

Simultaneous Selection and Adaptation of Source Data via Four-Level Optimization

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Abstract

In many NLP applications, to mitigate data deficiency in a target task, source data is collected to help with target model training. Existing transfer learning methods either select a subset of source examples that are close to the target domain or try to adapt all source examples into the target domain, then use selected or adapted source examples to train the target model. These methods either incur significant information loss or bear the risk that after adaptation, source examples which are originally already in the target domain may be outside the target domain. To address the limitations of these methods, we propose a four-level optimization based framework which simultaneously selects and adapts source data. Our method can automatically identify in-domain and out-of-domain source examples and apply example-specific processing methods: selection for in-domain examples and adaptation for out-of-domain examples. Experiments on various datasets demonstrate the effectiveness of our proposed method.

1 Introduction

Transfer learning (TL) (Zhuang et al., 2020), which aims at improving a model in a target domain by utilizing data from a source domain, has been broadly studied in natural language processing. One paradigm of TL methods (Sun et al., 2011; Song et al., 2012; Wang et al., 2017b; Patel et al., 2018; Qu et al., 2018; Liu et al., 2019b) focuses on selecting a subset of source examples that are close to the target domain and using selected examples as additional training data for the target model. Another paradigm of TL methods (Pan et al., 2010; Ganin et al., 2016; Bousmalis et al., 2017) focuses on adapting the entire set of source examples into the target domain and using adapted source examples as additional training data for the target model. The problem of selec-

tion based methods is that unselected examples are discarded. Though having a domain discrepancy with target data, unselected examples still contain useful information that can be leveraged to improve the target model. Discarding them would lead to information loss. The problem of adaptation based methods is that some source examples may already be in the target domain; performing domain adaptation on these source examples is a waste of effort; and, even worse, after adaptation, these source examples may be outside the target domain.

To address the limitations of both paradigms of methods, we propose a new approach which simultaneously performs selection and adaptation of source examples: Our method automatically identifies which source examples are in the same domain as target data (referred to as in-domain source data) and which are not (referred to as out-of-domain source data); for in-domain source data, they are directly used to train the target model; for out-of-domain source data, they are first adapted, then utilized to train the target model. Compared with previous methods, our approach has the following advantage: Instead of using a single way to deal with all source examples (either performing selection or adaptation), our method applies example-specific ways to deal with different examples based on whether they are in-domain or out-of-domain.

Our method is based on four-level optimization, which performs the following four stages end-to-end. At the first stage, a domain distance network is trained based on self-supervised learning. At the second stage, the domain distance network is used to identify out-of-domain source examples and adapt them into the target domain. At the third stage, in-domain source examples selected by the domain distance network and adapted source examples are used to train a target model. At the fourth stage, the trained target model is evaluated on a validation set and data weights

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of the domain distance network are updated by minimizing the validation losses. Experiments on a variety of datasets demonstrate the effectiveness of our proposed method. To summarize, the major contributions of this work is that we propose a four-level optimization based framework for simultaneous selection and adaptation of source examples in transfer learning and demonstrate the effectiveness of our proposed method on two NLP applications.

2 Related Works

Domain Adaptation (DA). DA (Pan et al., 2010; Ganin et al., 2016; Bousmalis et al., 2017; Sun and Saenko, 2016; Long et al., 2017b; Ben-David et al., 2010; Hoffman et al., 2018; Long et al., 2017a; Kang et al., 2019; Long et al., 2015; Long et al., 2017b; Hoffman et al., 2018; Mitsuzumi et al., 2021) considers the problem of transferring knowledge from a label-rich source domain to a label-deficient target domain where the two domains have distributional discrepancies. There are mainly two paradigms of approaches. One paradigm (Sun and Saenko, 2016; Long et al., 2017b; Ben-David et al., 2010; Kang et al., 2019) is based on metric learning, where a distance metric is defined to measure the distribution discrepancy between domains and domain-invariant representations are learned by minimizing the distance. The other paradigm is based on adversarial learning (Hoffman et al., 2018; Long et al., 2017a; Tzeng et al., 2017; Sankaranarayanan et al., 2018; Motiian et al., 2017), which learns a domain discriminator and a feature learning network adversarially. The domain discriminator is trained to tell whether an instance is from source domain or target domain, while the feature learning network learns domain-invariant features by fooling the domain discriminator. CDAN (Long et al., 2017a) uses multilinear conditioning to capture the cross-covariance between feature representations and classifier predictions, and leverages entropy conditioning to control the uncertainty of classifier predictions. MME (Saito et al., 2019) performs adaptation by maximizing the conditional entropy of unlabeled target data w.r.t the classifier and minimizing it w.r.t the feature encoder. SRDC (Tang et al., 2020) performs deep discriminative clustering with source regularization for unsupervised domain adaptation. These

methods perform adaptation on all source data, which leads to waste of efforts and incurs a risk of moving in-domain source data outside the target domain.

Data Selection in Transfer Learning. Many methods (Jiang and Zhai, 2007; Foster et al., 2010; Moore and Lewis, 2010; Axelrod et al., 2011; Ge and Yu, 2017; Ruder and Plank, 2017; Sivasankaran et al., 2017; Zhang et al., 2017; Guo et al., 2019; Liu et al., 2019a; Tang and Jia, 2019; Wang et al., 2019a,b; Bateson et al., 2020) have been developed for selecting source data that is suitable for training target models, based on reinforcement learning (Patel et al., 2018; Qu et al., 2018; Liu et al., 2019b), adversarial learning (Wang et al., 2019a), curriculum learning (Zhang et al., 2017; Wang et al., 2019b), entropy (Song et al., 2012; Wang et al., 2017b), Bayesian optimization (Ruder and Plank, 2017), multi-task learning (Ge and Yu, 2017), and bi-level optimization (BLO) (Ren et al., 2018, 2020; Hu et al., 2019; Shu et al., 2019; Wang et al., 2020a,b). BLO based approaches select data by minimizing a validation loss, where the lower-level optimization problem trains network weights on a training dataset and the upper-level optimization problem learns data selection variables on a validation set. These methods select part of source data and discard the rest, which incurs information loss.

Transfer Learning (TL). TL (Pratt, 1993; Mihalkova et al., 2007; Niculescu-Mizil and Caruana, 2007; Pan and Yang, 2009; Luo et al., 2017; Zhuang et al., 2020) aims at training a better target model by utilizing source data. Many TL methods have been developed, including those based on 1) distribution alignment (Huang et al., 2006; Foster et al., 2010; Wang et al., 2017b; Ngiam et al., 2018), 2) regularization (Luo et al., 2008; Tommasi et al., 2010; Duan et al., 2012), 3) adversarial domain-invariant representation learning (Ganin et al., 2016; Long et al., 2017a; Hoffman et al., 2018; Zhang et al., 2019), and 4) latent space projection (Borgwardt et al., 2006; Pan et al., 2010; Long et al., 2013; Wang et al., 2017a). These methods either select part of source data or adapt all source data, which leads to information loss, waste of efforts, and the risk of adapting in-domain source data outside the target domain.

