Consistency Regularization on Clean Samples for Learning with Noisy Labels

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SUMMARY In the recent years, deep learning has achieved significant results in various areas of machine learning. Deep learning requires a large amount of data to train a model, and data collection techniques such as web crawling have been developed. However, there is a risk that these data collection techniques may generate incorrect labels. If a deep learning model for image classification is trained on a dataset with noisy labels, the generalization performance significantly decreases. This problem is called Learning with Noisy Labels (LNL), and there have been several approaches proposed to tackle the problem of learning with noisy labels (LNL) [8]–[10]. In particular, DivideMix [1] achieves excellent results on baseline datasets with noisy labels by modeling the loss distribution to select clean samples and training a classification model in a semi-supervised manner. Following the results of DivideMix’s successful selection of clean samples, we propose to adopt a consistency regularization on the selected samples during training for preventing a classification model from overfitting to the simple patterns of the selected samples. The consistency regularization is one of the machine learning techniques developed in the field of semi-supervised learning. In our problem setting, the consistency regularization encourages a model to make consistent predictions on the perturbed images that match the predictions to the original images. Since the selected samples tend to be easy to classify, the sample selection may overlook samples that have the correct label but are difficult to classify. The consistency regularization on clean samples mitigates this drawback by adding noise to samples with simple patterns to reduce overfitting. In recent years, several approaches for data augmentation have been proposed, and we use one of them, RandAugment [11], to perturb the selected samples. An overview of our method is shown in Fig. 1.

The contributions of this study are summarized as follows. (1) We introduce the consistency regularization on the samples selected as clean as an extension of DivideMix. (2) Extensive evaluation on CIFAR-10 and CIFAR-100 with synthetically generated label noise is performed to confirm that DivideMix with the consistency regularization yields comparably or better than state-of-the-art methods. (3) We performed the ablation study on the value of hyperparameter for consistency regularization based on dataset with noisy labels and confirmed that consistency regularization contributes to the generalization performance of model.
2. Related Works

2.1 Learning with Noisy Labels

Robust-loss approaches train the classification model with a new loss function that mitigates the effect of label noise. Mean Absolute Error (MAE) is a more noise tolerant loss than widely used cross-entropy loss [12] and Generalized Cross Entropy [13] is a method that leverages the advantages of both the MAE and cross entropy. Loss-correction approaches [8], [14] estimate the noise transition matrix to correct loss function and mitigate the effect of noisy labels while training. Robust-loss and loss-correction techniques do not utilize the robustness of deep learning in the early learning phase [6].

Sample selection [9] exploits the robustness in the early learning phase to determine if the samples are labeled correctly or not. The value of loss function on the mislabeled data tends to be higher than ones with correctly labeled. Then samples with lower loss value are selected as clean samples. One drawback of the sample selection approach is that samples with low loss values and easy classification are always selected over samples that are correctly labeled but difficult to classify [15].

Label correction approaches also utilize the robustness of deep learning in the early learning stage to update the original labels with the prediction output from the softmax layer of the training model [10], [16], [17]. Some other approaches use both label correction and iterative sample selection [1], [18], [19]. Most notably, DivideMix [1] uses two networks for sample selection with a two-component mixture model, and applies the semi-supervised learning technique called MixMatch [20]. Recently, AugDesc [21] has been proposed which models the loss distribution with weakly augmented samples for sample selection and trains the model with strongly augmented samples.

More recently, two studies similar to our method have been published by R. Yi et al. [22] and K. Nishi et al. [21]

In the former work, the baseline model is a self-ensemble network, and the sample selection is based on the total classification loss of the model for the original images and transformed images by scaling rotation, and flipping. Our method differs from that method in that it selects images based on the loss distribution of the original image and transforms the selected images with simple patterns by RandAugment. Furthermore, the baseline model of our method is DivideMix. In the latter work, K. Nishi et al. [21] proposed AugDesc which utilizes weak augmentation and strong augmentation to images for sample selection and parameter update. The images transformed by the weak augmentation are fed to the model and calculate the loss values for each image. The GMM is fitted to these values, and divide the training image data into clean samples and noisy samples. Based on these binary classification, the dataset with strong augmentation is divided into labeled samples and unlabeled samples, and the model is trained using a semi-supervised learning method.

