Measuring Stochastic Data Complexity with Boltzmann Influence Functions

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Abstract

Estimating the uncertainty of a model's prediction on a test point is a crucial part of ensuring reliability and calibration under distribution shifts. A minimum description length approach to this problem uses the predictive normalized maximum likelihood (pNML) distribution, which considers every possible label for a data point, and decreases confidence in a prediction if other labels are also consistent with the model and training data. In this work we propose IF-COMP, a scalable and efficient approximation of the pNML distribution that linearizes the model with a temperature-scaled Boltzmann influence function. IF-COMP can be used to produce well-calibrated predictions on test points as well as measure complexity in both labelled and unlabelled settings. We experimentally validate IF-COMP on uncertainty calibration, mislabel detection, and OOD detection tasks, where it consistently matches or beats strong baseline methods.

1 Introduction

Safely deploying machine learning models in real world settings requires making accurate predictions, as well as quantifying the uncertainty in those predictions. This is particularly important in high-stakes settings such as healthcare (Ghassemi & Mohamed, 2022), medical imaging (Esteva et al., 2017), and self-driving cars (Bojarski et al., 2016), where uncertainty estimates can help accurately assess the risk in utilizing a model's decision and decide when to ignore it altogether (Ovadia et al., 2019). Common methods for quantifying uncertainty rely on Bayesian principles, which require defining a prior distribution and sampling from a posterior distribution. However, specifying a good prior is difficult in the context of modern deep learning, and Bayesian methods face significant scalability challenges.

The Minimum Description Length (MDL) Principle (Rissanen, 1996) provides an alternative approach to uncertainty estimation which does not require explicitly defining a prior or even a notion of ground truth. MDL instantiates a version of Occam's Razor by favoring models that minimize the combined codelength of both the model and the observed data under a suitable coding scheme. This code can then be used to make a prediction for an unseen test point by selecting the label with the shortest code. However, this code may perform poorly compared to a hindsight-optimal code that is allowed to observe the test point and label. We can attempt to minimize this difference in codelengths, or regret, by using a universal code such as the predictive normalized maximum likelihood (pNML) (Roos & Rissanen, 2008; Fogel & Feder, 2018) distribution, which minimizes the maximum possible regret for all labels.

Intuitively, given a query data point \mathbf{x} , pNML considers augmenting the training set with the pair (\mathbf{x}, \mathbf{y}) for some label \mathbf{y} and fitting a hindsight-optimal model to the augmented training set. If one can just as easily fit the augmented dataset with arbitrary labels, then one is highly uncertain about the label; if one choice of labels is much easier to fit than the others, then one has low uncertainty. This intuition is captured

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by the minimax optimality of pNML. The universal codelength pNML defines for a data point is called the stochastic data complexity (Rissanen, 1986), and consists of two terms. The error term captures the best possible loss a hindsight-optimal model could achieve by observing the true label. The second parametric complexity term quantifies how many essentially different distributions the model class could assign to the test data. Minimizing the pNML codelength then requires both learning an accurate model of the data as well as avoiding overfitting outlier observations.

Calculating the pNML distribution requires optimizing a hindsight-optimal model over the training set and the additional example for each possible label, which is both computationally intractable and misspecified for overparameterized neural networks that can easily fit random labels (Zhang et al., 2017). A simple way to restrict the hindsight-optimal model subclass is with a proximal objective that penalizes movement in function and weight space. Approximating this proximal objective by linearizing the network and applying a second order approximation of the proximal terms has been shown to correspond to the influence function (IF) in neural networks (Bae et al., 2022).

However, when models are overconfident on training examples, the proximal terms can become too restrictive, making the hindsight-optimal model unable to fit low probability labels. In this work we propose to temperature-scale the proximal objective and approximate it with a Boltzmann influence function (BIF), allowing the hindsight-optimal model to better accommodate arbitrary labels. We then propose Influence Function Complexity (IF-COMP), a complexity measure and associated pNML code which uses the BIF to produce estimates of the hindsight-optimal output probabilities. These estimates can then be used to produce calibrated output distributions as well as measure the stochastic complexity of both labelled and unlabelled examples.

We validate IF-COMP's ability to reliably estimate ground truth pNML complexity, then investigate its use on three tasks that test its different capabilities. We consider (1) uncertainty calibration, which requires producing reliable uncertainty estimates under distribution shifts, (2) mislabel detection, which requires estimating complexity for labelled training examples, and (3) OOD detection, which requires estimating complexity for unlabelled test examples. Across all three tasks, IF-COMP displays strong performance, consistently matching or beating Bayesian and optimization tracing approaches that often use more information than is available to our method, as well as a similar pNML approximation method that explicitly takes steps in parameter space (Zhou & Levine, 2021). Compared to this baseline, IF-COMP also provides a 7-15 times speedup in computational efficiency. IF-COMP demonstrates the potential of MDL-based approaches for uncertainty and complexity estimation in deep neural networks, enabling better calibrated decision making.

2 Background and Preliminaries

2.1 Minimum Description Length and Stochastic Complexity

In this work we consider a supervised classification task from an input space \mathcal{X} to a discrete output space \mathcal{Y} , where we are given a finite training set $\mathcal{D}_{\text{train}} = \{(\mathbf{x}_i, \mathbf{y}_i)\}_n$ consisting of pairs of examples $(\mathbf{x}_i, \mathbf{y}_i)$. We consider a hypothesis class of possible models $\{\boldsymbol{\theta} \in \boldsymbol{\Theta}\}$, each of which defines a conditional probability distribution $p_{\boldsymbol{\theta}}(\mathbf{y}|\mathbf{x}) = \sigma(f_{\boldsymbol{\theta}}(\mathbf{x}))$ where σ is the softmax function. We can define a *prefix code* for \mathbf{y} using the Kraft inequality (Cover & Thomas, 1991), such that the codelength $\mathcal{L}(\boldsymbol{\theta}, \mathbf{x}, \mathbf{y})$, or number of bits required to describe \mathbf{y} given \mathbf{x} , is:

$$\mathcal{L}(\boldsymbol{\theta}, \mathbf{x}, \mathbf{y}) = -\log p_{\boldsymbol{\theta}}(\mathbf{y}|\mathbf{x}), \tag{1}$$

or the log loss.

The Minimum Description Length (MDL) Principle (Rissanen, 1978; Grunwald, 2004) dictates that we should choose the model in our hypothesis class whose code can describe the training labels given the

training inputs in the fewest number of bits. For parametric hypothesis classes such as neural networks, the MDL principle reduces to the maximum likelihood principle (Grunwald, 2004; Dwivedi et al., 2023). Then, a given learner q will aim to find the parameters $\hat{\theta} \in \Theta$ that minimizes the standard training objective:

$$\mathcal{J}(\boldsymbol{\theta}, \mathcal{D}_{\text{train}}) = \sum_{i=1}^{n} \mathcal{L}(\boldsymbol{\theta}, \mathbf{x}_i, \mathbf{y}_i)$$
 (2)

We call the resulting model the base model and the codelength it defines for an example the base codelength.

