# Self-Training: A Survey

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#### **Abstract**

Semi-supervised algorithms aim to learn prediction functions from a small set of labeled observations and a large set of unlabeled observations. Because this framework is relevant in many applications, they have received a lot of interest in both academia and industry. Among the existing techniques, self-training methods have undoubtedly attracted greater attention in recent years. These models are designed to find the decision boundary on low density regions without making additional assumptions about the data distribution, and use the unsigned output score of a learned classifier, or its margin, as an indicator of confidence. The working principle of self-training algorithms is to learn a classifier iteratively by assigning pseudo-labels to the set of unlabeled training samples with a margin greater than a certain threshold. The pseudo-labeled examples are then used to enrich the labeled training data and to train a new classifier in conjunction with the labeled training set. In this paper, we present self-training methods for binary and multiclass classification as well as their variants and two related approaches, namely consistency-based approaches and transductive learning. We examine the impact of significant self-training features on various methods, using different general and image classification benchmarks, and we discuss our ideas for future research in self-training. To the best of our knowledge, this is the first thorough and complete survey on this subject.

# 1 Introduction

Semi-supervised learning has risen to prominence within the machine learning domain, tackling the core challenge of making inference from partially labeled data. The framework is particularly useful in scenarios where there are limited labeled examples but an abundance of unlabeled data available for training. This is highly relevant in a range of applications, such as image classification and segmentation, where the acquisition of labeled data can be a costly endeavor.

**Central hypothesis** The basic assumption in semi-supervised learning, called *smoothness*, stipulates that two examples in a high density region should have identical class labels [14, 2]. This means that if two points are a part of the same group or cluster, their class labels will most likely be the same. If they are separated by a low density zone, on the other hand, their desired labels should be different. Hence, if the examples of the same class form a partition, unlabeled training data might aid in determining the partition boundary more efficiently than if just labeled training examples were utilized.

**Three main semi-supervised learning families** The implementation of this assumption in various contexts results in three main families of semi-supervised learning techniques, each with its own adaption of the smoothness hypothesis.

Data clustering uses a mixture model and assigns class labels to groups using the labeled data they include; and it constitutes the working principle of generative approaches [57, 41]. The cluster assumption, which underpins these approaches, asserts that if two examples are in the same group, they are likely to be in the same class. This hypothesis may be explained as follows: if a group is formed by a large number of instances, it is rare that they belong to different classes. This does not imply that a class is constituted by a single group of examples, but rather that two examples from distinct classes are unlikely to be found in the same cluster.

If we consider the partitions of instances to be high density areas, a form of the cluster assumption known as low density separation entails determining the decision boundary over low density regions, and it constitutes the basis of discriminant techniques. The main difference between generative and discriminant techniques is that discriminant approaches find directly the prediction function without making any assumption on how the data are generated [3, 59].

Density estimation is often based on a notion of distance, which for high dimensional spaces may also become meaningless. A third hypothesis, known as the manifold assumption, stipulates that in high-dimensional spaces, instances reside on low-dimensional topological spaces that are locally Euclidean, which is supported by a variety of semi-supervised models called graphical approaches [7, 18].

Compatibility Although semi-supervised algorithms have been successfully applied in many situations, there have been cases where unlabeled data have been shown to have no effect on the performance of a learning task [76]. Several attempts have been made in recent years to investigate the value of unlabeled data in the training process [12, 49], and the capacity of semi-supervised learning approaches to generalize [68, 52]. The bulk of these studies are founded on the notion of *compatibility* defined by [5], and they strive to exhibit the connection between the marginal data distribution and the target function to be learned. According to these findings, unlabeled data will be beneficial for training only if such a relationship exists.

In generative approaches, the marginal distribution is viewed as a mixture of class conditional distributions, and when compared to the supervised case, semi-supervised learning has been shown to achieve lower finite-sample error bounds in some general cases, or a faster rate of error convergence in others [12, 68, 52, 76]. In this line, [8] showed that accessing the marginal distribution on unlabeled training data would

not provide sample size guarantees superior to those obtained by supervised learning unless very strong assumptions about conditional distribution on class labels are made.

For graph-based approaches, [58] provided a context in which such algorithms may be studied and perhaps justified; the key finding of the study is that unlabeled data can help learning in some situations by explicitly defining the manifold.

Finally, discriminant approaches mostly embed a margin maximization method that searches the decision boundary in low-density regions by pushing it from the unlabeled data [37]. In this survey we focus on self-training algorithms that follow this principle by assigning pseudo-labels to high-confidence unlabeled training examples and include these pseudo-labeled samples in the learning process.

**Paper structure** The reminder of this paper is organized as follows. In Section 2, we present the framework and notations used throughout the paper. In Section 3, we go over the self-training method in detail, covering pseudo-labeling and its variants, as well as providing some insights into current theoretical studies. Other related approaches are described in Section 4. Section 5 reviews application of self-training methods in different domains. In Section 6, we investigate the influence of important self-training features on different approaches using a variety of general and image classification benchmarks. The views and future prospects are discussed in Section 7.

### 2 Framework and notations

We consider classification problems where the input and the output spaces are respectively  $\mathcal{X} \subseteq \mathbb{R}^d$  and  $\mathcal{Y}$ . We further suppose available a set of labeled training examples  $S = (\mathbf{x}_i, y_i)_{1 \leqslant i \leqslant m} \in (\mathcal{X} \times \mathcal{Y})^m$  generated from a joint probability distribution  $\mathbb{P}(\mathbf{x}, y)$  (denoted as  $\mathcal{D}$ ) and a set of unlabeled training examples  $X_{\mathcal{U}} = (\mathbf{x}_i)_{m+1 \leqslant i \leqslant m+u} \in \mathcal{X}^u$  supposed to be drawn from the marginal distribution  $\mathbb{P}(\mathbf{x})$ .

The classic case corresponds to when  $m \ll u$ , and the issue is thrown into the unsupervised learning framework if S is empty. The opposite extreme scenario is when  $X_{\mathcal{U}}$  is empty and the problem is reduced to supervised learning. Given a hypothesis set of functions  $\mathcal{H}$  mapping  $\mathcal{X}$  to  $\mathcal{Y}$ , the learner receives a labeled set S and an unlabeled set  $X_{\mathcal{U}}$  and outputs a hypothesis  $h \in \mathcal{H}$  which is assumed to have a generalization error  $R(h) = \mathbb{E}_{(\mathbf{x},y) \sim \mathcal{D}}[\mathbb{1}_{h(\mathbf{x}) \neq y}]$  smaller than if just S was used to find the prediction function, where by  $\mathbb{1}_{\pi}$  we denote the indicator function equal to 1 if the predicate  $\pi$  is true and 0 otherwise.