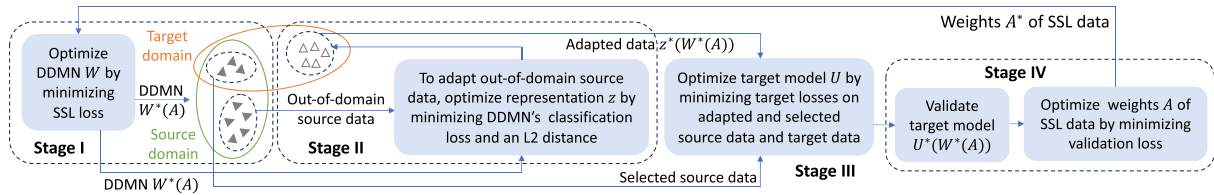


Figure 1: The overall framework.

Bi-level Optimization (BLO). BLO finds broad applications in hyperparameter tuning (Feurer et al., 2015), data selection (Shu et al., 2019; Ren et al., 2020; Wang et al., 2020b), training data generation (Such et al., 2019), neural architecture search (Liu et al., 2018), learning rate adaptation (Baydin et al., 2017), meta learning (Finn et al., 2017), etc. In BLO-based methods, model weights are learned by solving an inner optimization problem and meta parameters are learned by solving an outer optimization problem. The two optimization problems are nested. Different from existing BLO-based methods, our method is based on four-level optimization.

3 Methods

In this section, we present the method for simultaneous selection and adaptation of source examples based on four-level optimization. We aim to train a target model for a specific target domain using dataset D_t . In practical scenarios, the target domain often suffers from a lack of labeled training data. This lack can lead to overfitting on the training data and poor generalization on test data. To address this issue, we utilize a dataset from a source domain, D_s , which has an abundance of labeled examples. However, there is a notable discrepancy between the source and target data. We categorize D_s examples into two types: those that belong to the same domain as D_t (in-domain source data) and those that do not (out-of-domain source data). We aim to select in-domain source data and adapt out-of-domain source data into the target domain, and use selected and adapted source data to train the target model. The overall framework is shown in Figure 1. The notations are shown in Table 1.

3.1 A Four-Level Optimization Framework

We propose a four-level optimization framework to perform simultaneous selection and adaptation of source data. The framework consists of four learning stages which are performed end-to-end.

Notation	Meaning
D_t	Target dataset.
N_t	The number of examples in D_t .
$D_t^{(tr)}$	Training dataset of D_t .
$D_t^{(val)}$	Validation dataset of D_t .
$d_{t,i}^{(tr)}$	The i -th example in $D_t^{(tr)}$.
D_s	Source dataset.
N_s	The number of examples in D_s .
$d_{s,i}$	The i -th example in D_s .
q	Query example.
R	A set of examples.
o	Binary label regarding whether q and R are from the same domain.
M	The number of SSL training examples.
W	The DDMN's weight parameters.
a	The weight of an SSL training example.
A	The weights of all SSL training examples.
$W^*(A)$	The optimal solution of W , which depends on A .
$f(q, D_t; W^*(A))$	A binary label regarding whether q is out of the target domain.
$W_1^*(A)$	The encoder in $W^*(A)$.
$W_2^*(A)$	The rest of layers in $W^*(A)$ besides $W_1^*(A)$.
$\hat{z}(q; W_1^*(A))$	The latent representation of q extracted by $W_1^*(A)$.
$z(q)$	The adapted representation of q .
$z^*(q, W^*(A))$	The optimal solution of $z(q)$, which depends on $W^*(A)$.
γ	A tradeoff parameter in Eq.(2).
λ	A tradeoff parameter in Eq.(6).
U	Target model.
\hat{U}	A sub-network of U , which takes $z^*(q, W^*(A))$ as input.
ℓ_{tgt}	The target model's training loss.
L_t	The target model's training loss on target training data.
L_s	The target model's training loss on selected source data.
L_a	The target model's training loss on adapted source data.

Table 1: Notations.

Stage I. At the first stage, we learn a domain distance metric network via self-supervised learning (SSL) (He et al., 2019; Chen et al., 2020). This network takes a query example q and a set of examples R as inputs and predicts a binary label representing whether q is in the domain of

R. The architecture of this network is as follows. For q and each example in R , an encoder is used to generate a latent representation for the example. Then self-attention (Vaswani et al., 2017) is performed on these latent representations to generate attentive representations. Finally, the attentive representation of q and the averaged attentive representation of examples in R are concatenated and fed into a feedforward layer to predict the binary label.

To learn this domain distance metric network (DDMN), we construct self-supervised training examples. We randomly sample a subset of examples R from D_t , and randomly sample a query example q_t from D_t and a query example q_s from D_s . We label (q_t, R) as a positive pair since q_t and R are both from target domain, and label (q_s, R) as a negative pair since q_s is from source domain and R is from the target domain. This procedure repeats M times, yielding $2M$ training examples denoted by $D_{ssl} = \{(q_i, R_i, o_i)\}_{i=1}^{2M}$ where o_i is a binary label representing whether q_i and R_i are from the same domain. Let W denote weight parameters of the DDMN. We learn W by minimizing the binary classification loss: $\sum_{i=1}^{2M} \ell(q_i, R_i, o_i; W)$ where ℓ is a cross-entropy loss.

Note that these self-supervised examples may be noisy since the binary labels are given based on heuristics without human scrutiny. It could be the case that q_s happens to be in the same domain as R while q_t is not. It is necessary to automatically identify and remove these noisy examples. To achieve this goal, we associate each example (q_i, R_i, o_i) with a weight $a_i \in [0, 1]$. The smaller a_i is, the more likely that (q_i, R_i, o_i) is noisy. We multiply a_i to the loss $\ell(q_i, R_i, o_i; W)$. If a_i is close to zero, $a_i \ell(q_i, R_i, o_i; W)$ is made close to zero, which effectively removes (q_i, R_i, o_i) from the training set. This stage amounts to solving the following optimization problem:

$$W^*(A) = \operatorname{argmin}_W \sum_{i=1}^{2M} a_i \ell(q_i, R_i, o_i; W), \quad (1)$$

where $A = \{a_i\}_{i=1}^{2M}$. $W^*(A)$ denotes that W^* is a function of A . This is because W^* is a function of the loss which is a function of A . A is tentatively fixed at this stage and will be updated at a later stage. A cannot be updated at this stage. Otherwise, all the values in A would be zero.

