Flowing on Riemannian Manifold: Domain Adaptation by Shifting Covariance

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Abstract—Domain adaptation has shown promising results in computer vision applications. In this paper, we propose a new unsupervised domain adaptation method called domain adaptation by shifting covariance (DASC) for object recognition without requiring any labeled samples from the target domain. By characterizing samples from each domain as one covariance matrix, the source and target domain are represented into two distinct points residing on a Riemannian manifold. Along the geodesic constructed from the two points, we then interpolate some intermediate points (i.e., covariance matrices), which are used to bridge the two domains. By utilizing the principal components of each covariance matrix, samples from each domain are further projected into intermediate feature spaces, which finally leads to domain-invariant features after the concatenation of these features from intermediate points. In the multiple source domain adaptation task, we also need to effectively integrate different types of features between each pair of source and target domains. We additionally propose an SVM based method to simultaneously learn the optimal target classifier as well as the optimal weights for different source domains. Extensive experiments demonstrate the effectiveness of our method for both single source and multiple source domain adaptation tasks.

Index Terms—Domain adaptation, riemannian manifold, support vector machine.

I. INTRODUCTION

In many real world applications, the domain of interest (i.e., the target domain) contains very few or even no labeled samples because the collection of labeled samples is generally expensive and time consuming. The goal of domain adaptation is to learn robust target classifiers in this scenario by leveraging a large number of labeled training samples from other domains (i.e., auxiliary/source domains). Domain adaptation methods have been successfully used in many computer vision applications (please refer to Section II for more details).

In domain adaptation, the samples from the source domain and the target domain often have different data distributions. To reduce the mismatch of data distributions, the technique which introduces intermediate domains (see Fig. 1) can be employed to extract domain invariant features to bridge the distribution mismatch between the source and target domain. Specifically, Gopalan et al. [1] proposed an unsupervised domain adaptation method [called sampling geodesic flow (SGF) in [2]] by representing each domain as one subspace spanned by principal components of samples. Then they constructed a few intermediate domains between the source and target domains by interpolating subspaces between the two subspaces. Gong et al. [2] further extended [1] and proposed a new kernel based approach called geodesic flow kernel (GFK) by integrating an infinite number of subspaces. However, modeling one domain as a subspace is not sufficient to represent the distribution difference between two domains, especially when the two subspaces intersect in a common subspace. We give two examples in Fig. 2, where we plot the samples from two domains (i.e., red points and blue triangles) in a 2-D intersected subspace of two domains. It can be observed that the distributions of the samples from two domains are different. However, existing subspace-based methods such as
[1], [2] cannot be used to reduce the distribution mismatch in the intersected subspace, since the principal angles between the two subspaces of source and target domains are zero degrees in the intersected subspace (see Section III-A for more details). In Section III, we propose a new unsupervised domain adaptation approach for object recognition, which can better reduce the domain distribution mismatch in all the feature dimensions without the aid of the subspaces as in [1] and [2]. According to [3], any symmetric positive-definite (SPD) matrix resides on a Riemannian manifold. Thus, we assume the covariance matrices of the samples from two domains can be represented as two distinct points on a Riemannian manifold. Then we propose a simple but effective approach to seek for a geodesic flow curve connecting these two points. Specifically, we first embed the two distinct points into the log-Euclidean space and interpolate a few isometric points in this space. By inversely mapping these interpolated points back into the original Riemannian manifold, we obtain a set of interpolated covariance matrices. Then, we learn the projection matrices from these covariance matrices by using PCA, with which the samples from both domains can be projected into a set of intermediate feature representations that bridge the two domains. Linear discriminant analysis (LDA) [4] can be further used to extract the discriminant features. Finally, we concatenate all the features to form domain-invariant features and employ the nearest neighbor (NN) or support vector machine (SVM) classifier for the single source domain adaptation task.