2.2 DivideMix

DivideMix [1] selects training samples with lower loss value as a set of labeled data and the rest of the samples are treated as a set of unlabeled data, and train the model in a semi-supervised learning manner. DivideMix fits a two-components Gaussian Mixture Model (GMM) [23] to the loss distribution of all training samples to find the clean probability of each sample then the samples, and divides the samples based on that probability.

Let \( \mathcal{D} = \{(x_i, y_i)\}_{i=1}^{N} \) denote the training samples where \( x_i \) is an image and \( y_i \) is an one-hot vector represents label over \( L \) classes. Suppose the parameters of a deep learning model are denoted as \( \theta \) and the objective function for training is the cross entropy loss \( \ell(\theta) \) as follows:

\[
\ell(\theta) = \{ e_{i}\}_{i=1}^{N} = \{ - \sum_{l=1}^{L} y_{il} \log(p'_{\text{model}}(x_i, \theta))\}_{i=1}^{N} \quad (1)
\]

where \( p'_{\text{model}} \) is a softmax output from the model for class...
l. After \( \ell \) is computed for all training samples, a two-component GMM is fitted to \( \ell \) using the Expectation-Maximization algorithm. Probability that a sample being clean is defined as \( w_i \) and equals to the posterior probability \( p(g | \ell_i) \) for each sample from the Gaussian component \( g \) with smaller mean.

At each epoch, training data is divided into a labeled set \( X \) and an unlabeled set \( U \) by setting a threshold \( \tau \) on \( w_i \) given by GMM of one network, and the other network is trained on the divided set in a semi-supervised manner. By alternating the roles of the two networks, they teach an estimated set of clean samples each other and avoid accumulating confirmation bias.

After dataset is divided into two sets, and a mini-batch of labeled set \( \{(x_b, y_b, w_b); b \in (1, \cdots, B)\} \) is given, label refinement is performed as follows:

\[
\bar{y}_b = y_b y_b + (1 - w_b) \hat{p}_b
\]

where \( \bar{y}_b \) is a refined label, \( p_b \) is a network’s prediction (averaged across multiple augmentations of \( x_b \)) and \( \hat{y}_b \) is a clean probability given by the other network. Given a mini-batch of unlabeled set \( \{u_b; b \in (1, \cdots, B)\} \), predictions on unlabeled samples from two networks are averaged to estimate their labels \( \hat{y}_b \). Each \( \bar{y}_b \) and \( \hat{y}_b \) are transformed by sharpening function to reduce their temperature and obtain \( \tilde{y}_b \) and \( \hat{y}_b \).

Given \( \bar{X} \) and \( \bar{U} \), MixUp [24] is applied to them where each sample is interpolated with another sample randomly chosen from the combined mini-batch of \( \bar{X} \) and \( \bar{U} \). The transformed sets are denoted as \( \bar{X}' \) and \( \bar{U}' \). Finally, semi-supervised method called MixMatch [20] is applied to the augmented dataset \( \bar{X}' \) and \( \bar{U}' \).

The loss on \( \bar{X}' \) is the cross entropy loss \( L_X \) and the loss on \( \bar{U}' \) is the mean squared error \( L_U \) as follows:

\[
L_X = -\frac{1}{|\bar{X}'|} \sum_{x \in \bar{X}'} p \log(p_{\text{model}}(x; \theta))
\]

\[
L_U = -\frac{1}{|\bar{U}'|} \sum_{x \in \bar{U}'} \|p - p_{\text{model}}(x; \theta)\|_2^2
\]

where \( p \) is a mixed label for input \( x \). With the addition of another regularization term \( L_{\text{reg}} \) which prevents assigning all samples to a single class, the final total error is:

\[
L = L_X + \lambda_U L_U + \lambda_r L_{\text{reg}}
\]

where \( \lambda_r \) is set to 1 for all experiments and \( \lambda_U \) is set to the same value as used in the experiment of DivideMix [1].

