Sample-Dependent Adaptive Temperature Scaling
for Improved Calibration

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Abstract
It is now well known that neural networks can be wrong with high confidence in their predictions, leading to poor calibration. The most common post-hoc approach to compensate for this is to perform temperature scaling, which adjusts the confidences of the predictions on any input by scaling the logits by a fixed value. Whilst this approach typically improves the average calibration across the whole test dataset, this improvement typically reduces the individual confidences of the predictions irrespective of whether the classification of a given input is correct or incorrect. With this insight, we base our method on the observation that different samples contribute to the calibration error by varying amounts, with some needing to increase their confidence and others needing to decrease it. Therefore, for each input, we propose to predict a different temperature value, allowing us to adjust the mismatch between confidence and accuracy at a finer granularity. Our method is applied post-hoc, enabling it to be very fast with a negligible memory footprint and is applied to off-the-shelf pre-trained classifiers. We test our method on the ResNet50 and WideResNet28-10 architectures using the CIFAR10/100 and Tiny-ImageNet datasets, showing that producing per-data-point temperatures improves the expected calibration error across the whole test set.

1 Introduction
For neural networks (NNs) to be employed in real-world safety-critical applications, we do not only require them to produce correct predictions, but also provide reliable confidence estimates in their predictions (i.e. they are calibrated). Limiting our scope to neural classifiers, using the maximum probability of the predictive distribution as a confidence measure, literature has established that a mismatch exists between such notion of confidence and the expected accuracy. Indeed, such models generally suffer from being on average overconfident over the test set.

A simple approach to rectify this issue is to perform temperature scaling (Guo et al. 2017), a post-hoc method which scales the logits by a single scalar value, obtained through cross validation. This approach improves the classifier’s performance on standard calibration metrics across a test set. However, from a per-sample point of view there are significant issues. Since the temperature is found by minimising the calibration error (in expectation) over the entire validation set, and since neural networks are overconfident on average, practically speaking, the effect of temperature scaling is to reduce the confidence for every prediction. However, as we will discuss, different samples contribute by varying amounts to the calibration error.

This issue can be seen in Fig. 1, which shows the histogram of the individual contributions to the calibration errors; i.e. the distribution of \( p(y|p_i) - p_i \), where \( p(y|p_i) \) is the accuracy and \( p_i \) is the softmax probability for the data point \( i \), the calibration error can be obtained by taking the weighted average over all the values.¹ Here the mismatch between per-data-point confidence and accuracy is not constant across all the data-points, and hence miscalibration cannot be fixed by scaling the logits by a single fixed value, a key assumption in vanilla temperature scaling. The calibration error varies significantly, with a small (but not insignificant) number of samples on which the network is overconfident. Consequently, scaling the predictions with a single temperature value will adjust all of the errors in the same way. Typically, the temperature values obtained are greater than 1, resulting in a reduction of confidence of all predictions, regardless of whether they are correct with low confidence or incorrect with high confidence.

To combat this, we propose a method which produces per-data-point predictions of the temperature, permitting an adequate decrease in the confidence on samples which the classifier is likely to get wrong, and an increase in the confidence on predictions it is likely to get correct. As a result, we obtain better test Expected Calibration Error (ECE) (Guo et al. 2017) both on in-distribution sets (i.e. the test set is i.i.d. with respect to the training set) and under covariate-shifted sets (i.e. the test set shares the same set of labels of the training set, but the inputs are not i.i.d. with respect to the training set).

Like temperature scaling, our method is applied post-hoc and is very fast to train and test. We extensively test the calibration of ResNet50 (He et al. 2016) and WideResNet28 (Zagoruyko and Komodakis 2016) when using our method on CIFAR10/CIFAR100 and TinyImageNet, in-

¹Here \( p(y|p_i) \) is obtained through histogram binning and represents the accuracy of each bin, and the weights are proportional to the number of samples in each bin.
ing the softmax, i.e., scaling the logits by a constant factor than their expected accuracy (Guo et al. 2017).

