Biased Representation Learning for Domain Adaptation

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Abstract

Representation learning is a promising technique for discovering features that allow supervised classifiers to generalize from a source domain dataset to arbitrary new domains. We present a novel, formal statement of the representation learning task. We argue that because the task is computationally intractable in general, it is important for a representation learner to be able to incorporate expert knowledge during its search for helpful features. Leveraging the Posterior Regularization framework, we develop an architecture for incorporating biases into representation learning. We investigate three types of biases, and experiments on two domain adaptation tasks show that our biased learners identify significantly better sets of features than unbiased learners, resulting in a relative reduction in error of more than 16% for both tasks, with respect to existing state-of-the-art representation learning techniques.

1 Introduction

Supervised natural language processing (NLP) systems have been widely used and have achieved impressive performance on many NLP tasks. However, they exhibit a significant drop-off in performance when tested on domains that differ from their training domains. (Gildea, 2001; Sekine, 1997; Pradhan et al., 2007) One major cause for poor performance on out-of-domain texts is the traditional representation used by supervised NLP systems (Ben-David et al., 2007). Most systems depend on lexical features, which can differ greatly between domains, so that important words in the test data may never be seen in the training data. The connection between words and labels may also change across domains. For instance, “signaling” appears only as a present participle (VBG) in WSJ text (as in, “signaling that...”), but predominantly as a noun (as in “signaling pathway”) in biomedical text.

Recently, several authors have found that learning new features based on distributional similarity can significantly improve domain adaptation (Blitzer et al., 2006; Huang and Yates, 2009; Turian et al., 2010; Dhillon et al., 2011). This framework is attractive for several reasons: experimentally, learned features can yield significant improvements over standard supervised models on out-of-domain tests. Moreover, since the representation-learning techniques are unsupervised, they can easily be applied to arbitrary new domains. There is no need to supply additional labeled examples for each new domain.

Traditional representations still hold one significant advantage over representation-learning, however: because features are hand-crafted, these representations can readily incorporate the linguistic or domain expert knowledge that leads to state-of-the-art in-domain performance. In contrast, the only guide for existing representation-learning techniques is a corpus of unlabeled text.

To address this shortcoming, we introduce representation-learning techniques that incorporate a domain expert’s preferences over the learned features. For example, out of the set of all possible distributional-similarity features, we might prefer those that help predict the labels in a labeled training data set. To capture this preference, we might bias a representation-learning algorithm towards features with low joint entropy with the labels in the training data. This particular biased form of
representation learning is a type of semi-supervised learning that allows our system to learn task-specific representations from a source domain’s training data, rather than the single representation for all tasks produced by current, unsupervised representation-learning techniques.

We present a novel formal statement of representation learning, and demonstrate that it is computationally intractable in general. It is therefore critical for representation learning to be flexible enough to incorporate the intuitions and knowledge of human experts, to guide the search for representations efficiently and effectively. Leveraging the Posterior Regularization framework (Ganchev et al., 2010), we present an architecture for learning representations for sequence-labeling tasks that allows for biases. In addition to a bias towards task-specific representations, we investigate a bias towards representations that have similar features across domains, to improve domain-independence; and a bias towards multi-dimensional representations, where different dimensions are independent of one another. In this paper, we focus on incorporating the biases with HMM-type representations (Hidden Markov Model). However, this technique can also be applied to other graphical model-based representations with little modification. Our experiments show that on two different domain-adaptation tasks, our biased representations improve significantly over unbiased ones. In a part-of-speech tagging experiment, our best model provides a 25% relative reduction in error over a state-of-the-art Chinese POS tagger, and a 19% relative reduction in error over an unbiased representation from previous work.

The next section describes background and previous work. Section 3 introduces our framework for learning biased representations. Section 4 describes how we estimate parameters for the biased objective functions efficiently. Section 5 details our experiments and results, and section 6 concludes and outlines directions for future work.

