Active Bayesian Assessment of Black-Box Classifiers

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Abstract
Recent advances in machine learning have led to increased deployment of black-box classifiers across a wide variety of applications. In many such situations there is a crucial need to both reliably assess the performance of these pre-trained models and to perform this assessment in a label-efficient manner (given that labels may be scarce and costly to collect). In this paper, we introduce an active Bayesian approach for assessment of classifier performance to satisfy the desiderata of both reliability and label-efficiency. We begin by developing inference strategies to quantify uncertainty for common assessment metrics such as accuracy, misclassification cost, and calibration error. We then propose a general framework for active Bayesian assessment using inferred uncertainty to guide efficient selection of instances for labeling, enabling better performance assessment with fewer labels. We demonstrate significant gains from our proposed active Bayesian approach via a series of systematic empirical experiments assessing the performance of modern neural classifiers (e.g., ResNet and BERT) on several standard image and text classification datasets.

Introduction
Complex machine learning models, particularly deep learning models, are now being applied to a variety of practical prediction problems ranging from diagnosis of medical images (Kermany et al. 2018) to autonomous driving (Du et al. 2017). Many of these models are black boxes from the perspective of downstream users, such as models developed remotely by commercial entities and hosted as a service in the cloud (Yao et al. 2017; Sanyal et al. 2018). For a variety of reasons (legal, economic, competitive), users will often have no direct access to the detailed workings of the model, how the model was trained, or the training data. In this context, careful attention needs to be paid to have accurate, detailed and robust assessments of the quality of a model’s predictions, such that the model can be held accountable by users. In real-world deployment scenarios, acquiring labeled data for assessment is likely to be scarce and costly to collect, e.g., for a model being deployed in a diagnostic imaging context in a particular hospital. With this in mind we develop a framework for active Bayesian assessment of black-box classifiers, using techniques from active learning to efficiently select instances to label so that uncertainty of assessment can be reduced, and deficiencies of models such as low accuracy, high calibration error or high cost mistakes can be quickly identified. Our primary contributions are:
- We propose a general Bayesian framework to assess black-box classifiers with uncertainty for quantities such as class-wise accuracy, expected calibration error (ECE), confusion matrices, and performance comparison across groups.
- We propose a general framework for active Bayesian assessment for an array of fundamental tasks including (1) estimation of model performance; (2) identification of model deficiencies; (3) performance comparison between groups.
- We demonstrate that our proposed approaches need significantly fewer labels than baselines, via a series of experiments assessing the performance of modern neural classifiers (e.g., ResNet and BERT) on several standard image and text classification datasets.

Notation
We consider classification problems with a feature vector x and a class label y ∈ {1, ..., K}, e.g., classifying image pixels x into one of K classes. We are interested in assessing the performance of a pretrained prediction model M that makes predictions of y given a feature vector x, where M produces K numerical scores per class in the form of a set of estimates of class-conditional probabilities $p_M(y = k|x), k = 1, \ldots, K$. $\hat{y} = \arg \max_k p_M(y = k|x)$ is the classifier’s label prediction for a particular input x. $s(x) = p_M(y = \hat{y}|x)$ is the score of a model, as a function of x, i.e., the class probability that the model produces for its predicted class $\hat{y} \in \{1, \ldots, K\}$ given input x. This is also referred to as a model’s confidence in its prediction and can be viewed as a model’s own estimate of its accuracy. The model’s scores in general need not be perfectly calibrated, i.e., they need not match the true probabilities p(y = $\hat{y}$|x).

We focus in this paper on assessing the performance of a model that is a black box, where we can observe the inputs x and the outputs $p_M(y = k|x)$, but don’t have any other information about its inner workings. Rather than learning a model itself we want to learn about the characteristics of a fixed model that is making predictions in a particular environment characterized by some underlying unknown distribution p(x, y).
Performance Assessment

Performance Metrics and Tasks: We will use θ to indicate a performance metric of interest, such as classification accuracy, true positive rate, expected cost, calibration error, etc. Our approach to assessment of a metric θ relies on the notion of disjoint groups (or partitions) g = 1, . . . , G of the input space x ∈ R_g, e.g., grouping by predicted class ỹ. For any particular instantiation of groups g and metric θ, there are three particular assessment tasks we will focus on in this paper: (1) estimation, (2) identification, and (3) comparison.

Estimation: Let θ_1, . . . , θ_G be the set of true (unknown) values for some metric θ and some grouping g. The goal of estimation is to assess the quality of a set of estimates Ÿ_1, . . . , Ÿ_G relative to the true values. In this paper we will focus on RMSE loss \((\sum_g p_\theta(\theta_g - \hat{\theta}_g)^2)^{1/2}\) to measure estimation quality, where \(p_\theta = p(x \in R_g)\) (e.g., as estimated from unlabeled data).

Identification: Here the goal is to identify extreme groups, e.g., g* = arg min\_g θ\_g, such as the predicted class with the lowest accuracy (or the highest cost, swapping max for min). We will investigate methods for finding the \(m\) groups with highest or lowest values of a metric θ. To compare the set of identified groups to the true set of \(m\)-best/worst groups, we can use (for example) ranking measures to evaluate and compare the quality of different identification methods.