overconfident as the confidence of the predictions are higher than their accuracy. Typically, neural networks are expected to be 80% on the set of points. Typically, neural networks are expected to be 80% on the set of points. Then, the accuracy should be 80% on the set of points. Typically, neural networks are expected to be 80% on the set of points. One might suggest introducing the temperature as one of the outputs of the network and jointly learning it as part of the training process. Here we outline why this approach of learning to predict the temperature values \( T \) and the predictive probabilities \( p \) might result in suboptimal performance. Consider the last layer of a NN with parameters \( \boldsymbol{w} \in \mathbb{R}^{D \times K} \) for a feature space of size \( D \) and the cross entropy loss \( \mathcal{L} : \mathbb{R}^{K} \rightarrow \mathbb{R} \). The gradient for the layer is given as

\[
\frac{\partial \mathcal{L}}{\partial \bar{w}} = \frac{\partial s}{\partial \bar{w}} (\sigma(s) - y),
\]

where \( y \) denotes the one-hot ground-truth label and \( \sigma(s) - y = \{\sigma(s_j) - y_j : j \in \{1 \ldots K\}\} \), \( \bar{w} \) indicates the network weights are flattened to column vector form. Inspecting the gradients indicates that the gradient starts to vanish when \( s_k \rightarrow \infty \) and \( s_{\setminus k} \rightarrow -\infty \), where \( k \) is the correct class. Or to put it simply, the optimisation does not converge until the network produces one-hot logits.

This forces the magnification of the network weights (Mukhoti et al. 2020), which subsequently leads to an overconfident network and hence miscalibrated predictions. A mechanism to achieve the desired one-hot prediction without magnifying the weights could be instead to naively learn the temperature alongside the logits, assuming the model is trainable and converges. In this case, gradient updates would decrease the value of \( T \), resulting in a lower-entropy distribution that is more "peaky". We now discuss why this approach might not work well in practice.

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If we consider the gradient of the temperature, which is
given as
\[
\frac{\partial \mathcal{L}}{\partial T} = \sum_k y_k \frac{\exp \left( \frac{s_k}{T} \right)}{T^2} - \sum_i \sum_k \frac{\exp \left( \frac{s_i}{T} \right)}{T^2},
\]
(2)

which decreases the value of \( T \) for a correct prediction \( \hat{k} = \arg \max_k s_k \), leading to more confident predictions. Typically, the train accuracy will approach 100%, meaning that gradient updates to \( T \) cause it to decrease without any moderation, preventing the network from learning how to predict \( T \) appropriately. In short, there is essentially only data for correct predictions, preventing crucial information on how the network should behave when it's wrong being incorporated in the learning process. Consequently, learning \( T \) naïvely is not a feasible option as the network just learns to be confident everywhere.

**Learning to Calibrate**

Our approach involves a new temperature prediction module (a small neural network) that operates on each input sample independently and whose objective is to extract information from the trained model itself in order to calibrate the confidences of each prediction. We call this learning to calibrate.

Doing so requires learning a temperature prediction module on a data-set consisting of data-points \( \mathcal{X}_{cal} = \{ x_n \}_N \), \( \mathcal{X}_{train} \cap \mathcal{X}_{cal} = \emptyset \), neural network predictions \( p_{cal} = \{ p_n \}_N \) and labels \( y_{cal} = \{ y_n \}_N \), \( y \in \{ \ldots , K \} \) for \( K \) classes. It is important to note that the objective here is to learn to assign low confidences to data points which are likely to be incorrect and high confidences to those which are likely to be correct.

Specifically for a given data-point \( x \in \mathcal{X}_{cal} \), we propose to optimise the temperature prediction module over \( T \) by maximising the log probability of the label \( y \) under the categorical probability distribution parametrised by the \( T \)-scaled logits \( s \), i.e. \( T^* = \arg \max_T \log \text{Cat}(y; \text{softmax}(s/T))^2 \). Here we do not optimise \( s \) but keep it fixed; we are only optimising w.r.t \( T \).

In situations where \( y = \arg \max_k p_k \) (i.e. correct prediction), the target function is maximised when \( T \to 0 \), as we want the predicted probabilities to match the one-hot logits, e.g. see \( T = 0.1 \) in Fig. 2. This is equivalent to minimising the entropy of the predictive distribution by only manipulating \( T \), which is the desired outcome for a correct prediction.

