Cleaning Crowdsourced Labels Using Oracles For Supervised Learning

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ABSTRACT

Nowadays, crowdsourcing is being widely used to collect training data for supervised learning. However, crowdsourced labels are often noisy, and there is a performance gap between learning with noisy labels and learning with true labels. In this paper, we consider how to apply oracle-based label cleaning to reduce the gap. We propose TARS, a label-cleaning advisor that can provide two pieces of valuable advice for data scientists when they need to train or/and test a model using noisy labels. Firstly, in the model testing stage, given a test dataset with noisy labels, and a classification model, TARS can use the test data to estimate how well the model will perform w.r.t. true labels. Secondly, in the model training stage, given a training dataset with noisy labels, and a supervised-learning algorithm, TARS can determine which label should be sent to an oracle to clean such that the model can be improved the most. For the first advice, we propose an effective estimation technique, and study how to compute confidence intervals to bound its estimation error. For the second advice, we propose a new cleaning strategy along with two optimization techniques, and illustrate that it is superior to the existing cleaning strategies. We evaluate the effectiveness of TARS on both simulated and real-world datasets. The results show that (1) TARS can use noisy test data to accurately estimate a model’s true performance for various evaluation metrics (e.g., Accuracy, F-Score); and (2) TARS can improve the model accuracy by a larger margin than the existing cleaning strategies, for the same cleaning budget.

1 INTRODUCTION

Supervised learning, which can solve a large variety of real-world problems, such as spam detection, entity resolution, sentiment analysis, and image classification, has recently attracted significant attention in both industry and academia. Its basic idea is to learn a function from a collection of (instance, label) pairs such that the function can predict the labels of unseen instances. The resulting function is also known as a model or a classifier. Typically, the collection of labeled pairs is split into two parts: training data and test data, where the training data is for model training and the test data is for model evaluation.

Despite the great success that supervised learning has achieved, a fundamental limitation of using it in practice is the high cost of data labeling. Crowdsourcing is a promising way to solve this problem. Crowdsourcing platforms such as Amazon Mechanical Turk have hundreds of thousands of crowd workers available. These workers can be used to label data at low cost and fast speed. But at the same time, crowd workers are not accurate. They often provide noisy labels with certain probabilities to be wrong. Although this issue can be mitigated by assigning an instance to multiple workers and then infer the instance’s true label using a truth-inference algorithm like Majority Vote, the state-of-the-art truth-inference algorithms are still far from perfect [39].

Having noisy labels in training data will negatively affect the performance of a supervised-learning algorithm because the algorithm tries to predict noisy labels rather than true labels. In the Machine Learning community, there has already been some work that studies how to clean noisy labels to solve this problem [3]. However, the noisy labels are cleaned by heuristic algorithms, which have no guarantee on cleaning accuracy and may even mess up a lot of correct labels [12].

Unlike the existing work, our work draws some inspiration from the recent progress in the data cleaning community [1, 11, 16], and focuses on a different data-cleaning scenario. We consider that there exists an oracle who can be queried to clean noisy labels. Each query is to ask the oracle to verify whether a training example, (instance, label), is correctly labeled or not. If not, replace its label with the true label. This scenario more often than not holds in reality. Imagine a data scientist needs to train a good model on noisy data. In this situation, she can ask internal experts from her company to serve as oracles to clean the noisy labels.

It is worth noting that the goal of this paper is not to develop yet another supervised learning algorithm for noisy labels. Instead, we aim to develop a label cleaning advisor, named as TARS\(^1\), that can advise a data scientist on how to use the best use of oracle-based cleaning for supervised learning. As shown in Figure 1, suppose a data scientist has already trained a model on a noisy dataset. The first piece of advice she may ask for is how accurate the model is. If the current model is already good enough, there is no need to further spend effort on label cleaning. However, if she finds that the model does not meet her need, the next question she may ask is which label should be cleaned such that the model can be improved the most. TARS can provide the two pieces of advice:

- Advice 1. Model Evaluation. Given a model, and a test dataset labeled by the crowd, TARS can tell a data scientist how well

\(^1\)TARS is named after an intelligent robot in Interstellar who can provide insightful advice for human beings.
the model will perform w.r.t. true labels (rather than w.r.t. noisy labels).

• Advice 2. Cleaning Strategy. Given a learning algorithm, and a training dataset labeled by the crowd, TARS can tell a data scientist which label in the training dataset should be cleaned such that the new model, re-trained on the cleaned training dataset using the learning algorithm, has the best performance.

There are some straightforward solutions to the above two problems. In the following, we will use the simple example in Figure 2 to illustrate the limitations of these solutions, and then demonstrate the contributions made in this paper to overcome the limitations.

Let us first consider Advice 1. Figure 2(b) shows a test dataset labeled by two crowd workers w1 and w2. For simplicity, we assume that the workers w1 and w2 have the same noise rate = 0.2, which means that each of them has a probability of 0.2 to give a different label from the true label. We apply a given model to the test dataset and obtain the predicted label of each instance (see Table (c)). Since true labels are unknown, the model’s true accuracy cannot be directly derived. One naive approach is to treat noisy labels as true labels and then compute the accuracy based on the noisy labels. However, this approach is biased because it ignores the workers’ noise rates. In this example, the accuracies computed based on the noisy labels and the true labels are $\frac{3}{5}$ and $\frac{4}{5}$, respectively, and the difference is $\frac{1}{5}$.

To overcome the limitation, we present a new estimator that estimates the model’s true performance (e.g., accuracy, precision, recall, or F-score) by considering not only noisy labels but also noise rates. We prove that our estimator is unbiased (i.e., in expectation the estimator’s estimated value is equal to the true value). We further study how to compute a confidence interval for the estimator in order to bound its estimation error (i.e., bound the difference between the estimated value and the true value). This turns to be a challenging problem because the estimator’s error comes from two sources: sampling and labeling. We theoretically analyze how each source contributes to the overall estimation error, and show that the contribution of the first (second) source is controlled by test data size N (a noise rate $\beta$). Interestingly, they will not be affected by each other: (i) as $N$ increases, the overall estimation error will decrease at a rate of $O(\frac{1}{\sqrt{N}})$, regardless of what $\beta$ is; and (ii) as $\beta$ decreases, the overall estimation error will decrease at a rate of $O(\frac{1}{\sqrt{N} \cdot \frac{1}{1-\beta}})$, regardless of what $N$ is. In other words, the estimation error can be decreased by either increasing test data size or improving worker quality; the above theoretical results show the trade-off of each choice.

Next, let us consider Advice 2. Suppose Figure 2(b) represents a training dataset, where w1’s noise rate is 0.4 and w2’s noise rate is 0.01. There are five noisy labels in the dataset, and TARS needs to decide which one should be sent to an oracle to clean. Please note that this problem is different from active learning [29] because active learning assumes that data is unlabeled but here data has been labeled by crowd workers. We need to incorporate label noise into our cleaning strategy otherwise an oracle may clean many instances that have already been correctly labeled. For example, consider the data in Figure 2(b), the first four instances have the label noise of 0.4 and the label noise of the last instance is only 0.01. A good cleaning strategy should try to avoid sending the last instance to an oracle because it has a probability of 0.99 to be correct. (In addition to active learning, there are some other cleaning strategies proposed in the literature. Please refer to Section 5.1 for a more detailed discussion.)

To this end, we propose a new cleaning strategy, called expected model improvement (EMI). EMI estimates the expected model improvement of cleaning each noisy label and then selects the noisy label with the largest estimated value to clean. We illustrate the limitations of the existing cleaning strategies and explain why EMI can overcome the limitations. While the idea of EMI sounds promising in theory, we need to address some practical issues. The first issue is which data should be used to estimate the expected model improvement. If we choose the data improperly, EMI may end up training a model that performs well on the training data but not on the test data. The second issue is how to break the tie when two noisy labels have the same expected model improvement. We propose optimization techniques to address the issues and demonstrate their effectiveness experimentally.

Note that Figure 2 only shows a simplified version of our problem. In the paper, we study a more general version of the problem, where an instance can be labeled by multiple workers, a confusion matrix is used to model worker quality, and various metrics such as accuracy, precision, recall, and F-score can be chosen for model evaluation. In summary, our paper makes the following contributions:

• To the best of our knowledge, we are the first to study how to use an oracle to clean crowdsourced labels for supervised learning. We identify two challenging problems (model evaluation and cleaning strategy) in this study and present the formal problem definitions.