Stage II. At the second stage, we use the learned domain distance network to identify out-of-domain source examples and adapt them into the target domain. For each data example q from the source dataset D_s , we feed q and the target dataset D_t into $W^*(A)$, which predicts a binary label $f(q, D_t; W^*(A))$. If $f(q, D_t; W^*(A)) = 0$, it means q is out of the target domain, and we use the domain distance network to adapt it into the target domain. The adaptation is performed in the following way. In $W^*(A)$, let $W_1^*(A)$ denote the encoder (containing all layers before self-attention) and $W_2^*(A)$ denote the rest of layers (including self-attention and feedforward layers) used for predicting the binary label. Let $\hat{z}(q; W_1^*(A))$ denote the latent representation of q extracted by $W_1^*(A)$ —the layer before self-attention. We learn another representation $z(q)$ of q which falls into the target domain (in the latent space) and is close to $\hat{z}(q; W_1^*(A))$. Closeness is measured using L_2 distance. To encourage $z(q)$ to fall into the target domain, we encourage $W_2^*(A)$ to predict that $z(q)$ is in the target domain, i.e., minimizing the binary classification loss $l(z(q), D_t, t_q = 1; W_2^*(A))$. At this stage, we solve the following optimization problem for each source example q that is predicted to be out-of-domain:

$$\begin{aligned} & z^*(q, W^*(A)) \\ &= \operatorname{argmin}_{z(q)} \underbrace{l(z(q), D_t, t_q = 1; W_2^*(A))}_{\text{Encourage } z(q) \text{ to fall into the target domain}} \\ & \quad + \gamma \underbrace{\|z(q) - \hat{z}(q; W_1^*(A))\|_2^2}_{\text{Encourage } z(q) \text{ to be close to } \hat{z}(q; W_1^*(A))} \end{aligned} \quad (2)$$

where γ is a tradeoff parameter.

Stage III. At the third stage, we use selected and adapted source examples, together with the target training data $D_t^{(tr)}$, to train the target model U . Let ℓ_{tgt} denote the loss function of the target task, N_s and N_t denote the number of examples in D_s and D_t , and $f(q, D_t; W^*(A))$ denote the binary label predicted by the domain distance network $W^*(A)$ regarding whether a source example q is in the target domain. $d_{t,i}^{(tr)}$ denotes the i -th example in $D_t^{(tr)}$, $d_{s,i}$ denotes the i -th example in D_s . For a source example which is predicted to be in the target domain (where $f(d_{s,i}, D_t; W^*(A)) = 1$), it is directly used to train the target model U by minimizing the loss $\ell_{tgt}(U, d_{s,i})$. For a source

example which is predicted to be out of the target domain (where $f(d_{s,i}, D_t; W^*(A)) = 0$), its adapted representation $z^*(d_{s,i}, W^*(A))$ obtained in the second stage is used to train the target model. Note that since $z^*(d_{s,i}, W^*(A))$ is already a latent representation, only part of weight parameters (denoted by \widehat{U}) in the target model U is needed to make prediction on $z^*(d_{s,i}, W^*(A))$. We define the loss on target training data as:

$$L_t(U) = \sum_{i=1}^{N_t} \ell_{tgt}(U, d_{t,i}^{(tr)}), \quad (3)$$

the loss on selected source data as:

$$L_s(U, W^*(A)) = \sum_{i=1}^{N_s} f(d_{s,i}, D_t; W^*(A)) \ell_{tgt}(U, d_{s,i}), \quad (4)$$

and the loss on adapted data as:

$$L_a(\widehat{U}, W^*(A)) = \sum_{i=1}^{N_s} (1 - f(d_{s,i}, D_t; W^*(A))) \ell_{tgt}(\widehat{U}, z^*(d_{s,i}, W^*(A))). \quad (5)$$

At this stage, we solve the following optimization problem:

$$U^*(W^*(A)) = \operatorname{argmin}_U L_t(U) + \lambda (L_s(U, W^*(A)) + L_a(\widehat{U}, W^*(A))), \quad (6)$$

where λ is a tradeoff parameter.

Stage IV. At the fourth stage, we use the trained target model to make predictions on the validation dataset $D_t^{(val)}$ of the target task. We update the weights A of self-supervised training examples in the first stage by minimizing the validation loss:

$$\min_A L(U^*(W^*(A)), D_t^{(val)}). \quad (7)$$

A Four-level Optimization Based Framework. Putting all these pieces together, we have the following four-level optimization based framework.

$$\underbrace{\min_A L(U^*(W^*(A)), D_t^{(val)})}_{\text{Stage IV}} \\ \text{s.t.} \\ \underbrace{U^*(W^*(A)) = \operatorname{argmin}_U L_t(U) + \lambda (L_s(U, W^*(A)) + L_a(\widehat{U}, W^*(A)))}_{\text{Stage III}} \\ \underbrace{z^*(q, W^*(A)) = \operatorname{argmin}_{z(q)} l(z(q), D_t, t_q = 1; W_2^*(A)) + \gamma \|z(q) - \hat{z}(q; W_1^*(A))\|_2^2}_{\text{Stage II}} \\ \underbrace{W^*(A) = \operatorname{argmin}_W \sum_{i=1}^{2M} a_i \ell(q_i, R_i, o_i; W)}_{\text{Stage I}} \quad (8)$$

To make the objective at the third stage differentiable, we approximate $f(d_{s,i}, D_t; W^*(A))$ using the probability calculated by $W^*(A)$ regarding whether $d_{s,i}$ and D_t are in the same domain.

3.2 Optimization Algorithm

We leverage a gradient-based method (Liu et al., 2018) to solve the problem in Eq.(8). Convergence of this algorithm has been analyzed (Ghadimi and Wang, 2018; Grazi et al., 2020; Ji et al., 2021; Liu et al., 2021; Yang et al., 2021). At each level of the optimization problem, the exact value of the optimal solution (on the left-hand side of the equal sign, marked with $*$) is computationally expensive to compute. To address this problem, following Liu et al. (2018), we approximate the optimal solution using a one-step gradient descent update and plug the approximation into the next level of optimization problem. In the sequel, $\frac{\partial \cdot}{\partial \cdot}$ denotes partial derivative. $\frac{d \cdot}{d \cdot}$ denotes an ordinary derivative. Following Liu et al. (2018), we approximate $W^*(A)$ using one-step gradient descent update of W :

$$W^*(A) \approx W' = W - \eta_w \nabla_W \sum_{i=1}^{2M} a_i \ell(q_i, R_i, o_i; W). \quad (9)$$

We plug $W^*(A) \approx W'$ into the loss function at the second stage and get an approximated objective. Let W'_2 and W'_1 denote the approximations of $W_2^*(A)$ and $W_1^*(A)$, respectively. The approximated objective is $O(z(q), W'_2, W'_1) = l(z(q), D_t, t_q = 1; W'_2) + \gamma \|z(q) - \hat{z}(q; W'_1)\|_2^2$. We approximate $z^*(q, W^*(A))$ using one-step gradient descent update of $z(q)$ w.r.t the approximated objective:

$$z^*(q, W^*(A)) \approx z'(q) = z(q) - \eta_z \nabla_{z(q)} O(z(q), W'_2, W'_1). \quad (10)$$

We plug $z^*(q, W^*(A)) \approx z'(q)$ and $W^*(A) \approx W'$ into the objective at the third stage and get an approximated objective. Let $g(d_{s,i}, D_t; W^*(A))$ denote the probability that $d_{s,i}$ and D_t are in the same domain. We approximate $U^*(W^*(A))$ using one-step gradient descent update of U w.r.t the approximated objective:

$$U^*(W^*(A)) \approx U' = U - \eta_u \nabla_U (L_t(U) + \lambda (L_s(U, W') + L_a(\widehat{U}, W'))). \quad (11)$$

Finally, we plug the approximation $U^*(W^*(A)) \approx U'$ into the validation loss at the fourth stage and update A by minimizing the approximated loss using gradient descent:

$$A \leftarrow A - \eta_a \nabla_A L(U', D_t^{(val)}). \quad (12)$$

while not converged do

1. Update the approximation W' of $W^*(A)$ using Eq.(9)
2. Update the approximation $z'(q)$ of $z^*(q, W^*(A))$ using Eq.(10)
3. Update the approximation U' of $U^*(W^*(A))$ using Eq.(11)
4. Update A using Eq.(13)

end

Algorithm 1: Optimization algorithm.