2.3 RandAugment

Data augmentation is a widely used technique to increase the number of training samples to enhance the generalization performance of image classification model. Typical data augmentation methods for images include rotation, flipping, cropping, etc. In general, data augmentation methods require expertise in each domain to apply plausible transformations to each sample. Recently, several approaches [25], [26] have been developed that learn optimal policies on a small proxy task for automatically designing augmentation strategies without prior knowledge of each domain. However, these approaches require a huge computational cost to find the optimal hyperparameters in their search space.

Instead of searching for hyperparameters in a proxy task, RandAugment [11] performs a grid search on the validation set to determine the best hyperparameters with drastically reduced computational cost. RandAugment is a simple data augmentation method using hyperparameters \( n \) and \( m \), where \( n \) controls the number of transformations to be applied to a single sample and \( m \) controls the magnitude of each transformation. RandAugment selects \( n \) transformation from the following transformations with uniform probability for every image in every minibatch:

- identity
- autoContrast
- equalize
- rotate
- solarize
- color
- posterize
- contrast
- brightness
- sharpness
- shear-x
- shear-y
- translate-x
- translate-y

3. Consistency Regularization on Selected Samples

Our proposed method is based on DivideMix [1] which is an excellent approach for learning with noisy labels by modeling the loss distribution and selecting clean samples. Since recent studies show that a deep learning model in the early learning phase is robust against the noisy labels [6], [15], DivideMix selects samples with small loss values that are easy to classify as clean samples. In other words, DivideMix may overfit to the samples with simple patterns and fail to select samples with true labels but hard to classify.

Our method prevents DivideMix from overfitting to the samples with simple patterns by introducing a consistency regularization, which is widely used in semi-supervised learning (SSL). In SSL, consistency regularization encourages the training model to output the same values to the perturbed version of the unlabeled sample as to the original sample. In our method, consistency regularization is applied to selected samples by GMM and it encourages predictions on perturbed selected samples to be consistent with predictions on the original ones. This prevents training model from overfitting to the simple patterns, and encourages the model to learn samples with true labels but hard to classify by transforming easier samples by perturbation. While DivideMix uses only random cropping and horizontal flipping as data augmentation methods, our method uses RandAugment [11] as a method for adding perturbation to training images. The abstract of our method is summarized in Fig. 1.

At first, we train two networks \( \theta_1, \theta_2 \) on the original noisy dataset \( D = \{X, Y\} = \{(x_i, y_i)\}_{i=1}^N \) for a few epochs. This training period comes from the belief about robustness of deep learning in the early learning phase. After the
warmup period, all training samples are fed into $\theta_1$ and $\theta_2$ and a two-components GMM is fit to these loss distributions. Then select a set of clean samples as a set of labeled samples $D_{b1}, D_{b2}$ and define a complementary set as a set of unlabeled samples $\bar{D}_{b1}, \bar{D}_{b2}$ based on the output from a two-components GMM. If a clean probability $w_i$ of each sample $i$ exceeds $\tau$, it is selected as clean. As shown in DivideMix, compute the refined labels $\bar{y}_b$ and $\hat{y}_b$ of each sample from $D_k$ and $\bar{D}_k$ for $k \in \{\theta_1, \theta_2\}$. After transforming each label, MixUp and MixMatch transforms each dataset into $D'_{b1}$ and $D'_{b2}$. The loss on $D'_{b1}$ is the cross entropy loss $L_X$ and the loss on $\bar{D}'_{b2}$ is the mean squared error $L_{U}$ as follows:

$$L_X = -\frac{1}{|D'_{b1}|} \sum_{x \in D'_{b1}} \sum_{l} p_l \log(p_{\text{model}}(x; \theta))$$

$$L_{U} = -\frac{1}{|\bar{D}'_{b2}|} \sum_{x \in \bar{D}'_{b2}} \| p - p_{\text{model}}(x; \theta) \|_2^2$$

where $p$ is a mixed label for input $x$. In addition to these two terms, the loss for MixMatch consists of a regularization term as in Eq. (5).