• We propose an estimator that can estimate a model’s true accuracy based on noisy data. We prove that the estimator is unbiased and we compute a confidence interval to bound its estimation error. We also discuss how to extend our solution to other evaluation metrics such as precision, recall, F-score.

• We develop a new cleaning strategy, called EMI, that can effectively decide which label should be cleaned. We explain why EMI is superior to the existing cleaning strategies to solve our problem. We further improve the effectiveness of EMI by developing two optimization techniques.

• We evaluate TARS on both simulated and real-world datasets. The results show that (1) TARS can use noisy labels to accurately estimate how well a model will perform w.r.t. true labels; and (2) TARS can improve the model accuracy by a larger margin than the existing cleaning strategies, for the same cleaning budget.

The remainder of this paper is organized as follows. Section 2 formally defines the model evaluation and cleaning strategy problems. Since each instance may be labeled by multiple workers, we introduce how label consolidation works in Section 3. After that, we discuss how to solve the model evaluation problem in Section 4 and the cleaning strategy problem in Section 5. Experimental results are presented in Section 6, followed by related work (Section 7) and conclusion (Section 8).

2 BACKGROUND AND PROBLEM

FORMALIZATION

We first provide some background knowledge in Section 2.1, and then formally define our problems in Sections 2.2 and 2.3.
2.1 Background

Learning With True Labels. Let $\mathcal{G}$ be the joint distribution on $(x, y) \in X \times Y$, where $x$ represents an input instance (typically a vector) and $y \in \{-1, +1\}$ represents the true label. Denote a sample drawn i.i.d. from $\mathcal{G}$ as $S = \{(x_i, y_i)\}_{i=1}^{N}$. A supervised learning algorithm aims to train a predictive model $f$ (i.e., a classifier) based on $S$ and then make predictions over the unseen instances in $\mathcal{G}$, where $f$ can be thought of as a decision function that takes an instance $x$ as input and outputs a real value $t = f(x)$. The instance will be classified as $+1$ if $t > 0$, and $-1$ otherwise.

Crowdsourced Data. Obtaining ground truth labels can be expensive, but labeling instances with imprecise crowd workers can be cheap. Let $\mathcal{W} = \{w_j\}_{j=1}^{K}$ denote a set of workers. We assume that each instance $x_i$ is labeled by a subset of $k_i$ workers; let $\mathcal{L}^j = \{(w_j, i) \mid w_j \text{ labels } x_i\}$ denote the corresponding labels, where $i, j \in \{-1, +1\}$ represents the label given by worker $w_j$. Let $C = \{(x_i, \mathcal{L}^j)\}_{i=1}^{N}$, which we call the crowdsourced data.

Worker Model. Since crowd workers are not perfect, existing crowdsourcing work typically uses confusion matrix to model worker quality. The confusion matrix of each worker $w_j$ is a $2 \times 2$ matrix,

$$q^{(j)} = \begin{bmatrix} q^{(j)}_{1_1, -1} & q^{(j)}_{1_1, +1} \\ q^{(j)}_{1_2, -1} & q^{(j)}_{1_2, +1} \end{bmatrix},$$

where each row represents a true label, each column represents a worker’s provided label, and $q^{(j)}_{y_1, y_2} (y_1 \in \{\pm 1\}, y_2 \in \{\pm 1\})$ means that an instance with true label $y_2$ is labeled by a subset of $k_i$ workers; let $\mathcal{L}^j = \{(w_j, i) \mid w_j \text{ labels } x_i\}$ denote the corresponding labels, where $i, j \in \{-1, +1\}$ represents the label given by worker $w_j$. Let $C = \{(x_i, \mathcal{L}^j)\}_{i=1}^{N}$, which we call the crowdsourced data.

Prior Probability. Each cell in a confusion matrix is a conditional probability, $q^{(j)}_{y_1, y_2} = P(L = y_1 \mid Y = y_2)$. To compute the joint probability distribution $P(L, Y)$, we also need to know a prior probability (or simply called the prior), denoted by $P(Y)$. Intuitively, the prior is the probability of an instance having a label of $+1$ or $-1$.

Computing Confusion Matrices and Prior. An important problem is how to compute workers’ confusion matrices $q^{(j)}$ (for all $j \in [1, K]$) and the prior $P(Y)$ in practice. This problem has been extensively studied in the crowdsourcing literature [18]. One simple idea is to manually label a small sample of instances upfront, and then mix these instances with other unlabeled instances and ask workers to label them all. Since workers do not know which instances have been pre-labeled, these pre-labeled instances can be used to compute workers’ confusion matrices as well as the prior. Another common idea is to leverage label redundancy. To improve quality, each instance is often labeled by multiple workers. If a worker often provides inconsistent labels with the majority of other workers, then the worker is very likely to be a low-quality worker. Based on this idea, existing work treats confusion matrices and the prior as unknown parameters and adopts an EM algorithm [8] to iteratively estimate their values.

2.2 Advice 1: How Good is a Model?

The first piece of advice from TARS is focused on the model testing stage. It considers the situation when a user has already trained a model and wants to evaluate the model’s performance using crowdsourced data.

Evaluation Metrics. To evaluate a model’s performance, people often compute a confusion matrix for the model and then derive different types of evaluation metrics from the matrix. Figure 3(a) illustrates a model’s confusion matrix. We can see that it is similar to a worker’s confusion matrix, where each row also represents a true label, but the difference is that each column represents a model’s predicted label rather than a worker’s provided label. The matrix has four cells:

- **True Positive (TP):** the number of positive instances that are correctly predicted by a model;
- **False Positive (FP):** the number of positive instances that are falsely predicted by a model;
- **True Negative (TN):** the number of negative instances that are correctly predicted by a model;
- **False Negative (FN):** the number of negative instances that are falsely predicted by a model.

TARS aims to estimate the value of each cell. In this way, any evaluation metric computed based on these cells can be derived in a straightforward manner. Figure 3(b) shows the definitions of four representative evaluation metrics.

Let $\text{eval}$ denote a user-specified evaluation metric. The model’s true performance is denoted by $\text{eval}(\mathcal{G}, f)$. For example, suppose $\text{eval}$ is accuracy, then $\text{eval}(\mathcal{G}, f)$ computes the $f$’s accuracy on $\mathcal{G}$. Since we do not have access to $\mathcal{G}$ but only crowdsourced data $\mathcal{C}$, TARS aims to use $C$ to estimate $\text{eval}(\mathcal{G}, f)$. Let $\text{eval}(\mathcal{C}, f)$ denote an estimation of $\text{eval}(\mathcal{G}, f)$. We say $\text{eval}(\mathcal{C}, f)$ to be unbiased if the expected value of the estimation is equal to the true value, i.e., $E[\text{eval}(\mathcal{C}, f)] = \text{eval}(\mathcal{G}, f)$.

In addition, TARS computes a confidence interval to bound the estimation error of $\text{eval}(\mathcal{C}, f)$. Suppose the estimated value is $\text{eval}(\mathcal{C}, f) = 0.8$. Given a confidence level (e.g., 95%), a confidence interval (e.g., 0.8 ± 0.01) indicates that the difference between the estimated value
and the true value is within ±0.01 with 95% confidence probability. The wider the confidence interval, the larger the estimation error.

**Problem 1 (Model Evaluation).** Given crowdsourced data \( C \), a model \( f \), and an evaluation metric \( \text{eval} \), TARS aims to determine (1) an unbiased estimator, \( \hat{\text{val}}(C, f) \), of the model’s true performance, and (2) a confidence interval, \( [\hat{\text{val}}(C, f) - \epsilon_1, \hat{\text{val}}(C, f) + \epsilon_2] \), at a given confidence level.

### 2.3 Advice 2: Which Label Should Be Cleaned?

The second piece of advice that TARS can provide is focused on the model training stage. It considers the situation when a user trains a model using noisy data, but the model is not good enough. The user wants to know which instance-label pair should be sent to an oracle to clean such that the model’s true performance can be improved the most.

**Oracle Labeller.** Cleaning, or ground truth labeling, can be thought of as querying a perfect worker \( w_0 \) called an oracle. It follows that the noise rates for \( w_0 \) are \( P(L = -1 | Y = +1) = P(L = +1 | Y = -1) = 0 \). We assume that queries are expensive, and thus calls to the oracle are constrained by a budget.