2 Background and Previous Work

2.1 Terminology and Notation

A representation is a set of features that describe data points. Formally, given an instance set \( \mathcal{X} \), it is a function \( \mathcal{R} : \mathcal{X} \rightarrow \mathcal{Y} \) for some suitable space \( \mathcal{Y} \) (often \( \mathbb{R}^d \)), which is then used as the input space for a classifier. For instance, a traditional representation for POS tagging over vocabulary \( V \) would include (in part) \( |V| \) dimensions, and would map a word to a binary vector with a 1 in only one of the dimensions. By a structured representation, we mean a function \( R \) that incorporates some form of joint inference. In this paper, we use Viterbi decoding of variants of Hidden Markov Models (HMMs) for our structured representations, although our techniques are applicable to arbitrary (Dynamic) Bayes Nets. A domain is a probability distribution \( D \) over the instance set \( \mathcal{X} \); \( R(D) \) denotes the induced distribution over \( \mathcal{Y} \).

In domain adaptation tasks, a learner is given samples from a source domain \( D_S \), and is evaluated on samples from a target domain \( D_T \).

2.2 Theoretical Background

Ben-David et al. (2010) give a theoretical analysis of domain adaptation which shows that the choice of representation is crucial. A good choice is one that minimizes error on the training data, but equally important is that the representation must make data from the two domains look similar. Ben-David et al. show that for every hypothesis \( h \), we can provably bound the error of \( h \) on the target domain by its error on the source domain plus a measure of the distance between \( D_S \) and \( D_T \):

\[
E_{x \sim D_T} \mathcal{L}(x,R,f,h) \leq E_{x \sim D_S} \mathcal{L}(x,R,f,h) + d_1(R(D_S),R(D_T))
\]

where \( \mathcal{L} \) is a loss function, \( f \) is the target function, and the variation divergence \( d_1 \) is given by

\[
d_1(D,D') = 2 \sup_{B \in \mathcal{B}} |Pr_D[B] - Pr_{D'}[B]| \tag{1}
\]

where \( \mathcal{B} \) is the set of measurable sets under \( D, D' \).

2.3 Problem Formulation

Ben-David et al.’s theory provides learning bounds for domain adaptation under a fixed \( R \). We now reformulate this theory to define the task of representation learning for domain adaptation as the following optimization problem: given a set of unlabeled instances \( U_S \) drawn from the source domain and unlabeled instances \( U_T \) from the target domain, as well as a set of labeled instances \( L_S \) drawn from...
the source domain, identify a function $R^*$ from the space of possible representations $\mathcal{R}$:

$$
R^* = \arg\min_{R \in \mathcal{R}} \left\{ \min_{h \in \mathcal{H}} \left( E_{x \sim D_S} \mathcal{L}(x, R, f, h) \right) + d_1(R(D_S), R(D_T)) \right\}
$$

Unlike most learning problems, where the representation $R$ is fixed, this problem formulation involves a search over the space of representations and hypotheses. The equation also highlights an important underlying tension: the best representation for the source domain would naturally include domain-specific features, and allow a hypothesis to learn domain-specific patterns. We are aiming, however, for the best general classifier, that happens to be trained on training data from one or a few domains. Domain-specific features would contribute to distance between domains, and to classifier errors on data taken from unseen domains. By optimizing for this combined objective function, we allow the optimization method to trade off between features that are best for classifying source-domain data and features that allow generalization to new domains.

Naturally, the objective function in Equation 2 is completely intractable. Just finding the optimal hypothesis for a fixed representation of the training data is intractable for many hypothesis classes. And the $d_1$ metric is intractable to compute from samples of a distribution, although Ben-David et al. propose some tractable bounds (2007; 2010). We view Equation 2 as a high-level goal rather than a computable objective. We leverage prior knowledge to bias the representation learner towards attractive regions of the representations space $\mathcal{R}$, and we develop efficient, greedy optimization techniques for learning effective representations.

2.4 Previous Work

There is a long tradition of research on representations for NLP, mostly falling into one of three categories: 1) vector space models and dimensionality reduction techniques (Salton and McGill, 1983; Turney and Pantel, 2010; Sahlgren, 2005; Deerwester et al., 1990; Honkela, 1997) 2) using structured representations to identify clusters based on distributional similarity, and using those clusters as features (Lin and Wu, 2009; Candito and Crabbé, 2009; Huang and Yates, 2009; Ahuja and Downey, 2010; Turian et al., 2010; Huang et al., 2011) 3) and structured representations that induce multi-dimensional real-valued features (Dhillon et al., 2011; Emami et al., 2003; Morin and Bengio, 2005). Our work falls into the second category, but builds on the previous work by demonstrating how to improve the distributional-similarity clusters with prior knowledge. To our knowledge, we are the first to apply semi-supervised representation learning techniques for structured NLP tasks.

