# Unbiased Risk Estimator to Multi-Labeled Complementary Label Learning

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# Abstract

Multi-label learning (MLL) usually requires assigning multiple relevant labels to each instance. While a fully supervised MLL dataset needs a large amount of labeling effort, using complementary labels can help alleviate this burden. However, current approaches to learning from complementary labels are mainly designed for multi-class learning and assume that each instance has a single relevant label. This means that these approaches cannot be easily applied to MLL when only complementary labels are provided, where the number of relevant labels is unknown and can vary across instances. In this paper, we first propose the unbiased risk estimator for the *multi-labeled complementary label* learning (MLCLL) problem. We also provide an estimation error bound to ensure the convergence of the empirical risk estimator. In some cases, the unbiased estimator may give unbounded gradients for certain loss functions and result in overfitting. To mitigate this problem, we improve the risk estimator by minimizing a proper loss function, which has been shown to improve gradient updates. Our experimental results demonstrate the effectiveness of the proposed approach on various datasets.

# 1 Introduction

*Multi-label learning* (MLL) is a method of training a classifier that can predict multiple labels for an unseen instance simultaneously [Zhang and Zhou, 2014a]. It has been widely used in open-environment [Zhou, 2022] and various real-world applications such as text categorization [Maltoudoglou *et al.*, 2022; Zhang *et al.*, 2022] and image retrieval [Xu *et al.*, 2022a; Ma *et al.*, 2022]. However, collecting accurate multi-labeled data can be challenging due to the difficulty of identifying small objects or complex semantic labels in complex images, as well as the unknown number of relevant labels. This requires a significant amount of labeling effort as annotators need to exercise caution and possess specialized knowl-edge. Precisely annotating images, such as the one shown in



Figure 1: An example. The image is labeled with a variety of relevant labels including "cloud", "plant", "house", "balloon", "people", "Europe", and others. The complementary label of this image is "river". Labeling the image with the label "Europe" requires specialized knowledge as it is a complex semantic label that is challenging to annotate. On the other hand, the label "people" refers to a small object that is difficult to spot, as it is highlighted by a blue box in the image.

Fig. 1, requires a high level of attention and expertise, particularly when it comes to identifying the easily overlooked label "people" or specific geographic location label "Europe". Additionally, accurate identification of other relevant labels involves a thorough examination of each label within the entire label space.

To release the laborious of collecting multi-labeled data, we study the setting of *multi-labeled complementary label learning* (MLCLL) that could significantly reduce labeling effort, whose goal is to learn a multi-labeled classifier that can assign relevant labels for unseen instances. In MLCLL, each instance is associated with a single complementary label that specifies an irrelevant label of the instance. Obviously, providing weakly supervised information – complementary labels – releases annotation costs since selecting a complementary label does not need one-by-one checking of the entire label space and prior knowledge. For example, selecting the label "river" as the complementary label for the image in Fig. 1 is much easier than selecting all relevant labels.

Complementary label learning (CLL) is a method previously proposed for multi-class scenarios by Ishida *et al.* [2017]. Ishida *et al.* [2019] proposed an unbiased risk estimator by deriving a transition matrix using the fact that the complementary label is uniformly sampled besides the only one relevant label, such that any loss function can be

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used. With such an unbiased risk estimator, the classification risk of fully-supervised learning can be evaluated on CLL training data, enabling empirical risk minimization [Vapnik, 1991]. CLL problems are further studied for biased complementary labels [Yu *et al.*, 2018b], generative adversarial network [Xu *et al.*, 2020], multiple complementary labels [Feng *et al.*, 2020], improved gradient estimation [Chou *et al.*, 2020], or easing the dependence on estimating the transition matrix [Gao and Zhang, 2021]. These existing works are based on the fact that only one relevant label exists for multi-class cases, and most solutions are strongly dependent on the known number of irrelevant labels. They cannot be used to solve the MLCLL problem, due to the unknown number of relevant labels in MLL given only a complementary label.

To address the MLCLL problem, we first propose an unbiased risk estimator that utilizes the distribution of complementary data to approximate the distribution of fullysupervised data under certain assumptions. We also establish an estimation error bound for this estimator, guaranteeing that the classifier learned from complementary labeled data will converge to the optimal one from fully-supervised MLL. Furthermore, we analyze the impact of cross-entropy loss on learning when a crucial assumption is not met and propose a gradient-friendly loss function to prevent overfitting. Our experimental results demonstrate the effectiveness of the proposed risk minimization methods. Our contributions are summarized as follows:

- The proposal of an unbiased risk estimator for MLCLL and the derivation of an estimation error bound to ensure that the classifier learned from complementary labeled data is similar to the optimal one learned from fullysupervised MLL
- The improvement of the risk estimator by using a gradient-friendly loss function to prevent overfitting, which can occur when a crucial assumption does not hold.

The rest of this paper is organized as follows. Section 2 briefly reviews related work. We introduce our proposed approach and the gradient-friendly loss function Section 3 and 4 respectively. Section 5 describes experimental results, and Section 6 gives the conclusion.