We now consider an unseen test example $\mathbf{z} = (\mathbf{x}, \mathbf{y})$, for which our base model may not achieve the optimal codelength. If q is additionally given access to this test example, it will be able to find the parameters that minimize the total codelength on both the training set and the test example:

$$\boldsymbol{\theta}^{\star}(\mathbf{x}, \mathbf{y}, \epsilon) = \underset{\boldsymbol{\theta} \in \boldsymbol{\Theta}}{\arg \min} \, \mathcal{J}(\boldsymbol{\theta}, \mathcal{D}_{\text{train}}) - \epsilon \log p_{\boldsymbol{\theta}}(\mathbf{y}|\mathbf{x})$$
(3)

where ϵ represents the amount to weight the test point relative to the training set. We call the resulting model the *hindsight-optimal model* and the codelength it defines for an example the *hindsight-optimal codelength*. Typically we will always set $\epsilon = 1/n$ so that the test point is weighted equally to the other training points, but we include this additional parameter for reasons that will become clear later. For ease of notation, we define the hindsight-optimal parameters as $\theta^*(\mathbf{x}, \mathbf{y}) := \theta^*(\mathbf{x}, \mathbf{y}, 1/n)$.

The regret of the learner q for a test point \mathbf{z} is the difference between the base codelength and the hindsight-optimal codelength:

$$R(q, \mathcal{D}_{\text{train}}, (\mathbf{x}, \mathbf{y})) = -\log p_{\hat{\boldsymbol{\theta}}}(\mathbf{y}|\mathbf{x}) + \log p_{\boldsymbol{\theta}^{\star}(\mathbf{x}, \mathbf{y})}(\mathbf{y}|\mathbf{x}). \tag{4}$$

The predictive normalized maximum likelihood (pNML) distribution (Shtar'kov, 1987; Roos et al., 2008; Roos & Rissanen, 2008) is then defined as the universal model which minimizes the worst case regret across all possible labels:

$$p_{\text{pNML}}(\mathbf{y}|\mathbf{x}) = \frac{p_{\boldsymbol{\theta}^{\star}(\mathbf{x},\mathbf{y})}(\mathbf{y}|\mathbf{x})}{\sum_{\mathbf{y}' \in \mathcal{Y}} p_{\boldsymbol{\theta}^{\star}(\mathbf{x},\mathbf{y}')}(\mathbf{y}'|\mathbf{x})}.$$
 (5)

The codelength of the pNML distribution is

$$\Gamma(\mathbf{z}) = -\log p_{\boldsymbol{\theta}^{\star}(\mathbf{x}, \mathbf{y})}(\mathbf{y}|\mathbf{x}) + \log \sum_{\mathbf{y}' \in \mathcal{Y}} p_{\boldsymbol{\theta}^{\star}(\mathbf{x}, \mathbf{y}')}(\mathbf{y}'|\mathbf{x}),$$

$$(6)$$
parametric complexity

also known as the stochastic complexity (Rissanen, 1996; Barron et al., 1998) of z relative to the model class Θ . The first error term measures the hindsight-optimal codelength or log loss for z. The second parametric complexity term quantifies how many distinguishable distributions the model class Θ could assign to the data in hindsight. A model that is expressive enough to overfit its training data and any arbitrary test data will be able to achieve low error but at the cost of high parametric complexity. A less expressive model that underfits its training data and any arbitrary test data will have low parametric complexity but will incur high error. Choosing a model with a small description length under the pNML code necessitates trading off between these two extremes.

2.2 The Infinity Problem and Proximal Bregman Objective

For many overparameterized model classes, the denominator in (5) is either infinite (for continuous label spaces) or constant (for discrete label spaces). To solve this *infinity problem*, the hindsight-optimal model

must be restricted to a subclass of models that "comply" with the original training data (Fogel & Feder, 2019). This can be done by restricting the label space (Stine & Foster, 2001), enforcing moment matching (Farnia & Tse, 2017), confidence intervals, or minimum training likelihood (Fogel & Feder, 2019).

Another approach directly alters the objective with a proximal term, such as in a ridge estimator (Dwivedi et al., 2023). In this work we propose to use a proximal bregman objective (PBO) (Bae et al., 2022) that restricts movement in function and weight space away from the base model optimum while training on the additional unseen test example:

$$Q_{\beta}(\boldsymbol{\theta}, \mathcal{D}_{\text{train}}, \mathbf{z}) = -\log p_{\boldsymbol{\theta}}(\mathbf{y}|\mathbf{x}) + \sum_{i=1}^{n} D_{KL}(p_{\boldsymbol{\theta}}(\mathbf{y}_{i}|\mathbf{x}_{i}) \parallel p_{\hat{\boldsymbol{\theta}}}(\mathbf{y}_{i}|\mathbf{x}_{i})) + \frac{\lambda}{2} ||\boldsymbol{\theta} - \hat{\boldsymbol{\theta}}||_{2}^{2}.$$
(7)

Since the base model may not have reached convergence, the PBO penalizes movement in function space using the KL divergence between the base and hindsight model output distributions rather than using the true training labels.

2.3 Influence Functions

Influence functions (Cook, 1979; Hampel, 1974) are a classical method from robust statistics that attempt to measure the sensitivity of an estimator to individual datapoints (Fisher et al., 2023). We can formulate this alternative objective using Eq. 3 which we rewrite as a response function $r_{(\mathbf{x},\mathbf{y})}: \mathbb{R} \to \Theta$

$$r_{(\mathbf{x},\mathbf{y})}(\epsilon) = \boldsymbol{\theta}^{\star}(\mathbf{x},\mathbf{y},\epsilon)$$

where we assume that the objective (3) is strongly convex and hence the optimum is unique given some factor ϵ . Under these assumptions, note that $r_{(\mathbf{x},\mathbf{y})}(0) = \hat{\boldsymbol{\theta}}$ and the response function is differentiable at $\epsilon_0 = 0$ by the Implicit Function Theorem (Griewank & Walther, 2008; Krantz & Parks, 2002). This allows us to approximate the response function with a first order Taylor expansion about ϵ_0 :

$$r_{(\mathbf{x}, \mathbf{y})}(\epsilon) = \hat{\boldsymbol{\theta}} + \mathbf{G}^{-1} \nabla_{\boldsymbol{\theta}} \mathcal{L}(\hat{\boldsymbol{\theta}}, \mathbf{x}, \mathbf{y}) \epsilon,$$

where G is the Hessian of (2) or a positive definite approximation such as the Generalized Gauss-Newton Hessian or Fisher Information.