In practice, classifiers are defined based on a scoring function f from a class of functions  $\mathcal{F} = \{f : \mathcal{X} \times \mathcal{Y} \to \mathbb{R}\}$ , and for an example  $\mathbf{x}$  the corresponding classification function h outputs the class for which the score of f is the highest:

$$h(\mathbf{x}) = \operatorname{argmax}_{y \in \mathcal{Y}} f(\mathbf{x}, y).$$

We define the margin  $\rho_f(\mathbf{x},y)$  of a function f for an example  $\mathbf{x} \in \mathcal{X}$  and a class  $y \in \mathcal{Y}$  as

$$\rho_f(\mathbf{x}, y) = f(\mathbf{x}, y) - \max_{y' \neq y} f(\mathbf{x}, y').$$

In the binary case,  $\mathcal{Y} = \{-1, +1\}$ , we define the unsigned margin of a classification function  $f \in \mathcal{F}$  over an example  $\mathbf{x}$  [19, 1] as

$$m_f(\mathbf{x}) = |\rho_f(\mathbf{x}, +1)|.$$

In the multi-class classification case,  $\mathcal{Y} = \{1, \dots, K\}$ , the unsigned margin [19, 27] is defined as

$$m_f(\mathbf{x}) = \sum_{y \in \mathcal{Y}} f(\mathbf{x}, y) \rho_f(\mathbf{x}, y).$$

The maximization of the unsigned margin tends to find a decision boundary that passes through low density regions and hence follows the low density separation assumption.

Based on this idea, self-training algorithms define a pseudo-labeling strategy for assigning pseudo-labels to the examples of  $X_{\mathcal{U}}$ . This strategy can be characterized by a function, called *pseudo-labeler*:

$$\Phi_{\ell}: \mathcal{X} \times \mathcal{F} \to \mathcal{X} \times \mathcal{Y}.$$

We denote  $\tilde{y}$  the pseudo-label of an unlabeled  $\mathbf{x} \in X_{\mathcal{U}}$  for a score function  $f \in \mathcal{F}$  assigned by  $\Phi_{\ell}$  and  $X_{\mathcal{U}}$  the set of pseudo-labeled examples.

# 3 Self-Training

Self-training, also known as decision-directed or self-taught learning machine, is one of the earliest approach in semi-supervised learning [72, 28] that has risen in popularity in recent years.

This wrapper algorithm starts by learning a supervised classifier on the labeled training set S. Then, at each iteration, the current classifier selects a part of the unlabeled data,  $X_{\mathcal{U}}$ , and assigns pseudo-labels to them using the classifier's predictions. These pseudo-labeled unlabeled examples are removed from  $X_{\mathcal{U}}$  and a new supervised classifier is trained over  $S \cup X_{\mathcal{U}}$ , by considering these pseudo-labeled unlabeled data as additional labeled examples. To do so, the classifier  $h \in \mathcal{H}$  that is learned at the current iteration is the one that minimizes a regularized empirical loss over S and  $X_{\mathcal{U}}$ :

$$\frac{1}{m} \sum_{(\mathbf{x}, y) \in S} \ell(h(\mathbf{x}), y) + \frac{\gamma}{|X_{\mathcal{U}}|} \sum_{(\mathbf{x}, \tilde{y}) \in X_{\mathcal{U}}} \ell(h(\mathbf{x}), \tilde{y}) + \lambda ||h||^2$$

where  $\ell:\mathcal{Y}\times\mathcal{Y}\to[0,1]$  is an instantaneous loss most often chosen to be the crossentropy loss,  $\gamma$  is a hyperparameter for controlling the impact of pseudo-labeled data in learning, and  $\lambda$  is the regularization hyperparameter. This process of pseudo-labeling and learning a new classifier continues until the unlabeled set  $X_{\mathcal{U}}$  is empty or there is no more unlabeled data to pseudo-label. The pseudo-code of the self-training algorithm is shown in Algorithm 1.

## 3.1 Pseudo-labeling strategies

At each iteration, the self-training selects just a portion of unlabeled data for pseudolabeling, otherwise, all unlabeled examples would be pseudo-labeled after the first iteration, which would actually result in a classifier with performance identical to the initial classifier [14]. Thus, the implementation of self-training arises the following question: how to determine the subset of examples to pseudo-label?

A classical assumption, that stems from the low density separation hypothesis, is to suppose that the classifier learned at each step makes the majority of its mistakes on observations close to the decision boundary. In the case of binary classification, preliminary research suggested to assign pseudo-labels only to unlabeled observations for which the current classifier is the most confident [79]. Hence, considering thresholds  $\theta^-$  and  $\theta^+$  defined for respectively the negative and the positive classes, the pseudo-labeler assigns a pseudo-label  $\tilde{y}$  to an instance  $\mathbf{x} \in X_{\mathcal{U}}$  such that:

$$\tilde{y} = \begin{cases} +1, & \text{if } f(\mathbf{x}, +1) \geqslant \theta^+, \\ -1, & \text{if } f(\mathbf{x}, -1) \leqslant \theta^-, \end{cases}$$
 (1)

and  $\Phi_{\ell}(\mathbf{x}, f) = (\mathbf{x}, \tilde{y})$ . An unlabeled example  $\mathbf{x}$  that does not satisfy the conditions (1) is not pseudo-labeled; i.e.  $\Phi_{\ell}(\mathbf{x}, f) = \emptyset$ .

Intuitively, thresholds should be set to high absolute values as pseudo-labeling examples with low confidence would increase chances of assigning wrong labels. However, thresholds of very high value imply excessive trust in the confidence measure underlying the model, which, in reality, can be biased due to the small labeled sample size. Using several iterations makes also the situation more intricate as at every iteration the optimal threshold might be different.

One way to select the thresholds is to set them equal to the average of respectively positive and negative predictions [79]. In this line, and in the context of multi-class classification [45] used Neural Networks as the supervised classifier and chose the most confident class to infer pseudo-labels for unlabeled data using the current model' outputs. The pseudo-labeled examples were then added to the labeled training set and treated similarly as labeled examples.

[97] adapted the idea of [79] for multi-class classification by not choosing thresholds but rather fixing a proportion p of the most confident unlabeled data to be pseudo-labeled and then increasing this proportion at each iteration of the algorithm until p=0.5 was reached.