# 2 Related Work

In MLL, the main challenge is that the number of the output space grows exponentially as the number of labels increases [Zhang and Zhou, 2014b]. There are three routes to cope with MLL problems: *first-order approach* [Zhang *et al.*, 2018; Zhang and Zhou, 2007], *second-order approach* [Elisseeff and Weston, 2001; Fürnkranz *et al.*, 2008] and *high-order approach* [Read *et al.*, 2011; Tsoumakas *et al.*, 2011]. The first-order approaches decompose MLL problems into a series of binary classification problems to solve [Zhang *et al.*, 2018; Boutell *et al.*, 2004]. Subsequently, Zhang et al. [Zhang and Zhou, 2014b; Zhang and Wu, 2015] revealed that label correlations exist in multi-labeled data, more and more studies consider label correlations to address MLL problems [Gerych

*et al.*, 2021; Li *et al.*, 2017]. Among them, the secondorder approaches consider label correlations between label pairs [Fürnkranz *et al.*, 2008; Zhang and Zhou, 2006; Li *et al.*, 2017]. Beyond the second-order relationship, the highorder approaches pay attention to exploring label correlations among label sets [Read *et al.*, 2011; Gerych *et al.*, 2021; Zhao *et al.*, 2021].

In practice, collecting precisely multi-labeled data is difficult because labeling information is often incomplete [Xu *et al.*, 2022b]. A weakly supervised framework – *partial multilabel learning* (PML) – is designed to alleviate the pain of collecting precisely labeled data [Zhou, 2018]. PML was firstly proposed by Xie and Huang [2018], where each instance is associated with a set of *candidate labels* that consists of relevant labels and irrelevant (noisy) labels. Existing PML approaches handle PML problems according to the assumption that noisy labels only compose a small portion of candidate labels [Sun *et al.*, 2022; Xie and Huang, 2020; Sun *et al.*, 2019; Yu *et al.*, 2018a]. Then, these approaches use the matrix factorization technique to decompose the candidate label matrix into the low-rank multi-label matrix and the sparse noisy label matrix to solve PML problems.

CLL was first applied in multi-class learning [Ishida *et al.*, 2017], whose emergence significantly reduces annotation costs in multi-class learning. CLL aims to recover relevant labels from complementary labels. Ishida *et al.* [2019] used uniformly sampled complementary labels to derive an unbiased risk estimator that is available for arbitrary loss functions to solve the CLL problem. In addition, studies of CLL problems further involve biased complementary labels, whose implementation depends on estimating a transition matrix [Yu *et al.*, 2018b]. To ease the dependence on estimating transition matrix, Gao and Zhang [2021] designed a discriminative way to directly model the probabilities of complementary labels. Moreover, multiple complementary labels were proposed to make the learning process have more labeling information compared with one complementary label [Feng *et al.*, 2020].

Compared with PML, MLCLL considers the hardest version of this problem – a high-noise PML problem, where the candidate label set of an instance is all labels other than its complementary label. In this case, existing PML approaches may be inapplicable for MLCLL since their implementations are based on the assumption of few noisy included in candidate labels. For CLL, existing approaches are accomplished by the fact that each instance has one relevant label in multiclass learning and irrelevant labels are known. However, the number of relevant labels is unknown and can vary across instances in MLL, which leads to CLL approaches that could not handle MLCLL problems.

## **3** The Proposed Approach

In this section, we first introduce notations and problem setting (Sec. 3.1) and the recovery of data distribution given complementary labels (Sec. 3.2). Then, we propose an unbiased risk estimator with its estimation error bound (Sec. 3.3).

### 3.1 Preliminaries

Let  $\mathcal{X} \subset \mathbb{R}^d$  be the feature space with d dimensions and  $\mathcal{Y} = \{1, 2, 3, \dots, K\}$  be the label space with K (K > 2)

possible labels. In MLL, an instance  $x \in \mathcal{X}$  with its relevant label set  $Y \subseteq \mathcal{Y}$  is drawn from the unknown joint probability distribution p(x, Y). Given fully supervised data, MLL aims to learn a classifier  $f : \mathcal{X} \mapsto [0, 1]^K$  by minimizing the following classification risk:

$$R(\boldsymbol{f}) = \mathbb{E}_{p(\boldsymbol{x},Y)}[\mathcal{L}(\boldsymbol{f}(\boldsymbol{x}),Y)], \qquad (1)$$

where

$$\mathcal{L}(\boldsymbol{f}(\boldsymbol{x}), Y) = \sum_{j=1, j \in Y}^{K} \ell_j(\boldsymbol{x}) + \sum_{j=1, j \notin Y}^{K} \bar{\ell}_j(\boldsymbol{x}).$$
(2)

Denoting the *j*-th prediction of f as  $f_j$  which estimates p(j = 1|x),  $\ell_j(x)$  and  $\bar{\ell}_j(x)$  calculate the loss of  $f_j(x)$  respectively on relevant and irrelevant labels. Specially, when  $\ell_j(x) = -\log(f_j(x))$  and  $\bar{\ell}_j(x) = -\log(1 - f_j(x))$ ,  $\mathcal{L}(f(x), Y)$  is the popular BCE loss.  $\mathcal{L}(f(x), Y)$  is not limited to BCE loss only. The *mean absolute error* (MAE) loss can also be used, resulting in  $\ell_j(x) = 1 - f_j(x)$  and  $\bar{\ell}_j(x) = f_j(x)$ .