For $\nabla_A L(U', D_t^{(val)})$, it can be computed as:

$$\begin{aligned} & \nabla_A L(U', D_t^{(val)}) \\ &= \frac{dW'}{dA} \left(\frac{\partial U'}{\partial W'} + \sum_{i=1}^{N_s} \frac{dz'(d_{s,i})}{dW'} \frac{\partial U'}{\partial z'(d_{s,i})} \right) \nabla_{U'} L(U', D_t^{(val)}), \end{aligned} \quad (13)$$

where

$$\begin{aligned} & \frac{\partial U'}{\partial z'(d_{s,i})} \\ &= -\eta_u \lambda (1 - g(d_{s,i}, D_t; W')) \nabla_{z'(d_{s,i}), U}^2 \ell_{tgt}(\hat{U}, z'(d_{s,i})), \end{aligned} \quad (14)$$

$$\begin{aligned} & \frac{dz'(d_{s,i})}{dW'} \\ &= -\eta_z \nabla_{W', z(q)}^2 (\ell(z(q), D_t, t_q = 1; W'_2) \\ & \quad + \gamma \|z(q) - \hat{z}(q; W'_1)\|_2^2), \end{aligned} \quad (15)$$

$$\begin{aligned} & \frac{\partial U'}{\partial W'} \\ &= -\eta_u \lambda \nabla_{W', U}^2 \sum_{i=1}^{N_s} (g(d_{s,i}, D_t; W') \ell_{tgt}(U, d_{s,i}) \\ & \quad + (1 - g(d_{s,i}, D_t; W')) \ell_{tgt}(\hat{U}, z'(d_{s,i}))) \end{aligned} \quad (16)$$

$$\frac{dW'}{dA} = -\eta_w \nabla_{A, W}^2 \sum_{i=1}^{2M} a_i \ell(q_i, R_i, o_i; W) \quad (17)$$

The gradient descent update of A in Eq.(13) can run one or more steps. After A is updated, the one-step gradient-descent approximations in Eq.(9), (10), and (11), which are functions of A , change with A and need to be re-updated. Then, the gradient of A , which is a function of one-step gradient-descent approximations, needs to be re-calculated and is used to refresh A . In sum, the update of A and the updates of one-step gradient-descent approximations mutually depend on each other. These updates are performed iteratively until convergence. Algorithm 1 shows the algorithm.

In the gradient of A calculated using the chain rule, the number of chains is the same as the number of levels in our proposed four-level optimization formulation. This shows that this optimization algorithm preserves the four-level nested optimization nature of the proposed formulation.

3.3 Reduce Computation and Memory Costs

To reduce computation and memory costs, we adopt the following methods.

- We reduced the frequencies of updating (including calculating hypergradients of) the weights A of self-supervised training examples. They were updated every 8 mini-batches (i.e., iterations) instead of on every mini-batch. We empirically found that this greatly reduced computational costs without significantly sacrificing accuracy. The rest of the parameters were updated on every mini-batch.
- We added a decorrelation regularizer (Cogswell et al., 2015) on W and U , which significantly speeds up convergence and allows reducing the number of epochs by half without sacrificing convergence quality.
- Parameter tying was performed to reduce the number of weight parameters and computation costs. We let W and U share the same feature learning layers. These layers account for >95% of parameters in each of these models. Sharing them across models significantly reduces the number of total parameters, which reduces the computational costs of updating these parameters.
- We optimized the implementation of our method to speed up computation by leveraging techniques including 1) automatic mixed precision (Micikevicius et al., 2017), 2) using multiple (4, specifically) workers and pinned memory in PyTorch DataLoader, 3) using cudNN autotuner, 4) kernel fusion, and so forth.

3.4 Applications

In this section, we apply the proposed four-level optimization framework for two NLP applications.

Text Classification. In many text classification problems, training data in a target domain is limited. To address the lack of target training data, one can leverage data from a source domain.

Visual Question Answering on Pathology Images. Pathology imaging (Mohan, 2015) is broadly used for identifying the causes and effects of diseases or injuries. Given a pathology image, being able to answer questions about the clinical findings contained in the image is very important

Domain	Dataset	Label Type	Train	Dev	Test	Classes
Biomedical	CHEMPROT	relation classification	4169	2427	3469	13
	RCT	abstract sent. roles	180040	30212	30135	5
Computer Science	ACL-ARC	citation intent	1688	114	139	6
	SciERC	relation classification	3219	455	974	7
News	HYPERPARTISAN	partisanship	515	65	65	2
	AGNEWS	topic	115000	5000	7600	4
Reviews	HELPFULNESS	review helpfulness	115251	5000	25000	2
	IMDB	review sentiment	20000	5000	25000	2

Table 2: Statistics of datasets used in Gururangan et al. (2020).

for medical decision-making (He et al., 2020). However, collecting a large-scale visual question answering (VQA) dataset is challenging, due to the lack of doctors for making questions and answers from pathology images. A dataset collected in He et al. (2020) has about 33K question-answer pairs generated from around 5K pathology images. Although largest of its kind, it is still relatively small compared with common VQA datasets. To mitigate the deficiency of training data, we collect an auxiliary source dataset. From the pathology literature, we collect 1792 pathology figures and create 36,471 VQA questions using the method proposed in He et al. (2020).

4 Experiments

In this section, we present experimental results on text classification and visual question answering on pathology images. Following the common data assumption in transfer learning, the amount of labeled source data in our experiments is significantly larger than that of target data. Every experiment runs 4 times with different random initializations. For all experiments, we performed significance tests using double-sided t-tests. The p-values of our methods against baselines are all less than 0.001, which shows that our methods are significantly better than baselines.