For multi-domain adaptation, the domain-invariant features from each pair of source and target domains are different, so the existing multiple source domain adaptation methods cannot be directly used to learn the optimal target classifier and select the most relevant source domains. In Section IV, we propose to simultaneously learn the optimal target classifier and the optimal weights for different source domains. In Section V, we conduct comprehensive experiments using the office [5] and the Caltech256 [6] datasets under both single source domain and multi-domain settings. The results clearly demonstrate the effectiveness of our DASC as well as our new SVM-based learning algorithm for domain adaptation tasks.

II. RELATED WORK

Domain adaptation methods have been used for document classification [7], object recognition [1], [2], [5], [8]–[10], object localization [11], face recognition [12], indoor location [13], event recognition [14], [15], and video concept detection [7], [16].

In general, these methods can be categorized as feature (transform)-based approaches and classifier-based approaches. The feature (transform)-based approaches try to learn domain-invariant features for domain adaptation. Pan et al. [1] proposed a metric learning method by enforcing the samples that are from the same class but different domains to be closer with each other, which was further generalized in [8] by learning an asymmetric nonlinear transformation. However, their methods utilized the labeled data from the target domain, and hence cannot be applied to unsupervised domain adaptation. Gopalan et al. [1] and Gong et al. [2] proposed two methods to reduce the domain distribution mismatch based on the Grassmann manifold assumption. Recently, Zhu proposed a probabilistic graphic model based method [18], and Shi et al. [10] proposed an information theory-based method for domain adaptation.

The classifier-based approaches [16], [19], [20] directly seek target classifiers (e.g., SVM-based classifiers) for domain adaptation. Yang et al. [19] developed Adaptive SVM (A-SVM) by leveraging the existing source classifiers. Duan et al. [16] proposed a new multiple kernel learning (MKL)-based approach for domain adaptation by simultaneously learning the optimal linear weight coefficients of base kernels and the target classifier. Besides the two categories, other methods were also proposed, such as structure correspondence learning [21], sample reweighting [22], and feature replication [23].

Multiple source domain adaptation methods [7], [15], [24], [25] have also been studied. Schweikert et al. [24] proposed to match the means of different domains based on kernel mean matching (KMM) [22]. Duan et al. [7], [15] proposed two multi-domain adaptation methods by leveraging or selecting source domains. Chattopadhyay et al. [25] also proposed to weight different source domains based on a smoothness regularizer. Hoffman et al. [9] proposed a new clustering-based approach to partition a dataset into latent domains and extended [8] for multi-domain adaptation by learning multiple transformations. Moreover, some theoretical results can be found in [26]–[28], and [29]. Please refer to [30] for a comprehensive survey on transfer learning.
III. PROPOSED METHOD

In this paper, we denote a vector/matrix by a lower-case/uppercase letter in boldface. The transpose of a vector or matrix is denoted by the superscript ‘T’. Moreover, we use $O_{m \times n}$ to represent an $m$-by-$n$ matrix with all zeros and $I_n$ to represent an $n$-by-$n$ identity matrix, respectively. We also define $1$ as the vector with all ones. The inequality $u = [u_1, \ldots, u_n] \geq 0$ means that $u_i \geq 0$ for $i = 1, \ldots, n$.

In the common setting of the domain adaptation (DA) problem, the source domain samples are labeled and the target domain contains no labeled samples or only a limited number of labeled samples, which are referred to as unsupervised DA and semi-supervised DA, respectively. In this paper, we focus on the unsupervised DA. However, the proposed approach can be easily applied to semi-supervised DA. Formally, let us denote $X = [x_1, \ldots, x_N] \in \mathbb{R}^{D \times N_s}$ as the source domain data and denote $y_i^s \in \{1, 2, \ldots, K\}$ as the label of the $i$th sample $x_i$, where $D$ is the feature dimension, $N_s$ is the total number of source samples and $K$ is the total number of classes. Similarly, we denote $X' = [x_1', \ldots, x_{N_t}'] \in \mathbb{R}^{D \times N_t}$ as the target domain data from the same $K$ classes, where $N_t$ is the total number of target samples.