To approximate the effects on the loss at a specific test point $\mathbf{z}' = (\mathbf{x}', \mathbf{y}')$, we can linearize the model about $\hat{\boldsymbol{\theta}}$ and apply the chain rule:

$$\mathcal{L}(\boldsymbol{\theta}^{\star}(\mathbf{x}, \mathbf{y}, \epsilon), \mathbf{x}', \mathbf{y}') = \mathcal{L}(\hat{\boldsymbol{\theta}}, \mathbf{x}', \mathbf{y}') + \epsilon \nabla_{\boldsymbol{\theta}} \mathcal{L}(\hat{\boldsymbol{\theta}}, \mathbf{x}', \mathbf{y}')^{\mathsf{T}} \mathbf{G}^{-1} \nabla_{\boldsymbol{\theta}} \mathcal{L}(\hat{\boldsymbol{\theta}}, \mathbf{x}, \mathbf{y})$$

If the train and test point are the same, the quantity

IF
$$(\boldsymbol{\theta}, \mathbf{x}, \mathbf{y}) = \nabla_{\boldsymbol{\theta}} \mathcal{L}(\hat{\boldsymbol{\theta}}, \mathbf{x}, \mathbf{y})^{\mathsf{T}} \mathbf{G}^{-1} \nabla_{\boldsymbol{\theta}} \mathcal{L}(\hat{\boldsymbol{\theta}}, \mathbf{x}, \mathbf{y})$$
 (8)

is often referred to as the self-influence of z relative to the model parameterized by θ (Koh & Liang, 2017).

Although influence functions were meant to approximate the effects of true retraining (Basu et al., 2020), recent work has shown that they more closely approximate an alternate objective (Bae et al., 2022; Grosse et al., 2023), which coincides with the proximal bregman objective (7) described in Section 2.2.

3 IF-COMP: Measuring Complexity with Boltzmann Influence Functions

Although the pNML distribution and stochastic complexity are useful for calibrating the uncertainty of a given base model, exactly calculating them is intractable because they require finding the hindsight-optimal

model for each possible label. In addition, overparameterized neural networks can achieve arbitrarily low log loss on the additional test point regardless of its label, introducing the infinity problem. We empirically verify this behavior in Appendix A.1. Methods such as ACNML (Zhou & Levine, 2021) which take explicit steps in parameter space are also computationally expensive and can become unreliable for overconfident models.

To solve these issues, we begin by defining a temperature-scaled proximal Bregman objective that softens the local curvature, allowing the hindsight-optimal model to fit low probability labels. Linearizing the model produces a corresponding Boltzmann influence function (BIF) which can directly approximate the hindsight-optimal output distribution. We then propose Influence Function Complexity (IF-COMP), a complexity measure and associated pNML code that uses the BIF to produce calibrated output distributions as well as approximate the stochastic complexity for labelled and unlabelled data points. Finally, we describe how to efficiently compute IF-COMP and then validate it against the ground truth pNML complexity.

3.1 Boltzmann Influence Functions

Since the PBO (Eq. 7) regularizes movement in function space using the model's own output distribution, overconfident predictions can make this objective *too* restrictive for training the hindsight-optimal model with very low probability labels. Inspired by the use of temperature scaling to reduce model overconfidence and improve calibration for predictions (Guo et al., 2017), as well as the alternative Boltzmannian MDL formulation (Perotti et al., 2018), we propose to directly temperature scale our codelength as:

$$E_{\beta,\theta}(\mathbf{x}, \mathbf{y}) = -\log p_{\beta,\theta}(\mathbf{y}|\mathbf{x}) := -\log \sigma(\beta f_{\theta}(\mathbf{x})). \tag{9}$$

which gives us a corresponding minimax optimal Boltzmann pNML distribution

$$p_{\beta,\text{pNML}}(\mathbf{y}|\mathbf{x}) = \frac{p_{\beta,\theta^{\star}(\mathbf{x},\mathbf{y})}(\mathbf{y}|\mathbf{x})}{\sum_{\mathbf{y}'\in\mathcal{Y}} p_{\beta,\theta^{\star}(\mathbf{x},\mathbf{y}')}(\mathbf{y}'|\mathbf{x})}.$$
(10)

and a stochastic complexity

$$\Gamma_{\beta}(\mathbf{z}) = -\log p_{\beta, \boldsymbol{\theta}^{\star}(\mathbf{x}, \mathbf{y})}(\mathbf{y} | \mathbf{x}) + \log \sum_{\mathbf{y}' \in \mathcal{Y}} p_{\beta, \boldsymbol{\theta}^{\star}(\mathbf{x}, \mathbf{y}')}(\mathbf{y}' | \mathbf{x})$$
(11)

We can replace the loss in the original PBO with our temperature-scaled loss to define a corresponding hindsight-optimal model with a Boltzmann PBO (BPBO):

$$Q(\boldsymbol{\theta}, \mathcal{D}_{\text{train}}, \mathbf{z}) = E_{\beta, \boldsymbol{\theta}}(\mathbf{x}, \mathbf{y}) + \sum_{i=1}^{n} D_{KL}(p_{\beta, \boldsymbol{\theta}}(\mathbf{y}_{i} | \mathbf{x}_{i}) \parallel p_{\beta, \hat{\boldsymbol{\theta}}}(\mathbf{y}_{i} | \mathbf{x}_{i})) + \frac{\lambda}{2} ||\boldsymbol{\theta} - \hat{\boldsymbol{\theta}}||_{2}^{2}.$$
(12)

If we take a first order approximation of the log loss and a second order approximation of the proximal terms about $\hat{\theta}$, we can formulate the influence function as

$$IF_{\beta}(\hat{\boldsymbol{\theta}}, \mathbf{x}, \mathbf{y}) = \nabla_{\boldsymbol{\theta}} E_{\beta, \hat{\boldsymbol{\theta}}}(\mathbf{x}, \mathbf{y})^{\mathsf{T}} \mathbf{G}_{\beta}^{-1} \nabla_{\boldsymbol{\theta}} E_{\beta, \hat{\boldsymbol{\theta}}}(\mathbf{x}, \mathbf{y}), \tag{13}$$

with a corresponding Fisher information:

$$\mathbf{G}_{\beta} = \frac{1}{n} \sum_{i=1}^{n} \underset{\mathbf{y}_{i}^{\prime} \sim p_{\beta, \hat{\boldsymbol{\theta}}}(\mathbf{y}_{i} | \mathbf{x}_{i})}{\mathbb{E}} \left[\nabla_{\boldsymbol{\theta}} E_{\beta, \hat{\boldsymbol{\theta}}}(\mathbf{x}_{i}, \mathbf{y}_{i}^{\prime}) \nabla_{\boldsymbol{\theta}} E_{\beta, \hat{\boldsymbol{\theta}}}(\mathbf{x}_{i}, \mathbf{y}_{i}^{\prime})^{\intercal} \right].$$

Intuitively, temperature scaling softens the function space distance loss and allows the model to better accommodate arbitrarily labelled data points. We call the influence function calculated with this temperature-scaled loss the **Boltzmann influence function (BIF)**.

Different choices for β define different BIFs. At $\beta = 1$, we recover the standard influence function. As $\beta \to 0^+$, the KL divergence loss becomes an MSE loss over logits (Kim et al., 2021) and we recover a BIF resembling TRAK (Park et al., 2023). In Section 4.3 we will see that tuning β directly within the influence function is an important part of achieving high quality complexity estimates.