**Cleaning Data.** Suppose we query an oracle to clean an instance \( x_i \) and obtain ground truth label \( y_i \). One could update crowdsourced data \( C \) with this new knowledge by replacing \( L^i \) with \( \{(w_0, y_i)\} \). We say that we are cleaning instance \( x_i \), because we are substituting a set of imprecise labels with the ground truth label provided by the oracle.

**Cleaning Strategy.** Recall that a learning algorithm \( A \) takes crowdsourced data \( C \) as input and outputs a model \( f \). By cleaning the labels in \( C \), we hope \( A \) can produce a better model. It is expensive to query an oracle to get ground truth labels, thus the goal is to strategically choose \( x_i \) to clean. The best choice of \( x_i \) would result in a dataset which leads to a model more accurate than cleaning any other label. Based on the notation above, we have a formal definition of the problem below:

**Problem 2 (Cleaning Strategy).** Given crowdsourced data \( C \), a learning algorithm \( A \), and an evaluation metric \( \text{eval} \), let \( f_i \) be the model resulting from cleaning instance \( x_i \), then training on \( C \) with \( A \). TARS aims to determine which instance should be cleaned such that the model’s true performance can be improved the most:

\[
\hat{T}^* = \arg\max_{i=1, \ldots, N} \hat{\text{val}}(C, f_i).
\]

### 3 PRELIMINARY: LABEL CONSOLIDATION

Given crowdsourced data, if there are instances in the data having multiple worker labels, TARS will first consolidate these label into a single label. In other words, after this process, each instance in the crowdsourced data will have a single consolidated label along with a confusion matrix that quantifies the uncertainty of the consolidated label. In this section, we will start by showing why there is a need for this process, and then present how to get the consolidated label as well as the consolidated confusion matrix.

#### 3.1 The Need For Label Consolidation

Both Problems 1 and 2 require estimating the performance of some model which is defined based on the true labels in \( G \). Rather than true labels, we only have worker labels, which are noisy. The trick to bridging the gap between worker labels and true labels is noticing that there are some relationships between them, which are captured by the workers’ confusion matrices.

Therefore, we can use worker labels along with workers’ confusion matrices to infer the most likely true label for each instance, and then figure out how to quantify the uncertainty of each inferred label. Specifically, given a crowdsourced dataset \( C = \{(x_i, L^i)\}_{i=1}^N \), we aim to get a new dataset, denoted by \( D = \{(x_i, y_i', r_i)\}_{i=1}^N \), where \( y_i' \) and \( r_i \) represent the inferred label and the uncertainty of the inferred label mentioned above. We will present how to compute \( y_i' \) and \( r_i \) in Sections 3.2 and 3.3, respectively.

#### 3.2 Computing Consolidated Labels

Given an instance \( x_i \) with worker labels \( L^i \), the basic idea of getting the \( x_i \)’s most likely label is to compare the values of two conditional probabilities: \( P(Y_i = +1 | L^i) \) and \( P(Y_i = -1 | L^i) \). The probability that instance \( x_i \) has a true label of +1 (resp. -1), conditioned on the labels that the workers provide. If the former (latter) is larger, it means that the instance’s label is more likely to be +1 (resp. -1).

We use the work of Dawid and Skene [8] to compute the conditional probabilities. Below is an explanation of their approach, adapted to our notation. Assuming that workers provide labels independently of one another, we have:

\[
P(Y_i = +1 | L^i) = \frac{P(L^i = +1 | Y_i = +1) P(Y_i = +1)}{P(L^i = +1)}
\]

\[
= \frac{P(L^i = +1)}{P(L^i = +1)} \prod_{l_i \in L^i} P(L_j = l_{i,j} | Y_i = +1)
\]

\[
= P(Y = +1) \prod_{l_{i,j} \in L^i} q_{l_{i,j}}(j)
\]

Similarly, we can compute \( P(Y_i = -1 | L^i) \). By comparing their values, we obtain the consolidated label \( y_i' \) (if there is a tie, we break the tie randomly).

\[
y_i' = \arg\max_{y \in \{-1, 1\}} \prod_{l_{i,j} \in L^i} q_{l_{i,j}}(j)
\]

Thus, the chance of instance \( x_i \) having a ground truth label of, say, +1 is influenced by two main factors. If the instances for which \( Y = +1 \) are extremely common (i.e. \( P(Y = +1) \) is very close to 1), this increases our belief that \( Y_i = +1 \). Likewise, if the labels that workers provide are likely to happen provided that \( Y = +1 \) were true (i.e. \( q_{l_{i,j}}^{l_{i,j}} \) is very close to 1), then this also increases our belief that \( Y_i = +1 \).

#### 3.3 Quantifying Consolidated Label’s Uncertainty

Suppose an instance \( x_i \) is labeled by a group of \( k_i \) workers, denoted by \( W_i \). Let \( y_i' \) denote the consolidated label inferred from worker labels using the above method. Like the definition of a worker’s confusion matrix, we define the consolidated confusion matrix associated with a group of workers as a \( 2 \times 2 \) matrix:

\[
r^{(W_i)} = \begin{pmatrix}
(\text{prob}(W_i))_{+1,-1} & (\text{prob}(W_i))_{+1,+1} \\
(\text{prob}(W_i))_{-1,+1} & (\text{prob}(W_i))_{-1,-1}
\end{pmatrix}
\]

where each row represents a true label, each column represents a consolidated label, and \( r_{y_i', y_i'} \) (\( y_i' \in \{\pm 1\} \)) means that given an instance with true label \( y_i \), the consolidated label is \( y_i' \) with probability of \( r_{y_i', y_i'} \). In other words, each cell in the matrix is a
conditional probability \( r(W_i) = P(Y_i^\prime = y^\prime_i \mid Y = y) \) (\( y^\prime_i, y \in \{ \pm 1 \} \)). If the context is clear, \( r(W_i) \) will be abbreviated as \( r(i) \) or \( r \).

We use the Law of Total Probability to compute the conditional probability.

\[
P(Y_i^\prime \mid Y) = \sum_n P(Y_i^\prime \mid \hat{L}^i_n, Y) P(\hat{L}^i_n \mid Y),
\]

(3)

where \( \hat{L}^i_n : n = 1, 2, \cdots, 2^k \) represents all combinations of the labels that the group of \( k \) workers from \( W_i \) can provide. For example, suppose there are two workers, \( w_1, w_2 \). Then, there will be four combinations of worker labels: \( \hat{L}^i_1 = \{(w_1, -1), (w_2, -1)\} \), \( \hat{L}^i_2 = \{(w_1, -1), (w_2, +1)\} \), \( \hat{L}^i_3 = \{(w_1, +1), (w_2, -1)\} \), and \( \hat{L}^i_4 = \{(w_1, +1), (w_2, +1)\} \), where, e.g., \( \hat{L}^i_1 \) means that \( w_1 \) provides -1 and \( w_2 \) provides +1.

The equation depends on two forms of conditional probabilities: \( P(Y^\prime \mid \hat{L}^i_n, Y) \) and \( P(\hat{L}^i_n \mid Y) \). For the latter, we have already discussed how to compute it in Equation 1.

\[
P(\hat{L}^i_n \mid Y) = \prod_{(w_j, l) \in \hat{L}^i_n} g_{y_i(l)}
\]

(4)

For the former, since \( y_i^\prime \) depends only on \( \hat{L}^i_n \) (i.e., the workers and the labels that they provide), we have that:

\[
P(Y^\prime = y^\prime_i \mid \hat{L}^i_n, Y) = \begin{cases} 1 & \text{if } y^\prime_i = \tilde{y} \\ 0 & \text{otherwise} \end{cases}
\]

(5)

where \( \tilde{y} \) represents the consolidated label inferred from \( \hat{L}^i_n \).

4 MODEL EVALUATION

Now we have a noisy dataset \( D = \{(x_i, y_i^\prime, r(i))\}_{i=1}^N \), where each instance is associated with a single noisy label \( y_i^\prime \) along with a noise rate \( r(i) \). There are a number of challenging problems that need to be addressed: (1) how to develop a unified estimation framework that works for different evaluation metrics (Section 4.1); (2) how to bound the difference between the estimated value and the true value under the new framework (Section 4.2); (3) how to give a quantitative analysis on how each factor (e.g., sample vs. population, noisy labels vs. true labels) contributes to the bound (Section 4.2.1). In this section, we propose novel solutions to these non-trivial problems.