Following this idea, [11, 93] proposed an adaptation of curriculum learning to pseudo-labeling, which entails in learning a model using easy-to-learn observations before moving on to more complex ones. The principle is that at the step k of the algorithm, the pseudo-labeler selects unlabeled examples having predictions that are in the  $(1 - \alpha_k)^{th}$  percentile of the distribution of the maximum probability predictions assumed to follow a Pareto distribution, and where  $\alpha_k \in [0,1]$  is an hyperparameter that varies from 0 to 1 in increments of 0.2

Considering the distribution of predictions over unlabeled data, and the majority vote classifiers, such as Random Forest or Adaboost [70], it is possible to automatically choose a threshold for pseudo-labeling. Formally, the learning of a majority vote

#### Algorithm 1. Self-Training

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\begin{aligned} & \textbf{Input:} \ S = (\mathbf{x}_i, y_i)_{1 \leqslant i \leqslant m}, X_{\mathcal{U}} = (\mathbf{x}_i)_{m+1 \leqslant i \leqslant m+u}. \\ & k \leftarrow 0, X_{\mathcal{U}} \leftarrow \emptyset. \\ & \textbf{repeat} \\ & \text{Train } f^{(k)} \text{ on } S \cup X_{\mathcal{U}} \\ & \Pi_k \leftarrow \{\Phi_\ell(\mathbf{x}, f^{(k)}), \mathbf{x} \in X_{\mathcal{U}}\} \quad \triangleright \text{Pseudo-labeling} \\ & X_{\mathcal{U}} \leftarrow X_{\mathcal{U}} \cup \Pi_k \\ & X_{\mathcal{U}} \leftarrow X_{\mathcal{U}} \setminus \{\mathbf{x} \mid (\mathbf{x}, \tilde{y}) \in \Pi_k\} \\ & k \leftarrow k+1 \\ & \textbf{until } X_{\mathcal{U}} = \emptyset \vee \Pi_k = \emptyset \\ & \textbf{Output:} \ f^{(k)}, X_{\mathcal{U}}, X_{\mathcal{U}} \end{aligned}
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classifier with partially labeled data can be defined as follows. After observing the training set  $S \cup X_{\mathcal{U}}$ , the task of the learner is to choose a posterior distribution Q over a set of hypothesis  $\mathcal{H}$  such that the Q-weighted majority vote classifier  $B_Q$  defined by:

$$\forall \mathbf{x} \in \mathcal{X}, B_Q(\mathbf{x}) = \operatorname{argmax}_{y \in \mathcal{Y}} \mathbb{E}_{h \sim Q} \left[ \mathbb{1}_{h(\mathbf{x}) = y} \right],$$

will have the smallest possible risk on examples of  $X_{\mathcal{U}}$ . The associated *Gibbs* classifier,  $G_Q$ , is defined as the random choice of a classifier h according to the distribution Q, and its error over an unlabeled set  $X_{\mathcal{U}}$  is given by:

$$\hat{R}_u(G_Q) = \frac{1}{u} \sum_{\mathbf{x}' \in X_{\mathcal{U}}} \mathbb{E}_{h \sim Q}[\mathbb{1}_{h(\mathbf{x}') \neq y'}],$$

where, for every unlabeled example  $\mathbf{x}' \in X_{\mathcal{U}}$  we refer to y' as its true unknown class label. For binary and multi-class classification respectively, [1, 27] showed that a tight upper-bound on the Gibbs classifier's risk that holds with high probability over the random choice of S and  $X_{\mathcal{U}}$ , guarantees a tight bound on the error of the Bayes classifier over the unlabeled training set:

$$\hat{R}_u(B_Q) = \frac{1}{u} \sum_{\mathbf{x}' \in X_M} \mathbb{1}_{B_Q(\mathbf{x}') \neq y'}.$$

This bound is mainly based on the distribution of predictions over unlabeled data and the derivations can be extended to bound the risk of voted majority classifiers having margins greater than a threshold  $\theta$ ,  $\hat{R}_{u \wedge \theta}(B_Q)$ , defined as:

$$\hat{R}_{u \wedge \theta}(B_Q) = \frac{1}{u} \sum_{\mathbf{x}' \in X_U} \mathbb{1}_{B_Q(\mathbf{x}') \neq y' \wedge m_{B_Q}(\mathbf{x}') > \theta},$$

with a slight abuse of notation for  $m_{B_Q}$ . One of the practical aspects that arises from this result is the possibility to specify a threshold  $\theta$  which minimizes an upper-bound

of the conditional risk of a voted majority classifier  $B_Q$  over the unlabeled training set,  $X_U$ , defined as:

$$\hat{R}_{u|\theta}(B_Q) = \frac{\hat{R}_{u \wedge \theta}(B_Q)}{\frac{1}{u} \sum_{\mathbf{x} \in X_U} \mathbb{1}_{m_{B_Q}(\mathbf{x}) \geqslant \theta}},$$

where the denominator is the proportion of the unlabeled examples with the confidence higher than the threshold  $\theta$ , and the numerator is the joint Bayes risk on  $X_{\mathcal{U}}$ . Thus, the criterion can be interpreted as a trade-off between the number of examples going to be pseudo-labeled and the error they induce. Furthermore, these bounds are shown to be tight in the case where the majority vote classifier makes its error mostly on low margin regions. [27] demonstrated that this technique outperforms conventional fixed-threshold pseudo-labeling strategies on different multi-class classification problems. [15] highlighted two major issues with self-training: the snowball effects of cascading pseudo-labeling mistakes and random sampling of tiny samples (called data bias). The authors suggest two-phase solutions to address these problems for image classification using deep learning. First, they proposed a classification head to separate the creation and use of pseudo-labels in order to reduce training errors. An additional head is utilized to receive the pseudo-labels and carry out training on unlabeled data while the default head is used for classification and pseudo-labeling.

#### 3.2 Self-training with two classifiers

In the wake of works utilizing only a single classifier in self-training algorithms, new studies have been proposed with the use of two classifiers, where each model learns on the output of the other [86, 16, 40]. Most of these techniques are based on the idea of consensus in predictions between two classifiers and were inspired by the seminal work of [9] who proposed the co-training algorithm.