3.2 IF-COMP

We now describe **IF-COMP**, which estimates the Boltzmann pNML distribution and stochastic complexity using our Boltzmann influence function. Recall from (10) and (11) that our goal is to approximate $p_{\beta,\theta^*(\mathbf{x},\mathbf{y})}(\mathbf{y}|\mathbf{x})$ for all $\mathbf{y} \in \mathcal{Y}$. Using the BIF formulation above, we can apply a first order Taylor expansion of the output probability about $\hat{\boldsymbol{\theta}}$ to approximate

$$p_{\beta, \boldsymbol{\theta}^{\star}(\mathbf{x}, \mathbf{y})}(\mathbf{y}|\mathbf{x}) = p_{\beta, \hat{\boldsymbol{\theta}}}(\mathbf{y}|\mathbf{x}) + \frac{1}{n} p_{\beta, \hat{\boldsymbol{\theta}}}(\mathbf{y}|\mathbf{x}) \text{ IF}_{\beta}(\hat{\boldsymbol{\theta}}, \mathbf{x}, \mathbf{y}).$$

Similar to other work (Ilyas et al., 2022; Park et al., 2023), we choose to linearize our model in the log probability space, where we have used the identity

$$\nabla p_{\beta,\hat{\boldsymbol{\theta}}}(\mathbf{y}|\mathbf{x}) = p_{\beta,\hat{\boldsymbol{\theta}}}(\mathbf{y}|\mathbf{x}) \nabla \log p_{\beta,\hat{\boldsymbol{\theta}}}(\mathbf{y}|\mathbf{x})$$

to change the log probability gradient into a probability space gradient.

We can then rewrite the Boltzmann pNML parametric complexity as

$$\log \sum_{\mathbf{y}' \in \mathcal{Y}} p_{\beta, \boldsymbol{\theta}^{\star}(\mathbf{x}, \mathbf{y}')}(\mathbf{y}' | \mathbf{x}) = \log \left(1 + \frac{1}{n} \mathbb{E}_{\mathbf{y}' \sim p_{\beta, \hat{\boldsymbol{\theta}}}(\mathbf{y} | \mathbf{x})} \left[\mathrm{IF}_{\beta}(\hat{\boldsymbol{\theta}}, \mathbf{x}, \mathbf{y}') \right] \right).$$

For large enough n we can further simplify $\log(1+x) \approx x$, which gives us a final complexity

$$\Gamma_{\beta}(\mathbf{z}) = \underbrace{-\log p_{\beta,\boldsymbol{\theta}^{\star}(\mathbf{x},\mathbf{y})}(\mathbf{y}|\mathbf{x})}_{\text{error}} + \underbrace{\frac{1}{n} \mathbb{E}_{\mathbf{y}' \sim p_{\beta,\hat{\boldsymbol{\theta}}}(\mathbf{y}|\mathbf{x})}}_{\text{parametric complexity}} \left[\text{IF}_{\beta}(\hat{\boldsymbol{\theta}}, \mathbf{x}, \mathbf{y}') \right].$$
(14)

We leave the error term in its original form, since we assume access to true labels only for training data, for which our model is already hindsight-optimal. For unlabelled data that may not have a valid label, the error term is undefined and we compute only the parametric complexity term.

$$\Gamma_{\beta}(\mathbf{x}) = \mathbb{E}_{\mathbf{y}' \sim p_{\beta,\hat{\boldsymbol{\theta}}}(\mathbf{y}|\mathbf{x})} \left[\text{IF}_{\beta}(\hat{\boldsymbol{\theta}}, \mathbf{x}, \mathbf{y}') \right]$$
(15)

If we are interested in the calibrated probabilities of the Boltzmann pNML distribution, we can calculate the distribution directly as

$$p_{\beta,\text{pNML}}(\mathbf{y}|\mathbf{x}) = \frac{p_{\beta,\hat{\boldsymbol{\theta}}}(\mathbf{y}|\mathbf{x}) + \alpha/n \ p_{\beta,\hat{\boldsymbol{\theta}}}(\mathbf{y}|\mathbf{x}) \ \text{IF}_{\beta}(\hat{\boldsymbol{\theta}},\mathbf{x},\mathbf{y})}{1 + \alpha/n \ \mathbb{E}_{\mathbf{y}' \sim p_{\beta,\hat{\boldsymbol{\theta}}}(\mathbf{y}|\mathbf{x})} \left[\text{IF}_{\beta}(\hat{\boldsymbol{\theta}},\mathbf{x},\mathbf{y}') \right]}$$
(16)

where α controls the weighting of the test point relative to the training set. A value of $\alpha = 0$ corresponds to the original model output distribution. Compared to ACNML (Zhou & Levine, 2021), which explicitly optimizes the hindsight-optimal parameters using a weighted approximate posterior, IF-COMP linearizes the model directly which allows us to easily control this weighting *after* computing the BIF for each label.

3.3 Efficiently Computing IF-COMP

Although we have simplified the calculation of stochastic data complexity considerably, calculating the BIF still requires estimating and inverting the Fisher information matrix. Similar to previous work (Grosse et al., 2023), we use an eigenvalue-corrected Kronecker-factored approximation to the Fisher information matrix (EKFAC) (George et al., 2018) which produces an eigendecomposition $\mathbf{G}_{\beta} \approx Q \lambda Q^{T}$. We can then calculate the influence function as the squared L2 norm in the inverse eigenspace:

$$IF_{\beta}(\hat{\boldsymbol{\theta}}, \mathbf{x}, \mathbf{y}) = ||\nabla_{\boldsymbol{\theta}} E_{\beta, \boldsymbol{\theta}}(\mathbf{x}, \mathbf{y})^{\mathsf{T}} Q(\lambda + \delta)^{-1/2}||_{2}^{2}, \tag{17}$$

Table 1: Average pNML output distribution (16) inference time per input (in seconds). On smaller models such as LeNet, IF-COMP can be computed almost as efficiently as a gradient norm while ACNML is nearly 7 times slower. On larger ResNet18 models, IF-COMP maintains a greater than 10 times speedup compared to ACNML.

Method	MNIST LeNet	CIFAR-10 ResNet18	CIFAR-100 ResNet18
GradNorm	0.000226s	0.00336s	0.03783s
ACNML-EKFAC	0.001770s	0.23183s	2.33628s
IF-COMP (ours)	0.000241s	0.01576s	0.18883s
IF-COMP vs. ACNML Speedup	7.34 ×	14.71 ×	12.4 ×

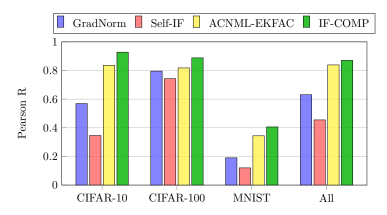


Figure 1: Pearson R correlation of different methods of approximating hindsight-optimal outputs with ground truth parametric complexity on in-domain (CIFAR-10) and out-of-domain datasets. IF-COMP achieves the highest correlation across all datasets, beating ACNML, a computationally more expensive alternative.

where δ is a damping term added to ensure invertibility. Since we need only calculate and invert the EKFAC once for a given model and training set, calculating the IF-COMP then requires only one jacobian vector product (JVP) per label per sample.