4.1 Text Classification

Dataset. Following Gururangan et al. (2020), we experiment with four domains: biomedical, computer science, news, and reviews. For the biomedical domain, we use two target datasets: CHEMPROT (Kringelum et al., 2016) and RCT (Dernoncourt and Lee, 2017), and one source dataset which contains 2.68 million full-text papers from S2ORC (Lo et al., 2019) with 7.55 billion tokens. For the computer science domain,

we use two target datasets: ACL-ARC (Jurgens et al., 2018) and SciERC (Luan et al., 2018), and one source dataset which contains 2.22 million full-text papers from S2ORC (Lo et al., 2019) with 8.1 billion tokens. For the news domain, we use two target datasets: HYPERPARTISAN (Kiesel et al., 2019) and AGNEWS (Zhang et al., 2015), and one source dataset which contains 11.9 million articles from RealNews (Zellers et al., 2019) with 6.66 billion tokens. For the reviews domain, we use two target datasets: HELPFULNESS (McAuley et al., 2015) and IMDB (Maas et al., 2011), and one source dataset which contains 24.75 million articles from Amazon Reviews (He and McAuley, 2016) with 2.11 billion tokens. Statistics of the target datasets are summarized in Table 2. In our method, we split the original target training set into a new training set and a validation set, with a ratio of 1:1. The new training set is used as $D_{cls}^{(tr)}$ and the validation set is used as $D_{cls}^{(val)}$. Note that baseline methods are trained on the combination of $D_{cls}^{(tr)}$ and $D_{cls}^{(val)}$.

Baselines. We compare our method with the following baselines. In baseline methods, the target data used for training the target model includes both training and validation sets.

- Domain adaptive pretraining (DAPT) (Gururangan et al., 2020): Given an RoBERTa-base model which has been pretrained on large amounts of corpora (used in Liu et al., 2019c), we continue to pretrain it on source data, then finetune it on target data.
- Task adaptive pretraining (TAPT) (Gururangan et al., 2020): Given an RoBERTa-base model which has been pretrained on large amounts of corpora (used

in Liu et al., 2019c), we continue to pretrain it on input texts of each target dataset.

- Data selection methods for transfer learning, based on Bayesian optimization (BO) (Ruder and Plank, 2017), minimax game (MMG) (Wang et al., 2019a), learning to select instance (LSI) (Huan et al., 2021).
- Domain adaptation methods, including DANN (Ganin et al., 2016), CDAN (Long et al., 2017a), MME (Saito et al., 2019), SRDC (Tang et al., 2020), SSDA (Kim and Kim, 2020), GDA (Mitsuzumi et al., 2021), and ATDOC (Liang et al., 2021).
- SimCSE (Gao et al., 2021): A contrastive learning method. The same input sentence is fed into a pretrained RoBERTa encoder twice by applying different dropout masks, to get two different embeddings. These two embeddings are labeled as being “similar”. Embeddings of different sentences are labeled as being “dissimilar”. Contrastive learning is performed on these “similar” and “dissimilar” pairs.

Implementation Details. In the domain distance network, the hyperparameters of self-attention and feed-forward layer are the same as those in Transformer (Vaswani et al., 2017). The subset R has varying cardinality (sampled uniformly). M is set to 10k. The tradeoff parameter γ and λ is set to 0.1 and 0.5 respectively. Baselines and our method receive a similar amount of tuning time and efforts. F1 is used as the evaluation metric. We use RoBERTa-base as a data encoder. For a fair comparison, most of our hyperparameters are the same as those in Gururangan et al. (2020). The maximum text length was set to 512. For all datasets, we used a batch size of 16 with gradient accumulation. We used the AdamW optimizer (Loshchilov and Hutter, 2017) with a warm-up proportion of 0.06, a weight decay of 0.1, and an epsilon of 1e-6. In AdamW, β_1 and β_2 are set to 0.9 and 0.98, respectively. The maximum learning rate was 2e-5. For the reader’s convenience, we summarize the hyperparameters as follows.

- The number M of self-supervised training examples: 10k
- Tradeoff parameters γ and λ : 0.1, 0.5
- Maximum text length: 512
- Batch size: 16

- Optimizer: AdamW
- Warm-up proportion, weight decay, and epsilon in AdamW: 0.06, 0.1, and 1e-6
- β_1 and β_2 in AdamW: 0.9, 0.98
- Maximum learning rate: 2e-5

To tune the hyperparameters, we randomly split the validation set into two equal-sized subsets, denoted by A and B . For each configuration of hyperparameters, we use the validation set A to learn the importance weights of self-supervised training examples. Then we measure the performance of the trained model on validation set B . Hyperparameter values yielding the best performance on validation set B are selected. Each baseline method received an equal amount of tuning effort as that for our method.

Results and Analysis. Table 3 shows the results. From this table, we make the following observations. **First**, our method outperforms source data selection methods including BO, MMG, and LSI. In these baseline methods, out-of-domain source examples are discarded, which incurs information loss. In contrast, our method adapts out-of-domain source examples into the target domain and uses the adapted examples to train the target model. **Second**, our method works better than domain adaptation methods including DANN, CDAN, MME, SRDC, SSDA, GDA, and ATDOC. The reason is: These methods perform adaptation on all source examples without identifying which ones are already in the target domain; as a result, some in-domain source examples may be adapted out of the target domain. In contrast, our method first identifies which source examples are already in-domain and only performs adaptation on out-of-domain examples. **Third**, our method outperforms vanilla RoBERTa-base which does not leverage source data to learn representations. This demonstrates that leveraging source data is helpful for improving the target model. **Fourth**, our method performs better than DAPT. In DAPT, all source examples are leveraged to pretrain the target encoder without considering the fact that some source examples have large domain discrepancy with the target domain and are not suitable for pretraining the target encoder. **Fifth**, our method outperforms TAPT and SimCSE. These methods do not leverage auxiliary source data or select/adapt source data.