4.1 Estimating Model’s True Performance

We first present our unified estimation framework. Recall that a model’s performance (e.g., accuracy, F-score) is determined by its confusion matrix: \( \begin{bmatrix} TN & FN \\ FP & TP \end{bmatrix} \). To estimate a model’s performance, the key is to figure out how to estimate the values of the four cells in the confusion matrix. We will use TP as an example to illustrate this estimation process.

**Overview.** The estimation framework consists of two steps. The first step is to write TP in a form of the sum of loss, and the second step is to use an existing approach \([23]\) to estimate the loss.

**Step 1.** A loss function, denoted by \( \text{Loss}(t, y) \), measures the difference between a model’s prediction \( t \) and a true label \( y \). When true labels are accessible, we can represent TP as follows:

\[
TP = \sum_{i=1}^N \text{Loss}(t_i, y_i).
\]

(6)

where \( \text{Loss}(t_i, y_i) = 1 \) if \( t_i \) and \( y_i \) are both positive; 0, otherwise.

**Step 2.** In reality, however, we do not have access to true labels but only noisy labels. Thus, we need to use a noisy label \( y^\prime \) along with its noise rate \( r \) to estimate the true loss \( \text{Loss}(t, y) \). Natarajan et al. \([23]\) proposed an unbiased estimator:

\[
\tilde{\text{Loss}}(t, y^\prime) = \frac{(1 - r \cdot y^\prime \cdot y) \cdot \text{Loss}(t, y^\prime) - r y^\prime \cdot \text{Loss}(t, -y^\prime)}{1 - r \cdot 1 - r \cdot 1 + 1}.
\]

(7)

Please note that this estimator works for any bounded loss function. It is defined based on a noisy label \( y^\prime \), thus does not require knowing the true label \( y \).

**Unbiased Estimator of TP.** By plugging \( \tilde{\text{Loss}}(t, y^\prime) \) into Equation 6, we obtain an estimator of TP:

\[
\hat{TP} = \sum_{i=1}^N \tilde{\text{Loss}}(t_i, y_i^\prime).
\]

(8)

Since \( \tilde{\text{Loss}}(t, y^\prime) \) is unbiased, due to the linearity of expectation, we can easily prove that \( \hat{TP} \) is unbiased, i.e., \( \mathbb{E}[\hat{TP}] = TP \).

**Unbiased Estimator of TN, FN, FP.** Using a similar approach, we can get an unbiased estimator for TN, FN, and FP. The only difference from TP is that they need to choose a different loss function. For example, suppose we want to estimate TN. Then, the loss function w.r.t. TN should be defined as \( \text{Loss}(t_i, y_i) = 1 \) if \( t_i \) and \( y_i \) are both negative; 0, otherwise.

**Estimating Accuracy, Precision, Recall, F-Score.** Now we have known how to estimate each value in the model’s confusion matrix. These estimated values can be composed to get the model’s performance w.r.t. each evaluation metric that is defined in Figure 3. For example, by plugging the estimated values of TP and TN into the accuracy’s definition, we can get the estimated value of accuracy: \( \hat{\text{Accuracy}} = \frac{\hat{TP} + \hat{TN}}{\hat{TP} + \hat{TN} + \hat{FP} + \hat{FN}} \).

Similarly, we can get the estimated values for precision: \( \hat{\text{Precision}} = \frac{\hat{TP}}{\hat{TP} + \hat{FP}} \), and F-score: \( \hat{\text{F-score}} = \frac{2\hat{TP}}{2\hat{TP} + \hat{FP} + \hat{FN}} \).

For accuracy, we can prove that the estimation is unbiased, i.e.,

\[
\mathbb{E}[\frac{\hat{TP} + \hat{TN}}{\hat{N}}] = \mathbb{E}[\frac{\hat{TP}}{\hat{N}}] + \mathbb{E}[\frac{\hat{TN}}{\hat{N}}] = \frac{TP + TN}{N}.
\]

However, for the other three evaluation metrics, their estimators are not unbiased. This is because that when both numerator \( X \) and denominator \( Y \) are random variables, we do not have \( \mathbb{E}[[X]] = \mathbb{E}[[X]] \). This type of estimator is often called conditionally unbiased given \( Y \). Despite that they are not unbiased in theory, we validate their effectiveness in the experiments, and find that they perform very well on both synthetic and real-world datasets.

4.2 Bounding Estimation Error

In this section, we study how to compute confidence intervals for these estimators. This problem is challenging because there are two sources of error involved and a confidence interval has to take both of them into consideration.

**Sample vs. Population.** The first source of error comes from sampling. Since the entire population is not accessible, our estimator can only look at a sample of data and use it to estimate how well a model will perform over the entire population.

**Noisy Labels vs. True Labels.** The other source of error comes from noisy labels. Since true labels are not accessible, our estimator can only look at noisy labels and use them to estimate how well a model will perform w.r.t. true labels.
To address this challenge, we develop an analytical confidence interval based on the central limit theorem (CLT). From the analytical confidence interval, we can easily see how each source of error contributes to the overall estimation error (i.e., half the width of the confidence interval). However, the analytical confidence interval only works for accuracy. For the other evaluation metrics, we show how to compute their empirical confidence intervals using bootstrapping and conduct experiments to explore the impact of the two sources of error on the overall estimation error in various situations.

4.2.1 Analytical Confidence Interval. We first introduce some background knowledge about CLT, then present an analytical confidence interval for accuracy, and finally dive into the confidence interval to gain more insights.

Central Limit Theorem (CLT). Consider a population with mean \( \mu \) and variance \( \sigma^2 \). Given a random sample of size \( N \) from the population, \( \{X_1, X_2, \cdots, X_N\} \), CLT states that the sample mean \( \bar{\mu} = \frac{1}{N} \sum_{i=1}^{N} X_i \) follows a normal distribution with mean \( \mu \) and variance \( \frac{\sigma^2}{N} \). Note that CLT does not require that the original population has a normal distribution.

Suppose that we treat the sample mean as an estimator of the population mean. Based on CLT, the confidence interval for the estimator is

\[
\mu \pm \lambda \sqrt{\frac{\sigma^2}{N}},
\]

where \( \lambda \) is a parameter determined by a confidence level (e.g., \( \lambda = 1.96 \) for 95% confidence interval, \( \lambda = 2.38 \) for 99% confidence interval). Since the population mean \( \mu \) and variance \( \sigma^2 \) are unknown, they can be replaced by the estimated mean \( \bar{\mu} \) and variance \( \bar{\sigma}^2 \) based on a sample.

Analytical Confidence Interval. The estimator of accuracy can be represented as follows:

\[
\text{accuracy} = \frac{T_P + T_N}{N} - \frac{1}{N} \sum_{i=1}^{N} \text{Loss}_{0/1}(t_i, y_i'),
\]

where \( \text{Loss}_{0/1}(t_i, y_i') \) is an unbiased estimator of \( \text{Loss}_{0/1}(t_i, y_i) \), and \( \text{Loss}_{0/1}(t_i, y_i) = 1 \) if \( t_i \) and \( y_i \) have the same sign (i.e., either both positive or both negative); \( \text{Loss}_{0/1}(t_i, y_i) = 0 \), otherwise. We can see that the estimator is in the form of mean. Let \( X_i = \text{Loss}_{0/1}(t_i, y_i') \) for each \( i \in [1, N] \). The confidence interval for the estimator can be directly derived from Equation 10.

\[
E[X] \pm \lambda \sqrt{\frac{\text{var}(X)}{N}}
\]

In-Depth Analysis. We now provide an in-depth analysis of the confidence interval. As mentioned in the beginning of this section, there are two sources of error. Our analysis aims to answer two questions: (1) how does sample size affect the confidence interval? (2) how does label noise affect the confidence interval?

For simplicity, we assume that each instance has the same label noise of \( r_{-1, +1} = r_{+1, -1} = \beta \). Based on Equation 7, we find that \( \text{Loss}_{0/1}(t, y') \) can only take two possible values:

\[
\text{Loss}_{0/1}(t, y') = \begin{cases} 
\frac{1-\beta}{1-2\beta} & \text{if } t \text{ and } y' \text{ have the same sign} \\
\frac{-\beta}{1-2\beta} & \text{otherwise}
\end{cases}
\]

Figure 4: The relationships between sample size (\( N \)), label noise (\( \beta \)), and estimation error (half the width of the 95% confidence interval), for model accuracy \( \theta = 0.8 \). For simplicity, we do not show the estimation error for \( \beta \in (0.5, 1) \) because based on Equation 15, it will be the same as \( 1 - \beta \).