We verify the computational efficiency of this approach by comparing the time to calculate the final pNML output distribution (16) for a single example with IF-COMP, ACNML, and gradient norm methods. Specific experimental details are provided in Appendix B. We present our results in Table 1. For small LeNet models, we find that the additional overhead of the EKFAC multiplication performed in IF-COMP is relatively small (6%) compared to a simple gradient norm, while ACNML is nearly 7 times slower. For larger ResNet18 models, IF-COMP becomes slower than the gradient norm, but still provides a 12-15 times speedup over ACNML.

3.4 pNML Validation

To verify that IF-COMP can accurately approximate the ground truth pNML parametric complexity on both in-distribution (ID) and out-of-distribution (OOD) samples, we fine-tune a CIFAR-10 (Krizhevsky, 2009) pre-trained ResNet-18 (He et al., 2016) model with the BPBO (12) on 20 random test images each from CIFAR-10, CIFAR-100, and MNIST (Deng, 2012). We label each example with every possible output class for a total of 600 individual training example pairs. After training, we can use the hindsight-optimal output probabilities to calculate the complexity as in Eq. 15.

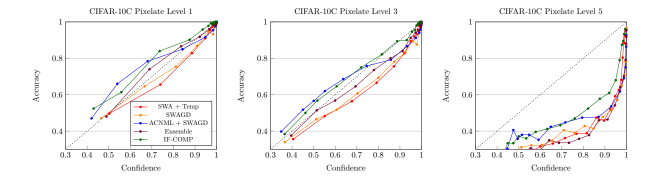


Figure 2: Reliability diagrams for Pixelate corruptions on CIFAR-10C. IF-COMP outperform ACNML as well as Bayesian methods and ensembles even as corruptions increase in severity. Although IF-COMP and ACNML perform similarly on lower confidence examples, IF-COMP maintains this reliability on higher confidence examples. Dotted lines represent perfect calibration.

We compare IF-COMP with other baselines for approximating hindsight-optimal training. Specifically, we consider averaging gradient norms across classes, self influence calculated with EKFAC, and ACNML with an EKFAC posterior. We present our results in Figure 1. We find that IF-COMP consistently achieves the highest Pearson correlation with true complexity on all datasets, beating ACNML which explicitly takes steps in parameter space. For ID CIFAR-10 examples, IF-COMP achieves a strong Pearson R of 0.928, which it maintains on CIFAR-100 examples. All methods degrade considerably on MNIST, which indicates that complexity becomes more difficult to estimate for further OOD examples.

4 Experiments

To experimentally validate IF-COMP, we consider three tasks that test different capabilities. The first, uncertainty quantification, requires producing calibrated outputs across distribution shifts. The second, mislabel detection, requires measuring complexity on labelled training examples. We use this task as a case study to understand how the error and parametric complexity terms trade off during training, as well as the effects of temperature scaling. Finally, OOD detection requires measuring complexity on unlabelled out-of-distribution test examples. Across all three tasks, IF-COMP consistently exhibits strong performance compared to baseline methods. We provide additional experiments on data pruning and hindsight-optimal retraining in Appendix A as well as full experimental details in Appendix B.

4.1 Uncertainty Calibration

We begin by evaluating the uncertainty calibration of IF-COMP output distributions under distribution shifts. Following the experimental setup in Ovadia et al. (2019), we measure the expected calibration error (ECE) (Naeini et al., 2015) of ResNet18 models trained on CIFAR-10 and tested on the CIFAR10-C datasets (Hendrycks & Dietterich, 2019). CIFAR-10C applies 19 corruptions with 5 severity levels across the test images of CIFAR-10, allowing us to compare calibration on a wide range of distribution shifts. We calculate ECE by dividing model predictions sorted by confidence into 20 equal sized bins (Zhou & Levine, 2021).

We compare against a wide range of Bayesian and NML baselines, including ensembling (Lakshminarayanan et al., 2017), Stochastic Weight Averaging (SWA) (Izmailov et al., 2019), SWA Gaussian Diagonal (SWAG-D) (Maddox et al., 2019), Monte-Carlo dropout (Gal & Ghahramani, 2016), and ACNML (Zhou

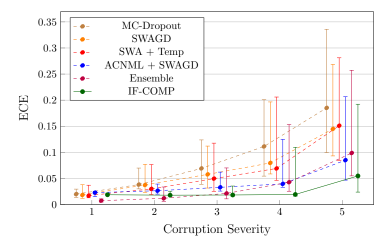


Figure 3: Expected calibration error (ECE) for various methods across increasing levels of CIFAR-10C corruptions. We plot medians and inter-quartile ranges. IF-COMP achieves lower ECE across almost all corruption levels compared to both Bayesian methods and other NML-based methods.

& Levine, 2021) with a SWAG-D posterior. For a fair comparison, all methods except ensembling and MC-Dropout use the same SWA base model. Additionally, we apply the same IF-COMP temperature scaling to the SWA output logits, which is equivalent to using a value of $\alpha = 0$. We produce 30 model samples for all Bayesian methods.

We present median ECE and inter-quartile ranges across corruptions for each severity level in Figure 3. On more severe corruptions (3-5), IF-COMP produces better calibrated uncertainties than all baselines, beating the most relevant ACNML baseline as well as a Bayesian ensemble of 30 trained models. At corruption severity 2 IF-COMP outperforms all other baselines except the ensemble. At the lowest corruption level all methods perform similarly.

The reliability diagrams in Figure 2 show a more detailed look at the calibration for varying Pixelate corruption severity levels. For low corruption levels all methods perform similarly. As severity increases, IF-COMP maintains strong calibration compared to Bayesian baselines and improves over ACNML for high confidence outputs. Although all methods degrade considerably at the highest corruption severity, IF-COMP still improves over the strong ACNML baseline.

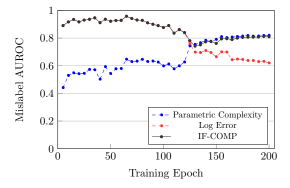
4.2 Mislabel Detection

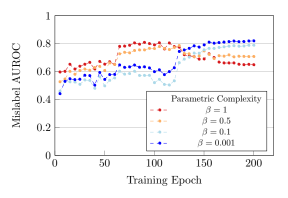
Next, we investigate IF-COMP's ability to measure the complexity of labelled training examples. We follow the mislabel detection setup in Zhu et al. (2021); Srikanth et al. (2023) and use CIFAR-10 and CIFAR-100 datasets corrupted with various types of label noise and noise rates. We train a ResNet-18 model on this corrupted dataset then attempt to identify which examples were mislabelled using only the trained model and noised dataset. We consider 4 types of label noise. Human noise replaces all labels with labels from a single human annotator (Zhu et al., 2021; Wei et al., 2022). Data-dependent noise is generated by jointly modelling the clean label and a class-dependent projection of the feature vector (Zhu et al., 2021). Symmetric noise changes the label to another uniformly at random. Asymmetric noise changes each label to another fixed similar label.