In situations where the prediction is incorrect, \( y \neq \arg \max_k p_k \), to maximise the target function we need to maximise \( p_{\hat{k}} \) and minimise \( p_k \), where \( \hat{k} = \arg \max_k p_k \). As the temperature prediction module cannot change the predicted label, the optimization accepts the incorrect prediction and maximise the target function by flattening the Softmax outputs with \( T \gg 1 \), which is equivalent to maximising the entropy of the predictive distribution. This effect can be seen by considering the case where predicting class 2 in Fig. 2 is the incorrect prediction; among the three cases shown, \( T = 10 \) maximises \( \text{Cat}(y \neq 2; \text{softmax}(s/T)) \).

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1Which is equivalent to the cross entropy loss.

**4 Adaptive Temperature Scaling**

We are now ready to outline the specifics of our proposed temperature prediction module. Given a data-set \( \mathcal{X}_{cal} \), we want to learn which samples the classifier should be confident about and which it should not. Rather than acting on the image space, we instead use the feature extractor of the classifier, as it has already learnt how to extract the information needed for class prediction and also contains a notion of the associated confidence. This has been demonstrated in (Guo et al. 2017), where working only on the feature space has already provided highly promising results in calibrating models for a variety of tasks; suggesting that the feature space already contains sufficient information for calibration. What we now need is a method to extract this confidence information from the feature space and leverage it appropriately to calibrate the predictions.

**Representing Uncertainty with the Variational Autoencoder** variational autoencoders (VAEs) (Kingma and Welling 2013) act as an efficient model to obtain representations of data; the representations encapsulate the generative factors in a lower-dimensional subspace and are rich enough to reconstruct the data sample. In the specification of the generative model, the user has to specify a prior over the latent variables (typically an isotropic Gaussian) where the KL distance between the prior and approximate posterior is minimised during training. Unlike a standard autoencoder (Hinton and Zemel 1994), there is now a mechanism to obtain a likelihood on the latent codes. In reality this value forms part of the importance weight and can be used as a proxy to the true likelihood but avoids issues associated with deep generative models (Nalisnick et al. 2019).

From a mechanistic point of view, we expect samples which are much more common to be placed in the centre of the prior. Here we leverage this idea and use the latent likelihoods as a basis to predict the temperature value. Indeed, we find empirically that this approach works well in practice. Rather than using an isotropic Guassian as the prior, we instead introduce a Gaussian mixture prior, with component for each class specified by the learnable parameters \( \lambda_k = \{ \mu_k, \sigma_k \} \in \mathbb{R}^{D_k} \), such that \( p_k(z) = \frac{1}{K} \sum_k p_{\lambda_k}(z \mid y) \).

This allows for an individual unimodal prior for each class; preventing any issue with clusters for individual classes being placed in lower likelihood regions of the latent space, as would be the case if an isotropic prior is used. With this mixture prior, the evidence lower bound is given as

\[
\log p(\Phi(x), y) \geq \mathbb{E}_{q_\psi(\Phi(x))} \log \frac{p_{\Theta}(\Phi(x) \mid z) p_{\lambda_k}(z \mid y)}{q_\psi(\Phi(x))},
\]

where \( q_\psi(\Phi(x)) \), \( p_{\Theta}(\Phi(x) \mid z) \) and \( p_{\lambda_k}(z \mid y) \) represent the encoder, decoder and mixture prior component, the parameters of the VAE are given as \( \Theta = \{ \theta, \psi, \lambda_1, \ldots , \lambda_K \} \). The parameters of the mixture prior are learnt alongside the parameters of the encoder and decoder(Tomczak and Welling 2018). The use of this mixture prior forces the aggregate posterior for each class to match a Gaussian distribution, i.e. \( \sum_{x \in X_k} q(z \mid \Phi(x)) \approx \mathcal{N}(z; \mu_k, \sigma_k) \). This encourages the
Temperature Prediction Network