Given a set of selected samples $D_{\bar{b}} = \{X_{\bar{b}}, Y_{\bar{b}}\}$ where $k = \{1, 2\}$ be a set of clean samples for each network $(\theta_1, \theta_2)$ at each epoch, each label $y_b$ is refined to $\bar{y}_b$ by label refinement and sharpening of DivideMix. Then for each $(x_b, \bar{y}_b) \in D_{\bar{b}}$ for $b \in \{1, \cdots, B\}$, RandAugment is applied to $x_b$ to convert the easy to classify samples into the hard to classify samples and a modified sample and modified sets of samples are denoted as $x'_{b}$ and $D'_{\bar{b}} = \{X'_{\bar{b}}, Y'_{\bar{b}}\} = \{X'_{\bar{b}}, Y_{\bar{b}}\}$ where $y'_{b} \in Y'_{\bar{b}}$ of each sample is equal to the $\bar{y}_b$ of original samples $D_{\bar{b}}$. In terms of the consistency regularization, a sample $(x_b, \bar{y}_b) \in D_{\bar{b}}$ corresponds to original sample and $(x'_{b}, y'_{b}) \in D'_{\bar{b}}$ corresponds to the perturbed sample. Then the consistency on selected samples for $k$-th network is defined as follows:

$$L_c = -\frac{1}{|D'_{\bar{b}}|} \sum_{x' \in D'_{\bar{b}}} \sum_{l} \hat{y}_l \cdot \log(p'_{\text{model}}(x'; \theta))$$

For the consistency regularization, we did not apply MixUp for further modification of inputs. $L_X$ modifies $D_{\bar{b}}$ by MixUp, where MixUp is a linear interpolation between a sample of $D_{\bar{b}}$ and the other sample. Therefore $L_X$ maintains information that the label of $x_b$ is $\bar{y}_b$. Then the total loss for each network is a sum of the loss of MixMatch and the consistency regularization:

$$L = L_X + \lambda_L L_{U} + \lambda_c L_{ceg} + \lambda_c L_c$$

where $\lambda_c$ is a hyperparameter.

In Algorithm 1, we summarized the entire computational procedure. In later chapters, in addition to the comparison between the proposed method and existing methods, we will discuss the change in the value of the error function for the selected samples and the change in the accuracy of sample selection by consistency regularization.

### 3.1 Extension of Consistency Regularization

For further improvement of our method, we introduced a consistency regularization averaged over multiple inputs. The new consistency regularization is defined as follows:

$$L_{c} = -\frac{1}{I} \cdot \frac{1}{|D'_{\bar{b}}|} \sum_{i=1}^{I} \sum_{x' \in D'_{\bar{b}}^{(i)}} \sum_{l} \hat{y}_l \cdot \log(p'_{\text{model}}(x'; \theta))$$

where $I$ is the number of trials that create augmented version of $D_{\bar{b}}$ by RandAugment. $D'_{\bar{b}}^{(i)}$ is the augmented dataset at $i$-th trial. $L_{c}$ encourages model to receive different representations of $x'$ with consistent label $\hat{y}$ over $I$ trials.

### 4. Experiments

First, we performed an evaluation on a dataset containing synthetically generated label noise, and then conducted a comparison experiment with state-of-the-art approaches. In later subsections, we evaluate the sensitivity of the model to
Table 1 Comparison with baseline methods and current state-of-the-art approaches on CIFAR-10 and CIFAR-100 with symmetric label noise in test accuracy (%). DivideMix [1] and AugDesc-SAW/WAW [21] was reimplemented using public code. The mean accuracy and its standard deviation are computed over five noise realizations.