For each example, we calculate the full stochastic complexity (14) by combining the model's log error on the true label as well as the parametric complexity across all labels. For our baselines, we consider

Table 2: Mislabel detection AUROC for CIFAR-10 and CIFAR-100 with various types and rates of label noise (best method bolded). IF-COMP achieves strong detection across all CIFAR-10 noise types without requiring extra checkpoints, even on the difficult data-dependent noise that other methods fail to perform better than random on. On CIFAR-100, IF-COMP achieves strong performance on symmetric and asymmetric noise, although it fails to detect mislabelled data with human and data-dependent noise, similar to other methods.

Extra		CIFAR10			CIFAR100				
Method	Checkpoints?	Human	Data	Asym 0.3	Sym 0.6	Human	Data	Asym 0.3	Sym 0.6
Trac-IN	√	90.50	49.13	70.98	64.40	50.01	50.25	58.25	57.95
EL2N	\checkmark	73.38	52.18	51.53	51.61	49.79	50.51	56.30	55.45
GraNd	\checkmark	70.53	51.30	50.48	50.52	50.52	50.07	52.26	50.16
Self-IF	×	95.38	55.77	87.30	43.16	49.81	50.29	72.15	58.40
IF-COMP	Х	96.86	88.07	95.69	97.83	49.52	50.30	79.39	95.21





- (a) AUROC of the individual components of IF-COMP.
- (b) Parametric complexity AUROC at varying temperatures

Figure 4: IF-COMP accurately trades off between log error and parametric complexity, maintaining strong AUROC throughout training. Tuning the temperature is critical to achieving accurate complexity estimates near convergence.

the self-influence score (Self-IF) (Koh & Liang, 2017), Trac-IN (Pruthi et al., 2020), EL2N (Paul et al., 2021), and GraNd (Paul et al., 2021) and measure performance using AUROC. Trac-IN requires evaluating intermediate checkpoints, and EL2N and GraNd require multiple short training runs, meaning they utilize more information than is available to IF-COMP.

We present our results in Table 2. For CIFAR-10 datasets, IF-COMP consistently achieves the highest AUROC across all types of noise, beating even the baselines that utilize additional checkpoint information. On the difficult data-dependent noise where all other methods achieve close to random AUROC, IF-COMP maintains strong performance. For CIFAR-100 datasets where all baselines consistently achieve close to random AUROC across all noise types, IF-COMP is still able to detect both asymmetric and symmetric noise. On human and data-dependent noise no method performs better than random.

4.3 Analyzing the Components of IF-COMP

We take a brief detour and use mislabel detection as a case study to understand how each component of IF-COMP contributes to the final complexity estimate. We consider CIFAR-10 human mislabel noise and

Table 3: AUROC for OOD detection methods on near and far distribution shifts. Best methods are bolded, and second best methods are starred. We show only the most common baseline methods, although we compare against all methods in the OpenOOD benchmark. IF-COMP achieves a new state of the art on MNIST and CIFAR-10 benchmarks, beating out all 20 other baselines.

	MNIST		CIFAR-10		CIFAR-100	
Method	Near	Far	Near	Far	Near	Far
MSP	91.45	98.51	88.03	90.73	80.27	77.76
ODIN	92.38	99.02	82.87	87.96	79.90	79.28
Energy	90.77	98.77	87.58	91.21	80.91*	79.77
MLS	92.49	99.08	87.52	91.10	81.05	79.67
KNN	96.52	96.66	90.64*	92.96	80.18	82.40*
$\operatorname{GradNorm}$	76.55	96.39	54.90	57.55	70.13	69.14
RMDS	98.00*	98.12	89.80	92.20	80.15	82.92
VIM	94.63	98.99	88.68	93.48*	74.98	81.70
Gram	73.90	99.75*	58.66	71.73	51.66	73.36
IF-COMP	99.40	99.97	92.23	95.63	79.40	79.41

train a model to convergence, taking checkpoints every 5 epochs. For each model, we calculate IF-COMP, log error, and temperature-scaled parametric complexity. We plot the mislabel detection AUROC across training for these values in Figure 4.

Figure 4a shows that mislabelled examples are ignored early in training, increasing their error and making them easy to distinguish. Later in training the model learns to memorize these examples and decrease their error, increasing parametric complexity accordingly. Choosing only one of these values to detect mislabelled examples with would fail in different scenarios, while IF-COMP accurately captures the tradeoff between the two throughout all stages of training. Figure 4b shows that temperature-scaling is a crucial component of IF-COMP. With a standard temperature $\beta = 1$, parametric complexity estimates are inaccurate and mirror log error, providing no additional signal for mislabel detection. As we increase the temperature we soften the second order restrictions, making complexity estimates more accurate and providing a complementary signal to the error.

4.4 OOD Detection

Finally, we investigate IF-COMP's ability to measure complexity on unlabelled data by detecting OOD examples. We use the OpenOOD benchmark (Yang et al., 2022; Zhang et al., 2023) and consider the ID MNIST, CIFAR-10, and CIFAR-100 datasets using the provided pretrained LeNet (Lecun et al., 1998) and ResNet-18 models. For each dataset, the benchmark provides a set of Near-OOD datasets which exhibit similar visual features to the ID training set, and a set of Far-OOD datasets that do not. Given a test example from one of these datasets, an OOD detection method aims to assign a high score to OOD examples and a low score to ID examples. We use the IF-COMP parametric complexity (15) as our score function, and measure performance using AUROC. We compare our method against the full suite of 20 baseline approaches implemented in OpenOOD, although we show only 9 of the most common or most similar methods in our results. We refer readers to Zhang et al. (2023) for more details on baseline methods.

Our results are shown in Table 3. On MNIST, IF-COMP achieves a new state of the art AUROC, beating all 20 baselines with near perfect detection on both Near and Far OOD datasets. IF-COMP similarly achieves a new state of the art AUROC for CIFAR-10, beating the next best method by over 2 AUROC on Far OOD datasets. On CIFAR-100 IF-COMP ranks in the middle of the baselines. For this dataset all baselines perform quite similarly, with only small AUROC gaps between the best and second best method.

Strong baselines exhibit variance in performance across datasets, with the best method, RMDS (Ren et al., 2021), beating all other baselines only on 2 datasets. In contrast, IF-COMP achieves the best AUROC across 4 of the 6 datasets.