Given that the VAE structures the latent in space in a way which makes it amenable to confidence prediction, we learn a very simple MLP parameterised by $\theta$, which predicts the temperature based on the latent embeddings, using the cross entropy loss as an objective. Rather than using the latent samples as input to the MLP, given the observations in Fig. 3, we choose to predict the temperature as a function of the vector of log-likelihoods on all of the conditional priors, specifically $T = g_\theta(\bar{q})$ where $g : \mathbb{R}^K \rightarrow \mathbb{R}$ is the MLP which predicts the temperature and $\bar{q} = \{\log p_{\lambda_y}(z \mid y)\}_{y \in \mathcal{Y}}$, i.e., each element $\bar{q}_y$ contains the log-likelihood of $z$ on the corresponding conditional prior $p_{\lambda_y}(z \mid y)$. Evaluating $\log p_{\lambda_y}(z \mid y)$ can be viewed as a pseudo likelihood of $x$, consequently the module predicts the temperature as a non-linear transform of a pseudo-likelihood of the sample. It is also important to point out that due to the use of feature space as the input, we are able to use small architectures, making this approach very fast during training and at test time. We represent a high level overview and the graphical model in Fig. 4.

Calibrated Training Details

The overall post-hoc learning algorithm is very simple and the module can be trained in under a minute on an 8Gb Titan Xp for most datasets, depending on the validation set and feature space size, we give an overview of the procedure in Alg. 1. We combine learning the VAE and the temperature prediction network into one objective. Specifically, we maximise the following objective

$$\mathcal{L}(x, y) = \mathcal{L}_{ELBO}(\Phi(x)) + \log \text{Cat}(y \mid \text{softmax}(s / g_\theta(\bar{q})))$$

(4)

with $\bar{q} = \{\log p_{\lambda_y}(z \mid y)\}_{y \in \mathcal{Y}}$ and using Normal distributions for $z$; Laplace distribution over $\Phi(x)$ (L1 loss); and a Categorical over $y$. To train the VAE, we use a held-out dataset from training the network, i.e., $\mathcal{X}_{\text{train}} \cap \mathcal{X}_{\text{cal}} = \emptyset$. We used the Adam optimiser with a learning rate of 0.001 and trained for 50 epochs. This is an additional benefit of training the VAE on the $\Phi(x)$, as the feature space has a lower dimensionality and simpler structure than the image space, leading to much faster training and the ability to use simpler networks.

Calibration at Test Time During test time, the features of the data point $\Phi(x)$ and predicted logits are computed using the classifier $s = f(\Phi(x))$, the temperature can then be predicted through $T = g_\theta(\bar{q})$, where $\bar{q} = \{\log p_{\lambda_y}(z \mid y)\}_{y \in \mathcal{Y}}$ with $z = q_\phi(z \mid x)$. The calibrated predictions are then computed as $p = \sigma(s/T)$. 

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Algorithm 1: Learning Adaptive Temperature

Require: \( X_{\text{cal}}, Y_{\text{cal}}, P_{\text{cal}} \)

1: while not converged do
2: \( x, y \leftarrow \text{Random batch} \)
3: \( \nabla_{VAE} \leftarrow \nabla_{\text{ELBO}}(\Phi(x)) \)
4: \( q = \{ \log p_{x}(z \mid y) \mid y \} \quad z \sim \alpha(z|\Phi(x)) \)
5: \( \nabla_T \leftarrow \nabla \log \text{Cat}(y; \text{softmax}(s/g_{\theta}(q))) \)
6: \( \{\Theta, \theta\}_{t+1} \leftarrow \{\Theta, \theta\}_t - \alpha(\nabla_{VAE} + \nabla_T) \)
7: end while

5 Results

Before evaluating the model, we define the hypothesis we are trying to test. Specifically, we want to evaluate if predicting the temperature on a per-data-point basis leads to improved calibration over vanilla temperature scaling. Secondly, we wish to investigate how adaptive temperature performs under dataset shift.

We performed our experiments on the WideResNet28-10 (Zagoruyko and Komodakis 2016) architecture. We report calibration results on CIFAR10/CIFAR100 (Krizhevsky, Hinton et al. 2009) and Tiny-ImageNet (Torralba, Fergus, and Freeman 2008). We conducted distribution-shift experiments using the variants CIFAR10-C/CIFAR100-C to test for domain shift (Hendrycks and Dietterich 2019). We used the following as models for our evaluation:

• Cross Entropy Loss, due to its popularity and wide adoption.
• Brier Score (Brier et al. 1950), due to its ability to obtain well calibrated predictions (Mukhoti et al. 2020).
• Deep Ensembles (Lakshminarayanan, Pritzel, and Blundell 2016), as it achieves state of the art results.