For the MLCLL problem studied in this paper, fully supervised data is unavailable. Instead,  $\overline{D} = \{(\boldsymbol{x}_i, \overline{y}_i)\}_{i=1}^n$  containing *n* training instances is given, where  $\overline{y}_i \in \{\mathcal{Y} - Y_i\}$ is the complementary label of  $\boldsymbol{x}_i \in \mathcal{X}$ . Even given the complementary training set, the goal of MLCLL is still to learn a multi-labeled classifier  $\boldsymbol{f} : \mathcal{X} \mapsto [0, 1]^K$ , i.e., the same goal as fully supervised MLL. As  $(\boldsymbol{x}, Y)$  follows a distribution  $p(\boldsymbol{x}, Y)$ ,  $(\boldsymbol{x}, \overline{y})$  also follows a distribution, which is denoted as  $\overline{p}(\boldsymbol{x}, \overline{y})$  on which  $\overline{D}$  is drawn. In the next subsection, we will construct the relationship between  $p(\boldsymbol{x}, Y)$  and  $\overline{p}(\boldsymbol{x}, \overline{y})$  under some assumptions, and propose one scenario which guarantees assumptions to hold.

#### **3.2 Recovering the Distribution**

Without any additional knowledge, it is naturally difficult to construct the relationship between p(x, Y) and  $\bar{p}(x, \bar{y})$ , especially when the number of unlabeled data is unknown. To enable the construction, below we provide some assumptions.

**Assumption 1.** Instance-Independent Assumption: Given the complementary label  $\bar{y}$ , the relevant label set Y is independent of  $\boldsymbol{x}$ , i.e.  $p(Y|\bar{y}) = p(Y|\boldsymbol{x}, \bar{y})$ .

Assumption 1 is motivated by existing works on complementary labels [Yu *et al.*, 2018b; Feng *et al.*, 2020; Ishida *et al.*, 2019], which assumes that the complementary label is conditionally independent of x given relevant labels. This assumption does not necessarily lead to the prediction of p(Y|x), because the information on complementary labels in MLL is not complete with only one complementary label provided. Information on other complementary labels and learning on x are still essential. The following result provided a conclusion based on Assumption 1.

**Lemma 1.** Under Assumption 1,  $\sum_{\bar{y}=1, \bar{y}\notin Y}^{K} \frac{\bar{p}(\boldsymbol{x}, \bar{y})}{2^{K-1}-1}$  is a valid probability mass function with respect to  $\boldsymbol{x}$  and Y, i.e., it is non-negative and

$$\int_{\mathcal{X}} \int_{\mathcal{Y}} \sum_{\bar{y}=1, \bar{y} \notin Y}^{K} \frac{\bar{p}(\boldsymbol{x}, \bar{y})}{2^{K-1} - 1} \mathrm{d}\boldsymbol{x} \mathrm{d}Y = 1.$$
(3)

The proof is stated in Appendix A. We then give another assumption, which describes the relationship between Y and  $\bar{y}$ .

Assumption 2. Uniform Generation Assumption:

$$p(Y|\bar{y}) = \begin{cases} \frac{1}{2^{K-1}-1}, & \bar{y} \notin Y\\ 0, & \bar{y} \in Y \end{cases}.$$
 (4)

Assumption 2 provides the information on how complementary labels can contribute to the prediction of Y, i.e., Y can be any subset of  $\mathcal{Y} - \bar{y} - \emptyset$  with uniform probability. This is a strong assumption for the difficult MLCLL problem, which implies that the complementary label  $\bar{y}$  will contribute to learning no more than an irrelevant label. Such an assumption will be a necessary ground to give the unbiased risk estimation for MLCLL in Section 3.3, and will discuss an alternative solution in Section 4 if such an assumption does not hold.

With these assumptions, the following result gives the relationship between p(x, Y) and  $\bar{p}(x, \bar{y})$ .

Theorem 1. Under Assumption 1 and Assumption 2,

$$p(\boldsymbol{x}, Y) = \sum_{\bar{y}=1, \bar{y} \notin Y}^{K} \frac{\bar{p}(\boldsymbol{x}, \bar{y})}{2^{K-1} - 1}.$$
(5)

It has been verified in Lemma 1 that the right side of Eq. (5) forms a valid probability mass function under Assumption 1.

#### 3.3 Unbiased Risk Estimator

An unbiased risk estimator enables the classification risk of fully-supervised MLL to be evaluated on the given MLCLL training data. The following result shows an unbiased risk estimator of the fully supervised MLL classification risk R(f) defined in Eq. (1).

**Theorem 2.** With  $p(\mathbf{x}, Y)$  defined in Eq. (5) and  $R(\mathbf{f})$ defined in Eq. (1),  $R(\mathbf{f}) = \overline{R}(\mathbf{f})$ , where  $\overline{R}(\mathbf{f}) = \mathbb{E}_{\overline{p}(\mathbf{x},\overline{y})}[\overline{\mathcal{L}}(\mathbf{f}(\mathbf{x}),\overline{y})]$  is the expected risk on complementary data, and

$$\bar{\mathcal{L}}(\boldsymbol{f}(\boldsymbol{x}), \bar{y}) = \frac{2^{K-2}}{2^{K-1} - 1} \sum_{j=1, j \neq \bar{y}}^{K} \ell_j(\boldsymbol{x}) +$$
(6)  
$$2^{K-2} - 1 \sum_{j=1, j \neq \bar{y}}^{K} \ell_j(\boldsymbol{x}) +$$

$$\frac{2^{K-2}-1}{2^{K-1}-1}\sum_{j=1,j\neq\bar{y}}\bar{\ell}_j(\bm{x})+\bar{\ell}_{\bar{y}}(\bm{x}).$$

The proof is proved in Appendix B. Theorem 2 shows that the fully-supervised classification risk can be estimated using the complementary labels by the unbiased estimator  $\bar{R}(f)$ , with the corresponding complementary loss function defined in Eq. (6). Eq. (6) shows that the complementary loss gives low confidence in treating other labels as either relevant or irrelevant with smaller-than-one weights for them.