In co-training, examples are defined by two modalities that are comparable but not entirely correlated. Each view of an example is expected to contain complementary information about the data and if there are enough labeled training data, each of them is supposed to be sufficient for learning. The main principle is to learn a classifier on each view, taking initially the available labeled examples as the training set. Then, one of the classifiers assigns pseudo-labels to unlabeled data, and the other one uses them to retrain the model by including them into its training set. At each iteration, the classifiers alternately switch their roles, thereby co-training each other. As for self-training algorithms with a single classifier, this procedure continues until there are no more unlabeled instances to be pseudo-labeled. In practice, several studies artificially generated the two modalities for classification problems where examples are *monoviewed* and described by a vector representation. These approaches create the two modalities out of one by selecting at random the set of features that should correspond to each modality from the initial set of features; and their efficiency was empirically proved on various applications [88].

Co-training can thus be thought of as a form of self-training: rather than training a classifier on its own previous predictions, each classifier is trained on the predictions of another classifier that was trained on the predictions of the former. Without splitting the input feature set, [16] proposed Cross Pseudo Supervision for semantic segmentation

	Base classifier		Clas	Classification		Threshold	
	Single	Double	Binary	Multi-class	Fixed	Optimized	Account
Scudder [[72]]	<b>√</b>	_	✓	_	<b>√</b>	_	_
Blum et al. [[9]]	_	✓	✓	_	✓	_	✓
Joachims. [[37]]	✓	_	✓	_	✓	_	_
Tur et al. [[79]]	✓	_	_	✓	✓	_	_
Amini et al. [[1]]	✓	_	✓	_	_	✓	_
MeanTeacher [[78]]	_	✓	_	✓	✓	_	_
Feofanov et al. [[27]]	✓	_	_	✓	_	✓	_
Xie et al. [[85]]	✓	_	_	✓	✓	_	_
Chen et al. [[16]]	_	✓	_	✓	✓	_	_
Cascante et al. [[11]]	✓	_	_	✓	✓	_	_
Zhang et al. [[93]]	✓	_	_	✓	_	✓	_
Du et al. [[22]]	_	✓	_	✓	✓	_	_
Chen et al. [[15]]	$\checkmark$	_	_	✓	✓	_	✓
Hadjadj et al. [[33]]	✓	_	✓		_	✓	✓

Table 1: A summary of principal self-training algorithms presented in this survey.

in images. This method employs two neural networks as supervised classifiers having the same images as inputs. Each neural-network is learned at every mini-batch by considering the pseudo-labels generated by the other network for unlabeled instances as ground-truths.

The learnability of co-training was studied under the PAC framework [80], which also accounts for noise in the class labels of unlabeled examples caused by pseudo-labeling. The injection of noisy labels in the pseudo-labeling step is in fact inherent to any self-training algorithm. Taking into account noisy labels in training a model was first considered in supervised learning, following the paradigm of learning with an imperfect supervisor in which training data contains an unknown portion of imperfect labels [55, 34]. Most of these studies tackle this problem from an algorithmic point of view, employing regularization [54] or estimating mislabeling errors by modeling the transition probability between noisy and true labels [61].

Table 1 summarizes the main self-training approaches presented in this survey by emphasizing their key aspects.

#### 3.3 Theoretical studies

Several studies have recently looked into the theoretical properties of self-training algorithms.

In this line, [83] suggest a new concept of *expansion* defined as the quantity of data dispersion in an example's neighbor, where the term *neighbor* refers to adding adversarial perturbations [54] or augmentations [77] to the example. The study establishes distributional guarantees of self-training when the label distribution meets such expansion properties and classes are suitably separated according to neighbors. The study generates finite sample bounds for Deep Neural Networks (DNNs) by combining generalization bounds with DNN generalization bounds. Experiments with a Generative Adversarial Network (GAN) are also used to verify the expansion assumption.

[29] examine a self-training algorithm with linear models for the binary classification using gradient-based optimization of the cross-entropy loss after supervised

learning with a small number of samples. The classifier is a mixture model with concentration and anti-concentration properties. The authors show that utilizing  $O(d/\epsilon^2)$  unlabeled observations in the self learning algorithm, with d the number of input variables, suffices to achieve the classification error of the Bayes-optimal classifier up to an  $\epsilon$  error if the initial pseudo-labeling strategy has a classification error smaller than an absolute constant  $C_{err}$ . Furthermore, the authors demonstrate that a constant number of labeled examples is sufficient for optimal performance in a self-training algorithm by demonstrating that using only O(d) labeled examples, the standard gradient descent algorithm can learn a pseudo-labeling strategy with a classification error no more than  $C_{err}$ .

[94] study the generalization ability of self-training in the case where the base classifier is a two-layer neural network with the second layer weights all fixed to one, and assuming that the ground truth is realisable, the labels are observed without noise, and the labeled and unlabeled instances are drawn from two isotropic Gaussian distributions. The authors show that, given some plausible assumptions about the initial point and the amount of unlabeled training examples, the algorithm converges to the ground truth with fewer observations than needed when no unlabeled data is provided. The reader can refer [95] for a broader context. [94] extend their main result to a more general setting, where it is shown that the model still converges towards a given convex combination of the ground truth and the initial point, and is guaranteed to outperform the initial supervised model, without fixing any requirement on the number of labeled training examples.

[33] propose a first bound over the misclassification error of a self-training method which utilizes half-spaces as the base classifier in the case where class labels of examples are supposed to be corrupted by a Massart noise model. Under this assumption, it is shown that the use of unlabeled data in the proposed self-training algorithm does not degrade the performance of the first half-space trained over the labeled training data.

# 4 (Un)related approaches

In semi-supervised learning, there are probably two main other areas of research that are related to self-training. The first, referred to as *consistency learning*, uses classifier predictions over unlabeled data as a confidence indicator and constrains model outputs to be comparable for similar unlabeled examples without assigning pseudo-labels. The second method, known as *transductive learning*, is based on the low density separation assumption and tends to give class labels for only the set of unlabeled training samples. In this Section, we will also briefly discuss *self-supervised learning*, which, despite its similar name with self-training, is an entirely separate technique that employs unlabeled data to train or pre-train a model.

## 4.1 Consistency-based approaches

Early studies in this line, see for example [96] for binary classification, were proposed to learn a single classifier defined from a scoring function  $f: \mathcal{X} \times \mathcal{Y} \to \mathbb{R}$  penalized for quick changes in its predictions. The similarity matrix  $\mathbf{W} = [W_{ij}]_{\substack{1 \le i \le u \\ 1 \le j \le u}}$ 

over the unlabeled training data, is used to measure the similarity between instances. The penalization is mostly expressed as a regularization term in the learning objective. As an example, adapting the work of [96] to multi-class classification, the penalization term can be written as:

$$\Omega_{\mathbf{W}}(X_{\mathcal{U}}) = \sum_{i,j=1}^{u} W_{ij} \| f(\mathbf{x}_{m+i},.) - f(\mathbf{x}_{m+j},.) \|^{2}$$

where for a given example  $\mathbf{x}$ ,  $f(\mathbf{x}, \cdot) = (f(\mathbf{x}, k))_{k \in \mathcal{Y}}$  is the vector class predictions of f. In terms of learning,  $\Omega_{\mathbf{W}}$  can be seen as a regularization term, constraining the model to have the same predictions on similar unlabeled instances.