Results are obtained for multiple seeds for Cross Entropy and Brier Score, but only one seed for Deep Ensembles due to the number of models needed.

Calibration

Here we evaluate how adaptive temperature scaling affects standard calibration metrics compared to vanilla temperature scaling. We report results using the ECE, which divides the probability into equally sized bins and then computes the absolute difference between confidence and accuracy for each bin before taking the average. However, the ECE is known to be a biased estimator (Ding et al. 2020), with its performance depending on the binning size and on the distribution of samples in each bin. For this reason, the reliability of the ECE as a miscalibration metric is being questioned and several alternatives have been proposed (e.g. Nixon et al. 2019; Roelofs et al. 2022; Mukhoti et al. 2020). Among these, we choose to also use the AdaECE (Mukhoti et al. 2020), which uses adaptive bin sizes to ensure each bin contains the same number of samples.

We report the results in Tab. 1 where it can be seen that adaptive temperature scaling improves calibration compared to standard temperature scaling. In all cases our method is able to outperform vanilla temperature scaling, with large improvements obtained when using the cross entropy loss, e.g. 0.93 → 0.76 and 3.76 → 2.95 ECE for CIFAR10 and CIFAR100 when using the WideResNet2810 Network.

| Model | Scaling | Accuracy | ECE | AdaECE |
|-------|---------|----------|-----|--------|
| CIFAR10 | CE | 95.52 ± 0.4 | 2.15 ± 0.1 | 2.13 ± 0.1 |
| | TS | 95.52 ± 0.4 | 0.93 ± 0.2 | 0.98 ± 0.3 |
| | AdaTS | 95.52 ± 0.4 | 0.76 ± 0.1 | 0.86 ± 0.2 |
| | Brier | 95.84 ± 0.1 | 0.92 ± 0.2 | 1.50 ± 0.1 |
| | TS | 95.84 ± 0.1 | 1.88 ± 0.2 | 1.94 ± 0.1 |
| | AdaTS | 95.84 ± 0.1 | 1.65 ± 0.1 | 1.61 ± 0.1 |
| Ensm | VTS | 96.35 | 1.68 | 1.61 |
| | AdaTS | 96.37 | 0.51 | 0.46 |
| CIFAR100 | CE | 80.71 ± 0.1 | 5.76 ± 0.1 | 5.70 ± 0.1 |
| | TS | 80.71 ± 0.1 | 3.76 ± 0.2 | 3.68 ± 0.3 |
| | AdaTS | 80.71 ± 0.1 | 2.95 ± 0.4 | 2.90 ± 0.4 |
| | Brier | 79.25 ± 0.1 | 4.19 ± 0.2 | 4.13 ± 0.2 |
| | TS | 79.25 ± 0.1 | 3.87 ± 0.6 | 3.90 ± 0.6 |
| | AdaTS | 79.25 ± 0.1 | 3.67 ± 0.8 | 3.64 ± 0.7 |
| Ensm | VTS | 83.19 | 4.24 | 4.21 |
| | TS | 83.18 | 3.71 | 3.55 |
| | AdaTS | 83.22 | 2.95 | 2.66 |
| Tiny-ImageNet | CE | 60.47 ± 0.1 | 7.54 ± 4.0 | 7.53 ± 4.0 |
| | TS | 60.47 ± 1.1 | 6.28 ± 2.4 | 6.15 ± 2.4 |
| | AdaTS | 60.94 ± 1.2 | 5.18 ± 1.4 | 5.17 ± 1.3 |
| | Brier | 50.23 ± 0.4 | 5.56 ± 0.6 | 5.52 ± 0.6 |
| | TS | 50.23 ± 0.4 | 4.55 ± 0.2 | 4.43 ± 0.6 |
| | AdaTS | 50.23 ± 0.4 | 4.43 ± 0.4 | 4.21 ± 0.5 |
| Ensm | VTS | 66.16 | 6.21 | 6.19 |
| | TS | 66.16 | 5.12 | 5.06 |
| | AdaTS | 66.00 | 4.58 | 4.41 |

Table 1: Calibration results, here we can see that adaptive temperature scaling is able to improve calibration on a variety of models. Bold indicates the best result, or within one standard deviation of the best result.