Comparison: The goal here is to determine if the difference between two groups \(g_1\) and \(g_2\) is statistically significant, e.g., to assess if accuracy or calibration for one group is significantly better than another group for some black-box classifier. A measure of the quality of a particular assessment method in this context is to compare how often, across multiple datasets of fixed size, a method correctly identifies if a significant difference exists and, if so, its direction.

There are multiple definitions of groups that are of interest in practice. One grouping of particular interest is where groups correspond to a model’s predicted classes, i.e., \(g = k\), and the partition of the input space corresponds to the model’s decision regions \(x \in R_k\), i.e., \(g(x) = k\). If θ refers to classification accuracy, then Ÿ_k is the accuracy per predicted class. For prediction problems with costs, Ÿ_k can be the expected cost per predicted class, and so on.

Another grouping of interest for classification models are groups \(g\) that correspond to bins \(b\) of a model’s score \(^1\), i.e., \(s(x) \in b\), \(b = 1, . . . , B\), or equivalently \(x \in R_b\) where \(R_b\) is the region of the input space where model scores lie in score-bin \(b\). \(\hat{\theta}_b\) can be defined as the accuracy per score-bin, which in turn can be related to the well-known expected calibration error (ECE, e.g., Guo et al. (2017)) as we will discuss in more detail later in the paper.

In an algorithmic fairness context, for group fairness (Hardt, Price, and Srebro 2016) the \(g\) groups can correspond to categorical values of a protected attribute such as gender or race, and θ can be defined (for example) as accuracy or true positive rate per group.

In the remainder of the paper we focus on developing and evaluating the effectiveness of different methods for assessing groupwise metrics \(\theta_g\). In the two sections below we first describe a flexible Bayesian strategy for assessing performance metrics θ in the context of the discussion above, and then outline a general active assessment framework that uses the Bayesian strategy to address the three assessment tasks in a label-efficient manner.

Bayesian Assessment

We outline below a Bayesian approach to make posterior inferences about performance metrics given labeled data, where the posteriors on \(\theta\) can be used to support the three assessment tasks (estimation, identification, and comparison). For simplicity we begin with the case where \(\theta\) is assumed to be accuracy and then extend to other metrics such as ECE. The accuracy for a group \(g\) can be treated as an unknown Bernoulli parameter \(\theta_g\). Labeled observations \((x_i, y_i)\), \(i = 1, . . . , N_g\) are sampled randomly per group conditioned on \(x_i \in R_g\), leading to a binomial likelihood with binary accuracy outcomes \(1(y_i, \hat{y}_i) \in \{0, 1\}\). The standard frequency-based estimate is \(\hat{\theta}_g = \frac{1}{N_g} \sum_{i=1}^{N_g} 1(y_i, \hat{y}_i)\).

It is natural to consider Bayesian inference in this context, especially in situations where there is relatively little labeled data available per group. With a conjugate prior \(\theta_g \sim \text{Beta}(\alpha_g, \beta_g)\) and a binomial likelihood on binary outcomes \(1(y_i, \hat{y}_i)\), we can update the posterior distribution of \(\theta_g\) in closed-form to \(\text{Beta}(\alpha_g + r_g, \beta_g + N_g - r_g)\) where \(r_g\) is the number of correct label predictions \(\hat{y} = y\) by the model given \(N_g\) trials for group \(g\).

For other metrics, we sketch the basic idea here for Bayesian inference for ECE and provide additional discussion in the Supplement. ECE is defined as \(\frac{\sum_k p_b |s_b - \theta_b|}{B}\) where \(B\) is the number of bins (corresponding to groups \(g\)), \(p_b\) is the probability of each bin \(b\), and \(\theta_b\) and \(s_b\) are the accuracy and average confidence per bin respectively. We can put Beta priors on accuracies \(\theta_b\), model the likelihood of outcomes for each bin \(b\) as binomial resulting again in closed form Beta posteriors for accuracy per bin \(b\). The posterior density for the marginal ECE itself is not available in closed form, but can easily be estimated by direct Monte Carlo simulation from the \(B\) posteriors for the \(B\) bins. Being Bayesian about ECE per group \(ECE_g\) (e.g., per class, with \(g = k\)) follows in a similar manner by defining two levels of grouping, one at the class level and one at the bin level (see Supplement for details).

Illustrative Example: To illustrate these ideas, we trained a standard ResNet-110 classifier on the CIFAR-100 training data set and performed Bayesian inference about accuracy and ECE performance on the 10,000 labeled examples in the test set. The groups \(g = k\) correspond to \(K\) predicted classes by the model, \(y = k \in \{1, . . . , K\}\). We used Beta priors with \(\alpha_k = \beta_k = 1, k = 1, . . . , K\) for classwise accuracy, and \(\alpha_0 = 2s_0, \beta_0 = 2(1 - s_0), b = 1, . . . , K\) for binwise accuracy. Figure 1 shows the resulting mean posterior estimates (MPEs) and 95% credible intervals (CIs) for accuracy

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1The score-bins can be defined in any standard way, e.g., equal width \(1/B\) or equal weight \(p(s(x) \in b) = 1/B\)

2We use ECE for illustration in our results since it is widely used in the recent classifier calibration literature, but other calibration metrics could also be used, e.g., see Kumar, Liang, and Ma (2019).
and ECE values for each of the \( K = 100 \) classes. The accuracies and ECE values of the model vary substantially across classes, and classes with low accuracy tend to be less calibrated. There is also considerable posterior uncertainty for these metrics even using the whole test set of CIFAR-100. For example, while there is confidence that the least accurate class is “lizard” (top left point), there is much less certainty about the most accurate class (bottom right).