| Model          | Noise | CIFAR-10 20% | CIFAR-10 50% | CIFAR-10 80% | CIFAR-100 20% | CIFAR-100 50% | CIFAR-100 80% |
|----------------|-------|--------------|--------------|--------------|---------------|--------------|--------------|
| Cross Entropy  | Best  | 86.39 ± 0.51 | 79.49 ± 0.48 | 59.48 ± 1.19 | 61.99 ± 0.39 | 47.28 ± 0.59 | 22.35 ± 0.80 |
|                | Last  | 83.63 ± 0.17 | 58.49 ± 0.31 | 26.44 ± 0.53 | 61.73 ± 0.37 | 38.58 ± 0.47 | 10.72 ± 0.20 |
| DivideMix [1]  | Best  | 96.07 ± 0.07 | 94.72 ± 0.06 | 92.72 ± 0.27 | 76.38 ± 0.16 | 72.98 ± 0.13 | 56.02 ± 0.62 |
|                | Last  | 95.79 ± 0.07 | 94.44 ± 0.11 | 92.47 ± 0.31 | 75.93 ± 0.14 | 72.46 ± 0.07 | 55.65 ± 0.64 |
| AugDesc-SAW [21]| Best | 96.13 ± 0.07 | 94.55 ± 0.43 | 93.50 ± 0.40 | 78.86 ± 0.04 | 76.72 ± 0.17 | 55.44 ± 0.36 |
|                | Last  | 95.96 ± 0.09 | 94.39 ± 0.44 | 93.33 ± 0.38 | 78.61 ± 0.04 | 76.44 ± 0.22 | 55.20 ± 0.36 |
| AugDesc-WAW [21]| Best | 96.17 ± 0.08 | 95.24 ± 0.13 | 93.56 ± 0.48 | 78.86 ± 0.09 | 76.67 ± 0.20 | 64.53 ± 0.50 |
|                | Last  | 95.99 ± 0.09 | 95.04 ± 0.16 | 93.38 ± 0.49 | 78.61 ± 0.12 | 76.38 ± 0.20 | 64.30 ± 0.50 |
| Ours (I=1)     | Best  | 96.73 ± 0.08 | 96.20 ± 0.09 | 93.99 ± 0.13 | 80.83 ± 0.09 | 77.17 ± 0.27 | 61.19 ± 0.58 |
|                | Last  | 96.47 ± 0.12 | 95.96 ± 0.09 | 93.78 ± 0.14 | 80.28 ± 0.14 | 76.73 ± 0.41 | 60.81 ± 0.68 |
| Ours (I=2)     | Best  | 96.67 ± 0.05 | 96.37 ± 0.07 | 93.83 ± 0.11 | 81.11 ± 0.17 | 76.50 ± 0.38 | 61.05 ± 0.45 |
|                | Last  | 96.39 ± 0.04 | 96.11 ± 0.06 | 93.64 ± 0.09 | 80.51 ± 0.12 | 76.20 ± 0.35 | 60.68 ± 0.51 |

Table 2 The value of RandAugment hyperparameters $N$ and $M$ used in Table 1 with CIFAR-10 and CIFAR-100 at different levels of noise rate.

| Noise Rate | CIFAR-10 20% | CIFAR-10 50% | CIFAR-10 80% | CIFAR-100 20% | CIFAR-100 50% | CIFAR-100 80% |
|------------|--------------|--------------|--------------|---------------|--------------|--------------|
| $N$        | 2            | 2            | 1            | 1             | 2            | 2             |
| $M$        | 2            | 2            | 4            | 4             | 2            | 2             |
| $a_c$      | 0.5          | 0.5          | 0.1          | 0.5           | 0.5          | 0.1           |
| $a_H$      | 0            | 25           | 25           | 150           | 150          | 150           |