**Estimation Error Bound.** With Theorem 2, the expected risk  $\bar{R}(f)$  can be approximated by its empirical estimation  $\bar{R}_n(f)$ , which is

$$\bar{R}_n(\boldsymbol{f}) = \frac{1}{n} \sum_{i=1}^n \bar{\mathcal{L}}(\boldsymbol{f}(\boldsymbol{x}_i), \bar{y}_i).$$
(7)

Denote  $\mathcal{F}$  as the hypothesis class and  $\mathcal{G}_j = \{g : x \mapsto f_j(x) | f \in \mathcal{F}\}$  as the functional space for the label  $j \in \mathcal{Y}$ .  $\mathfrak{R}_n(\mathcal{G}_j)$  gives the *Rademacher Complexity* [Bartlett and Mendelson, 2002] of  $\mathcal{G}_j$ , defined as  $\mathfrak{R}_n(\mathcal{G}_j) = \mathbb{E}_{x,\sigma} \left[ \sup_{g \in \mathcal{G}_j} \frac{1}{n} \sum_{i=1}^n g(x_i) \right]$ . We further denote  $f_n \in \mathcal{F}$ 

#### Algorithm 1 MLCLL with the GDF loss

Input:  $\overline{D} = \{(\boldsymbol{x}_i, \overline{y}_i)\}_{i=1}^n$ : the training data; E: the number of epochs;  $\mathcal{A}$ : an external stochastic optimization algorithm **Output:**   $\boldsymbol{\theta}$ : model parameter for  $\boldsymbol{f}(\boldsymbol{x}; \boldsymbol{\theta})$ 1: for t = 1 to E do 2: Let  $\mathcal{L}$  be the risk,  $\mathcal{L} = \frac{1}{n} \sum_{i=1}^n \overline{\mathcal{L}}_{GDF}(\boldsymbol{f}(\boldsymbol{x}_i), \overline{y}_i) = \frac{1}{n} \sum_{i=1}^n \left\{ -\sum_{j=1, j \neq \overline{y}_i}^K \log f_j(\boldsymbol{x}_i) - \log(1 - f_{\overline{y}_i}(\boldsymbol{x}_i)) \right\};$ 3: Set gradient  $- \nabla \boldsymbol{\theta} \mathcal{L};$ 4: Update  $\boldsymbol{\theta}$  by  $\mathcal{A};$ 

5: end for

and  $f^* \in \mathcal{F}$  as the minimizers of  $\overline{R}_n(f)$  and R(f) respectively. Based on these notations, we give the estimation error bound.

**Theorem 3.** Suppose  $M = \sup_{\boldsymbol{x} \in \mathcal{X}, \boldsymbol{f} \in \mathcal{F}} \mathcal{L}(\boldsymbol{f}(\boldsymbol{x}), \bar{y})$ . For any  $j \in \mathcal{Y}$ , assuming  $\ell_j(\boldsymbol{x})$  and  $\bar{\ell}_j(\boldsymbol{x})$  are  $\beta^+$ -Lipschitz and  $\beta^-$ -Lipschitz with respect to  $\boldsymbol{f}(\boldsymbol{x})$  respectively. For any  $\delta$ , with the probability at least  $1 - \delta$ ,

$$R(\boldsymbol{f}_{n}) - R(\boldsymbol{f}^{*}) \leq M \sqrt{\frac{\log 2/\delta}{2n} + 4\sqrt{2}} \left[ \frac{(K-1)2^{K-2}}{2^{K-1} - 1} \beta^{+} + \frac{(K-1)2^{K-2} - K}{2^{K-1} - 1} \beta^{-} \right] \sum_{j=1}^{K} \Re_{n}(\mathcal{G}_{j}).$$
(8)

The proof is shown in Appendix C. Theorem 3 demonstrates that the empirical risk minimizer converges to the true risk minimizer as  $n \to \infty$ . It also demonstrates the impact of K on learning complementary labeled problems.

## 4 Gradient Descent Friendly Loss

In the above section, necessary assumptions are made for giving an unbiased risk estimator for the MLCLL problem, especially Assumption 2, which assumes that there exists no additional bias in learning besides that  $\bar{y}$  is an irrelevant label. In this section, we will investigate the situation that Assumption 2 does not hold from the perspective of the loss function and give a moderated solution.

Our proposed complementary loss function Eq. (6) is eligible to accommodate any loss functions. If we use the popular BCE loss, we have

$$\bar{\mathcal{L}}_{BCE}(\boldsymbol{f}(\boldsymbol{x}), \bar{y}) = -\frac{2^{K-2}}{2^{K-1} - 1} \sum_{j=1, j \neq \bar{y}}^{K} \log(f_j(\boldsymbol{x}_i)) \qquad (9)$$

$$-\frac{2^{K-2}-1}{2^{K-1}-1}\sum_{j=1,j\neq\bar{y}}\log(1-f_j(\boldsymbol{x}))-\log(1-f_{\bar{y}}(\boldsymbol{x})).$$