It is straightforward to apply this type of Bayesian inference to other metrics and to other assessment tasks, such as estimating a model’s confusion matrix, ranking group performance with uncertainty (Marshall and Spiegelhalter 1998), analyzing significance of differences in performance across groups, and so on. For the CIFAR-100 dataset, based on the test data we can, for example, say that with 96% probability the ResNet-110 model is less accurate when predicting the superclass “human” than it is on “trees”; and that with 82% probability, the accuracy when the model predicts “woman” is lower than when it predicts “man.” Given constraints on space, details and examples for these approaches are provided in the Supplement.

**Active Bayesian Assessment**

Rather than relying on a random sample of labeled instances for inference, we propose to improve data efficiency by extending our Bayesian framework to support **active assessment** by actively selecting examples \( x \) for labeling in a data-efficient manner. We develop below active assessment approaches for the following three tasks: estimation, identification, and comparison. Efficient active selection of examples for labeling is particularly relevant when we have a potentially large pool of unlabeled examples \( x \) available, and have limited resources for labeling (e.g., a human labeler).

The Bayesian framework described in the last section readily lends itself to be used in Bayesian active learning algorithms, by considering model assessment as multi-armed bandit problems where each group \( g \) corresponds to an arm or a bandit. In Bayesian assessment, there are two key building blocks: (i) the assessment algorithm’s current beliefs (prior or posterior distribution) for the metric of interest \( \theta_g \sim p(\theta_g) \), and (ii) a generative model (likelihood) of the labeling outcome \( z \sim q(z|g) \), \( \forall g \).

Instead of labeling randomly sampled data points from a pool of unlabeled data, we propose instead to actively select data points to be labeled by iterating between: (1) **labeling:** actively select a group \( \hat{g} \) based on the assessment algorithms current beliefs about \( \theta_g \), randomly select a data point \( x_i \sim \mathcal{R}_g \) and then query its label; (2) **assessment:** update the assessment model given the outcome \( z_i \). This active selection approach requires defining a **reward function** \( r(z|g) \) for the revealed outcome \( z \) for the \( g \)-th group. For example, if the assessment task is to generate low variance estimates of groupwise accuracy, \( r(z|g) \) can be formulated as the reduction in uncertainty about \( \theta_g \), given an outcome \( z \), to guide the labeling process. Our goal in this paper is to demonstrate the utility of active assessment in general for performance assessment rather than comparing different active selection methods. With this in mind, we focus in particular on the framework of Thompson sampling (Thompson 1933; Russo et al. 2018) since we found it to be more reliable in terms of reliability and efficiency compared to other active selection methods such as epsilon-greedy and upper-confidence bound (UCB) approaches (additional discussion in the Supplement).

Algorithm 1 describes a general active assessment algorithm based on Thompson sampling. At each step \( i \), a set of metrics \( \theta_g, 1 \ldots, G \) are sampled from the algorithm’s current beliefs, i.e., \( \theta_g \sim p_{i-1}(\theta_g) \) (line 4). As an example, when assessing groupwise accuracy, \( p_{i-1}(\theta_g) \) represents the algorithm’s belief (e.g., in the form of a posterior Beta distribution) about the accuracy for group \( g \) given \( i-1 \) labeled examples observed so far. The sampling step is a key difference between Thompson sampling and alternatives that use a point estimate to represent current beliefs (such as greedy

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**Figure 1:** Scatter plot of estimated accuracy and expected calibration error (ECE) per class of a ResNet-110 image classifier on the CIFAR-100 test set, using our Bayesian assessment framework, with posterior means and 95% credible intervals per class. Red and blue for the top-10 least and most accurate classes, gray for the other classes.

**Figure 2:** An outline of the algorithm for active Bayesian assessment using multi-arm bandit Thompson sampling with arms corresponding to groups \( g \).

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**Algorithm 1** Thompson Sampling\((p, q, r, M)\)

1: Initialize the priors on metrics \( \{p_0(\theta_1), \ldots, p_0(\theta_g)\} \)
2: for \( i = 1, 2, \ldots \) do
3:  \# Sample parameters for the metrics \( \theta \)
4:  \( \theta_g \sim p_{i-1}(\theta_g), \, g = 1, \ldots, G \)
5:  \# Select a group \( g \) (or arm) by maximizing expected reward
6:  \( \hat{g} \leftarrow \arg \max_g E_{\theta_g}[r(z|g)] \)
7:  \# Randomly select an input data point from \( \hat{g} \)-th group and compute its predicted label
8:  \( x_i \sim \mathcal{R}_g \)
9:  \( \hat{y}_i(x_i) = \arg \max_k p_M(y = k|x_i) \)
10: \# Query to get a true label (pull arm \( \hat{g} \))
11: \( z_i \leftarrow f(y_i, \hat{y}_i(x_i)) \)
12: \# Update parameters of the \( g \)-th metric
13: \( p_i(\theta_g) \propto p_{i-1}(\theta_g) q(z_i|\theta_g) \)
14: end for