Most previous work on domain adaptation has focused on the case where some labeled data is available in both the source and target domains (Daumé III, 2007; Jiang and Zhai, 2007; Daumé III et al., 2010). Learning bounds are known (Blitzer et al., 2007; Mansour et al., 2009). A few authors have considered domain adaptation with no labeled data from the target domain (Blitzer et al., 2006; Huang et al., 2011) by using features based on distributional similarity. We demonstrate empirically that incorporating biases into this type of representation-learning process can significantly improve results.

3 Biased Representation Learning

As before, let $U_S$ and $U_T$ be unlabeled data, and $L_S$ be labeled data from the source domain only. Previous work on representation learning with Hidden Markov Models (HMMs) (Huang and Yates, 2009) has estimated parameters $\theta$ for the HMM from unlabeled data alone, and then determined the Viterbi-optimal latent states for training and test data to produce new features for a supervised classifier. The objective function for HMM learning in this case is marginal log-likelihood, optimized using the Baum-Welch algorithm:

$$
\mathcal{L}(\theta) = \sum_{x \in U_S \cup U_T} \log \sum_y p(x, Y = y | \theta)
$$

where $x$ is a sentence, $Y$ is the sequence of latent random variables for the sentence, and $y$ is an instance of the latent sequence. The joint distribution in an HMM factors into observation and transition distributions, typically mixtures of multinomials:

$$
p(x, y | \theta) = P(y_1)P(x_1 | y_1) \prod_{i \geq 2} P(y_i | y_{i-1})P(x_i | y_i)
$$
Innocent bystanders are often the victims. For example, in Figure 1, we want to learn the best HMM distribution for the sentence “Innocent bystanders are often the victims” for POS tagging task. The hidden sequence \( y_1, y_2, y_3, y_4, y_5, y_6 \) can have any distribution \( p_1, p_2, p_3, ..., p_m, ..., p_n \) from the latent space \( \mathcal{Y} \). Since we are doing POS tagging, we want the distribution to learn the information encoded in the original POS labels “JJ NNS RB VBP DT NNS”. Therefore, by calculating the entropy between the hidden sequence and real labels, we can identify a subset of desired distributions that have low entropy, shown in the dotted oval. By minimizing the KL divergence between the learned distribution and the set of desired distributions, we can find the best distribution which is the closest to our desire.

The following subsections describe the specific properties we investigate; here we show how to incorporate them into the objective function. Let \( z \) be the sequence of labels in \( L_S \), and let \( \phi(x,y,z) \) be a property of the completed data that we wish the learned representation to minimize, based on our prior beliefs. Let \( Q \) be the subspace of the possible distributions over \( Y \) that have a small expected value for \( \phi: Q = \{ q(Y) | E_{Y \sim q}[\phi(x,Y,z)] \leq \xi \} \), for some constant \( \xi \). We then add penalty terms to the objective function (3) for the divergence between the HMM distribution \( p \) and the “good” distributions \( q \), as well as for \( \xi \):

\[
\mathcal{L}(\theta) - \min_{q,\xi} \left[ KL(q(Y)||p(Y|x,\theta)) + \sigma|\xi| \right] \quad (4)
\]

s.t. \( E_{Y \sim q}[\phi(x,Y,z)] \leq \xi \) \quad (5)

where \( KL \) is the Kullback-Leibler divergence, and \( \sigma \) is a free parameter indicating how important the bias is compared with the marginal log likelihood.

To incorporate multiple biases, we define a vector of properties \( \phi \), and we constrain each property \( \phi_i \leq \xi_i \). Everything else remains the same, except that in the penalty term \( \sigma|\xi| \), the absolute value is replaced with a suitable norm: \( \sigma ||\xi|| \). To allow ourselves to place weights on the relative importance of the different biases, we use a norm of the form \( \|x\|_A = \sqrt{x^TAx} \), where \( A \) is a diagonal matrix whose diagonal entries \( A_{ii} \) are free parameters that provide weights on the different properties. For our
experiments, we set the free parameters σ and A_{ii}
using a grid search over development data, as de-
scribed in Section 5.1

3.1 A Bias for Task-specific Representations

Current representation learning techniques are unsu-
servised, so they will generate the exact same represen-
tation for different tasks. Yet it is exceedingly rare that two state-of-the-art NLP systems for differ-
et tasks share the same feature set, even if they do
tend to share some core set of lexical features.