Suppose that \( \text{Loss}_{0/1}(t, y') \) has a probability of \( p \) being \( a = \frac{1-\beta}{1-2\beta} \) and \( 1-p \) being \( b = \frac{\beta}{1-2\beta} \). It is easy to see that \( a + b = 1 \). The expected value of \( \text{Loss}_{0/1}(t, y') \) is:

\[
E[X] = pa + (1-p)b
\]

The variance of \( \text{Loss}_{0/1}(t, y') \) is:

\[
\text{Var}[X] = \text{E}[X^2] - (\text{E}[X])^2
\]

\[
= pa^2 + (1-p)b^2 - \text{E}[X]^2
\]

\[
= pa^2 + (1-p)b^2 - (pa + (1-p)b - \text{E}[X]) - \text{E}[X]^2
\]

\[
= -ab + \text{E}[X] - \text{E}[X]^2
\]

Let \( \theta \) denote a model’s true accuracy. Since \( \frac{1}{N} \sum_{i=1}^{N} \text{Loss}_{0/1}(t_i, y_i') \) is an unbiased estimator of the true accuracy, then we have \( \text{E}[X] = \theta \).

\[
\text{Var}[X] = -ab + \theta - \theta^2
\]

By plugging Equation 14 into Equation 10, we obtain a closed form confidence interval of our estimator:

\[
\theta \pm \lambda \sqrt{-ab + \theta - \theta^2} \frac{1}{\sqrt{N}}
\]

where \( ab = -\beta(1-\beta) \), \( \beta \) is label noise, \( \theta \) is a model’s true accuracy, \( N \) is sample size, and \( \lambda \) is constant determined by a confidence level.

From this equation, we can analyze how each source of error contributes to the overall estimation error.

Insight 1. The first source of error (sample vs. population) is controlled by sample size \( N \). It only affects the denominator of the confidence interval. Figure 4(a) shows the relationship between sample size and estimation error, for different label noise \( \beta \). We can see that as sample size \( N \) increases, regardless of what \( \beta \) is, estimation error will decrease at a rate of \( O\left(\frac{1}{\sqrt{N}}\right) \). For example, when sample size is increased from \( N = 100 \) to 1000, estimation error will decrease by about \( O\left(\sqrt{\frac{1000}{100}}\right) = 3 \) times.

Insight 2. The second source of error (noisy label vs. true label) is controlled by label noise \( \beta \). It only affects the numerator of the confidence interval. Figure 4(b) demonstrates the relationship between label noise and estimation error, for different sample size \( N \). We can see that as noise decreases, regardless of what \( N \) is, estimation
error will decrease at a rate of $O\left(\frac{1}{\sqrt{n}}\right)$. For example, when $\beta$ is decreased from $\beta = 0.4$ to $\beta = 0.1$, estimation error will decrease by about $O\left(\frac{0.5-0.1}{0.5-0.4}\right) = 4$ times.

### 4.2.2 Empirical Confidence Interval

We first introduce some background knowledge about bootstrapping, and then present our approach to compute empirical confidence intervals for other estimators than accuracy.

**Bootstrapping.** Consider a population $P$ and a random sample of the population $S$. Given an estimator $\text{est}$, suppose the estimator can use the sample to estimate a complex parameter of the population. Once the estimate $\text{est}(S)$ is derived, we want to compute its confidence interval. If we could create multiple samples from the population, $S_1, S_2, \ldots, S_n$, then we would get a distribution of the estimates, $\text{est}(S_1), \text{est}(S_2), \ldots, \text{est}(S_n)$. Based on the distribution, we can find an interval that covers 95% of the estimates and use it as a 95% confidence interval of the estimation. In practice, however, the population is not available, thus it is impossible to draw another sample from $P$.

The basic idea of bootstrapping is to simulate this procedure by constructing a number of resamples of $S$. Specifically, to get a resample, it samples the data from $S$ with replacement. Let $S(1), S(2), \ldots, S(n)$ denote $n$ resamples. Then, it can get a distribution of the estimates based on resamples, $\text{est}(S(1)), \text{est}(S(2)), \ldots, \text{est}(S(n))$, from which we can compute a 95% confidence interval.

**Empirical Confidence Interval.** Now we present how to use the bootstrap to compute empirical confidence intervals for our estimators. Consider a crowdsourced dataset $D$. We can think of $D$ as a random sample of a population. While it is feasible to get multiple crowdsourced datasets from the population, the monetary cost and the time for doing so can be quite high. For example, suppose each instance needs the time for doing so can be quite high. For example, suppose each instance needs $0.5$ dollar and 5 seconds to label on average. Getting one crowdsourced dataset of size $|D| = 1000$ will cost us $500$ dollars and 1.4 hours. Repeat this for 1000 times will cost us as high as $0.5$ million dollars and 58 days.

We use bootstrapping to avoid the need to repeatedly draw samples from the population. Figure 5 illustrates the procedure. Given $D$ and an estimator $\text{eval}(D, f)$, the first step is to construct $n$ resamples of $D$, denoted by $D(1), D(2), \ldots, D(n)$. Then, we apply the estimator to each resample to get $n$ estimates,\[ \text{eval}(D(1), f), \text{eval}(D(2), f), \ldots, \text{eval}(D(n), f). \]

Given a confidence level (e.g., 95%), let $\text{eval}_{2.5\%}$ and $\text{eval}_{97.5\%}$ denote the 2.5th and 97.5th percentile of the distribution, respectively. Then, the 95% confidence interval is denoted by $[\text{eval}_{2.5\%}, \text{eval}_{97.5\%}]$.

This approach works for all the estimators developed in Section 4.1 including precision, recall, F-score. Like accuracy, the estimation error of other estimators also come from two sources. We empirically study its relationship with sample size and label noise for these estimators in the experiments.

# 5 CLEANING STRATEGY

Now we present how TARS provides the second piece of advice: which label should be cleaned? Please note that, unlike the previous section, here we turn our focus to the model training stage.

Typically, in supervised learning, we are given $S = \{(x_i, y_i)\}_{i=1}^N$ drawn from $\mathcal{O}$, and aim to make predictions on the dataset $S_{\text{test}} = \{(x_j, y_j)\}_{j=1}^M$, also drawn from $\mathcal{O}$. We could train a model $f$ on $S$, predict labels on instances $x_j$ from $S_{\text{test}}$, and compare the predicted labels with each $y_j$. Ideally, the predicted labels should be “fairly close” to the actual labels $S_{\text{test}}$. One reason why this procedure works is because both $S$ and $S_{\text{test}}$ are drawn from $\mathcal{O}$; in other words, the data we train on is an acceptable representation of the data we are expected to make predictions on.

Labels in noisy dataset $D = \{(x_i, y_i, r_i)\}_{i=1}^N$ are not guaranteed to be correct, so $D$ might not adequately represent the data we ultimately have to make predictions on. This means if we train directly on the pairs $\{(x_i, y_i)\}_{i=1}^N$, we are asked to predict the labels from a testing set $S_{\text{test}}$, there could be unacceptably many errors. Therefore, we study the cleaning strategy problem, aiming to choose the “best” instance to clean that would bring the greatest benefit to the resulting model.

In the following, we first explain why the existing cleaning strategies do not work in Section 5.1. and then present the main idea of our cleaning strategy in Section 5.2. We find that the naive implementation of this idea did not work very well in the experiments. We discuss the issues and propose effective solutions in Section 5.3.

## 5.1 Limitations of Existing Cleaning Strategies

Below are three classes of existing cleaning strategies, and explanations as to why they are not the perfect solution to our problem.

**Active Learning.** Typical active learning settings start with a small pool of (cleanly) labeled data, and a large pool of unlabeled data. Similar to our problem setting, an oracle is available to obtain ground truth labels, and the goal is to choose the most informative instances to label under a budget [29]. Active learning literature refers to approaches to this problem as query strategies. There are many query strategies proposed in the literature, such as uncertain sampling [17], expected error reduction [28]. However, since active learning involves labeling unlabeled rather than noisy data, we believe that an effective cleaning strategy in our problem setting should leverage $y_j^r$ and $r_i$, rather than treat instances as unlabeled.