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*Slight modifications to the text for natural reading,结构调整和对英文内容的精炼。*
approaches). Conditioned on the sampled \( \theta \) values, the algorithm then selects the group \( g \) that maximizes the expected reward \( q_g = \arg \max_g \mathbb{E}_{q_g} [r(z|g)] \) (line 6) where \( r(z|g) \) is task-specific. The algorithm then draws an input datapoint \( x_t \) randomly from \( R_g \), and uses the model \( M \) to generate a predicted label \( \hat{y}_t \). The Oracle is then queried (equivalent to “pulling arm \( g \)" in a bandit setting) to obtain a label outcome \( z_t \) and the algorithm’s belief is updated (line 13) to update the posterior for \( \theta_g \), where \( z \sim q_g(z|g) \) is the likelihood for outcome \( z \). Note that this algorithm implicitly assumes that the \( \theta_g \)'s are independent (by modeling beliefs about \( \theta_g \)'s independently rather than jointly). In some situations there may be additional information across groups \( g \) (e.g., hierarchical structure) that could be leveraged (e.g., via contextual bandits) to improve inference but we leave this for future work.

We next discuss how specific reward functions \( r \) can be designed for different assessment tasks of interest, with a summary provided in Table 1.

**Estimation:** The MSE for estimation accuracy for \( G \) groups can be written in bias-variance form as \( \sum_{g=1}^{G} p_g \text{Bias}^2(\hat{\theta}_g) + \text{Var}(\hat{\theta}_g) \). Given a fixed labeling budget the bias term can be assumed to be small relative to the variance (e.g., see Sawade et al. (2010)), by using relatively weak priors for example. It is straightforward to show that to minimize \( \sum_{g=1}^{G} p_g \text{Var}(\hat{\theta}_g) \) the optimal number of labels per group \( g \) is proportional to \( \sqrt{\frac{p_g \text{Bias}(\hat{\theta}_g)}{p_g \text{Var}(\hat{\theta}_g)}} \), i.e., sample more points from larger groups and from groups where \( \theta_g \) is furthest from 0 or 1. While the group sizes \( p_g \) can be easily estimated from unlabeled data, the \( \theta_g \)'s are unknown, so we can’t compute the optimal weights a priori. Active assessment in this context allows one to minimize MSE (or RMSE) in an adaptive sequential manner. In particular we can do this by defining a reward function \( r(z|g) = p_g \cdot (\text{Var}(\hat{\theta}_g|\mathcal{L}) - \text{Var}(\hat{\theta}_g|\{\mathcal{L}, z\})) \), where \( \mathcal{L} \) is the set of labeled data seen to date, with the goal of selecting examples for labeling to minimize the overall posterior variance at each step. For confusion matrices, a similar argument applies but with multinomial likelihoods and Dirichlet posteriors on vector-valued \( \theta_g \)'s per group (see Table 1).

**Identification:** To identify the best (or worst performing) group, \( g = \arg \max_g \theta_g \), we can define a reward function using the sampled metrics \( \tilde{\theta}_g \) for each group. For example, to identify the least accurate class, the expected reward of the \( g \)-th group is \( \mathbb{E}_{q_g} [r(z_g|g)] = q_g(y=1)(-\tilde{\theta}_g) + q_g(y=0)(-\tilde{\theta}_g) = -\tilde{\theta}_g \). Similarly, because the reward functions of other identification tasks (Table 1) are independent of the value of \( y \), when the assessment tasks are to identify the group with the highest ECE or misclassification cost, maximization of the reward function corresponds to selecting the group with the greatest sampled ECE or misclassification cost.

To extend this approach to identification of the best-\( m \) arms, instead of selecting the arm with the greatest expected reward, we pull the top-\( m \)-ranked arms at each step, i.e. we query the true labels of \( m \) samples, one sample \( x \) randomly drawn from each of the top \( m \) ranked groups.

This method can be seen as an application of the general best-\( m \) arms identification method proposed by Komiyama, Honda, and Nakagawa (2015) for the problem of extreme arms identification. They proposed the multiple-play Thompson sampling (MP-TS) algorithm as a multiple-play multi-armed bandit problem, and proved that MP-TS has the optimal regret upper bound when the reward is binary. We also experimented with a modified version of Thompson sampling (TS) called top-two Thompson sampling (TTTS) (Russo 2016) but found that that TTTS and TS gave very similar results—so we just focus on TS in the results presented in this paper.