Motivated by [Chou *et al.*, 2020] which concluded that gradient estimation is important for weakly supervised learning, we further give the gradient of  $\overline{\mathcal{L}}_{BCE}$  with respect to  $\theta$ , which is learnable parameters for  $f_j(\boldsymbol{x})$ :

$$\frac{\partial \bar{\mathcal{L}}_{BCE}}{\partial \boldsymbol{\theta}} = \begin{cases} -(w^+ + w^-) \bigtriangledown_{\boldsymbol{\theta}} f_j(\boldsymbol{x}; \boldsymbol{\theta}), & \text{if } j \neq \bar{y}, \\ \frac{1}{1 - f_j(\boldsymbol{x}; \boldsymbol{\theta})} \bigtriangledown_{\boldsymbol{\theta}} f_j(\boldsymbol{x}; \boldsymbol{\theta}), & \text{if } j = \bar{y}, \end{cases}$$
(10)

where 
$$w^+ = \frac{2^{K-2}}{2^{K-1}-1} \frac{1}{f_j(\boldsymbol{x}; \boldsymbol{\theta})}$$
 and  $w^- = \frac{2^{K-2}-1}{2^{K-1}-1} \frac{1}{1-f_j(\boldsymbol{x}; \boldsymbol{\theta})}$ 

The calculation in Eq. (10) is divided into two parts: on the complementary label  $\bar{y}$  and non-complementary labels  $\mathcal{Y} - \bar{y}$ . On the complementary label, the gradient descent favors a prediction of zero for  $f_{\bar{y}}(\boldsymbol{x}; \boldsymbol{\theta})$ . However, serious overfitting can happen regarding the punishment on non-complementary labels  $\mathcal{Y} - \bar{y}$  if Assumption 2 does not hold, i.e.,  $\bar{y}$  provides some bias in learning other labels.

To give a simple example, imagine there is another label  $y_s$  which is similar to  $\bar{y}$ , such that its prediction  $f_{y_s}(x;\theta)$  will be close to the prediction on  $f_{\bar{y}}(x;\theta)$ , i.e., zero. When  $f_{y_s}(x;\theta) = 0$  or close to zero,  $w^+$  in Eq. (10) will become  $\infty$ , resulting in  $\infty$  gradient although the prediction  $f_{y_s}$  is already close to the groundtruth. In another example, when a label  $y_d$  is absolutely exclusive of  $\bar{y}$ , its close-to-groundtruth prediction should be close to one, resulting in an infinity  $w^-$ , still an  $\infty$  gradient despite that the prediction is close to groundtruth. These two naive examples could demonstrate that if Assumption 2 is violated, serious overfitting could happen when using the proposed unbiased risk estimator.

To mitigate the overfitting caused due to violating Assumption 2, we design a new gradient by using the  $\bar{y}$  part in Eq. (10), but a modification on the gradient of the  $Y - \bar{y}$  part. Such a gradient is potentially calculated from a Gradient-Descent-Friendly (GDF) loss function and shown as

$$\frac{\partial \bar{\mathcal{L}}_{\text{GDF}}}{\partial \boldsymbol{\theta}} = \begin{cases} -\frac{1}{f_j(\boldsymbol{x};\boldsymbol{\theta})} \bigtriangledown \boldsymbol{\theta} f_j(\boldsymbol{x};\boldsymbol{\theta}), & \text{if } j \neq \bar{y}, \\ \frac{1}{1 - f_j(\boldsymbol{x};\boldsymbol{\theta})} \bigtriangledown \boldsymbol{\theta} f_j(\boldsymbol{x};\boldsymbol{\theta}), & \text{if } j = \bar{y}. \end{cases}$$
(11)

Eq. (11) achieves a kind of gradient that will enable any label belonging to  $Y - \bar{y}$  to have a prediction higher than the prediction on the complementary label, i.e., achieving a good ranking effect. When the label absolutely exclusive from  $\bar{y}$  exists, such a gradient will favor the prediction to be close to one. It seems that the problem still remains if there is any label similar to  $\bar{y}$  exists, however, the learning on  $\bar{y}$  will provide a way of balancing the learning of these two labels, and a good ranking performance can still be achieved in MLL metrics such as *ranking loss* or *average precision*. In Eq. (11), we remove all weights related to K to make the gradients caused by individual labels have uniform weights in case one label dominates the learning.

The corresponding loss function of Eq. (11) is

$$\bar{\mathcal{L}}_{\text{GDF}}(\boldsymbol{f}(\boldsymbol{x}), \bar{y}) = -\sum_{j=1, j \neq \bar{y}}^{K} \log(f_j(\boldsymbol{x})) - \log(1 - f_{\bar{y}}(\boldsymbol{x})).$$
(12)

The procedure for optimizing the GDF loss is shown in Algorithm 1. We will show in the next section how optimizing Eq. (12) can mitigate overfitting and improve performance.

MAE loss is another widely used loss function, which is shown to be less likely to overfit but slower at convergence [Ghosh *et al.*, 2017]. Formally, the MLCLL corresponding MAE loss and its gradients are

$$\bar{\mathcal{L}}_{MAE}(\boldsymbol{f}(\boldsymbol{x}), \bar{y}) = \frac{2^{K-2}}{2^{K-1}-1} \sum_{j=1, j \neq \bar{y}}^{K} (1 - f_j(\boldsymbol{x}))$$

Table 1: Experimental results (mean $\pm$ std) on training data with first pre-processing way (each instance is given one complementary label and labels are kept under 15). The best performance of each dataset is shown in **boldface**, where  $\bullet/\circ$  denotes whether GDF is superior/inferior to baselines with pairwise *t*-test (at 0.05 significance level).