A. Subspace Analysis and Motivation

For domain adaptation, the key issue is to reduce the mismatch on the data distributions of two domains. Since the intermediate domains can contain more shared information of labeled samples, which are referred to as unsupervised DA and semi-supervised DA, respectively. In this paper, we focus on the unsupervised DA. However, the proposed approach can be easily applied to semi-supervised DA. Formally, let us denote $X = [x_1, \ldots, x_N] \in \mathbb{R}^{D \times N_s}$ as the source domain data and denote $y_i^s \in \{1, 2, \ldots, K\}$ as the label of the $i$th sample $x_i$, where $D$ is the feature dimension, $N_s$ is the total number of source samples and $K$ is the total number of classes. Similarly, we denote $X' = [x_1', \ldots, x_{N_t}'] \in \mathbb{R}^{D \times N_t}$ as the target domain data from the same $K$ classes, where $N_t$ is the total number of target samples.

However, when two subspaces $S_1$ and $S_2$ intersect into a $r$-dim subspace $\hat{S} = S_1 \cap S_2$, the first $r$ principal angles $\theta_1, \ldots, \theta_r$ must be equal to zero-degree, where we only need to assign the $r$ principal components of the common subspace $\hat{S}$ to $[u_1 = v_1, u_2 = v_2, \ldots, u_r = v_r]$ according to (1) and (2). As a result, the first $r$ columns of the intermediate point $S(t)$, corresponding to the common subspace $\hat{S}$, remain unchangeable when $t$ ranges from $0$ to $1$. In this case, after projecting these samples into the intermediate subspaces as used in SGF and GFK, the distributions of the samples from two domains in this intersected subspace are still unchanged. Empirically, given two domains, their intersection subspace is usually nonempty especially when $d$ is larger than $D/2$, and meanwhile the distributions of samples from two domains are usually different in the intersection subspace. Two examples are shown in Fig. 2, where the samples are projected into the 2-D space by using the first two dimensions of PCA on the common subspace.

To address the above problem, we directly represent each domain by a covariance matrix instead of a subspace (i.e., a column-orthogonal matrix). Due to the robustness of the covariance matrix for characterizing the data distribution, we then use the intermediate covariance matrices between two domains to reduce the data distribution mismatch.

B. Domain Adaptation by Shifting Covariance

We denote the covariance matrices of the source domain and the target domain as $C_s$ and $C_t$, respectively, which are symmetric positive-definite (SPD) matrices. Since $D$-by-$D$ SPD matrices lie on a Riemannian manifold, so each domain corresponds to a point on this manifold. To bridge the source and target domains, we seek a geodesic path $g(t)|_{t=0}$ from $C_s$ to $C_t$ on the Riemannian manifold in which we expect $g(0) = C_s$, $g(1) = C_t$, and $g(t)$ is a covariance matrix flowing on the geodesic path with gradually changing from $C_s$ to $C_t$ while $t$ increases from $0$ to $1$.

By flowing on the geodesic path $g(t)$, the covariance difference between two domains can be gradually reduced and finally it reaches the target covariance. Nevertheless, it is non-trivial to seek such a geodesic path on a Riemannian manifold. Partial differential equations (PDEs) may be used, however, it is usually computationally expensive. Below we employ a much more efficient approach called Log-Euclidean metric which can achieve the same excellent theoretical properties on a Riemannian manifold [3].

C. Computation With Log-Euclidean Metric

Before introducing the Log-Euclidean metric, we first give the definition of matrix exponential $\exp(\cdot)$ and logarithm $\log(\cdot)$ of a covariance matrix $C$ as follows:

$$\begin{align*}
\exp(C) &= U \exp(A)U', & \log(C) &= U \log(A)U' \\
&\text{where } C = U A U, & \text{and } \log(C) = U \log(A) U'
\end{align*}$$

where $A = \log(C)$ is from singular value decomposition (SVD). Then, we further define the logarithmic multiplication $\odot$ and