Traditional non-learned (i.e., manually-
engineered) representations essentially always
include task-specific features. In response, we pro-
pose to bias our representation learning such
that the learned representations are optimized for a
specific task. In particular, we propose a property
that measures how difficult it is to predict the labels
in training data, given the learned latent states.
Our entropy property uses conditional entropy of
the labels given the latent state as the measure of
unpredictability:

\[ \phi^{\text{entropy}}(y, z) = - \sum_i \tilde{P}(y_i, z_i) \log \tilde{P}(z_i|y_i) \] (6)

where \( \tilde{P} \) is the empirical probability and \( i \) indicates
the \( i \)th position in the data. We can plug this feature
into Equation 5 to obtain a new version of Equation
4 as an objective function for task-specific represen-
tations. We refer to this model as HMM+D+E. Un-
like previous formulations for supervised and semi-
supervised dimensionality reduction (Zhang et al.,
2007; Yang et al., 2006), our framework works effi-
ciently for structured representations.

3.2 A Bias for Domain-Independent Features

Following the theory in Section 2.2, we devise a bi-
ased objective to provide an explicit mechanism for
minimizing the distance between the source and tar-
gent domain. As before, we construct a property of
the completed data:

\[ \phi^{\text{distance}}(y) = d_1(\tilde{P}_S, \tilde{P}_T) \]

where \( \tilde{P}_S(Y) \) is the empirical distribution over lat-
tate space estimated from source-domain lat-
tate states, and similarly for \( \tilde{P}_T(Y) \). Essentially,
minimizing this property will bias the the represen-
tation towards features that appear approximately as
often in the source domain as the target domain. We
refer to the model trained with a bias of minimizing \( \phi^{\text{distance}} \) as HMM+D, and the model with both \( \phi^{\text{distance}} \) and \( \phi^{\text{entropy}} \) biases as HMM+D+E.

3.3 A Bias for Multi-Dimensional
Representations

Words are multidimensional objects. In English,
words can be nouns or verbs, singular or plural,
count or mass, just to name a few dimensions along
which they may vary. Factorial HMMs (FHMM-
s) (Ghahramani and Jordan, 1997) can learn multi-
dimensional models, but inference and learning are
complex and computationally expensive even in su-
pervised settings. Our previous work (2010) creat-
ed a multi-dimensional representation called an “I-
HMM” by training several HMM layers indepen-
dently; we showed that by finding several latent cat-
egories for each word, this representation can pro-
vide useful and domain-independent features for su-
pervised learners. In this work, we also learn a sim-
ilar multi-dimensional model (I-HMM+D+E), but
within each layer we add in the two biases described
above. While more efficient than FHMMs, the draw-
back of these I-HMM-based models is that there is
no mechanism to encourage the different HMM
models to learn different things. As a result, the lay-
ers may produce similar or equivalent features de-
scribing the dominant aspect of distributional sim-
ilarity in the data, but miss features that are less
strong, but still important, in the data.

To encourage learning a truly multi-dimensional
representation, we add a bias towards I-HMM mod-
els in which each layer is different from all previ-
sous layers. We define an entropy-based predictabili-
ty property that measures how predictable each pre-
vious layer is, given the current one. Formally, let \( y_i^l \)
denote the hidden state at the \( i \)th position in lay-
er \( l \) of the model. For a given layer \( l \), this property
measures the conditional entropy of \( y_m \) given \( y_i \),
summed over layers \( m < l \), and subtracts this from
the maximum possible entropy:

\[ \phi^{\text{predict}}_i(y) = \text{MAX} + \sum_{m < l} \tilde{P}(y_i, y_m) \log \tilde{P}(y_i|y_i^l) \]

The entropy between layer \( l \) and the previous layer-
s m measures how unpredictable the previous layers are, given layer l. By biasing the model such that MAX minus the entropy approaches zero, we encourage layer l towards completely different features from previous layers. We call the model with this bias P-HMM+D+E.

4 Efficient Parameter Estimation

Several machine learning paradigms have been developed recently for incorporating biases and constraints into parameter estimation (Liang et al., 2009; Chang et al., 2007; Mann and McCallum, 2007). We leverage the Posterior Regularization (PR) framework for our problem because of its flexibility in handling different kinds of biases; we provide a brief overview of the technique here, but see (Ganchev et al., 2010) for full details.