Data-Shift

A key hypothesis we want to test is how adaptive temperature scaling behaves under data-shift. Specifically, we use the widely used CIFAR10-C and CIFAR100-C datasets, which are corrupted versions of the CIFAR10 and CIFAR100 (Hendrycks and Dietterich 2019).

The dataset consists of standard CIFAR images which have undergone 15 synthetic corruptions (e.g. noise, weather conditions, image properties) at varying levels. Within this context, we use Deep Ensembles (Lakshminarayanan, Pritzel, and Blundell 2016), as it achieves state of the art results.

We include experiments for a standard autoencoder and MLP in the appendix.
Figure 5: Left: How temperature varies when interpolating between class feature means on CIFAR-10. Right: Histogram of temperature values for each image in CIFAR-10, here we can see that typically objects have a lower temperature than animals, indicating they are easier to classify.

Figure 6: How AdaECE changes with varying levels of motion-blur corruptions on CIFAR-10. Adaptive temperature consistently produces lower error rates.

scenario, the classifier should either be robust to such corruptions (retaining accuracy) or if the accuracy is compromised, reduce the confidences accordingly. As such, we report the test accuracy as well as ECE and AdaECE in Tab. 2, where adaptive temperature scaling shows improvements over temperature scaling.

We also expect to see adaptive temperature scaling provide improvement over temperature scaling as the intensity of corruptions are increased for CIFAR-10-C. We generate plots highlighting the AdaECE calibration metric as the level of the corruption intensity is increased; the plot for motion-blur is displayed in Fig. 6. Here despite a general increase in error for all methods adaptive temperature scaling consistently produces lower error rates than vanilla temperature scaling (orange) and vanilla predictions (blue). More examples are provided in the Appendix.

**Behaviour of the Temperature Prediction Module**

Can the temperature module predict high temperature in uncertain regions? If yes, then we should see a change in the temperature as we traverse the feature space. To conduct this experiment, inspired by the analysis provided in (Pinto et al. 2021, 2022), we obtain the average feature representation for each class $\Phi_k = \frac{1}{|X_k|} \sum_{x \in X_k} \Phi(x)$ and measure the temperature when interpolating between two classes. i.e. we predict the temperature for the features $\{\alpha \Phi_k + (1 - \alpha) \Phi_j \}$ $\alpha \in [0, 1]$. We plot the interpolation results in Fig. 5 (Left) for the classes in CIFAR-10, where the horizontal axis represent $\alpha$ and the vertical axis represents the temperature. For some classes we see a significant rise in the temperature as we interpolate between two classes, e.g. automobile and bird. This highlights the temperature prediction models ability to assign a low temperature in regions that the classifier is certain about, e.g. around the mean and a higher temperature in less certain regions, e.g. heavily interpolated regions.

Interestingly, this feature is not present for all class pairs; for some, e.g. cat and dog, where the temperature remains high between classes. We hypothesise that this is due to an interpolation between these classes being a plausible realisation of an image, unlike for automobile and bird.

**Misclassification Rejection**

Calibrated uncertainty estimates should render that the models are able to reject samples in order to preserve the accuracy. In this setting we report results for AURRA, which

| Model     | Scaling | Accuracy | ECE  | AdaECE |
|-----------|---------|----------|------|--------|
| CE        | -       | 75.07 ± 1.4 | 15.70 ± 1.1 | 15.68 ± 1.1 |
| CE       | TS      | 75.07 ± 1.4 | 12.19 ± 0.9 | 12.17 ± 0.9 |
| CE       | AdaTS   | 75.07 ± 1.4 | 12.03 ± 1.3 | 12.02 ± 1.3 |
| Brier    | -       | 75.27 ± 0.7 | 16.21 ± 0.8 | 16.45 ± 0.7 |
| Brier    | TS      | 75.27 ± 0.7 | 15.87 ± 0.4 | 15.86 ± 0.4 |
| Brier    | AdaTS   | 75.27 ± 0.7 | 14.84 ± 0.8 | 14.81 ± 0.9 |
| Ensm     | -       | 77.28      | 13.45 | 13.43  |
| Ensm     | TS      | 77.28      | 10.12 | 10.09  |
| Ensm     | AdaTS   | 77.21      | 9.29  | 9.25   |

Table 2: Corrupted calibration results. Here we can see that adaptive temperate scaling is able to improve calibration on a variety of models. Bold indicates best results, or within one standard deviation of best results.