Other types of penalization have been studied in the literature. [52] suggested an approach that partitions partially labeled data and then uses labeled training samples to identify dense clusters having predominant classes with a fraction of non-predominant classes below a given threshold extending earlier results on supervised classification [38]. In this situation, the proposed penalization term measures the learner's inability to predict the predominant classes of the identified clusters which in turn constrains the supervised classifier to be consistent with the structure of the dense clusters.

In this line, [66] consider non-decomposable metrics with consistency regularization by giving a cost-sensitive framework that consists of minimizing a cost-sensitive error on pseudo labels and consistency regularization. They demonstrate theoretically that they can build classifiers that can maximize the required non-decomposable measure more effectively than the original model used to produce pseudo-labels under comparable data distribution assumptions.

Without explicitly stating a penalization term, consistency learning was extended to cases with two classifiers. The Mean-Teacher approach [78] is perhaps one of the earliest popular techniques that have been proposed in this context. This method employs Neural Networks (NNs) as supervised classifiers, and it is based on the assumption that two close models with the same input should make the same prediction. One of the models is called the *teacher*, while the other is referred to as the *student*. These two NN models are structurally identical, and their weights are related in that the teacher's weights are an exponential moving average [44] of the student' weights. In this scenario, the student model is the only one that is trained over the labeled training set, and the consistency loss is computed between the teacher's probability distribution prediction and the student's one using the mean square error or the Kullback-Leibler divergence.

Other studies refined the Mean-Teacher approach with a data-augmentation technique by combining two images with random patches to improve prediction consistency [30, 85]. More recently, [22] provide a two-stage method to reduce label propagation errors; where in the first phase, the gradients of the student loss are computed and utilized to update the teacher. In the second stage, the teacher assigns pseudo-labels which are then utilized to train the current student.

## 4.2 Transductive learning

The goal of transductive learning, as previously stated, is to assign pseudolabels to samples from an unlabeled training set,  $X_{\mathcal{U}}$ . As this set is finite, the considered function class  $\mathcal{F}$ , for finding the transductive prediction function, is also finite.  $\mathcal{F}$  can be defined using a nested structure according to the structural risk minimization principle,  $\mathcal{F}_1 \subseteq \mathcal{F}_2 \subseteq \ldots \subseteq \mathcal{F}$  [81]. Transductive techniques often employ the distribution of unsigned margins of unlabeled examples to guide the search for a prediction function, limiting it to following the low density separation assumption in order to find the best function class among the current ones.

Transductive approaches also assume that the function class's structure should reflect prior knowledge of the learning problem at hand, and that it should be built in such a way that the correct prediction of class labels of labeled and unlabeled training examples is contained in a function class  $\mathcal{F}_j$  of small size with a high probability. In particular, [20] show that with high probability the error on the unlabeled training set of a function from a class function  $\mathcal{F}_j$  is bounded by its empirical loss on the labeled training set plus a complexity term that depends on the number of labeled examples m, the number of unlabeled examples u, and the size of the class function  $\mathcal{F}_j$ .

The Transductive Support Vector Machines (TSVM) [37] developed for the binary case is based on this paradigm, and is looking for the optimal hyperplane in a feature space that separates the best labeled examples while avoiding high density areas. TSVM does this by building a structure on a function class  $\mathcal{F}$  and sorting the outputs of unlabeled samples by their margins. The solutions to the associated optimization problem are the pseudo-labels of unlabeled examples for which the resulting hyperplane separates the examples of both labeled and unlabeled training sets with the largest margin.

[74] extended this idea to the multi-class classification case with Neural Networks. Similar to TSVM, class labels of unlabeled examples are treated as variables, and the algorithm tries to determine their optimal values, along with the optimal NNs parameter set get by minimizing a cross-entropy loss estimated over both labeled and unlabeled training sets through an iterative training process. The authors employ the MinMax Feature regularization to constrain the neural network to learn features of same-class images to be close, and features of different classes to be separated by a preset margin, in order to overcome incorrect label estimations on outliers and noisy samples.

## 4.3 Self-supervised Learning

Although similar in names, self-training is a completely different approach than self-supervised learning which has demonstrated encouraging results and has become an active area of research [60].

In self-supervised learning, a model acquires the ability to make predictions regarding various facets of its input data, all without the necessity of explicit labeled training data. Rather than depending on labeled data, self-supervised learning harnesses the inherent structure present in the input data and autonomously generates guidance to train the model. This procedure involves the formulation of a pretext task, also referred to as a proxy task, wherein the model is trained to make predictions concerning a specific

relationship inherent in the data. For instance, in the domain of computer vision, a pretext task might involve rotating images within a predefined range of angles, followed by training a supervised model to predict these angles.

Once the model has undergone training on the pretext task, the knowledge it has gained in this process can be applied to downstream tasks that do require labeled data. Consequently, by learning from extensive amounts of unlabeled data, self-supervised learning empowers the acquisition of robust data representations, capitalizing on the abundant, freely available unlabeled data resources.

Common approaches in self-supervised learning include predicting missing parts of an image [46], predicting the order of shuffled image patches or their orientation [75], reconstructing corrupted images [25], filling in missing words in a sentence [21], or predicting future frames in a video sequence [71]. These pretext tasks encourage the model to capture meaningful representations of the input data, which can then be used for various downstream tasks, such as image classification, object detection, or natural language processing.

# 5 Applications

In this section, we will concentrate on the most popular applications where self-training was employed, although this technique may be extended and used to a variety of additional machine learning tasks. The goal of our presentation here is not to be thorough, but rather to focus on the main features of self-training that were used in the literature among the selected applications.

#### 5.1 Natural Language Processing

Co-training is perhaps one of the preliminary self-training techniques which was applied to web pages classification [9]. In the paper, the content of a web page has been divided into two sets of words: those that appear on the page and those that appear in hyperlinks pointing to the page. The main hypothesis here is that each of the set of words contain sufficient information for the classification task and that there are enough labeled data to efficiently learn two supervised classifiers. Both theoretical and empirical studies of co-training show that if examples have two redundant but not entirely correlated views, then unlabeled data may be used to augment the original labeled training data to find more robust classifiers. However, the drawback of this strategy is that in general, text data is mono-view. For bag-of-word representation of texts, a solution was to split the set of words in two random sets, considered as two distinct views of a text [56], as mentioned in Section 3.2. However, this idea cannot be generalized to sequential models that could be used as base classifiers in co-training.