4.1 Overview of PR

PR introduces a modified EM algorithm to handle constrained objectives, like Equation 4. The modified E-step estimates a distribution \( q(Y) \) that is close to the current estimate of \( p(Y|x, \theta) \), but also close to the ideal set of distributions that (in expectation) have \( \phi = 0 \) for each property \( \phi \). The M step remains the same, except that it re-estimates parameters with respect to expected latent states computed with \( q \) rather than \( p \).

E step:

\[
q^{t+1} = \arg \min_{q} \min_{\xi} \text{KL}(q(Y)\|p(Y|x, \theta^t)) + \sigma \|\xi\| \quad \text{s.t. } \mathbb{E}[\phi(x, Y, z)] \leq \xi
\]

M step:

\[
\theta^{t+1} = \arg\max_{\theta} E_{q^{t+1}}[\log p(x, Y|\theta^t)]
\]

To make the optimization task in the E-step more tractable, PR transforms it to a dual problem:

\[
\max_{\lambda \geq 0, \|\lambda\|_1 \leq \sigma} -\log \sum_{Y} p(Y|x, \theta) \exp\{-\lambda \cdot \phi(x, Y, z)\}
\]

where \( \|\cdot\|_1 \) is the dual norm of \( \|\cdot\| \). The gradient of this dual objective is \( -E_{q}[\phi(x, Y, z)] \). A projected subgradient descent algorithm is used to perform the optimization.

4.2 Modifying \( \phi \) for Tractability

In unstructured settings, this optimization problem is relatively straightforward. However, for structured representations, we need to ensure that the dynamic programming algorithms needed for inference remain tractable for the biased objectives. For efficient PR over structured models, the properties \( \phi \) need to be decomposed as a sum over the cliques in the structured model. Unfortunately, the properties we mention above do not decompose so nicely, so we must resort to approximations.

In order to efficiently compute the expected value of the entropy property with respect to \( Y \sim q \), we need to be able to compute each component \( \mathbb{E}_{Y \sim q}[\phi_{\text{entropy}}(Y, z_i)] \) separately. Yet \( \tilde{P} \) depends on the setting of other latent states \( Y_j \) in the corpus. To avoid this problem, we pre-compute the expected empirical distributions over the completed data. For each specific value \( y \) and \( z \):

\[
\tilde{P}_q(y, z) = \frac{1}{|L_s|} \sum_{x} \sum_{i=1}^{|z|} 1[z_i = z] q(Y_i = y)
\]

\[
\tilde{P}_q(y) = \frac{1}{|L_s|} \sum_{x} \sum_{i=1}^{|z|} q(Y_i = y)
\]

These expected empirical distributions \( \tilde{P}_q \) can be computed efficiently using standard inference algorithms, such as the forward algorithm for HMMs. Note that \( \tilde{P}_q \) depends on \( q \), but unlike the original \( \tilde{P} \) from Equation 6, they do not depend on the data completions \( y \). Thus we can compute \( \tilde{P}_q \) once for each \( q^t \), and then substitute it for \( \tilde{P} \) for all values of \( Y \) in the computation of \( \mathbb{E}_{Y \sim q}[\phi_{\text{entropy}}(Y, z)] \), making this computation tractable. For the entropy-based predictability properties, the calculation is similar, but instead of using the label \( z \), we use the decoded states \( y^t_i \) from previous layers.

For the distance property, Ben-David et al.’s analysis depends on a particular notion of distance (Eqn. 1) that is computationally intractable. They also propose more tractable lower bounds, but these are again incompatible with the PR framework. Since no computationally feasible exact algorithm exists for this distance feature, we resort to a crude but efficient approximation of this measure: for each pos-
sible value $y$ of the latent states, we define:
\[
\phi_{dist}^y(y) = \sum_{i|\mathbf{x}_i \in U_g} 1\{y_i = y\} q(Y_i = y) / |U_g| - \sum_{i|\mathbf{x}_i \in U_T} 1\{y_i = y\} q(Y_i = y) / |U_T|
\]

Each of these individual properties is tractable for structured models. Combining these properties using the $||\cdot||_A$ norm results in a Euclidean distance (weighted by $A$) between the frequencies of features in each domain, rather than $d_1$ distance.