Method	CHEMPROT	RCT	ACL-ARC	SciERC	HYPERPARTISAN	AGNEWS	HELPLESSNESS	IMDB	Average
RoBERTa-base	81.9 _{1.0}	87.2 _{0.1}	63.0 _{5.8}	77.3 _{1.9}	86.6 _{0.9}	93.9 _{0.2}	65.1 _{3.4}	95.0 _{0.2}	81.3
TAPT	82.6 _{0.4}	87.7 _{0.1}	67.4 _{1.8}	79.3 _{1.5}	90.4 _{5.2}	94.5 _{0.1}	68.5 _{1.9}	95.5 _{0.1}	83.2
DAPT	84.2 _{0.2}	87.6 _{0.1}	75.4 _{2.5}	80.8 _{1.5}	88.2 _{5.9}	93.9 _{0.2}	66.5 _{1.4}	95.4 _{0.1}	84.0
SimCSE	83.2 _{0.2}	87.6 _{0.1}	69.5 _{2.6}	80.5 _{0.7}	90.9 _{2.7}	94.7 _{0.1}	68.7 _{1.7}	95.7 _{0.1}	83.9
BO	83.3 _{0.2}	87.5 _{1.0}	75.4 _{3.3}	79.1 _{1.3}	88.1 _{4.2}	94.0 _{0.2}	68.4 _{2.1}	95.3 _{0.2}	83.9
MMG	83.0 _{0.4}	87.4 _{1.0}	75.1 _{4.1}	79.6 _{0.8}	87.9 _{2.1}	94.2 _{0.1}	66.7 _{1.5}	95.7 _{0.1}	83.7
LSI	83.2 _{0.3}	87.5 _{1.0}	75.3 _{4.5}	80.2 _{0.6}	88.7 _{1.9}	94.5 _{0.1}	68.6 _{1.0}	95.4 _{0.1}	84.2
DANN	83.5 _{0.3}	87.7 _{0.1}	75.5 _{3.7}	80.5 _{1.1}	90.4 _{3.0}	94.3 _{0.1}	66.8 _{2.8}	95.2 _{0.1}	84.2
CDAN	83.8 _{0.2}	87.4 _{0.1}	75.9 _{2.4}	80.9 _{1.4}	88.7 _{5.5}	94.1 _{0.2}	67.3 _{3.4}	95.8 _{0.2}	84.2
MME	84.0 _{0.1}	87.4 _{0.1}	75.7 _{5.1}	80.5 _{0.8}	89.2 _{2.8}	94.6 _{0.2}	68.9 _{2.7}	95.4 _{0.1}	84.5
SRDC	84.3 _{0.3}	87.3 _{0.1}	75.7 _{3.4}	80.7 _{1.0}	88.9 _{3.5}	94.1 _{0.1}	67.6 _{3.0}	95.1 _{0.1}	84.2
SSDA	83.9 _{0.5}	87.9 _{0.1}	75.9 _{2.7}	81.0 _{1.4}	88.5 _{2.6}	94.4 _{0.1}	67.0 _{1.6}	95.7 _{0.2}	84.3
GDA	84.1 _{0.3}	87.3 _{0.1}	75.4 _{2.4}	80.8 _{0.9}	90.5 _{4.1}	94.2 _{0.2}	66.8 _{2.4}	95.5 _{0.1}	84.3
ATDOC	84.5 _{0.2}	87.5 _{0.1}	75.6 _{3.6}	81.1 _{1.2}	89.0 _{5.7}	93.9 _{0.2}	67.4 _{2.8}	95.1 _{0.1}	84.3
No-Adapt	84.1 _{0.2}	87.3 _{0.1}	75.9 _{2.2}	81.4 _{1.0}	90.3 _{4.2}	94.1 _{0.1}	66.9 _{3.3}	95.3 _{0.2}	84.4
No-In-Domain	84.5 _{0.1}	87.5 _{0.1}	75.5 _{4.5}	81.6 _{1.7}	88.8 _{3.9}	94.6 _{0.2}	67.1 _{3.1}	95.6 _{0.1}	84.4
Separate	85.3 _{0.5}	88.1 _{0.1}	75.8 _{2.9}	82.7 _{0.9}	90.1 _{3.6}	94.0 _{0.2}	68.3 _{1.5}	95.2 _{0.1}	84.9
MMD	85.9 _{0.4}	88.3 _{0.1}	75.5 _{4.9}	82.3 _{1.1}	89.5 _{1.8}	94.1 _{0.1}	68.9 _{1.2}	95.4 _{0.2}	85.0
AD	85.6 _{0.2}	88.7 _{0.1}	75.9 _{3.1}	82.6 _{0.8}	90.3 _{3.0}	94.7 _{0.2}	67.4 _{1.7}	95.8 _{0.1}	85.1
WMVL	85.5 _{0.4}	88.5 _{0.1}	75.7 _{2.8}	81.9 _{1.3}	90.6 _{4.6}	94.6 _{0.1}	68.1 _{2.3}	95.1 _{0.1}	85.0
H-Divergence	85.4 _{0.3}	88.7 _{0.1}	75.9 _{4.6}	82.1 _{0.9}	89.2 _{1.9}	94.2 _{0.1}	68.5 _{2.6}	95.3 _{0.1}	84.9
Our full method	87.1 _{0.2}	90.4 _{0.1}	77.6 _{2.3}	84.4 _{0.7}	92.4 _{1.2}	95.7 _{0.1}	70.9 _{0.8}	96.6 _{0.1}	86.9

Table 3: Text classification results. Following Gururangan et al. (2020), the results are micro-F1 for CHEMPROT and RCT, and macro-F1 for other datasets. For each x_y entry, x and y represent the mean and standard deviation of four random runs, respectively.

4.2 Visual Question Answering on Pathology Images

Datasets. For the target dataset, we use PathVQA (He et al., 2020), which contains 1,670 pathology images and 32,795 question-answer pairs. Of these, 16,466 questions are open-ended, with the following types: what, where, when, whose, how, why, how much/how many. The rest are close-ended ‘‘yes/no’’ questions. Based on images, the dataset is split into a training, validation, and test set with a ratio of 3:1:1 approximately. Note that baseline methods are trained on the combination of the training and validation sets. We collected a source dataset containing 1792 pathology figures extracted from papers in medRxiv where each figure has a caption. We create 36,471 VQA questions using the method proposed in He et al. (2020). This source dataset will be made available publicly.

Implementation Details. For the target model, we experimented with two state-of-the-art VQA models—LXMERT (Tan and Bansal, 2019) and bilinear attention networks (BAN) (Kim et al., 2018)—each containing an image encoder, a question encoder, and an answer generation head. Hyperparameters mostly follow those in previous work (Tan and Bansal, 2019; Kim et al., 2018; Yang et al., 2016). For LXMERT, the hidden size of the text encoder was set to 768.

The initial learning rate was set to $5e-5$ with the Adam (Kingma and Ba, 2014) optimizer used. The batch size was set to 256. The model was trained for 200 epochs. For BAN, words in questions and answers were represented using GloVe (Pennington et al., 2014) vectors. The initial learning rate was set to 0.005 with the Adamax optimizer (Kingma and Ba, 2014) used. The batch size was set to 512. The model was trained for 200 epochs.

From questions and answers in the PathVQA dataset, we create a vocabulary of 4,631 words that have the highest frequencies. Data augmentation is applied to images, including shifting, scaling, and shearing. We compare with baselines similar to those in Section 4.1. The Pretrain baseline works as follows: We pretrain the target encoder using source data, then finetune the target encoder using target data. Three metrics were used for evaluation, including BLEU (Papineni et al., 2002), macro-averaged F1 (Goutte and Gaussier, 2005), and accuracy (Malinowski and Fritz, 2014). We implement the methods using PyTorch and perform training on four GTX 1080Ti GPUs.

Results and Analysis. The results are shown in Table 4. From this table, we make similar observations as those in Table 3. The analysis of reasons is similar to that for results in Table 3. The