Methods	MLL		PML		CLL	Unbounded	Bounded	GDF				
	ML-KNN	LIFT	fpml	PML-LRS	L-UW	BCE	MAE	GDF				
One Error↓												
bookmark	.801±.006•	.649±.015●	.885±.019•	.584±.004•	.504±.008•	.568±.006•	.613±.011•	.483±.007				
Corel16k	.736±.054•	.789±.044●	.816±.023•	.730±.000●	$.703 {\pm} .058$	.798±.019●	.702±.053	.706±.053				
delicious	.592±.017•	.533±.014•	.617±.017•	.452±.007•	.482±.018●	.490±.011•	.467±.019•	.426±.012				
mediamill	.198±.020•	$.187 {\pm} .019$	.188±.019	.200±.002•	$.188 {\pm} .019$	.190±.014●	.188±.019	.184±.018				
eurlex_dc	.790±.033•	.759±.024●	.920±.024●	.517±.013●	.609±.024●	.575±.019•	.906±.010●	.459±.015				
eurlex_sm	.667±.017•	.674±.013•	.868±.032•	.457±.005•	.552±.011•	.534±.012•	.834±.008•	.400±.010				
scene	.692±.029•	.605±.022•	.815±.026•	.540±.022●	.764±.023●	.449±.031•	.805±.021•	.383±.020				
yeast	.297±.028•	.284±.027•	.251±.023	.738±.097●	$.253 {\pm} .022$	.277±.016●	.251±.023	.249±.024				
Ranking Loss												
bookmark	.348±.006•	.310±.007•	.468±.019•	.260±.004●	.196±.007	.260±.005•	.301±.008•	.196±.005				
Corel16k	.328±.045•	.392±.026•	.420±.031•	$.303 {\pm} .005$	.301±.039	.400±.016●	.316±.032	.307±.040				
delicious	.398±.004•	.383±.003•	.438±.008•	.305±.002•	.319±.005●	.336±.003•	.305±.004•	.292±.005				
mediamill	.200±.014●	.202±.015•	.206±.019•	$.160 {\pm} .001 {\circ}$	.193±.020●	.157±.009∘	.192±.019●	.174±.013				
eurlex_dc	.283±.008•	.294±.012•	.470±.037•	.179±.006●	.267±.015●	.216±.017•	.437±.008•	.161±.005				
eurlex_sm	.322±.007•	.333±.013•	.472±.034•	.238±.004•	.329±.005●	.273±.014●	.461±.006•	.209±.006				
scene	.340±.030●	.289±.019•	.504±.024•	.258±.006•	.457±.018●	.184±.019●	.492±.023●	.153±.009				
yeast	.247±.011•	.298±.012•	.233±.012•	.464±.018•	.252±.012●	.250±.015•	.238±.014•	.219±.013				
Average Precision <sup>↑</sup>												
bookmark	.383±.006•	.480±.010•	.267±.018•	.534±.004•	.604±.005•	.544±.005•	.494±.008•	.619±.006				
Corel16k	$.405 {\pm} .047$	.350±.033•	.325±.021•	$.423 {\pm} .006$	.434±.047	.343±.017•	$.426 {\pm} .040$	.429±.044				
delicious	.487±.006●	.511±.004●	.457±.006●	.580±.002•	.554±.006●	.544±.005●	.567±.006●	.586±.004				
mediamill	.711±.010●	.710±.009●	.709±.012•	.748±.0010	.717±.012●	.765±.005∘	.715±.012●	.732±.007				
eurlex_dc	.417±.018●	.429±.016●	.240±.028•	.616±.009•	.525±.017∙	.563±.014●	.267±.009●	.658±.010				
eurlex_sm	.426±.009•	.425±.012•	.285±.026•	.584±.004●	.475±.008●	.524±.012•	.288±.003•	.623±.009				
scene	.543±.023●	.600±.016●	.417±.020•	.637±.010●	.463±.018●	.717±.022•	.429±.018●	.759±.012				
yeast	.677±.018•	.636±.016•	.688±.016•	.459±.031•	.679±.016●	.679±.015•	.693±.017•	.712±.018				

$$+\frac{2^{K-2}-1}{2^{K-1}-1}\sum_{j=1,j\neq\bar{y}}^{K}f_{j}(\boldsymbol{x})+f_{\bar{y}}(\boldsymbol{x}),$$
(13)

and

$$\frac{\partial \bar{\mathcal{L}}_{\text{MAE}}}{\partial \boldsymbol{\theta}} = \begin{cases} -\frac{1}{2^{K-1}-1} \bigtriangledown_{\boldsymbol{\theta}} f_j(\boldsymbol{x}; \boldsymbol{\theta}), & \text{if } j \neq \bar{y}, \\ \bigtriangledown_{\boldsymbol{\theta}} f_j(\boldsymbol{x}; \boldsymbol{\theta}), & \text{if } j = \bar{y}. \end{cases}$$
(14)

Eq. (14) shows the reason why the  $\bar{\mathcal{L}}_{MAE}$  is less likely to overfit but slower in convergence compared to the  $\bar{\mathcal{L}}_{BCE}$ . By comparing Eqs. (10), (14) and (11), we can see that the GDF loss is a trade-off between the MAE one and the BCE one from the perspective of the gradient. The relationship between  $\bar{\mathcal{L}}_{GDF}$  and  $\bar{\mathcal{L}}_{MAE}$  can be further shown by the fact that  $\bar{\mathcal{L}}_{MAE} \leq \bar{\mathcal{L}}_{GDF}$ . The empirical comparison of optimizing these three loss functions is shown in Section 5.