Consider an example in Figure 6, where black (white) points represent positive (negative) instances and the red line represents the model. Since labels are noisy, there is a number associated with each point representing its label’s noise rate. If we ignore noise rates and simply apply an active-learning query strategy (e.g., uncertain sampling), $x_i$ will be selected because it is closest to the model’s decision boundary. However, $x_i$’s label only has a noise rate of 0.01, which is very unlikely to flip after cleaning. As shown in Figure 6 (b), if $x_i$’s label did not flip, the model would keep unchanged, thus it is a waste of cleaning budget. In comparison, $x_j$ has a much higher noisy rate, and cleaning it would be more likely to flip the label, leading to a big change of the model. Thus, $x_j$ should have a higher priority than $x_i$ to be selected, for this example.
As we can see, a good cleaning strategy should be eval
Nevertheless, it could be possible to compute the model’s expected to improve the model. For example, in Case 2, if
Case 2:
the model trained after cleaning instance
5.2 Main Idea: Expected Model Improvement
Computing expected model improvement (EMI) is defined as:
\[
\text{EMI}(i) = P(\text{case}_1) \cdot 0 + P(\text{case}_2) \cdot (\text{eval}(G, f_i) - \text{eval}(G, f))
\]
\[
= P(\text{case}_2) \cdot (\text{eval}(G, f_i) - \text{eval}(G, f))
\]
Our cleaning strategy computes EMI(i) for each instance \(x_i\), and then selects the instance \(x_i^*\) with the largest value and sends it to an oracle to clean.
\[
i^* = \arg\max_i \text{EMI}(i)
\]
Next, we discuss how to compute EMI(i), which consists of three parts:
Computing \(P(\text{case}_1)\). \(P(\text{case}_2)\) represents the probability that the noisy label is flipped after cleaning. Note that we have already known that the noisy label is \(y'_i\). Thus, \(P(\text{case}_1)\) represents the conditional probability of the true label being \(-y'_i\) given the noisy label \(y'_i\), i.e.,
\[
P(\text{case}_2) = P(Y_i = -y'_i | Y'_i = y'_i) \propto y_i | Y_i = -y'_i \] is equal to the noise rate \(\epsilon^{(i)}_{y'_i, y'_i}\) and \(P(Y_i = -y'_i)\) is equal to the prior \(P(Y = -y'_i)\). Therefore, we obtain
\[
P(\text{case}_2) \propto \epsilon^{(i)}_{y'_i, y'_i} \cdot P(Y = -y'_i)
\]
Similarly, we can obtain
\[
P(\text{case}_1) = \epsilon^{(i)}_{y_i, y_i} \cdot P(Y = y_i)
\]
Since \(P(\text{case}_1) + P(\text{case}_2) = 1\), we have
\[
P(\text{case}_2) = \frac{\epsilon^{(i)}_{y'_i, y'_i} \cdot P(Y = -y'_i)}{\epsilon^{(i)}_{y'_i, y'_i} \cdot P(Y = -y'_i) + \epsilon^{(i)}_{y_i, y_i} \cdot P(Y = y_i)}
\]
Computing \(\text{eval}(G, f_i)\). Since \(G\) is not available, we cannot compute \(\text{eval}(G, f)\) directly. Fortunately, in Section 4.1, we have discussed a way to estimate it based on noisy data \(D\), which is accessible. Thus, we can use \(\text{eval}(D, f)\) to approximate \(\text{eval}(G, f)\).
Computing \(\text{eval}(G, f_i)\). If we knew \(f_i\), \(\text{eval}(G, f_i)\) could be estimated similarly as above. Recall that \(f_i\) denotes the resulting model from training on \(D\) after cleaning instance \(x_i\). If \(x_i\)'s label is not flipped, we do not need to consider this case because the model stays the same as \(f\); if \(x_i\)'s label is flipped, we can retrain a model \(f_i\) on the new dataset \(D_i\), where \(D_i\) represents the dataset resulting from flipping the label of instance \(x_i\) of \(D\).
Remarks. EMI incorporates both noise rates and model changes, thus overcomes the limitations of the existing cleaning strategies. For example, consider \(x_i\) in Figure 6. Since it has a small noise...
rate, leading to a small value of \( P(\text{case} 2) \), EMI tends to not select \( x_i \). Consider \( x_i \) in Figure 7. Since the model would not change after flipping the label of \( x_i \), leading to a zero value of \( \text{eval}(\mathcal{G}, f) - \text{eval}(\mathcal{G}, f_i) \), EMI will not select \( x_i \).

However, the cost of having this effective cleaning strategy is that it needs to retrain \( N \) models, \( f_1, f_2, \ldots, f_N \), for each iteration, which could be highly inefficient when \( D \) is large. Fortunately, EMI is similar in spirit to an active learning query strategy called expected error reduction [28], which also needs to retrain \( N \) models. Many techniques have been proposed to reduce the retraining time, such as incremental training and subsampling techniques. In this paper, we treat this as an orthogonal problem and defer additional exploration to future work.

5.3 Further Optimization

We find that the naive implementation of EMI did not perform very well in the experiments. We discuss the reasons that cause the problem and propose effective techniques to optimize EMI.

Splitting the noisy data

The first reason is related to which noisy dataset should be used to estimate \( \text{eval}(\mathcal{G}, f) \) and \( \text{eval}(\mathcal{G}, f_i) \). One natural idea is to use \( D_t \) for \( \text{eval}(\mathcal{G}, f) \) and \( D \) for \( \text{eval}(\mathcal{G}, f_i) \) because \( f_i \) is trained on \( D_t \) and \( f \) is trained on \( D \):

\[
\text{eval}(\mathcal{G}, f) \approx \text{eval}(D_t, f), \quad \text{eval}(\mathcal{G}, f_i) \approx \text{eval}(D, f_i).
\]

However, there are two issues about this idea.

First, as shown in Equation 16, the goal is to estimate the difference between \( \text{eval}(\mathcal{G}, f) \) and \( \text{eval}(\mathcal{G}, f_i) \) as more accurate as possible. Let \( X \) and \( Y \) denote the estimators of \( \text{eval}(\mathcal{G}, f) \) and \( \text{eval}(\mathcal{G}, f_i) \), respectively. That is, we aim to minimize \( \text{var}(X - Y) = \text{var}(X) + \text{var}(Y) - \text{cov}(X, Y) \). In order to minimize \( \text{var}(X - Y) \), we need to increase \( \text{cov}(X, Y) \) as more as possible, i.e., making \( X \) and \( Y \) as more correlated as possible. If \( X \) and \( Y \) are estimated based on the same noisy data, it will make them much more more correlated than be estimated on two different ones. Another issue is about overfitting. If we train a model on a dataset and then use the same dataset to evaluate it, the model may suffer from overfitting. In other words, the model may perform well on the current dataset, but not learn to generalize to unseen data.

To address these issues, we split \( D \) into a training dataset \( D_{\text{train}} \) and a validation dataset \( D_{\text{val}} \). Only the instances in \( D_{\text{train}} \) can be cleaned and be used to train a model; the instances in \( D_{\text{val}} \) cannot be cleaned or train a model, and their job is to estimate \( \text{eval}(\mathcal{G}, f) \) and \( \text{eval}(\mathcal{G}, f_i) \):

\[
\text{eval}(\mathcal{G}, f) \approx \text{eval}(D_{\text{val}}, f), \quad \text{eval}(\mathcal{G}, f_i) \approx \text{eval}(D_{\text{val}}, f_i).
\]

It is worth mentioning that this idea has been widely adopted in machine learning, where a validation set is often used for hyperparameter tuning and has shown to be very effective to avoid overfitting.

Weighing with Model Uncertainty

Even after splitting noisy data into validation and training sets, another challenge that remains is that the values for EMI(\( i \)) for different instances \( i \) can be very similar. To see why this is the case, consider a simple situation where \( \text{eval}(\mathcal{G}, f) \) measures the percent of instances classified correctly by \( f \), and the data is labeled by a single worker. Then, the \( u^{(1)} \) constant across all instances \( i \), as is \( P(\text{case} 2) \).

This means if we have two instances \( i_1 \) and \( i_2 \), for which \( \text{eval}(\mathcal{G}, f_i) - \text{eval}(\mathcal{G}, f) \) and \( \text{eval}(\mathcal{G}, f_i) - \text{eval}(\mathcal{G}, f) \) are very similar, the resulting values \( \text{EMI}(\text{case} 1) \) and \( \text{EMI}(\text{case} 2) \) will be very similar. In the worst case, \( \text{EMI}(\text{case} 1) = \text{EMI}(\text{case} 2) \), which makes impossible to distinguish which instance would be a better candidate for cleaning.