Other current self-training techniques in NLP are mostly built on the concept of co-training and employ two base classifiers that are learned over each other's predictions. In this line, [84] proposed a Named Entity Recognition (NER) strategy that consists in automatically detecting and classifying named entities, with a first NER model trained on labeled training data serving as a teacher to predict the probability distribution of entity labels for each token in the unlabeled set. The pseudo-labeled

data with such soft labels are then used to train a student NER model for the unlabeled set and the process of pseudo-labeling and training is repeated until convergence as in co-training. For the task of Relation Extraction (RE) which consists in obtaining a predefined semantic relation between two entities in a given sentence, [89] proposed an approach which classifies the pseudo-labeled instances generated from a teacher into confident, ambiguous and hard sets. In the training of the student model, the confident and ambiguous instances are subsequently interpreted as positive and set-negatives observations, respectively.

Lately, [53] proposed an approach to leverage the power of language models that have been pre-trained on large corpora of text to generate pseudo-labels for unlabeled text data. The pseudo-labeled data along with a smaller set of labeled data are then used to train and fine-tune the text classifier, and the process of assigning pseudo-labels and retraining of the classifier is repeated until convergence. A challenge that arises when using a single base classifier in self-training for NLP tasks is to minimize the impact of label noise in the pseudo-labeling policy. To cope with this problem, [91] devised a bootstrapping technique for semantic role labeling that consists randomly selecting a subset of the most confident samples for pseudo-labeling. In the same vein, for the sentiment analysis task, [32] advocated selecting the top most confident samples for pseudo-labeling. However, as we shall see in the next section, the use of a fixed threshold in the pseudo-labeling policy may be suboptimal in general.

### **5.2** Computer Vision

As in NLP, the two variants of self-training with one or two classifiers, mainly referred to as student and teacher in the literature, are mainly considered for image classification. Most recent approaches use neural networks as base classifiers and rely on these models' ability to learn efficient representations of images, proposing various strategies to either improve the representation or reduce the effect of noise injection during the pseudo-labeling phase of self-training.

The most common strategy with student and teacher base classifiers is arguably the one proposed by [86], in which an EfficientNet model trained on labeled ImageNet images is used as a teacher to create pseudo labels on unlabeled ones. A larger EfficientNet is subsequently employed as a student model, being trained on a mix of labeled and pseudo-labeled images. This training involves altering the input images using various techniques like dropout, stochastic depth, and data augmentation. The objective is for the model to learn a representation of images that remains consistent despite these alterations. This procedure is repeated by reversing the roles of the student and the teacher. The input of the teacher model is not altered throughout the training process. The main motivation advanced is to ensure that the pseudo labels be as accurate as possible. Empirical evidence from various image collections demonstrates the effectiveness of this strategy.

[77] proposed a self-training approach called FixMatch that combines consistency regularization with a confidence-based mechanism to select high-confidence pseudo-labeled examples for training. The algorithm applies to the same image two different data augmentations procedures, called weak (flip-and-shift) and strong (more heavy distortions) augmentations. As in the previous case, these perturbations helps to in-

crease diversity and improve the model's robustness on the unlabeled images. The authors introduce a consistency loss term that encourages the consistency between the model's hard output of the weakly-augmented version and the model's soft output of the strongly-augmented version of the same unlabeled image. They demonstrate that the model learns to provide more trustworthy and accurate results by minimizing the discrepancy between these predictions. In order to decrease the influence of possibly inaccurate pseudo-labels on the learning process, the loss is evaluated only on those unlabeled data from the batch that have the confidence higher than a fixed threshold.

This idea has then been adapted to various correlated tasks, including object detection, image segmentation [17], remote sensing [36] and video anomaly detection [51], among others. [15] proposed an improvement of FixMatch by introducing two novel features. First of all, they introduce a separate classification head that is used to assign pseudo-labels and trained using labeled data only in order to avoid possible label noise from wrong pseudo-labels. Secondly, they improve the feature learning by introducing an adversarial classification head whose goal is to approximate the worst possible error on unlabeled data. All these approaches employ a constant predefined threshold across all classes to choose unlabeled data for training, disregarding varying learning conditions and complexities among different classes.

To tackle this concern, [93] introduced a curriculum learning technique to utilize unlabeled data based on the model's learning progress. The essence of this strategy involves dynamically adapting thresholds for distinct classes during each time step, enabling the inclusion of insightful unlabeled data and their corresponding pseudolabels. This approach has been successfully applied to many domains, including object detection [47], medical image classification [63], human action recognition [82] and facial expression identification [73].

## **5.3** Knowledge-driven applications

Through the incorporation of domain expertise, recent studies have developed more sophisticated self-training systems that reduce label noise in the pseudo-labeling phase across diverse applications. In the subsequent sections, we will consider advances made in this context in the domains of speech recognition, anomaly detection, genomics and proteomics.

#### 5.3.1 Speech Recognition

Newly developed methods have introduced filtering mechanisms that are congruent with domain knowledge for end-to-end speech recognition. These mechanisms establish rules that assess pseudo-labels using criteria specific to the domain. For example by using filters to verify if certain phonetic patterns that are common in the domain, are present in the pseudo-labels [31]. Similar techniques incorporate phonetic information relevant to the domain to validate pseudo-labels. In these approaches, incorrectly labeled examples that violate phonetic constraints are discarded from training the model [50].

Other approaches integrate domain-specific language models in the the pseudolabel generation process in order to ensure that the generated labels adhere to the linguistic nuances and terminologies of the domain. In this line, [39] introduced a self-training approach, with one base classifier combined with a language model for pseudo-labeling. Their approach involves implementing tailored filtering methods designed to address common errors arising from sequence-to-sequence models, alongside an inventive ensemble technique for enhancing the breadth of pseudo-label variations. Building upon this idea, [87] showcased that the synergy between self-training and unsupervised pre-training using wav2vec 2.0 [4] offers mutual benefits across diverse scenarios involving labeled and unlabeled data.

As in image classification, alternative methods for speech recognition apply dataaugmentation techniques, tailored to the unique aspects of the domain, to enhance the robustness of the model's predictions and consequently the quality of pseudo-labels. In this sense, [6] employed a text-to-speech system to generate audio training data from text-only sources.