## **5** Experiments

In this section, we conduct experiments to evaluate the performance of the unbiased risk estimator with various loss functions and our proposed GDF loss. Here, we adopt five MLL criteria, including *ranking loss, hamming loss, one error, coverage* and *average precision*, to measure the performance of approaches. For *average precision*, the greater the value, the better the performance, while for the remaining four criteria, the smaller the values the better the performance. We use PyTorch [Paszke *et al.*, 2019] and NVIDIA TITAN RTX to implement our experiments, where the code is available at https://github.com/GaoYi439/GDF.

### 5.1 Experimental Settings

**Datasets & pre-processing.** We use eight widely-used MLL datasets to experiments<sup>1</sup>, where we adopt two pre-processing ways to process datasets to verify the performance of the proposed approach. The first way follows Xie et al. [2018; 2020], rare labels and instances being relevant to these rare labels are removed for datasets with more than 15 labels, whose

<sup>&</sup>lt;sup>1</sup>Publicly available at https://mulan.sourceforge.net/datasetsmlc.html.

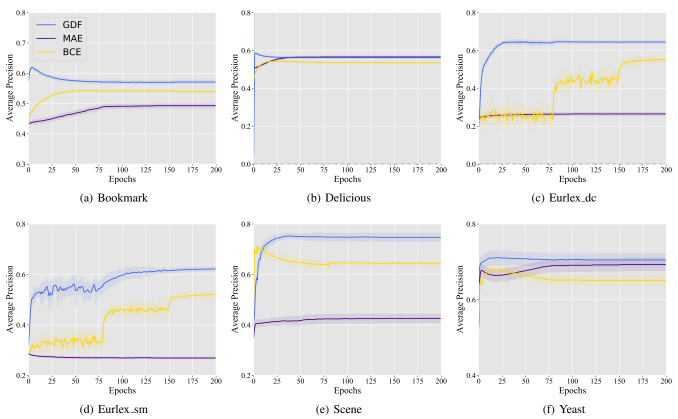


Figure 2: Average precision on various datasets adopted the first pre-processing way, where each instance is associated with one complementary label and labels are kept under 15. Dark colors show the mean of testing average precision and light colors are corresponding to the std.

labels are kept under 15. Datasets use the first way to process, where each instance is associated with one complementary label. The second way keeps the original number of labels for datasets, while the number of complementary labels per instance is half the original number of labels in datasets. Detailed descriptions of these datasets are shown in Appendix D.

**Baselines.** We adopt two PML approaches (fpml [Yu et al., 2018a] and PML-LRS [Sun et al., 2019]) as baselines, which use candidate labels  $(\mathcal{Y} \setminus \bar{y})$  to learn. Similarly, a CLL approach, L-UW [Gao and Zhang, 2021], is used in our experiments. Due to that L-UW is applied in multi-class learning, we use the Sigmoid layer and BCE loss to replace the Softmax layer and cross-entropy loss respectively, and help it suit the MLCLL problem. In addition, ML-KNN [Zhang and Zhou, 2007] and LIFT [Zhang and Wu, 2015] are two MLL approaches, which are absorbed to compare with our approach. They deal with MLCLL by taking all candidate labels  $(\mathcal{Y} \setminus \bar{y})$  of instances as possible labels. To verify the feasibility of theoretical analysis in Section 4, we use the BCE loss  $\bar{\mathcal{L}}_{\mathrm{BCE}}$  and MAE loss  $\bar{\mathcal{L}}_{\mathrm{MAE}}$  to our empirical risk estimator Eq. (7) as baselines. The proposed GDF loss is inserted into Eq. (7).

**Setup.** We adopt the linear model as the predictive model for fair comparisons and apply SGD with momentum 0.9 for optimization. We set batch-size and training epoch as 256 and 200 respectively. Weight decay is set as  $10^{-3}$  and learning rate is selected from  $\{10^{-1}, 10^{-2}, 10^{-3}\}$ , where the learning rate is multiplied by 0.1 at 100 and 150 epochs [Wu *et al.*,

2018]. We apply the same model and hyper-parameters of ours for L-UW,  $\bar{\mathcal{L}}_{BCE}$  and  $\bar{\mathcal{L}}_{MAE}$ . Ten-fold cross-validation is used to evaluate the performance of all approaches. Here, training data is only equipped with complementary labels and test data associated with the sets of relevant labels is used to verify the performance of approaches. We adopt the mean and *standard deviation* (std) of five criteria.  $\downarrow / \uparrow$  indicates that criteria are smaller/larger, the performance is better.

#### 5.2 Main Empirical Results

**Results.** Table 1 reports *one error*, *ranking loss*, and *average precision* of various approaches over eight datasets (the results of *hamming loss* and *coverage* are shown in Appendix D). Here, all datasets are processed by the first preprocessing way, where each instance is associated with one complementary label. As shown in Table 1, GDF outperforms the most approaches on eight datasets. Specifically, we improve upon the best baseline on *one error*, *ranking loss*, and *average precision* by 0.066, 0.031, and 0.099 respectively on scene and eurlex\_sm datasets. This indicates the effectiveness of our proposed approach GDF to solve the MLCLL problem.