### Table 1: Dataset statistics (SS: synthetic data with simulated noisy labels; RS: real-world data with simulated noisy labels; RR: real-world data with real-world crowdsourced noisy labels)

<table>
<thead>
<tr>
<th>Dataset</th>
<th>#Positive</th>
<th>#Negative</th>
<th>Dimension</th>
<th>Type</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gaussian</td>
<td>500</td>
<td>500</td>
<td>2</td>
<td>SS</td>
</tr>
<tr>
<td>Heart</td>
<td>120</td>
<td>150</td>
<td>13</td>
<td>RS</td>
</tr>
<tr>
<td>German</td>
<td>300</td>
<td>700</td>
<td>20</td>
<td>RS</td>
</tr>
<tr>
<td>Cancer</td>
<td>77</td>
<td>108</td>
<td>9</td>
<td>RS</td>
</tr>
<tr>
<td>Restaurant</td>
<td>102</td>
<td>1902</td>
<td>8</td>
<td>RR</td>
</tr>
</tbody>
</table>

To address this issue, we combine EMI with the uncertainty of model \( f \). More specifically, let \( u(x_i) = 1 - P(f(x_i) | x_i) \) measure the uncertainty of \( f \)’s prediction on \( x_i \). For example, we can interpret small \( P(f(x_i) | x_i) \) as a “less confident” prediction, which corresponds to large \( u(x_i) \).

Intuitively, if we have two instances whose EMI values are similar, we would like to defer the decision of “which is better” (i.e. which instance is better to clean) to the model’s uncertainty. In this case, the instance for which \( f \) more uncertain (i.e. larger \( u(x_i) \)) should be considered a better candidate for cleaning.

At first glance, it might be tempting to compare instances using \( u(x_i) \cdot \text{EMI}(i) \). Suppose \( x_i \) is the best instance to clean. Since \( \text{EMI}(i) \) is computed using estimators, it’s possible that \( \text{EMI}(i^*) < 0 \). Furthermore, if \( x_i^* \) is large, then the product \( u(x_i^*) \cdot \text{EMI}(i^*) \) could be very negative, which would rank \( x_i^* \) below other instances.

Before multiplying by \( u(x_i) \), we need to transform \( \text{EMI}(i) \) into a positive value, using some function \( \sigma: \mathbb{R} \rightarrow \mathbb{R}^+ \). In order to preserve the relative ordering of \( \text{EMI}(i) \), \( \sigma \) needs to be monotonically increasing. A convenient choice is the sigmoid function \( \sigma(t) = \frac{1}{1+e^{-t}} \). To weigh EMI with model uncertainty, we compute:

\[
\text{MU}(i) = u(x_i) \cdot \sigma(\text{EMI}(i)),
\]

Thus, with this optimization, we choose instance \( i^* \) to clean by computing:

\[
i^* = \arg\max_i \text{MU}(i)
\]
Noisy Labels. The noisy labels for the Gaussian, Heart, German, and Cancer datasets were randomly generated, controlled by two parameters, \( r_{-1,+1} \) and \( r_{+1,-1} \). For example, given \( r_{-1,+1} = 0.2 \) and \( r_{+1,-1} = 0.1 \), to generate the noisy labels for a dataset, we do the following for each instance. If the instance’s true label is -1, then it will be flipped with a probability of 0.2; if its true label is +1, then it will be flipped with a probability of 0.1. When an instance needs to be labeled by multiple workers, we will apply this process to generate multiple noisy labels for the instance and get its consolidated label using the method described in Section 3.

The noisy labels for the Restaurant dataset were collected from the real-world crowd workers in Amazon Mechanical Turk (AMT). Each instance was labeled by a single worker, and the entire dataset was labeled by 24 different workers in total. The noise rates of the workers were in the ranges of \( r_{-1,+1} \in [0, 0.4] \) and \( r_{+1,-1} \in [0, 0.2] \).

Cleaning Strategies. We compared TARS with three existing cleaning strategies. SortNoise cleans the instance whose noisy label is most likely to be wrong, without considering the impact of cleaning the instance to the current model. ActiveClean cleans the instance which, if the instance was cleaned, would impart the greatest change to the current model. Expected Error Reduction (ExpectError) [28] cleans the instance such that the current model’s error can be reduced the most, where the error is computed on noisy labels rather than estimated w.r.t. true labels. We chose ExpectError because it is similar (in spirit) to our strategy and has been shown to outperform other active-learning query strategies such as uncertain sampling [28].

All the code was written in Python 2.7. We trained logistic regression models on all datasets using scikit-learn\(^4\). Each dataset was randomly divided into a training set and a test set with the ratio of 2 to 1. We ran each experiment ten times and reported the average performance.

6.2 Evaluation of Advice 1
In this section, we first conduct sensitivity analysis on the Advice 1 provided by TARS in order to gain a deep understanding of its performance, and then examine its performance on real-world datasets.

6.2.1 Sensitivity Analysis. We evaluate the estimation error of TARS on the Gaussian dataset by varying the sample size, the noise rate, the number of votes, and the percentage of good workers, for accuracy, precision, recall, and F-score. When varying one parameter, we set the other parameters with their default values. By default, the sample size is 1000, the noise rate is 0.2, the number of votes is 1, and the percentage of good workers is 0%. We define the estimation error of TARS as half the width of the 95% confidence interval of its estimated value. We compared TARS with DirtyEval, which is a naive estimator presented in the Introduction section. It simply treats the noisy labels as the true labels without considering noise rates, thus leading to biased estimated results.

\(^4\)http://scikit-learn.org/
Sample Size. Figure 8 compares the estimation error of TARS and DirtyEval by varying the dataset size from 100 to 1000, w.r.t. different evaluation metrics. We can see that as the sample size was increased by 10 times, the estimation error of TARS was reduced by about 3 times, but DirtyEval did not change so much. The reason is that TARS takes into account noise rates, leading to larger variance. Increasing the sample size can reduce the estimator’s variance at the rate of \(O\left(\frac{1}{n}\right)\). In comparison, DirtyEval has a much larger bias but with a smaller variance. While the variance can still be decreased, since the overall estimation error is dominated by the bias, the improvement is marginal.

Noise Rate. Figure 9 compares the estimation error of TARS and DirtyEval by varying the noise rate from 0.45 to 0.1, w.r.t. different evaluation metrics. We can see that TARS follows a similar trend for all the figures: as the decrease of the noise rate, the estimation error will first decrease dramatically, quickly reaching an estimation error of less than 0.1 at the noise rate of about 0.35. After that, the decreasing speed tends to get slower. Our theoretical analysis in Section 4.2.1 has shown that for accuracy, the estimation error decreases at a rate of \(O\left(\frac{1}{0.5-p}\right)\). This experiment validated that it holds for the other evaluation metrics empirically.

Number of Votes and Percentage of Good Workers. We evaluated the performance of TARS by varying the number of votes and the percentage of good workers, respectively. The results can be found in Appendix A.

6.2.2 Performance on Real-world Datasets. We evaluate TARS on the four real-world datasets. We used the noise rate of 0.8 for Heart, German, and Cancer, and used the real-world crowd workers to label Restaurant. We aim to answer two questions in this experiment: (i) can TARS get an accurate estimate of the model’s true performance on real-world datasets? and (ii) can TARS bound the difference between the estimated value and the true value?

We used TARS and DirtyEval to estimate the model’s true accuracy on the Heart, German, and Cancer datasets. We estimated F-score for the Restaurant dataset because the class labels of the dataset is highly imbalanced. As a comparison, we computed the true accuracy (F-score) using the ground-truth labels. Note that since the entire clean population \(\mathcal{G}\) is not available, we can only compute the true accuracy (F-score) based on the clean sample \(S\) (i.e., the labeled datasets). Figure 10 shows the result. We can see that on the German and Cancer datasets, TARS returned almost the same accuracy as the true accuracy. On the Heart dataset, although the TARS’s performance was not as good as the one on the German and Cancer datasets, it was still better than DirtyEval. On the Restaurant dataset, TARS returned an estimated value of F-score with the error more than 2k smaller than DirtyEval. More importantly, on all the datasets, TARS was able to bound the estimation error. Being able to do so is essential for enabling reliable decisions in the real world.

6.3 Evaluation of Advice 2

In this section, we first examine how the proposed optimization techniques can improve the effectiveness of the naive implementation of EMI, and then compared TARS (i.e., EMI with all the optimization techniques) with the state-of-the-art cleaning strategies, ActiveClean, ExpectError, and SortNoise.