4.2 Comparison with State-of-the-Art Methods

We compared our method with DivideMix [1] and Augment Descent (AugDesc) [21], which is the current state-of-the-art for learning with noisy labels through sample selection and data augmentation, using the same network architecture. AugDesc defines the common random flip and crop image augmentation as weak data augmentation, and AutoAugment [25] as strong data augmentation. AugDesc models loss distribution on weakly or strongly augmented training samples by a two-component GMM to divide the dataset into a labeled set and an unlabeled set. Then AugDesc trains a classification model on strongly augmented training samples in a semi-supervised manner following training procedures of DivideMix. AugDesc-SAW (AugDesc-WAW) trains model with strong (weak) data augmentation during warmup period. The proposed method and AugDesc are quite similar in terms of sample selection and data augmentation strategy, but AugDesc is a method that learns only with perturbed samples for parameter updates, whereas our proposed method imposes constraints on classification model to ensure that the outputs for perturbed samples are similar to ones for original samples. This means that AugDesc does not obtain information of original samples and our proposed method receives samples selected as clean more frequently than AugDesc. In Table 1, Cross-Entropy denotes baseline model trained with Cross-Entropy loss using the same network architecture. We report the best test accuracy obtained during training and mean test accuracy averaged across the last 10 training epochs. The results with different levels of symmetric label noise on CIFAR-10 and CIFAR-100 are summarized in the Table 1 and the results given by our method (Ours ($I = 1$)) is compared with the above state-of-the-art approaches, where $I$ is the number of augmented samples used in the consistency regularization. We also report the results from a variants of our method: consistency regularization with 2 trials of data augmentation (Ours ($I = 2$)). As the number of $I$ increases, the model receives more augmented samples for stronger effect on pre-
vent preventing overfitting. At the same time, learning the original samples in the classification loss ($L_X$) becomes more difficult. We prepared these variants to explore the appropriate number of augmented samples in the consistency regularization. Our method (Ours ($I = 1$)) achieves superior results on CIFAR-10 with 50% label noise and CIFAR-100 with 20% label noise than the state-of-the-art approaches. When the noise rate of CIFAR-100 is 80%, AugDesc-WAW achieves better results than the other methods. Our method (Ours ($I = 1$)), which performs multiple trials of data augmentation to input, yields results comparative to or lower than ours with $I = 1$.

4.3 Analysis of the Effectiveness of Consistency Regularization

Table 3 shows the results of the analysis of the effect of the hyperparameter $\lambda_c$ on the performance of the model (Ours($I = 1$)) on datasets with label noise. The value of $\lambda_c$ is one of $\{0.1, 0.5, 1.0, 1.5\}$ in the experiment and Table 3 shows that the performance of model is sensitive to $\lambda_c$ when the noise rate is high (80%) on both datasets. For higher noise rate (80%), the model with smaller $\lambda_c$ yields better results. On the contrary, the results on the different noise rate are less sensitive to the value of $\lambda_c$. Figure 2 shows how the value of $L_X$ with different value of $\lambda_c$ changes as learning progresses on both datasets. We observe the value of $L_X$ to confirm the effect of the consistency regularization by changing the value of $\lambda_c$. Y-axis of each plot shows the average value of $L_X$ for each epoch. The value of $\lambda_c$ is one of $\{0.0, 0.1, 0.5, 1.0, 1.5\}$, and the top row is the result of CIFAR-10 and the bottom row is the result of CIFAR-100. $L_X$ is the loss of MixMatch on selected sample by GMM, and measures how well the model fits those samples. In all plots, the value of $L_X$ with larger $\lambda_c$ is greater than the ones with smaller $\lambda_c$. Since the best classification accuracy of our method is given when the value of $\lambda_c$ is nonzero, we can deduce that the larger value of $L_X$ is better for generalization performance, and the consistency regularization prevents model from overfitting to the selected samples. However, if the value of $\lambda_c$ is too large, $L_X$ also becomes large. This causes the learning model to underfit the selected sample, resulting in poor generalization performance as shown in Table 3.

Figure 3 shows the number of selected samples as clean and Area Under the Curve (AUC) for clean/noisy sample classification at each epoch on CIFAR-10 and CIFAR-100 training data with various $\lambda_c \in \{0.0, 0.1, 0.5, 1.0, 1.5\}$ when the noise rate is 20% or 80%. In results on both CIFAR-10 and CIFAR-100, if $\lambda_c > 0$, the number of selected sam-

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**Table 3** Test accuracy on CIFAR-10 and CIFAR-100 with noisy labels with different various on Attention module. The mean accuracy and its standard deviation are computed over three noise realizations.