**Comparison:** For the task of comparing differences in a performance metric \( \theta \) between two groups, an active assessment algorithm can learn about the accuracy of each group by sequentially allocating the labeling budget between them. Consider two groups \( g_1 \) and \( g_2 \) with a true accuracy difference \( \Delta = \theta_{g_1} - \theta_{g_2} \). Our approach uses the “rope” (region of practical equivalence) method of Bayesian hypothesis testing (e.g., Benavoli et al. (2017)) as follows. The cumulative density in each of three regions \( \mu = (P(\Delta < -\epsilon), P(-\epsilon \leq \Delta \leq \epsilon), P(\Delta > \epsilon)) \) represents the posterior probability that the accuracy of group \( g_1 \) is more than \( \epsilon \) lower than the accuracy of \( g_2 \), that the two accuracies are “practically equivalent,” or that \( g_1 \)'s accuracy is more than \( \epsilon \) higher than that of \( g_2 \), where \( \epsilon \) is user-specified.3

The assessment task is to identify the region \( \eta = \)
arg max(μ) in which ∆ has the highest cumulative density, where λ = max(μ) ∈ [0, 1] represents the confidence of the assessment. Using Thompson sampling to actively select labels from g1 and g2, at i-th step, when we get a z, for a data point from the g-th group, we update the Beta posterior of θg. The resulting decrease in uncertainty about θg depends on the realization of the binary variable z and the current distribution of θg. We use λ to measure the amount of evidence we gathered from the labeled data from both of the groups. Then we can select the group in a greedy manner that has the greater expected increase in the confidence of the g-th group, we update the Beta posterior of θg, which is equivalent to selecting the arm with the largest E[θg | L, (g, z)]. This approach of maximal expected model change strategy has also been used in prior work in active learning for other applications (Freytag, Rodner, and Denzler 2014; Vezhnovets, Buhmann, and Ferrari 2012).

### Experimental Settings

#### Datasets and Prediction Models

In our experiments we used a number of well-known image and text classification datasets, for both image classification (CIFAR-100 (Krizhevsky and Hinton 2009), SVHN (Netzer et al. 2011) and ImageNet (Russakovsky et al. 2015)) and text classification (20 Newsgroups (Lang 1995) and DBpedia (Zhang, Zhao, and LeCun 2015)). For models M we used well-known deep learning models such as ResNets (He et al. 2016) and BERT (Devlin et al. 2019) (additional details in the Supplement). Each model was trained on the standard training set used in the literature and assessment was performed on random samples from the test sets. Table 2 provides a summary of datasets, models, and test sizes.

Unlabeled data points X, from the test set were assigned to groups (such as predicted classes or score-bins) by each prediction model. Values for pgi (for use in active learning in reward functions and in evaluation of assessment methods) were estimated using the model-based assignments of test datapoints to groups. Ground truth values for θg were defined using the full labeled test set for each dataset.

#### Priors

We investigate both uninformative and informative priors to specify prior distributions over groupwise metrics. All of the priors we use are relatively weak in terms of prior strength, but as we will see in the next section, the informative priors can be very effective when there is little labeled data available. We set the prior strengths as αg + βg = N0 = 2 for Beta priors and ∑αg = N0 = 1 for Dirichlet priors in all experiments, demonstrating the robustness of the settings across a wide variety of contexts. For groupwise accuracy, the informative Beta prior for each group is Beta(N0s, N0(1 – s)), where s is the average model confidence (score) of all unlabeled test data points for group g. The uninformative prior distribution is α = β = N0/2.

For confusion matrices, there are O(K2) prior parameters in total for K Dirichlet distributions, each distribution parameterized by a K dimensional vector αj, As an informative prior for a confusion matrix we use the model’s own prediction scores on the unlabeled test data, αjk ∝ ∑x∈R N0s jk(x), where x is a K vector. The uninformative prior for a confusion matrix is set as αjk = N0/K, j, k. In experiments, we show that even though our models are not well-calibrated (as is well-known for deep models, e.g., Guo et al. (2017)), the model’s own estimates of class-conditional probabilities nonetheless contain valuable information about confusion probabilities.

### Experimental Results

We conducted a series of experiments across datasets, models, metrics, and assessment tasks, to systematically compare three different assessment methods: (1) non-active sampling with uninformative priors (UPrior), (2) non-active sampling with informative priors (IPrior), and (3) active Thompson sampling (Figure 2) with informative priors (IPrior+TS). Estimates of metrics (as used for example in computing RMSE or ECE) correspond to mean posterior estimates of metrics (as used for example in computing RMSE, ECE, and NDCG).

Table 2: Datasets and models used in experiments.

<table>
<thead>
<tr>
<th>Mode</th>
<th>Test Set Size</th>
<th>Number of Classes</th>
<th>Prediction Model</th>
</tr>
</thead>
<tbody>
<tr>
<td>CIFAR-100</td>
<td>Image 10K</td>
<td>100</td>
<td>ResNet-110</td>
</tr>
<tr>
<td>ImageNet</td>
<td>Image 50K</td>
<td>1000</td>
<td>ResNet-152</td>
</tr>
<tr>
<td>SVHN</td>
<td>Image 26K</td>
<td>10</td>
<td>ResNet-152</td>
</tr>
<tr>
<td>20 Newsgroups</td>
<td>Text 7.5K</td>
<td>20</td>
<td>BERTBASE</td>
</tr>
<tr>
<td>DBpedia</td>
<td>Text 70K</td>
<td>14</td>
<td>BERTBASE</td>
</tr>
</tbody>
</table>

Table 3: RMSE of classwise accuracy across 5 datasets. Each RMSE number is the mean across 1000 independent runs.