5 Related Work

The Minimum Description Length Principle MDL is one of many ways to measure complexity, including Kolmogorov complexity(Kolmogorov, 1965; Solomonoff, 1964), VC dimension (Vapnik & Chervonenkis, 1971) and Rademacher complexity (Bartlett & Mendelson, 2003). The Refined MDL coding scheme (Barron et al., 1998) which defines stochastic complexity (Rissanen, 1986) is based on the normalized maximum likliehood (NML) distribution (Shtar'kov, 1987), which has been modified for predictive settings (pNML) (Roos et al., 2008; Fogel & Feder, 2019). Refined MDL bears similarity to model selection criteria such as Akaike Information Criterion (AIC) (Akaike, 1973) and the Bayesian Information Criterion (BIC) (Schwarz, 1978). We point readers to Grunwald (2004) for further discussion of MDL.

MDL principles have been used to motivate methods to train autoencoders (Hinton & Zemel, 1993) and regularize weights (Hinton & van Camp, 1993). Applying MDL directly to overparameterized model classes is difficult, requiring subclass restrictions such as ridge estimators (Dwivedi et al., 2023). Recent work in MDL methods for neural networks have mainly attempted to approximate pNML distributions by performing additional gradient steps (Bibas et al., 2020) or optimizing with an approximate posterior (Zhou & Levine, 2021). Instead of explicitly calculating model weights, IF-COMP directly linearizes the model making optimization unnecessary and allowing post-hoc tuning of example weighting.

Quantifying Predictive Uncertainty A wide variety of methods have been proposed to quantify predictive uncertainty in deep neural networks. Bayesian approaches include Laplace approximations (Mackay, 1992; Ritter et al., 2018), variational inference (Graves, 2011), stochastic gradient MCMC (Welling & Teh, 2011), dropout (Gal & Ghahramani, 2016), and weight averaging (Izmailov et al., 2019; Maddox et al., 2019). Non-Bayesian methods include model ensembling (Lakshminarayanan et al., 2017; Osband et al., 2016), Platt scaling (Platt, 2000) with logit temperatures, (Guo et al., 2017) and more recently, MDL based approaches.

Bayesian and MDL methods have a deep connection. For parametric hypothesis classes, MDL reduces to standard maximum likelihood estimation (Grunwald, 2004). Refined MDL model selection coincides with Bayes factor model selection (Kass & Raftery, 1995) based on a Jeffreys prior (Bernardo & Smith, 1994), where codes correspond to MAP estimates. NML distributions over the full dataset can also be seen as a form of Bayesian marginal likelihood (Grunwald, 2004), although pNML does not coincide with any standard Bayesian interpretations.

Influence Functions Influence functions (Cook, 1979; Hampel, 1974) are a classical method from robust statistics that has found renewed interest in deep neural networks (Koh & Liang, 2017; Grosse et al., 2023). They have been applied for a wide range of tasks including detecting bias (Brunet et al., 2019), data poisoning (Koh et al., 2021), auditing predictions (Schulam & Saria, 2019), and fixing model mistakes (Tanno et al., 2022). The self-influence of a data point is related to its memorization (Feldman, 2021), which has been shown is necessary to cover the long tails of the data distribution and improve generalization (Feldman & Zhang, 2020). Although influence functions have been found to poorly match true retraining for neural networks (Basu et al., 2020), they correlate well with an alternate proximal objective (7) (Bae et al., 2022).

6 Conclusion

In this paper we have proposed IF-COMP, an efficient method for estimating stochastic data complexity in deep neural networks with temperature-scaled Boltzmann influence functions (BIFs). The BIF softens the second order curvature and linearizes the model, allowing us to efficiently approximate hindsight-optimal outputs even for low probability labels. Using these output estimates, IF-COMP can produce well calibrated output distributions as well as measure complexity on both labelled and unlabelled points. On tasks covering uncertainty calibration, mislabel detection, and OOD detection, IF-COMP consistently matches or outperforms strong baselines, including Bayesian and optimization tracing based approaches. Our results demonstrates the potential of MDL based approaches for improving uncertainty estimates in deep neural networks.

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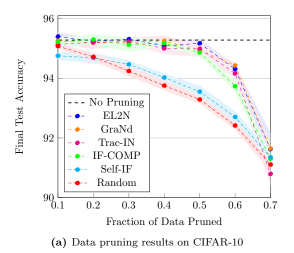
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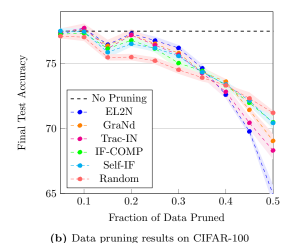


Figure 5: Data pruning results. Shaded regions correspond to standard deviations over 5 seeds.IF-COMP performs similarly to other methods that require access to additional checkpoints, including Trac-IN, GraNd, and EL2N. At the highest pruning levels for CIFAR-100, IF-COMP and Self-IF outperform baselines that perform worse than random.

A Additional Experiments

A.1 Hindsight-Optimal Retraining

We consider ground truth retraining with the original hindsight objective (3) on CIFAR-10 with a ResNet-18 model. Similar to our experiments in Section 3.4, we select 20 examples randomly from CIFAR-10, CIFAR-100, and MNIST test sets, then apply each of the possible 10 CIFAR labels for a total of 600 additional training examples.

For each example, we train the model from scratch on the full training set with the added example, and evaluate the probability of the true label class at convergence. We find that the model is able to effectively memorize all additional examples provided to it, achieving close to 1 probability consistently across all classes. Using this hindsight-optimal probability in the pNML formulation produces a constant uniform distribution that is useless for any tasks.

A.2 Dataset Pruning

We consider the task of dataset pruning, with the goal of removing training examples that do not provide essential training information to the model. Specifically, given a model trained on a dataset, we aim to rank training examples such that removing the least important examples degrades final test accuracy as little as possible. We consider ResNet-18 models trained on both CIFAR-10 and CIFAR-100 datasets, with varying levels of data pruning. We use IF-COMP to rank points, removing ones with the lowest complexity first and keeping those with higher complexity.

For baselines, we consider Trac-IN, GraNd, EL2N, and Self-IF, which follow our experimental setup for mislabel detection. We also consider a no pruning and random pruning baseline. Once points are ranked, we train 5 models with different seeds on each pruned dataset. Results are shown in Figure 5. We find that IF-COMP performs similarly to baseline methods across both datasets, even those that utilize additional training checkpoints. On CIFAR-100 at the highest pruning levels, IF-COMP and Self-IF outperform

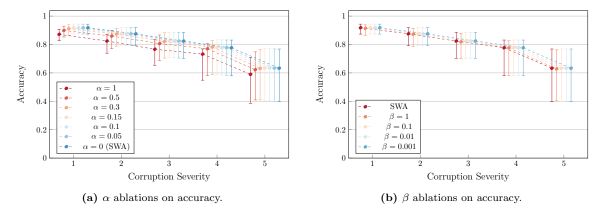


Figure 6: Ablations on the effects of the α and β hyperparameters on model accuracy. IF-COMP exhibits minimal differences from SWA baseline accuracy at all severity levels, except for particularly large values of α .

baselines, although all methods perform worse than random.