#### 5.3.2 Anomaly Detection

Leveraging domain knowledge to mitigate label noise in pseudo-labels within self-training approaches has also been considered in anomaly detection. In this case, the understanding of the anomaly patterns and characteristics specific to the domain are incorporated in the model. In this regard, [48] identified common anomaly types, their features and potential sources of noise and [65] performed time domain analysis. Also, [26] created features that capture domain-specific information for video anomaly detection. It was demonstrated that these features highlight the crucial elements for anomaly detection in videos, leading to their utilization for enhancing the pseudo-labeling phase within self-training. Alternate strategies focus on simulating anomalies within the unlabeled dataset using domain knowledge. This aids the model in learning from a broad spectrum of anomalies, mitigating the potential of becoming overly specialized in a particular anomaly type [64].

#### 5.3.3 Genomics and proteomics

Furthermore, datasets in the field of genomics and proteomics encompass a variety of characteristics including gene expression levels, epigenetic markers, and genetic variants. These characteristics have been shown to increase the effectiveness of features used in self-training approaches, together with the selection of important features and their physiologically coherent transformation.

[10] incorporated biological context into feature engineering that integrate unsupervised modeling of datasets relating to human disease with the supervised component that concentrated on training with mouse data. In this context, [67] amalgamated expression data from three distinct humanized mouse models that were subjected to live attenuated yellow fever vaccine challenges in self-training with different base classifiers. The results of this study show that self-training coupled with NRG-HIS/Fluc mice exhibited the most favorable outcomes across the tested human cohorts.

Additionally, [24] employed self-training in conjunction with bioinformatic tools in silico to anticipate secreted and protective proteins. This was done to eliminate

pseudo-label errors from the identified P. falciparum SEPs obtained through proteomics experiments and to anticipate new SEPs within the P. falciparum proteome.

[35] applied domain-specific quality control steps to clean and pre-process the data. This included filtering out low-quality samples, normalizing data to account for technical biases, and addressing batch effects that can introduce noise. By doing so, they ensured that the unlabeled data that is feed into the self-training pipeline is as accurate as possible. [13] utilized reference databases and annotation resources related to genomics. These resources provide information about genes, functional elements, pathways, and biological processes. Incorporating this information into the pseudolabeling process has been shown to lead to more accurate predictions by aligning them with known biological knowledge. [90] applied network analysis techniques to identify interactions between genes and proteins. The authors demonstrated that Pathway enrichment analysis can help identify genes that are functionally related and likely to be co-regulated. This information has been shown to guide the self-training process to produce more coherent and biologically plausible pseudo-labels.

# 6 Empirical Study

Within this section, we will evaluate the effectiveness and performance of the self-training algorithm. This assessment will be based on various key features presented in the preceding sections, and it will be conducted across multiple benchmark scenarios. Our primary focus will be on scenarios characterized by severely limited labeled training data, where the utilization of complex baseline classifiers like deep learning models is unfeasible. Additionally, we will address the prevalent scenario where there are sufficient labeled training data, enabling the development of an initial supervised complex model.

The impact of threshold selection. We first study the effect of selecting automatically the threshold for pseudo-labeling on 9 publicly available data sets proposed for semi-supervised learning [23]. The characteristics of these datasets are presented in Table 2. It is worth noting that certain datasets contain only a limited number of labeled training examples, comprising just a few hundred instances and accounting for less than 1% of the total training examples. This condition underscores the suitability of employing complex base classifiers.

In the experimentation, Random Forest was employed instead using the scikit-learn implementation [62] with 200 trees of maximum depth while leaving other parameters at their default values. The primary objective was to assess and contrast the classifier's performance in two scenarios: the supervised scenario (denoted by RF) and the self-training scenario where pseudo-labeling is automatically conducted following the approach introduced by  $[27]^1$  (denoted by PL\*). Additionally, we investigated the impact of setting the pseudo-labeling threshold at predefined values from the set  $\theta \in \{0.5, 0.7, 0.9\}$  (denoted by PL $_{\theta}$ ).

<sup>&</sup>lt;sup>1</sup>https://github.com/vfeofanov/trans-bounds-maj-vote

Data set	# of labeled examples $m$	# of unlabeled examples $u$	Dimension d	# of classes  K
Vowel	99	891	10	11
Protein	129	951	77	8
PageBlocks	1094	4379	10	5
Isolet	389	7408	617	26
HAR	102	10197	561	6
Pendigits	109	10883	16	10
Letter	400	19600	16	26
Fashion	175	69825	784	10
MNIST	175	69825	784	10

Table 2: Characteristics of data sets used in our experiments, d and K correspond to respectively the dimension of the input space and the number of classes.

The automatic pseudo-labeling strategy selects the threshold which minimizes the bound of the error of the Random Forest classifier over the unlabeled training samples.

Results are resumed in Table 3. Experiments are repeated 20 times by choosing randomly the labeled training examples, and  $\downarrow$  indicates that performance is statistically worse than the best result, shown in bold, according to the Wilcoxon rank-sum test.

These results suggest that the effectiveness of self-training heavily relies on the method used to determine the pseudo-labeling threshold. When the threshold is automatically determined, self-training (i.e. PL\*) can perform competitively, indicating that this approach has the potential to improve results compared to the supervised RF.

However, when a fixed threshold is applied, self-training tends to yield inferior results compared to the supervised learning approach. This suggests that an arbitrarily chosen threshold might not effectively capture the underlying patterns in the data for the pseudo-labeling process, leading to suboptimal performance.

Moreover, when the threshold is too low as for  $\theta \in \{0.5, 0.7\}$ , pseudo-labeling is likely to produce label noise and degrade the performance of self-training with respect to the supervised RF classifier in all cases. When the threshold it is too high (i.e.  $\theta = 0.9$ ), self-training becomes competitive compared to RF on Isolet and MNIST, but the quantity of pseudo-labeled unlabeled examples seems not to be sufficient to learn efficiently.

In summary, the findings emphasize the importance of a dynamic and adaptive threshold selection mechanism when implementing self-training.

**Noise Account.** We now consider the case where the initial labeled training set allows to train deep neural networks and examine the effects of taking into account noise in the pseudo-labeling process along with the dynamic selection of the threshold on CIFAR-10 and CIFAR-100 [42]. Both datasets contain 32x32 pixel RGB images belonging to respectively 10 and 100 classes; 50000 examples are used for training and 10000 samples for test.