5 Experiments

We tested the structured representations with biases on two NLP tasks: Chinese POS tagging and English NER. In both cases, we use a domain adaptation setting where no labeled data is available for the target domain — a particularly difficult setting, but one that provides a strong test for an NLP system’s ability to generalize. In our work (Huang and Yates, 2009), we used a plain HMM for domain adaptation tasks in which there is labeled source data and unlabeled source and target data, but no labeled target data for training. Therefore, here, we use the HMM technique as a baseline, and build on it by including biases.

5.1 Chinese POS tagging

We use the UCLA Corpus of Written Chinese, which is a part of The Lancaster Corpus of Mandarin Chinese (LCMC). The UCLA Corpus consists of 11,192 sentences of word-segmented and POS-tagged text in 13 genres. We use gold-standard word segmentation labels during training and testing. The LCMC tagset consists of 50 Chinese POS tags. Each genre averages 5284 word tokens, for a total of 68,695 tokens among all genres. We use the ‘news’ genre as our source domain and randomly select 20% of every other genre as labeled test data. To train our representation models, we use the ‘news’ text, plus the remaining 80% of the texts from the other genres. We use 90% of the labeled news text for training, and 10% for development. We replace *hapax legomena* in the unlabeled data with the special symbol *UNKNOWN*, and also do the same for word types in the labeled test sets that never appear in our unlabeled training texts.

Following our previous HMM setup in (Huang and Yates, 2009) for consistency, we use an HMM with 80 latent states. For our multi-layer models, we use 7 layers of HMMs. We tuned the free parameters $\sigma$ and $A$ on development data. We varied $\sigma$ from 0.1 to 1000. To tune $A$, we start by setting the diagonal entry for $\phi_{\text{entropy}}$ to 1, without loss of generality. We then tie all the entries in $A$ for $\phi_{dist}$ to a single parameter $\alpha$, and tie all of the entries for $\phi_{predict}$ to a parameter $\beta$. We vary $\alpha$ and $\beta$ over the set \{0.01,0.1,1,10,100\}. Figure 2 shows our results for $\sigma$ and $\alpha$ on news development data. A setting of $\alpha = 0.01$ and $\sigma = 100$ performs best, with all $\sigma = 100$ doing reasonably well. Results for each of these models on the general fiction test text confirm the general trends seen on development data — a comforting sign, since this indicates we can optimize the free parameters on in-domain development data, rather than requiring labeled data from the target domain. Our models tended to perform better with increasing $\beta$ on development data, though with diminishing returns. We pick the largest setting tested, $\beta = 100$, for our final models.

We use a linear-chain Conditional Random Field (CRF) for our supervised classifier. To incorporate the learned representations, we use the Viterbi Algorithm to find the optimal latent state sequence from each HMM-based model and then use the optimal states as features in the CRF. Table 1 presents the full list of features in the CRF. To handle Chinese, we add in two features introduced in previous work (Wang et al., 2009): radical features and repeated characters. A radical is a portion of a Chinese character that consists of a small number of pen or brush strokes in a regular pattern.
To evaluate on a second task, we turn to Named Entity Recognition. We use the training data from the General Fiction Domain (test data) to evaluate. We found that incorporating the multi-dimensional bias (P-HMM+D+E) adds an additional 0.6% improvement.

Our interpretation for the success of I-HMM+D+E and P-HMM+D+E is that the increase in the state space of the models yields improved performance. Because P-HMM+D+E biases against redundant states found in I-HMM+D+E, it effectively increases the state space beyond I-HMM+D+E. Ahuja and Downey (2010) and our own work with HMMs as representations (2010) have previously shown that increasing the state space of the HMM can significantly improve the representation, but memory constraints eventually prevent further progress this way. The I-HMM+D+E and P-HMM+D+E models can provide similar benefits, but because they split parameters across multiple HMMs, they can accommodate much greater state spaces in the same amount of memory.