<table>
<thead>
<tr>
<th>Mode</th>
<th>N/K</th>
<th>N</th>
<th>UPrior (baseline)</th>
<th>IPrior (our work)</th>
<th>IPrior+TS (our work)</th>
</tr>
</thead>
<tbody>
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<td>CIFAR-100</td>
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<td>2</td>
<td>28</td>
<td>137.572</td>
<td>0.144</td>
<td>0.025</td>
</tr>
<tr>
<td></td>
<td>5</td>
<td>70</td>
<td>33.617</td>
<td>0.019</td>
<td>0.003</td>
</tr>
<tr>
<td></td>
<td>10</td>
<td>140</td>
<td>0.000</td>
<td>0.004</td>
<td>0.001</td>
</tr>
</tbody>
</table>

Table 4: Mean relative RMSE for confusion matrix estimation. Same setup as Table 3.
As we will show below, our results clearly demonstrate that the Bayesian and active assessment frameworks are significantly more label-efficient and accurate across a wide array of assessment tasks. Best-performing values that are statistically significant, across the 3 methods, are indicated in bold in our tables. Statistical significance between the best value and next best was determined by a Wilcoxon signed-rank test with p=0.05. Results are statistically significant in all rows in all tables, except for SVHN results in Table 7. Code and scripts for all of our experiments are provided in the Supplemental Material.

**Estimation of Accuracy, Calibration, and Confusion Matrices:** We compared the estimation efficacy of each method as the labeling budget \( N \) increases, for classwise accuracy (Table 3), confusion matrices (Table 4), and ECE (Table 5). All reported numbers were obtained by averaging across 1000 independent runs, where a run corresponds to a sequence of sampled \( x_i \) values (and sampled \( \theta_y \) values for the TS method).

Table 3 shows the mean RMSE accuracy for the 3 methods on the 5 datasets. The results demonstrate that informative priors and active sampling have significantly lower RMSE than the baseline. e.g., reducing RMSE by a factor of 2 or more in the low-data regime of \( N/K = 2 \). Active sampling (IPrior+TS) improves on the IPrior method in 11 of the 15 results, but the gains are typically small. For other metrics and tasks below we will see much greater gains from using active sampling.

Table 4 reports the mean RMSE across runs of estimates of confusion matrix entries for 4 datasets\(^4\). RMSE is defined here as \( \text{RMSE} = \left( \sum_k p_k \sum_j (\hat{\theta}_{jk} - \theta_{jk})^2 \right)^{1/2} \), where \( \hat{\theta}_{jk} \) is the probability that class \( j \) is the true class when class \( k \) is predicted. To help with interpretation, we scaled the errors in the table by a constant \( \theta_0 \), defined as the RMSE of the confusion matrix estimated with scores from only unlabeled data, i.e. the estimate with IPrior when \( N = 0 \). Numbers greater than 1 mean that the estimate is worse than using \( \theta_0 \) (with no labels). The results show that using informed priors (IPrior and IPrior+TS) often produces RMSE values that are orders of magnitude lower than using simple uniform prior (UPrior). Thus, the model scores on the unlabeled test set (used to construct the informative priors) are highly informative for confusion matrix entries, even though the models themselves are (for the most part) miscalibrated. We see in addition that active sampling (IPrior+TS) provides additional significant reductions in RMSE over the IPrior method with no active sampling.

In our ECE experiments samples are grouped into 10 equal-sized bins according to their model scores. Table 5 reports the average relative ECE estimation error\(^2\), defined as \( \frac{(100/R) \sum_{r=1}^{R} |ECE_N - ECE_r|/ECE_N} \), where \( ECE_N \) is the ECE measured on the full test set, and \( ECE_r \) is the estimated ECE (using MPE estimates of \( \theta_i \)'s), for a particular method on the \( r \)th run, \( r = 1, \ldots, R = 1000 \). Both the IPrior and IPrior+TS methods have significantly lower percentage error in general in their ECE estimates compared to the naive UPrior baseline, particularly on the 3 image datasets (CIFAR-100, ImageNet, and SVHN). The bin-wise RMSE of the estimated \( \theta_i \)'s are reported in the Supplement and show similar gains for IPrior and IPrior+TS.

**Identification of Extreme Classes:** For our identification experiments, for a particular metric and choice of groups, we conducted 1000 different sequential runs. For each run, after each labeled sample, we rank the estimates \( \hat{\theta}_y \) obtained from each of the 3 methods, and compute the mean-reciprocal-rank (MRR) relative to the true top-\( m \) ranked groups (as computed from the full test set). The MRR of the predicted top-\( m \) classes is defined as \( MRR = \frac{1}{m} \sum_{i=1}^{m} \frac{1}{\text{rank}(i)} \), where \( \text{rank} \) is the predicted rank of the \( i \)th best class. Table 6 shows the mean percentage of labeled test set examples needed to correctly identify the target classes where “identify” means the minimum number of labeled examples required so that the MRR is greater than 0.99. For all 5 datasets the active method (IPrior+TS) clearly outperforms the non-active methods, with large gains in particular for cases where the number of classes \( K \) is large (CIFAR-100 and Imagenet). Similar gains in identifying the least calibrated classes are reported in the Supplement.