We consider the debiased self-training approach (DST) [15] to address the presence of noise in pseudo-labeling, in conjunction with the FlexMatch method [93] for the dynamic threshold determination in pseudo-labeling. As outlined in Section 3.1, DST

Data set	RF	$PL_{\theta=0.5}$	$PL_{\theta=0.7}$	$PL_{\theta=0.9}$	PL*
Vowel	<b>.586</b> ± .028	$.489^{\downarrow} \pm .016$	$.531^{\downarrow} \pm .034$	$.576^{\downarrow} \pm .028$	<b>.586</b> ± .026
Protein	$.764^{\downarrow} \pm .032$	$.653^{\downarrow} \pm .024$	$.687^{\downarrow}\pm.036$	$.724^{\downarrow}\pm.018$	<b>.781</b> ± .034
PageBlocks	$.965 \pm .003$	.931 <sup>↓</sup> ± .003	$.964 \pm .004$	$.965 \pm .002$	<b>.966</b> ± .002
Isolet	$.854^{\downarrow} \pm .016$	$.648^{\downarrow}\pm.018$	$.7^{\downarrow}\pm.04$	$.861^{\downarrow}\pm.08$	<b>.875</b> ± .014
HAR	.851 ±.024	$.76^{\downarrow} \pm .04$	$.81^{\downarrow} \pm .041$	$.823^{\downarrow} \pm .035$	<b>.854</b> ± .026
Pendigits	$.863^{\downarrow} \pm .022$	$.825^{\downarrow} \pm .022$	$.839^{\downarrow} \pm .036$	$.845^{\downarrow} \pm .024$	<b>.884</b> ± .022
Letter	$.711 \pm .011$	$.062^{\downarrow} \pm .011$	$.651^{\downarrow} \pm .015$	.673 $^{↓}$ ± .015	<b>.717</b> ± .013
Fashion	$.718 \pm .022$	$.625^{\downarrow} \pm .014$	$.64^{\downarrow}\pm .04$	$.68^{\downarrow}\pm .014$	<b>.723</b> ± .023
MNIST	.798 <sup>↓</sup> ± .015	$.665^{\downarrow}\pm .012$	$.705^{\downarrow} \pm .055$	$.823^{\downarrow}\pm .045$	<b>.857</b> ± .013

Table 3: Classification performance using the accuracy score on 9 publicly available data set. Best results are shown in bold and the sign  $^{\downarrow}$  shows if the performance is statistically worse than the best result on the level 0.01 of significance.

involves training a dedicated head on pseudo-labeled examples, allowing the model to implicitly capture and account for noise inherent in the pseudo-labels.

For FlexMatch, we followed the same experimental protocol than [93]. In this case, Wide ResNet (WRN) [92] was used as the base classifier in self-training. Parameter learning was accomplished using stochastic gradient descent (SGD) with a momentum coefficient of 0.9. The initial rate was set to  $\eta_0=0.03$  with a cosine learning rate decay schedule as  $\eta=\eta_0\cos(7\pi t/16T)$ , where t denotes the current training step and T is the total training step set at  $2^{20}$ . Additionally, exponential moving averaging with a momentum of 0.999 was implemented and the batch size for labeled data was fixed to 64. For DST, we used the code made available by the authors<sup>2</sup>.

We compared FlexMatch with and without the DST approach denoted respectively by FM and FM+DST. We also compared self-training with WRN trained in fully supervised manner. Each experiment was repeated 5 times by changing the seed at each time. Figure 1 presents the average accuracy of different models on the test set for the same number of initial labeled training samples per class within the set  $\{4, 10, 20, 50\}$  for both datasets. In both datasets, considering label noise within pseudo-labels leads to improved performance, with the improvement being more pronounced in the case of CIFAR-100.

In CIFAR-100, classes are structured into 20 superclasses, each comprising 5 related classes, addressing noise in this more complex task aids in class differentiation and enhances the model's ability to generalize. It is worth noting that with a greater number of initial labeled training examples, the gap between the FM and FM+DST approaches narrows, as the model becomes more proficient with the increased labeled data and makes fewer errors in pseudo-labeling.

<sup>&</sup>lt;sup>2</sup>https://github.com/thuml/Debiased-Self-Training

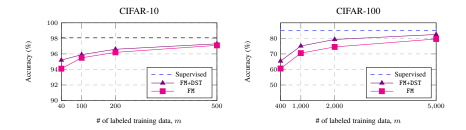


Figure 1: Comparisons in terms of Accuracy on CIFAR-10 and CIFAR-100 for a varying number of labeled training data. "Supervised" refers to the fully supervised learning (m=50000, u=0).

# 7 Conclusion and Perspectives

In this survey, we provided an overview of self-training approaches for semi-supervised learning that have received increasing attention in recent years.

First, we discussed the various strategies for selecting unlabeled samples for pseudolabeling that have been proposed. We emphasized the significance of considering margin distributions across unlabeled data as a pivotal factor in the development of these strategies. Next, we provided an overview of the diverse variants of self-training explored in the literature, along with relevant approaches. Furthermore, we examined recent theoretical advancements in this research domain and outlined the principal characteristics of self-training employed in several widely recognized applications. Lastly, we explored the impact of fundamental aspects of self-training on a range of benchmark datasets.

While the self-training approach is currently in widespread use, there are extensive opportunities for future research. Presently, the majority of studies have concentrated on perturbation-based deep learning, particularly in the domains of visual, text, and audio applications. However, there exist numerous other domains, such as industrial time-series or medical data, where the application of self-training could prove highly beneficial. As highlighted in recent research like [33], a promising avenue for future exploration lies in adopting a theoretical perspective to investigate self-training methods while explicitly considering the challenge of training a final classifier on data with noisy labels

Recently, self-training has expanded its scope beyond semi-supervised learning and has found extensive application in unsupervised domain adaptation [69, 97, 98]. In this context, the objective is to transfer knowledge from a labeled source domain to an unlabeled target domain. However, a notable challenge arises due to the distribution shift between the source and target domains. The prediction confidence provided by the training model may exhibit a strong bias toward the source domain, making it unreliable for generating pseudo-labels in the target domain.

In fact, substantial dissimilarity between the source and target domains can result in a significant drop in the performance of self-training, primarily because of the substantial errors introduced in pseudo-labels. Practical techniques like confidence regularization [98] can help mitigate the issue by reducing the occurrence of incorrect pseudo-labels [43]. However, from a theoretical standpoint, an important avenue for future research lies in the exploration of self-training methods under distribution shift conditions.

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