6assuming features of clean and corrupted inputs are not mapped exactly to the same point in the embedding space.
Table 3: AURRA scores for based on: confidence (AURRA-C), Demster-Schafer (Sensoy, Kaplan, and Kandemir 2018) (AURRA-DS) and entropy (AURRA-E). Unlike temperature scaling, adaptive temperature scaling does not suffer a reduction in rejection ability. Higher is better.

| Methods | AURRA-C       | AURRA-DS       | AURRA-E       |
|---------|---------------|---------------|---------------|
| None    | 93.07 ± 4.28  | 91.84 ± 5.24  | 92.95 ± 4.23  |
| Vanilla TS | 92.97 ± 4.25  | 91.70 ± 5.40  | 92.67 ± 4.41  |
| Adaptive TS | 93.20 ± 4.25  | 92.00 ± 5.39  | 92.99 ± 4.35  |
| None    | 84.21 ± 1.09  | 81.83 ± 0.64  | 83.69 ± 1.16  |
| Vanilla TS | 84.05 ± 1.09  | 81.63 ± 0.64  | 83.31 ± 1.11  |
| Adaptive TS | 84.68 ± 0.18  | 81.93 ± 0.28  | 84.09 ± 0.20  |

Table 4: CIFAR-10.1, here we see that adaptive temperature scaling is able to provide slightly improved calibration on the harder CIFAR-10.1 dataset.

| Methods | Accuracy (%) | ECE (%) | AdaECE (%) |
|---------|--------------|---------|------------|
| None    | 85.86 ± 2.48 | 8.35 ± 1.58 | 8.15 ± 1.68 |
| Vanilla TS | 85.86 ± 2.48 | 4.57 ± 0.32 | 4.24 ± 0.43 |
| Adaptive TS | 85.86 ± 2.48 | 3.67 ± 1.41 | 3.35 ± 1.39 |

Evaluating Hardness

Given our models ability to predict the temperature, it should naturally extract a notion of hardness, that is how difficult it is to classify. One would expect hard samples to have a high temperature and easy ones to have a low temperature. To conduct this experiment, we utilise the CIFAR-10.1 (Recht et al. 2018) dataset, which contains “harder”, but statistically similar images to CIFAR-10; consequently this experiment is not examining data-shift, but is instead measuring the performance on challenging samples. We report the standard metrics: accuracy, ECE and AdaECE in Tab. 4, where we see that adaptive temperature is able to obtain a lower calibration error than vanilla temperature scaling when the model is trained using cross entropy loss.

A key hypothesis we wish to test is “does the model assign higher temperatures to harder samples?”; harder samples should naturally contain a greater amount of uncertainty in their predictions. Consequently, we should see higher temperatures assigned to harder samples (CIFAR-10.1) than to easier ones (CIFAR-10). We test this hypothesis by plotting the histogram of temperature values for CIFAR-10 and CIFAR-10.1, for both correct and incorrect predictions in Fig. 7 (Right).

Here we see that generally, correct samples for CIFAR-10 (blue) are assigned a lower temperature than for CIFAR-10.1 (green), indicating that the adaptive temperature is able to recognise harder samples and assign a higher temperature increasing the uncertainty. We also see that adaptive temperature predicts higher temperatures for incorrect predictions on CIFAR-10 (orange), highlighting adaptive temperatures ability to reduce the confidence of samples which are likely to be incorrect. The same is not true for CIFAR-10.1, this is due to the fact that the samples from CIFAR-10.1 are by design harder, adaptive temperature predicts higher values of $T$ than for the easier CIFAR10 counterpart.

6 Discussion

Here we have presented Adaptive Temperature Scaling (AdaTS), a novel post-hoc method for predicting the temperature on per-sample basis. Given a data-point, our method can predict how confident the classifier should be about its prediction, providing the critical flexibility to increase of decrease the confidence, leading to improved calibration error. The adaptive temperature is also able to obtain better results under distribution shifts. This is achieved by leveraging the latent space of a VAE, which we found to naturally encapsulate and structure the information relating to confidence appropriately. As the model is applied post-hoc, training is very fast, requiring little computational overhead, furthermore it is very easy to implement.
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