Figure 3 compares our 3 assessment methods for identifying the predicted classes with highest expected cost, using data from CIFAR-100, with two different (synthetic) cost matrices. In this plot the x-axis is the number of labels \( L_x \) (queries) and the y-value is the average (over all runs) of the MRR conditioned on \( L_x \) labels. In the left column (Human) the cost of misclassifying a person (e.g., predicting tree when the true class is a woman, etc.) is 10 times more expensive than other mistakes. In the right column, costs are 10 times higher if a prediction error is in a different superclass than the superclass of the true class (for the 20 superclasses in CIFAR-100). The curves show the MRR as a function of the number of labels (on average, over 100 runs) for each of the 3 assessment methods. The active assessment (IPrior+TS) is

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\(^4\)ImageNet is omitted because 50K labeled samples is not sufficient to estimate a confusion matrix that contains 1M parameters.

\(^2\)We report error for overall ECE rather than error per score-bin since ECE is of more direct interest and more interpretable.

### Table 5: Mean percentage estimation error of ECE with bins as groups. Same setup as Table 3.

<table>
<thead>
<tr>
<th>N/K</th>
<th>N</th>
<th>UPrior (baseline)</th>
<th>IPrior (our work)</th>
<th>IPrior+TS (our work)</th>
</tr>
</thead>
<tbody>
<tr>
<td>CIFAR-100</td>
<td>2</td>
<td>20</td>
<td>76.1</td>
<td>26.4</td>
</tr>
<tr>
<td></td>
<td>5</td>
<td>50</td>
<td>50.3</td>
<td>23.4</td>
</tr>
<tr>
<td></td>
<td>10</td>
<td>100</td>
<td>25.7</td>
<td>21.5</td>
</tr>
<tr>
<td>ImageNet</td>
<td>2</td>
<td>20</td>
<td>198.7</td>
<td>51.8</td>
</tr>
<tr>
<td></td>
<td>5</td>
<td>50</td>
<td>122.0</td>
<td>55.3</td>
</tr>
<tr>
<td></td>
<td>10</td>
<td>100</td>
<td>66.0</td>
<td>40.8</td>
</tr>
<tr>
<td>SVHN</td>
<td>2</td>
<td>20</td>
<td>383.6</td>
<td>86.2</td>
</tr>
<tr>
<td></td>
<td>5</td>
<td>50</td>
<td>155.8</td>
<td>93.1</td>
</tr>
<tr>
<td></td>
<td>10</td>
<td>100</td>
<td>108.2</td>
<td>80.6</td>
</tr>
<tr>
<td>20 NewsGroups</td>
<td>2</td>
<td>20</td>
<td>54.0</td>
<td>39.7</td>
</tr>
<tr>
<td></td>
<td>5</td>
<td>50</td>
<td>32.8</td>
<td>28.9</td>
</tr>
<tr>
<td></td>
<td>10</td>
<td>100</td>
<td>24.7</td>
<td>22.3</td>
</tr>
<tr>
<td>DBpedia</td>
<td>2</td>
<td>20</td>
<td>900.3</td>
<td>118.0</td>
</tr>
<tr>
<td></td>
<td>5</td>
<td>50</td>
<td>249.6</td>
<td>130.5</td>
</tr>
<tr>
<td></td>
<td>10</td>
<td>100</td>
<td>169.1</td>
<td>125.9</td>
</tr>
</tbody>
</table>
clearly much more efficient at identifying the highest cost classes than the two non-active methods. The gains from active assessment were also robust to different settings of the relative costs of mistakes (details in the Supplement).

Comparison of Groupwise Accuracy: For comparison experiments, Table 7 shows the results for the number of labeled data points required by each method to reliably assess the accuracy difference of two predicted classes, averaged over independent runs for all pairwise combinations of classes. The labeling process terminates when the most probable region $\eta$ is identified correctly and the estimation error of the cumulative density $\lambda$ is within 5% of its value on the full test set. The results show that actively allocating a labeling budget and informative priors always improves label efficiency over uniform priors with no active assessment. In addition, active sampling (IPrior+TS) shows a systematic reduction of 5% to 35% in the mean number of labels required across datasets, over non-active sampling (IPrior).

Related Work

Bayesian and Frequentist Classifier Assessment: Prior work on Bayesian assessment of prediction performance, using Beta-Bernoulli models for example, has focused on specific aspects of performance modeling, such as estimating precision-recall performance (Goutte and Gaussier 2005), comparing classifiers (Benavoli et al. 2017), or analyzing performance of diagnostic tests (Johnson, Jones, and Gardner 2019). Welinder, Welling, and Perona (2013) used a Bayesian approach to leverage a classifier’s scores on unlabeled data for Bayesian evaluation of performance. Frequentist methods for label-efficient evaluation of classifier performance have included techniques such as importance sampling (Sawade et al. 2010) and stratified sampling (Kumar and Raj 2018), and low-variance sampling methods have been developed for evaluation of information retrieval systems (Aslam, Pavlu, and Yilmaz 2006; Yilmaz and Aslam 2006; Moffat, Webber, and Zobel 2007). Our paper significantly generalizes these earlier contributions, by addressing a broader range of metrics and performance tasks within a single coherent Bayesian assessment framework and by introducing the notion of active assessment for label-efficiency.

