleic Acids Res.*, 48, D9–D16.
- 29. Benson, D.A., Karsch-Mizrachi, I., Lipman, D.J., Ostell, J. and Sayers, E.W. (2010) GenBank. *Nucleic Acids Res.*, **38**, D46–D51.
- Paradis, E. and Schliep, K. (2019) ape 5.0: an environment for modern phylogenetics and evolutionary analyses in R. *Bioinformatics*, 35, 526–528.
- Ward, J.H. (1963) Hierarchical grouping to optimize an objective function. J. Am. Statist. Assoc., 58, 236–244.
- Sajek,M.P., Woźniak,T., Sprinzl,M., Jaruzelska,J. and Barciszewski,J. (2020) T-psi-C: user friendly database of tRNA sequences and structures. *Nucleic Acids Res.*, 48, D256–D260.
- 33. Kotikalapudi, R. (2017). keras-vis Github.

- 34. Jackman, J.E. and Alfonzo, J.D. (2013) Transfer RNA modifications: nature's combinatorial chemistry playground. *Wiley Interdiscip. Rev. RNA*, **4**, 35–48.
- Barraud, P. and Tisné, C. (2019) To be or not to be modified: miscellaneous aspects influencing nucleotide modifications in tRNAs. *IUBMB Life*, 71, 1126–1140.
- Machnicka, M.A., Olchowik, A., Grosjean, H. and Bujnicki, J.M. (2014) Distribution and frequencies of post-transcriptional modifications in tRNAs. *RNA Biol.*, **11**, 1619–1629.
- 37. Rose, S., Auxilien, S., Havelund, J.F., Kirpekar, F., Huber, H., Grosjean, H. and Douthwaite, S. (2020) The hyperthermophilic partners Nanoarchaeum and Ignicoccus stabilize their tRNA T-loops via different but structurally equivalent modifications. *Nucleic Acids Res.*, 48, 6906–6918.
- Kierzek, E., Malgowska, M., Lisowiec, J., Turner, D.H., Gdaniec, Z. and Kierzek, R. (2014) The contribution of pseudouridine to stabilities and structure of RNAs. *Nucleic Acids Res.*, 42, 3492–3501.
- Dalluge, J.J., Hashizume, T., Sopchik, A.E., McCloskey, J.A. and Davis, D.R. (1996) Conformational flexibility in RNA: the role of dihydrouridine. *Nucleic Acids Res.*, 24, 1073–1079.
- Machnicka, M.A., Dunin-Horkawicz, S., de Crécy-Lagard, V. and Bujnicki, J.M. (2016) tRNAmodpred: a computational method for predicting posttranscriptional modifications in tRNAs. *Methods*, 107, 34–41.
- Kerpedjiev, P., Hammer, S. and Hofacker, I.L. (2015) Forna (force-directed RNA): simple and effective online RNA secondary structure diagrams. *Bioinformatics*, 31, 3377–3379.