	Accuracy(%)	BLEU-1(%)	BLEU-2(%)	BLEU-3(%)	F1(%)	Average	Runtime(h)
<i>LXMERT (Tan and Bansal, 2019) based experiments</i>							
Vanilla LXMERT (Tan and Bansal, 2019)	57.6	57.4	3.1	1.3	9.9	25.9	29
Pretrain (He et al., 2016)	59.3	59.0	4.6	2.6	11.3	27.4	35
BO (Ruder and Plank, 2017)	59.5	58.9	3.8	2.7	11.4	27.3	41
MMG (Wang et al., 2019a)	59.3	58.4	3.7	2.6	10.9	27.0	38
LSI (Huan et al., 2021)	59.2	58.8	3.5	2.8	10.6	27.0	32
DANN (Ganin et al., 2016)	59.9	59.4	4.3	3.2	11.2	27.6	30
CDAN (Long et al., 2017a)	60.2	58.9	4.7	3.3	11.9	27.8	35
MME (Saito et al., 2019)	59.7	59.0	3.6	2.9	11.7	27.4	30
SRDC (Tang et al., 2020)	58.8	58.7	4.8	2.9	12.2	27.5	39
SSDA (Kim and Kim, 2020)	59.3	58.3	4.4	3.0	12.0	27.4	34
GDA (Mitsuzumi et al., 2021)	59.8	59.1	4.6	2.8	12.3	27.7	36
ATDOC (Liang et al., 2021)	59.6	59.6	4.3	3.1	12.2	27.8	30
Ours	62.5	61.1	5.2	3.7	12.9	29.1	29
<i>BAN (Kim et al., 2018) based experiments</i>							
Vanilla BAN (Kim et al., 2018)	55.1	56.2	3.2	1.2	8.4	24.8	25
Pretrain (He et al., 2016)	58.4	58.6	4.3	1.6	10.3	26.6	28
BO (Ruder and Plank, 2017)	58.3	58.5	4.2	2.0	10.9	26.8	32
MMG (Wang et al., 2019a)	58.7	58.1	4.6	2.3	11.2	27.0	29
LSI (Huan et al., 2021)	58.3	58.7	4.3	2.1	11.5	27.0	30
DANN (Ganin et al., 2016)	58.8	58.4	4.3	2.3	11.0	27.0	31
CDAN (Long et al., 2017a)	58.6	58.6	4.5	2.5	11.4	27.1	33
MME (Saito et al., 2019)	59.1	58.9	4.9	2.3	11.1	27.3	26
SRDC (Tang et al., 2020)	58.9	58.7	4.7	2.5	11.6	27.3	35
SSDA (Kim and Kim, 2020)	59.3	59.0	4.4	2.6	11.4	27.3	32
GDA (Mitsuzumi et al., 2021)	59.5	58.8	4.9	2.4	11.7	27.5	30
ATDOC (Liang et al., 2021)	59.1	59.1	5.0	2.7	11.5	27.5	29
Ours	62.4	61.7	5.6	3.2	12.5	29.1	26

Table 4: Results on the PathVQA dataset. Runtime (hours) for training is measured on a 1080TI GPU.

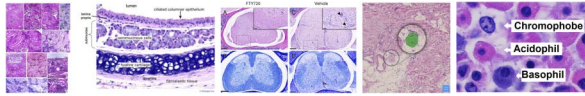


Figure 2: Randomly sampled source pathology figures identified by our method as being out of the target domain.

training time of our method is similar to that of baselines. Figure 2 shows some randomly sampled source pathology figures identified by our method as being out of the target domain. These images contain subfigures and texts, which are different from the target dataset.

4.3 Ablation Studies

4.3.1 Ablation by Removing Certain Components

To better understand the contributions of individual components in our framework, we perform the following ablation studies.

- **No-Adapt.** Out-of-domain source examples are discarded instead of being adapted. This is equivalent to removing stage II and the loss term $\mathbb{I}(f(d_{s,i}, D_t; W^*(A)) =$

$0)\ell_{tgt}(\hat{U}, z^*(d_{s,i}, W^*(A)))$ in Eq.(6) of stage III.

- **No-In-Domain.** In-domain source examples are discarded instead of being used for training the target model. This is equivalent to removing the loss term $\mathbb{I}(f(d_{s,i}, D_t; W^*(A)) = 1)\ell_{tgt}(U, d_{s,i})$ in Eq.(6) of stage III.
- **Separate.** Different stages are performed separately instead of jointly.

Table 3 shows ablation study results. From this table, we make the following observations. **First**, our full method works better than No-Adapt. This shows that it is beneficial to adapt out-of-domain source examples into the target domain and use adapted examples to train the target model, and our method is effective in achieving this goal. **Second**, our full method outperforms No-In-Domain, which demonstrates that the source examples selected by our method are useful for training the target model. **Third**, our full method achieves better performance than Separate. Our full method performs source data selection, adaptation, and target model training jointly, which enables these different tasks to mutually influence each other

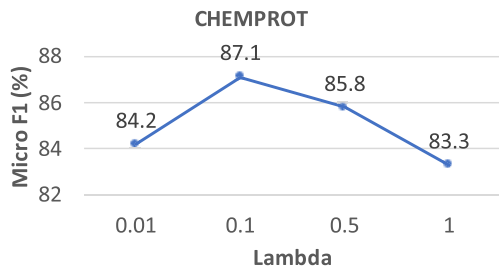


Figure 3: How the micro F1 on CHEMPROT changes with λ .

to achieve the best overall performance. Such a mechanism is lacking in Separate.

4.3.2 Ablation on the Adaptation Component

We perform an ablation study of the adaptation component in stage II of our framework by replacing it with the following baselines.

- Maximum mean discrepancy (MMD) (Kang et al., 2019), which is a broadly used metric for measuring discrepancies of two distributions. MMD-based adaptation method learns latent representations so that the selected out-of-domain source examples and the target examples have small MMD in the latent space.
- Adversarial adaptation (AD) (Ganin et al., 2016), which learns latent representations so that a domain discriminator cannot tell whether an example is from the source or from the target.

Table 3 shows the results. As can be seen, our adaptation method works better than the baselines. The reason is: The domain distance metric in our method is learned using many self-supervised training examples; it can better measure domain discrepancy and facilitate domain adaptation.

4.3.3 Ablation on the Selection Mechanism

We perform an ablation study of the selection mechanism in our framework by replacing it with the following baselines.

- Each source example is associated with a weight in $[0, 1]$. A larger weight indicates the example is more likely to be in the target domain. We learn these data weights by minimizing the validation loss (WMVL) of the target model.

- We use an \mathcal{H} -divergence (Elsahar and Gallé, 2019) based metric to measure domain similarity between a source example and the target dataset.

Table 3 shows the results. As can be seen, our selection mechanism works better than the baselines. The reason is: In our framework, the domain distance network (DSN) is learned in a discriminative way by performing classification on self-supervised training examples. Discriminative training can enable the DSN to better distinguish in-domain source examples from out-of-domain ones. In contrast, the selection mechanisms in the two baselines lack discriminability.

4.3.4 Ablation on the Tradeoff Parameter λ

We investigate how the performance of our method is affected by the tradeoff parameter λ . Figure 3 shows the test accuracy on ImageCLEF. As can be seen, when λ increases from 0.01 to 0.1, the accuracy improves. This is because the selected and adapted source data is used as additional training resources for the target model. However, as λ further increases, the accuracy drops. This is because the source data is not as reliable as the target data. An excessively large λ renders too much emphasis to be put on less-reliable source data.