We also tested the entropy and distance biases separately. Figure 4 shows the result of the distance-biased HMM+D on the general-fiction test text, as we vary \( \sigma \) over the set \{0.1,1,10,100,1000\} (we observed similar results for other domains). For all values of \( \sigma \), the biased representation outperforms the unbiased HMM. There is also a strong negative correlation between the expected value of \( \|\phi_{\text{distance}}\| \) and the resulting accuracy, as expected from Ben-David et al.’s theoretical analysis. The HMM+E model outperforms the HMM on the (source) news domain by 0.3%, but actually performs worse for most target domains. We suspect that the entropy feature, which is learned only from labeled source-domain data, makes the representation biased towards features that are important in the source domain only. However, after we add in the distance bias and a parameter to balance the weights from both biases, the representation is able to capture the label information as well as the target domain features. Thus, the representation won’t solely depend on source data. HMM+D+E, which combines both biases, outperforms HMM+D, suggesting that task-specific features for domain adaptation can be helpful, but only if there is some control for the domain-independence of the features.

5.2 English Named Entity Recognition

To evaluate on a second task, we turn to Named Entity Recognition. We use the training data from the
Table 2: POS tagging accuracy: The P-HMM+D+E tagger outperforms the unbiased HMM tagger and the Stanford tagger on all target domains. The ‘avg’ column includes source-domain development data results. Differences between the P-HMM+D+E and the Stanford tagger are statistically significant at $p < 0.01$ on average and on 11 out of 12 target domain. We used the two-tailed Chi-square test with Yates’ correction.

Table 3: English Named Entity recognition results

![Figure 4: Greater distance between domains correlates with worse target-domain tagging accuracy.](image)

CoNLL 2003 shared task for our labeled training set, consisting of 204k tokens from the newswire domain. We tested the system on the MUC7 formal run test data, consisting of 59k tokens of stories on the telecommunications and aerospace industries.

To train our representations, we use the CoNLL-L training data and the MUC7 training data without labels. We again use a CRF, with features introduced by Zhang and Johnson (2003) for our baseline. We use the same setting of free parameters from our POS tagging experiments.

Results are shown in Table 3. Our best biased representation P-HMM+D+E outperformed the unbiased HMM representation by 3.6%, and beats the I-HMM+D+E by 1.6%. The domain-distance and multi-dimensional biases help most, while the task-specific bias helps somewhat, but only when the domain-distance bias is included. The best system tested on this dataset achieved a slightly better F1 score (78.84) (Turian et al., 2010), but used a much larger training corpus (they use RCV1 corpus which contains approximately 63 million tokens). Other studies (Turian et al., 2010; Huang et al., 2011) have performed a detailed comparison between these types of systems, so we concentrate on comparisons between biased and unbiased representations here.

5.3 Does the task-specific bias actually help?

In this section, we test whether the task-specific bias (entropy bias) actually learns something task-specific. We learn the entropy-biased representations for two tasks on the same set of sentences, labeled differently for the two tasks: English POS tagging and Named Entity Recognition. Then we switch the representations to see whether they will help or hurt the performance on the other task. We randomly picked 500 sentences from WSJ section...
Table 4: Results of POS tagging and Named Entity recognition tasks with different representations. With the entropy-biased representation, the system has better performance on the task which the bias is trained for, but worse performance on the other task.

| Representation/Task | POS Accuracy | NER F1 |
|---------------------|--------------|--------|
| HMM                 | 88.5         | 66.3   |
| HMM+E(POS labels)   | 89.7         | 64.5   |
| HMM+E(NER labels)   | 86.5         | 68.0   |

0-18 as our labeled training data and 500 sentences from WSJ section 20-23 as testing data. Because WSJ data does not have gold standard NER tags, we manually labeled these sentences with NER tags. For simplicity, we only use three types of NER tags: person, organization and location. The result is shown in Table 4. When the entropy bias uses labels from the same task as the classifier, the performance is improved: about 1.2% in accuracy on POS tagging and 1.7% in F1 score on NER. Switching the representations for the tasks actually hurts the performance compared with the unbiased representation. The results suggest that the entropy bias does indeed yield a task-specific representation.

6 Conclusion and Future Work

We introduce three types of biases into representation learning for sequence labeling using the PR framework. Our experiments on POS tagging and NER indicate domain-independent biases and multi-dimensional biases significantly improve the representations, while the task-specific bias improves performance on out-of-domain data if it is combined with the domain-independent bias. Our results indicate the power of representation learning in building domain-agnostic classifiers, but also the complexity of the task and the limitations of current techniques, as even the best models still fall significantly short of in-domain performance. Important considerations for future work include identifying further effective and tractable biases, and extending beyond sequence-labeling to other types of NLP tasks.

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