Moreover, GDF achieves comparable results on three criteria against two MLL approaches over all datasets. For example, GDF on *average precision* are significantly superior to ML-KNN and LIFT by 0.241 and 0.229 respectively on eurlex\_dc dataset. This is because our approach is more suitable for dealing with complementary labeled data than MLL approaches. GDF achieves similar or better performance to

Table 2: Experimental results (mean $\pm$ std) on the training data adopted the second pre-processing way. The label space is original and the number of complementary labels per instance is half the original number of labels. The best performance of each dataset is shown in **boldface**, where  $\bullet/\circ$  indicates whether GDF is superior/inferior to baselines with pairwise *t*-test (at 0.05 significance level).

Methods	MLL	PML		CLL	Bounded	GDF						
Methous	ML-KNN	fpml	PML-LRS	L-UW	MAE	ODF						
Ranking Loss↓												
Corel16k	.276±.043	.286±.054	.281±.013	.293±.052•	.307±.051•	$.286 {\pm} .036$						
delicious	.235±.002•	.204±.002•	.261±.009•	.195±.002●	.201±.002●	$.186 {\pm} .002$						
eurlex_dc	.267±.004•	.263±.005•	.243±.003•	.252±.006•	.248±.008•	$.239 {\pm} .005$						
eurlex_sm	.173±.004●	.208±.007•	.170±.002•	.167±.004●	.167±.003●	$.159 {\pm} .004$						
Average Precision↑												
Corel16k	.224±.042•	.238±.045	.223±.009•	.244±.046	.245±.046	$.240 {\pm} .042$						
delicious	.205±.003•	.201±.002•	.232±.001•	.235±.003•	.234±.003•	$.299 {\pm} .004$						
eurlex_dc	$.300 {\pm} .011$	.179±.007●	.299±.003•	.278±.006●	.233±.008•	$.300 {\pm} .008$						
eurlex_sm	.407±.009•	.204±.004•	$.413 {\pm} .002$	.370±.006•	.321±.009•	$.413 {\pm} .008$						

PML approaches. Especially, the *average precision* of GDF is 0.418 higher than fpml on eurlex\_dc dataset, which shows that our approach can work well on MLCLL with the highnoisy problem compared with PML approaches.

Results of GDF shown in Table 1 outperform L-UW on all datasets, which indicates that the CLL approach can not enough handle the MLCLL problem. This is because their implementations depend on the assumption of one relevant label per instance, while the number of relevant labels for each instance is unknown in MLL. That leads to CLL approaches could not hold the relationship of multiple relevant labels and complementary labels in MLCLL. In addition, we observe that the unbounded loss BCE used to the empirical risk estimator Eq. (7) is inferior to our proposed GDF loss in most cases, which proves that GDF is indeed better than the unbounded one in MLCLL. Accordingly, experimental results of the empirical risk estimator Eq. (7) with MAE loss on three criteria are lower than ours in all datasets, which further proves the effectiveness of our proposed strategy - minimizing the empirical risk estimator by the GDF loss.

Effect of the gradient descent friendly loss GDF. Fig. 2 shows average precision of BCE loss, MAE loss and GDF loss on various datasets over 200 epochs, where six datasets both adopt the first pre-processing way to process and each instance is associated with one complementary label. As can be seen from Fig. 2, the curve of MAE loss is lower than that of BCE loss, but MAE loss is more stable than BCE loss in Fig. 2. It clearly illustrates the advantages and disadvantages of MAE loss: the bounded loss MAE is more robust for noisy data than unbounded loss BCE, while its convergence rate is inferior to the unbounded loss [Ghosh et al., 2017]. In addition, we observe that the curve of our proposed GDF loss is higher than these of the unbounded loss BCE and the bounded loss MAE, which demonstrates the effectiveness of our approach. Specifically, the curve of GDF loss shows that our approach has both the robustness of the bounded loss MAE and the superior convergence of the unbounded loss BCE. This demonstrates that removing the easily caused overfitting part of BCE loss to design GDF loss is a reasonable strategy in MLCLL to improve performance of our approach, which also confirms that GDF loss to optimize in MLCLL brings benefits to gradient update.

### 5.3 Additional Experiments

Table 2 shows ranking loss and average precision of different approaches on four datasets that are adopted the second preprocessing way to process, the remaining experimentations of three criteria are shown in Appendix D. Here, each dataset keeps the original number of labels and complementary labels given per instance is half of L(S) (the number of labels). In Table 2, the proposed approach achieves comparable performance to excellent baselines on four datasets, which indicates that GDF loss can also work on datasets with the original label space. Specifically, results of fpml and PML-LRS are inferior to GDF loss, which indicates that PML approaches depending on the assumption of sparse noisy labels can not deal with the high-noisy label problem of MLCLL. Table 2 illustrates that GDF loss outperforms L-UW, which proves that the proposal is better than the CLL approach in tackling the relationship between multiple relevant labels and complementary labels. GDF loss obtains better performance compared with MAE loss, which indicates that the proposed loss function designed by improving gradient updates can also deal with the multiple complementary-label problem.

# 6 Conclusion

In this paper, we study the setting of MLCLL, which aims to learn a multi-labeled classifier from complementary labeled data. To solve this problem, we propose an unbiased risk estimator with an estimation error bound, which supports that the learned risk minimizer from complementary labeled data converges to the optimal one of fully supervised MLL. Although our unbiased risk estimator has no restrictions on loss functions, it will produce unbounded gradients if certain unbounded loss functions are used and result in overfitting. To alleviate this issue, we design GDF loss to prevent the overfitting problem and find that it brings benefits to gradient updates. We verify the effectiveness of the proposed upper bound loss on various datasets.

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