6.3.1 Optimization Techniques. In Section 5.3, we identified the possible issues when applying EMI in practice, and proposed two optimization techniques, denote by \(\text{op1}\) and \(\text{op2}\), to address them, where \(\text{op1}\) represents the optimization of splitting the noisy data and \(\text{op2}\) represents the optimization of weighing with model uncertainty. Figure 11 compares the three variants of EMI on the Gaussian dataset in four settings of noise rates.

In Figure 11(a), we set the noise rates to \((0.4, 0)\) and \((0.2, 0.2)\). We can see that in both settings, EMI\( +\text{op1}\) improved the model’s accuracy from 0.77 to 0.94. This experiment validated that it avoided this kind of situation happen.

In Figure 11(b), we set the noise rates to \((0.4, 0)\) and \((0.2, 0.2)\). In this setting, we can see that the model’s accuracy was (almost) not affected by the noisy labels. Therefore, all three cleaning strategies started with a very accurate model. After sending some instances to an oracle to clean, EMI and EMI\( +\text{op1}\) sometimes led to a much worse model, but EMI\( +\text{op1}\) avoided this kind of situation happen.

In Figure 11(c), we set the noise rates to \((0.4, 0)\) and \((0.2, 0.2)\). We can see that in both settings, EMI\( +\text{op1}\) outperformed the other two variants, further validating the effectiveness of the proposed optimization techniques.

Since EMI\( +\text{op1}\) achieved the best performance, we will only use it in TARS and compare it with the existing cleaning strategies.

6.3.2 Cleaning Strategies. Like the previous experiments, we first set the noise rates to \((0.4, 0)\), leading to a big gap between learning with noisy labels and with true labels. We trained a model on the data and then asked an oracle to clean the data. Figure 12 compares the model’s accuracy w.r.t. different cleaning strategies on the Gaussian, Heart, German, and Cancer datasets for a cleaning budget of 100 instances.

We have two observations from the results. First, TARS outperformed SortNoise and ExpectError on all four datasets. The reason is that TARS considers both label noise and model changes in its cleaning strategy while SortNoise and ExpectError only considers one of them. Second, we find that the performance of ActiveClean may not be very stable for our problem setting (see Figure 12(b) and Figure 12(c)). This is because that ActiveClean is focused on a different cleaning scenario, where there is a large dirty dataset.
(both features and labels are dirty), and an oracle cleans the data in batches. It is common that each batch contains at least 100 instances, but in this experiment, the entire cleaning budget was only 100.

The above experiment demonstrated that when the initial model's accuracy was much lower than the final model's accuracy, the cleaning strategies were able to keep improving the model's accuracy through cleaning. A natural question is that when the initial model is already good enough, will the cleaning strategies hurt the model's performance? To answer this question, we set the noise rates to (0.2, 0.2) such that the noisy labels did not impose a significant negative impact on the model's performance. Figure 13 shows the result. We can see that TARS did a good job for keeping the model's accuracy as good as the initial model's accuracy.

Mix of Noise Rates. We also compared TARS with the existing cleaning strategies on the datasets with a mix of noise rates. The results can be found in Figures 16 and 17 in the Appendix. They further validated the observations described above.

7 RELATED WORK

Crowdsourcing. There are three research topics in crowdsourcing related to our work: task assignment [2, 4, 10, 21, 25, 40], truth inference [9, 13, 15, 27, 39], and active learning (from crowds) [14, 19, 22, 37]. Task assignment studies how to determine which task should be assigned to an incoming crowd worker. A number of task assignment algorithms have been proposed [2, 21, 40], but their objectives are not to maximize the performance of a supervised-learning model. Truth inference studies the problem of inferring the ground-truth label of each instance based on (inconsistent) labels from different workers. There are many interesting ideas proposed to solve this problem [13, 15, 27]. However, the goal of truth inference is to improve label quality rather than model quality. As discussed in Section 5.1, a good cleaning strategy should not only consider noise rates but also model changes. Active learning (from crowds) determines which unlabeled instance should be sent to a crowd worker to label. Since a crowd worker may make mistakes, there are some studies on the trade-off between asking another crowd worker to relabel an instance or label a new instance [19, 30]. In our problem, we consider that all the instances have been labeled by the crowd and an oracle can be used to clean the instances.

Data Cleaning. Algorithmic data cleaning approaches have been improving in quality, but still far from perfect [5]. In view of the challenge, human-guided data cleaning has recently attracted a lot of attention [1, 6, 7, 11, 16, 24, 33–36, 38]. The existing studies can be broadly divided into two categories. One category is to leverage humans (either crowd workers or experts) to solve a particular data-cleaning problem, such as entity resolution [7, 11, 33, 35], missing value imputation [24], and data repairing [6, 36]. The other category is to clean data for a particular data analysis task, such as building a machine-learning model [16] and answering SQL queries [1, 34]. Our work belongs to this category. In particular, we are focused on cleaning crowdsourced labels for supervised learning, which is a problem that has not been explored before.

Learning with Noisy Labels. There is a large body of work in the Machine Learning community on learning with noisy labels (see [12] for a survey). Some existing approaches aim to develop a robust algorithm to tolerate label noise [20, 31]. Liu et al. [20] use importance reweighting to ensure that any surrogate loss function can be used for classification, proving that the label noise involved in training will not affect the search for an optimal classifier. Sukhbaatar et al. [31] train neural networks on images with noisy labels directly, but add an extra layer designed to model the label noise. There are also some works [3, 32] that seek to leverage data cleaning for model training. Brodley and Friedl [3] use ensembles of classifiers to identify mislabeled instances and then remove them from the training data. Veit et al. [32] use a small sample of clean data to reduce the severity of label noise on the (much larger) noisy dataset. In contrast to these works, this paper is focused on a different data-cleaning scenario, i.e., oracle-based label cleaning.

8 CONCLUSION

In this paper, we have studied the problem of cleaning crowdsourced labels using oracles for supervised learning. We developed TARS, a label-cleaning advisor that can provide data scientists with two pieces of advice when they need to train or test a model using noisy labels. We formally defined the corresponding problems: model evaluation and cleaning strategy. For the first problem, we described effective techniques to estimate the model’s true performance as well as bound the estimation error, for different evaluation
metrics (accuracy, precision, recall, F-score). For the second problem, we devised a new cleaning strategy, called EMI, to overcome the limitations of the existing cleaning strategies, and developed two techniques to further optimize its effectiveness. The experimental results show that (1) TARS can accurately estimate the model’s true performance, with the estimation error up to 3× smaller than DirtyEval; (2) TARS can improve the model accuracy by a larger margin than ActiveClean, SortNoise, and ExpertError, for the same cleaning budget.

REFERENCES


APPENDIX

A ADDITIONAL EXPERIMENTS

Number of Votes

In Section 6.2.1, we assume that each instance has a single vote (i.e., labeled by a single worker). Next, we relax the assumption and investigate how adding the number of votes will affect the estimation error. We varied the number of votes from 1 to 15. Figure 14 reports the comparison results between TARS and DirtyEval w.r.t. different evaluation metrics. We can see that increasing the number of votes reduced the estimation error exponentially. This is because that as the increase of the number of votes, the noise rate of a consolidated label will decrease exponentially. We also see that TARS outperformed DirtyEval for not only a single-vote situation but also multiple-vote situations. Eventually, their estimation error will both converge to zero.

Percentage of Good Workers

In a real-world crowdsourcing setting, there could be a mix of high-quality and low-quality workers. We designed an experiment aiming to examine whether TARS can still achieve good performance in this situation. We consider that a “good” worker has the noise rate of 0.9, and a “bad” worker has the noise rate of 0.4. We varied the percentage of the instances labeled by good workers. Figure 15 reports the result. We can see that TARS achieved much smaller estimation error than DirtyEval for all situations. For example, when 50% of the instances were labeled by good workers, TARS got an estimation error of about 0.1 (w.r.t. accuracy and F-score), while the estimation error of DirtyEval was more than 2× larger.
Figure 14: Comparison of the estimation error between TARS and DirtyEval by varying the number of votes per instance (Gaussian).

Figure 15: Comparison of the estimation error between TARS and DirtyEval by varying the percentage of good workers (Gaussian).

Figure 16: Comparing the cleaning strategy of TARS with the existing cleaning strategies (Noise Rates: 50% of (0.4, 0) and 50% of (0.2, 0.2)).

Figure 17: Comparing the cleaning strategy of TARS with the existing cleaning strategies (Noise Rates: 90% of (0.4, 0) and 10% of (0.2, 0.2)).