A.3 α and β Ablations

In this section we provide additional ablations on the α weighting parameter and β inverse temperature parameter for the uncertainty calibration task. We measure their effect on accuracy in Figure 6 and on ECE in Figure 7. We compare all methods against the SWA baseline. When not specified, we set default values of $\beta = 0.66$ and $\alpha = 0.15$.

We find that accuracy is minimally affected by α , and only begins to slightly degrade as α becomes particularly large (> 0.5). Since a maximum value of only $\alpha = 0.3$ is required to achieve optimal ECE on severity level 5, even prioritizing calibration on the highest severity levels does not degrade accuracy. Similarly, β values have little effect on accuracy, even for extremely small values. This indicates that the BIF for each label provides as much information as the model output.

We find that the ECE for each corruption severity level changes with α , where smaller values of α are better for cleaner images, and larger values of α are better for noisier images. Tuning α allows practitioners to select what kinds of test images to prioritize strong calibration on. Compared to the raw outputs produced by temperature scaling SWA model outputs, IF-COMP consistently improves ECE across all severity levels for varying values of β . We do not tune a separate α for each β , and use the same value of 0.15.

B Experimental Details

B.1 Computing Environment

All experiments were implemented in PyTorch and were run on single RTX6000 or A40 GPUs.

B.2 EKFAC

We implement a version of the Eigenvalue Kronecker-Factored Approximate Curvature (EKFAC) (George et al., 2018) to efficiently approximate the inverse Fisher information matrix. For efficiency, in convolutional

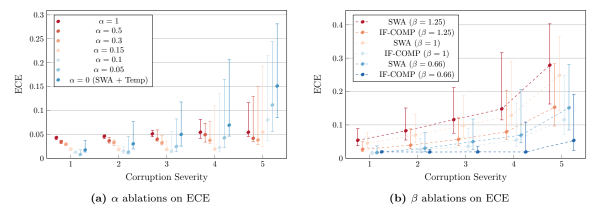


Figure 7: Ablations on the effects of the α and β hyperparameters on ECE. Different values of α are optimal for different severity levels. Selecting an α value allows practitioners to prioritize calibration on different images. Applying IF-COMP on top of SWA models improves ECE across all severity levels even for varying values of β (α is not tuned and held constant at 0.15).

layers we assume spatially uncorrelated activations (SUA), except for MNIST LeNet models, for which removing the SUA assumption does not introduce significant computational overhead. BatchNorm layers are implemented with a diagonal approximation which do not require eigendecomposition. We take a single pass through the full training set and sample a single label to calculate the activation and output gradient covariances, then perform the relevant eigendecompositions. We then take another full pass through the training set to estimate the second moments in the eigenbasis with a single label sample. Both passes perform no data augmentation and use the original training images. Calculating the BIF is performed by taking a temperature-scaled loss gradient, projecting into the eigenbasis, then taking the norm scaled by the inverse of the second moments. We apply a consistent damping value of $\delta = 1e-30$ for LeNet models and $\delta = 1e-12$ for ResNet-18 models.

To further improve efficiency, we use the PyTorch vmap operation to vectorize computations both across label spaces and across examples. We use an example chunk size of 8 for MNIST and CIFAR-10 experiments, and an example chunk size of 4 for CIFAR-100 experiments, with full chunking across the label space.

B.3 Uncertainty Calibration

All models are ResNet-18 models. CIFAR-10 ensemble models were trained with following standard training procedures using SGD with momentum of 0.9, weight decay of 0.0005, and a learning rate of 0.1 that decays by a factor of 5 at epochs 60, 120, and 160. To train SWA models we follow the setup from (Izmailov et al., 2019) and follow this setup for 160 epochs, then set a constant 0.01 learning rate and average checkpoints for another 140 epochs for a total of 300 epochs. For SWA and SWAG-D models, batch statistics were then updated on the full training set. For dropout models, we place dropout layers after the ReLU nonlinearities in each block before the residual connection, and test rates of 0.1, 0.2, 0.3, and 0.4. Empirically we find a rate of 0.3 to perform the best without significant accuracy degradation. We follow the setup described in Zhou & Levine (2021) for training ACNML models, including the recommended step size and weighting term. All Bayesian methods use 30 samples from the posterior which we average. IF-COMP uses a temperature of 1.5 and an α weight of 0.15, which we find by tuning on a held out validation set of additional perturbations of varying severity. Reliability diagrams and ECE values are calculated similar to Zhou & Levine (2021) by binning exampels sorted by confidence into 20 bins, then calculating the average L1 norm between the average confidence and accuracy within each bin.

B.4 Mislabel Detection

For both CIFAR-10 and CIFAR-100 datasets we use a ResNet-18 model trained with the standard training procedure detailed in the section above, with early stopping calculated on a clean validation set. We retrieve human and data-dependent noise directly from the relevant repositories, and generate asymmetric and symmetric noise locally. Asymmetric noise is applied to CIFAR-10 with a predefined label mapping, and for CIFAR-100 we perform a mapping to the next fine-grained class in the corresponding coarse model class.

For Trac-IN we take model checkpoints every 20 epochs for a total of 10 checkpoints, then compute gradient norms on the full model. EL2N and GraNd are computed by training a model for epoch 20 with 10 different seeds, then calculating average error L2 norm and gradient norms across all models. Self-IF is calculated using the same EKFAC estimated for IF-COMP with full gradients. IF-COMP is calculated at $\beta = 0.001$ for all CIFAR-10 experiments and $\beta = 1$ for all CIFAR-100 experiments. Note that we fit the EKFAC on the dataset with noised labels.

B.5 OOD Detection

All baseline results are provided in the OpenOOD benchmark (Yang et al., 2022; Zhang et al., 2023), as well as the pre-trained LeNet and ResNet-18 models used for MNIST and CIFAR-10/CIFAR-100 experiments. We use a temperature of $\beta=2$ for MNIST experiments, $\beta=0.001$ for CIFAR-10 experiments, and $\beta=1$ for CIFAR-100 experiments. Temperature values are found by tuning on a held out validation set of OOD examples that share no classes with the ones used for testing. This validation set is the same one used to tune hyperparameters for all benchmarks in the OpenOOD and are provided as a standard procedure from the framework (Zhang et al., 2023). We hypothesize that the extremely high temperatures necessary for strong CIFAR-10 results are a result of the stable and flat optima found with Batchnorm which prevent learning arbitrary labels without moving far away in weight and function space.

B.6 Timing Comparisons

For all experiments we use a single A40 GPU and attempt to use the same chunk size for all vmap operations. However, ACNML on CIFAR-100 runs out of memory, so we reduce the chunk size from 4 to 1, meaning we perform the vmap only over the label space. We follow the set of best practices provided in Zhou & Levine (2021) and perform 5 optimization steps. As in other experiments, we do not use the spatially uncorrelated activations (SUA) assumption for MNIST models, making the EKFAC computations slightly more expensive.