4.3.5 Ablation on the Train-Validate Ratio

In the next ablation study, we investigate how the performance of our method varies under different split ratios between target training and validation datasets. The study was performed on the text classification task. Table 5 shows the results. As can be seen, a more balanced ratio (e.g., 1:1) yields better performance. When the ratio is largely imbalanced (e.g., 1:9, 1:4, 1:0.1, 1:0.25), the performance is worse. If the target training dataset $D_t^{(tr)}$ is much smaller than the target validation dataset $D_t^{(val)}$, the target model’s weight parameters U , which are trained on $D_t^{(tr)}$, will not be sufficiently trained due to the lack of training data and thereby yield poorer performance. On the other hand, if $D_t^{(val)}$ is much smaller than $D_t^{(tr)}$, the weights A of SSL training examples, which are optimized by minimizing the loss on $D_t^{(val)}$, will not be sufficiently optimized due to the lack of data and thereby yield worse performance as well. Note that since $D_t^{(val)}$ and $D_t^{(tr)}$ are obtained

Train-val ratio	CHEMPROT	RCT	ACL-ARC	SciERC	HYPERPARTISAN	AGNEWS	HELPFULNESS	IMDB	Average
1:9	79.4 _{0.4}	86.9 _{0.1}	73.7 _{2.7}	78.5 _{1.3}	87.2 _{2.3}	93.0 _{0.2}	65.5 _{3.0}	93.3 _{0.1}	82.2
1:4	82.0 _{0.4}	87.8 _{0.1}	75.1 _{3.9}	81.0 _{1.5}	89.6 _{3.0}	94.1 _{0.1}	68.2 _{2.7}	94.1 _{0.2}	84.0
1:2	84.2 _{0.1}	89.1 _{0.1}	76.4 _{5.0}	83.2 _{0.9}	91.7 _{4.2}	94.5 _{0.1}	70.0 _{0.3}	95.9 _{0.2}	85.6
1:1	87.1 _{0.2}	90.4 _{0.1}	77.6 _{2.3}	84.4 _{0.7}	92.4 _{1.2}	95.7 _{0.1}	70.9 _{0.8}	96.6 _{0.1}	86.9
1:0.5	86.8 _{0.4}	90.6 _{0.1}	77.1 _{2.9}	84.0 _{1.2}	92.8 _{1.7}	95.6 _{0.1}	70.4 _{1.1}	96.1 _{0.1}	86.7
1:0.25	86.4 _{0.2}	89.9 _{0.1}	76.9 _{4.2}	83.8 _{1.6}	91.9 _{1.9}	95.2 _{0.2}	69.9 _{1.6}	95.3 _{0.2}	86.2
1:0.1	85.3 _{0.5}	88.5 _{0.1}	76.3 _{2.8}	82.6 _{0.9}	91.7 _{2.2}	94.9 _{0.1}	69.4 _{0.8}	95.0 _{0.1}	85.5

Table 5: Text classification results of our method under different split ratios between target training and validation sets.

Method	CHEMPROT	RCT	ACL-ARC	SciERC	HYPERPARTISAN	AGNEWS	HELPFULNESS	IMDB	Average
BLO	85.3 _{0.4}	88.7 _{0.1}	75.4 _{2.9}	83.1 _{1.1}	90.7 _{4.5}	94.9 _{0.2}	68.3 _{2.2}	95.0 _{0.1}	85.2
Ours	87.1 _{0.2}	90.4 _{0.1}	77.6 _{2.3}	84.4 _{0.7}	92.4 _{1.2}	95.7 _{0.1}	70.9 _{0.8}	96.6 _{0.1}	86.9

Table 6: Ablation study results of the BLO method on text classification.

by splitting the original target training dataset, it is always possible to obtain a balanced split.

4.3.6 A Bi-level Optimization Based Ablation Setting

We also perform an ablation study which reduces the proposed four-level optimization problem to a BLO problem. The study was conducted on the text classification task. For each source example $d_{s,i}$, we learn a weight $b_i \in [0, 1]$. A larger weight indicates that $d_{s,i}$ is more likely to be in the target domain. Let $B = \{b_i\}_{i=1}^{N_s}$ where N_s is the number of source examples. We use the Transformer (Vaswani et al., 2017) T to perform domain adaptation. It takes a source text t as input and generates an adapted text $f(t, T)$ which is expected to be in the target domain. The Gumbel-Softmax (Jang et al., 2017) trick is used to deal with the non-differentiability of generated texts. At the lower level in the BLO formulation, we train the target model U . We define the training loss on selected source data as:

$$L_s(U, B) = \sum_{i=1}^{N_s} b_i \ell_{tgt}(U, d_{s,i}), \quad (18)$$

and the training loss on adapted source data as:

$$L_a(U, B, T) = \sum_{i=1}^{N_s} (1 - b_i) \ell_{tgt}(U, f(d_{s,i}, T)). \quad (19)$$

The optimization problem at this level is:

$$U^*(B, T) = \operatorname{argmin}_U L_t(U) + \lambda (L_s(U, B) + L_a(U, B, T)), \quad (20)$$

where $L_t(U)$ is the training loss on target training data, as defined in Eq.(3). At the upper level,

Method	Accuracy
BO (Ruder and Plank, 2017)	79.9
MMG (Wang et al., 2019a)	82.1
LSI (Huan et al., 2021)	83.9
Separate	83.5
WMVL	81.7
H-Divergence	84.0
Our full method	88.2

Table 7: Human evaluation results.

we evaluate $U^*(B, T)$ on the target validation set $D_t^{(val)}$ and update B and T by minimizing the validation loss:

$$\min_{B, T} L(U^*(B, T), D_t^{(val)}). \quad (21)$$

The overall BLO formulation is:

$$\begin{aligned} & \min_{B, T} L(U^*(B, T), D_t^{(val)}) \\ & s.t. U^*(B, T) = \\ & \operatorname{argmin}_U L_t(U) + \lambda (L_s(U, B) + L_a(U, B, T)). \end{aligned} \quad (22)$$

Table 6 shows the results. As can be seen, this BLO method performs worse than our method. The reason is: Our method uses self-supervised learning to learn domain discrepancy (in stage I) and leverages the learned domain discrepancy metric to perform domain adaptation (in stage II). Such mechanisms are lacking in the BLO method. This further demonstrates the necessity of stage I and II in our method.

4.4 Human Evaluation

We perform a human evaluation on whether the identified in-domain and out-of-domain source examples are indeed in or out of the target domain.

The study is performed on CHEMPROT, RCT, and ACL-ARC. For each dataset, we randomly sample 200 source texts. Three undergraduates were asked to label whether these source texts are in-domain or out-of-domain. Majority vote is leveraged to decide the final label. The Kappa score among the annotations is 0.75, which indicates a strong level of agreement among the annotators. Different methods are applied to predict whether each source text is in-domain or not. Table 7 shows the results. Our method achieves the best accuracy in identifying in-domain and out-of-domain source examples, due to its mechanism of learning the domain distance network in a discriminative way (as analyzed in Section 4.3).

5 Conclusions and Discussion

We propose a framework for simultaneous selection and adaptation of source examples in transfer learning. Our method automatically identifies which source examples are in or out of the target domain, and performs example-specific operations (either selection or adaptation). This is different from previous methods, which 1) discard out-of-domain source examples, leading to information loss; or 2) try to adapt all source examples into the target domain, incurring a risk of moving in-domain source examples outside the target domain. Our framework is based on four-level optimization. Experiments on text classification and visual question answering demonstrate the effectiveness of our proposed method.

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