Active Assessment: While there is a large literature on active learning and multi-armed bandits (MAB) in general, (Settles 2012; Russo et al. 2018), our paper is the first that applies ideas from Bayesian active learning to general classifier assessment, building on MAB-inspired, pool-based active learning algorithms for data selection. Nguyen, Ramanan, and Fowlkes (2018) develop non-Bayesian active learning methods to select samples for estimating visual recognition performance of an algorithm on a fixed test set and similar ideas have been explored in the information retrieval literature (Sabharwal and Sedghi 2017; Li and Kanoulas 2017; Rahman et al. 2018; Voorhees 2018; Rahman, Kutlu, and Lease 2019). However, this prior work is significantly narrower in scope in terms of performance metrics and tasks compared to the more general approach we propose here.

Conclusions

In this paper we described a Bayesian framework for assessing performance metrics of black-box classifiers, developing inference procedures for an array of assessment tasks. In particular, we proposed a new framework called active assessment for label-efficient assessment of classifier performance, and demonstrated significant performance improvements across five well-known datasets. There are a number of interesting directions for future work, such as Bayesian estimation of continuous functions related to accuracy and calibration (rather than over regions). The framework can also be extended to assess a particular model operating in multiple environments using a Bayesian hierarchical approach, or to comparatively assess multiple models operating in the same environment.

Table 6: Percentage of labeled samples needed to identify the least accurate top-1 and top-\( m \) predicted classes across 5 datasets across 1000 runs.

<table>
<thead>
<tr>
<th>Dataset</th>
<th>Top m</th>
<th>UPrior (baseline)</th>
<th>IPrior (our work)</th>
<th>IPrior+TS (our work)</th>
</tr>
</thead>
<tbody>
<tr>
<td>CIFAR-100</td>
<td>1</td>
<td>81.1</td>
<td>83.4</td>
<td>24.9</td>
</tr>
<tr>
<td></td>
<td>10</td>
<td>99.8</td>
<td>99.8</td>
<td>55.1</td>
</tr>
<tr>
<td>ImageNet</td>
<td>1</td>
<td>96.9</td>
<td>94.7</td>
<td>9.3</td>
</tr>
<tr>
<td></td>
<td>10</td>
<td>99.6</td>
<td>98.5</td>
<td>17.1</td>
</tr>
<tr>
<td>SVHN</td>
<td>1</td>
<td>90.5</td>
<td>89.8</td>
<td>82.8</td>
</tr>
<tr>
<td></td>
<td>3</td>
<td>100.0</td>
<td>100.0</td>
<td>96.0</td>
</tr>
<tr>
<td>20 Newsgroups</td>
<td>1</td>
<td>53.9</td>
<td>55.4</td>
<td>16.9</td>
</tr>
<tr>
<td></td>
<td>3</td>
<td>92.0</td>
<td>92.5</td>
<td>42.5</td>
</tr>
<tr>
<td>DBpedia</td>
<td>1</td>
<td>8.0</td>
<td>7.6</td>
<td>11.6</td>
</tr>
<tr>
<td></td>
<td>3</td>
<td>91.9</td>
<td>90.2</td>
<td>57.1</td>
</tr>
</tbody>
</table>

Figure 3: MRR of 3 assessment methods for identifying the top 1 (top) and top 10 (bottom) highest-cost predicted classes, with 2 different cost matrices (right and left), averaged over 100 trials. See text for details.

Table 7: Average number of labels across all pairs of classes required to estimate $\lambda$ for randomly selected pairs of groups.

<table>
<thead>
<tr>
<th>Dataset</th>
<th>UPrior</th>
<th>IPrior</th>
<th>IPrior+TS</th>
</tr>
</thead>
<tbody>
<tr>
<td>CIFAR-100, Superclass</td>
<td>203.5</td>
<td>129.0</td>
<td>121.9</td>
</tr>
<tr>
<td>SVHN</td>
<td>391.1</td>
<td>205.2</td>
<td>172.0</td>
</tr>
<tr>
<td>20 Newsgroups</td>
<td>197.3</td>
<td>157.4</td>
<td>136.1</td>
</tr>
<tr>
<td>DBpedia</td>
<td>217.8</td>
<td>4.3</td>
<td>2.0</td>
</tr>
</tbody>
</table>
Ethics and Societal Impact Statement

Machine learning classifiers are currently widely used to make predictions and decisions across a wide range of applications in society: education admissions, health insurance, medical diagnosis, court decisions, marketing, face recognition, and more—and this trend is likely to continue to grow. When these systems are deployed in real-world environments it will become increasingly important for users to have the ability to perform reliable, accurate, and independent evaluation of the performance characteristics of these systems and to do this in a manner which is efficient in terms of the need for labeled data.

Our paper addresses this problem directly, providing a general-purpose and transparent framework for label-efficient performance evaluations of black-box classifier systems. The probabilistic (Bayesian) aspect of our approach provides users with the ability to understand how much they can trust performance numbers given a fixed data budget for evaluation. For example, a hospital system or a university might wish to evaluate multiple different performance characteristics of pre-trained classification models in the specific context of the population of patients or students in their institution. The methods we are proposing have the potential to contribute to an increase societal trust in AI systems based on machine learning classification models.

References