| Noise    | CIFAR-10 | CIFAR-100 |
|----------|----------|-----------|
| $\lambda_c = 0.1$ |        |           |
| Best     | 96.67    | 95.86     | 94.17     | 79.90     | 76.31     | 60.94     |
| Last     | 96.36    | 95.51     | 93.91     | 79.40     | 75.89     | 60.57     |
| $\lambda_c = 0.5$ |        |           |
| Best     | 96.79    | 96.35     | 93.77     | 80.69     | 77.18     | 60.14     |
| Last     | 96.48    | 96.09     | 93.52     | 80.38     | 76.79     | 60.01     |
| $\lambda_c = 1.0$ |        |           |
| Best     | 96.30    | 95.92     | 91.59     | 79.88     | 75.49     | 56.63     |
| Last     | 96.09    | 95.74     | 91.21     | 79.48     | 75.83     | 56.29     |
| $\lambda_c = 1.5$ |        |           |
| Best     | 96.16    | 95.79     | 90.38     | 79.14     | 74.51     | 54.87     |
| Last     | 95.76    | 95.52     | 90.19     | 78.73     | 74.02     | 54.45     |
Fig. 3  Results of the number of selected samples by GMM and AUC of classification accuracy for noisy/clean labels at each epoch on CIFAR-10 and CIFAR-100. Top row: Each plot shows number of samples selected as clean vs. Number of epochs. Bottom row: Each plot shows Area Under the Curve (AUC) for clean/noisy classification vs number of epochs.

samples is greater than the case with $\lambda_c = 0$ while AUC is comparable or better. This means that the loss distribution given by model trained with consistency regularization selects the more samples with correct labels more accurately. One possible explanation for less accurate sample selection with $\lambda_c = 0$ is that the model overfits to the samples with simple patterns and regards the samples with correct labels but hard to classify as corrupted data.

4.4 Discussion

Table 1 shows that our method achieves superior results than DivideMix [1] (baseline model) in all cases, and better than or comparative to the results given by AugDesc [21] except for 80% symmetric label noise on CIFAR-100 dataset. One possible explanation for this shortage is that the number of selected sample for each class is small by GMM for high noise rate, while AugDesc applies strong data augmentation to all training samples. Figure 4 shows that the number of selected samples by GMM at each epoch and the number of selected samples decreases as the noise rate in the dataset increases. Hyper-parameter setting used in this experiments are shown in Table 2. Therefore our method does not consider the consistency on the enormous unlabeled (not selected) samples during training when the noise rate is high. For further improvements of our methods, we should address the consistency on the unlabeled (not selected) samples during training. And we compared two variants of our method (Ours($I = 1$) and Ours($I = 2$)) in Table 1. As the value of $I$ increases, the model receives more augmented samples, which is expected to have a stronger effect on preventing over-fitting, but on the other hand, learning the original samples in $L_X$ becomes more difficult. Experimental results show that $I = 1$ is sufficient for learning.

Training model with consistency regularization receives the perturbed selected samples and their original ones at the same time. Then model does not overfit to the selected samples. As shown in Fig. 2 for both datasets with various noise rates, the classification loss on the selected samples...
\( L_X \) increases as the \( \lambda \) increases in both datasets with various noise rates, indicating that the model using consistency regularization does not overfit to the selected samples during training. Also our method with nonzero \( \lambda \) achieved the best classification accuracy, it’s provably showed that preventing the model from fitting to the selected samples is benefiting for higher generalization performance. Since the samples selected by fitting GMM to the loss distribution have simple patterns and tend to be easy to classify [6], our consistency regularization contributes to preventing the model from overfitting to the simple patterns and results in higher generalization performance.

Figure 3 showed that the number of selected samples and the value of AUC of classification accuracy for noisy/clean labels given by our method is larger than the baseline (\( \lambda = 0 \)) during training when noise rate is small (20%) in both datasets. This means that the model correctly outputs high loss value to the samples with noisy labels and the model was able to be trained with more correctly labeled samples during training. When the noise rate is 80% in CIFAR-100, the results are also better than the baseline. However, when the noise rate is 80% in CIFAR-10, the number of selected samples and AUC given by our method is almost equal to the baseline. As shown in Table 1, the classification accuracy is also equal to the baseline. These results also support the claim that our method is less effective when the noise rate is large, because the number of samples to which consistency regularization is applied is small. One possible improvement is to generate accurate pseudo-labels for unlabeled samples, and apply consistency regularization to them as well.

5. Conclusion

In this paper, we adopt a consistency regularization to the selected samples by fitting GMM to the loss distribution for learning with noisy labels. Our method selects clean samples and modify their representation by RandAugment, and prevents the training model from overfitting to the easy samples with simple patterns. Experimental results show that validity of our method to combat with noisy labels.

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