\textbf{1 We can always add a small positive value to the diagonal elements when a covariance matrix is not SPD.}
the logarithmic scalar multiplication $\otimes$ on the SPD matrices space as

\[ C_1 \otimes C_2 = \exp(\log(C_1) + \log(C_2)), \]
\[ \lambda \otimes C_1 = \exp(\lambda \log(C_1)) = C_1^\lambda \]

where $C_1$ and $C_2$ are two SPD matrices and $\lambda$ is a scalar. When the SPD matrix space is associated with the multiplication $\otimes$, it is actually a lie group [3], which may induce a bi-invariant metric, i.e., Log-Euclidean metric in [3] (referred as LEM here). Formally, the distance of two SPD matrices $C_1$ and $C_2$ based on LEM can be written as

\[ \text{DIST}(C_1, C_2) = \| \log(C_1) - \log(C_2) \|_F \]

where $\| \cdot \|_F$ is the matrix Frobenius norm. With LEM, the SPD matrix space is isomorphically and isometrically mapped to the Euclidean space of symmetric matrices [3], which means many calculations on Riemannian manifold can be simplified in the Euclidean space. Specifically, we can calculate the interpolated point on the geodesic path $g(t)$ using matrix exponential and logarithm as

\[ C(t) = \exp \left( (1-t) \log(C') + t \log(C') \right). \]

Therefore, to seek the geodesic path from the source domain to the target domain, we first map the covariance matrices $C'$ and $C$ into the Log-Euclidean space with the matrix logarithm. Given any $0 \leq t \leq 1$, we then obtain the interpolated point along the line between $\log(C')$ and $\log(C)$ in the Euclidean space. After that, we use matrix exponential to obtain the SPD matrix on Riemannian manifold. The procedure is illustrated in Fig. 3.

D. Performing Recognition with DASC

As shown in Fig. 3, we denote these interpolated covariance matrices as $C_i |_{i=1}^{n}$, and also denote the covariance matrices of the source domain and the target domain as $C_0$ and $C_{n+1}$, respectively. These $n+2$ covariance matrices actually represent $n+2$ intermediate domains that bridge the source domain and the target domain.

To obtain the domain-invariant features, a common way is to project the original features from the source/target domain into those intermediate domains. To this end, we perform principal component analysis (PCA) [4] on each covariance matrix to obtain the projection matrix. Formally, we denote $P_j |_{j=0}^{n+1} \in \mathbb{R}^{D \times d}$ as the projection matrices obtained from those $n+2$ covariance matrices using PCA. For any sample $x_i$ in the source/target domain (i.e., $x_i'$ or $x_i'$), the projected feature by using $P_j$ can be computed by $\tilde{x}_j = P_j x_i$. Then, we perform LDA [4] using the labeled samples in each intermediate domain to obtain the discriminative representation of each sample, denoted by $z_j$. The labeled data are from the source domain (resp., both the source and the target domain) for unsupervised DA (resp., semi-supervised DA). Finally, the $n+2$ features are concatenated to form the domain-invariant feature by $z_i = [z_{0,0}, \ldots, z_{(n+1),1}]^T$. After obtaining the domain-invariant features for the samples in both domains, any type of classifier (e.g., SVM classifier or NN classifier) can be employed to perform the recognition task.

IV. MULTIPLE SOURCE DOMAIN ADAPTATION

In this section, we first introduce the problem of multiple source DA, then we propose an SVM based method to learn the target classifier as well as the weights of source domains. Finally, we discuss the most related work with our proposed method.

A. Problem

When multiple source domains are available, multiple geodesic paths can be constructed on Riemannian manifold by connecting each source domain with the target domain. Formally, given $M$ source domains, $X_1^{s,1}, X_2^{s,2}, \ldots, X_r^{s,M}$, we can obtain $M$ pairs of domain-invariant features after performing feature extraction with our DASC method, that is $(Z_1^{s,1}, Z_1^{t,1}), (Z_2^{s,2}, Z_2^{t,2}), \ldots, (Z_1^{s,M}, Z_1^{t,M})$ where $Z_m^{s} = [z_1^{s,m}, \ldots, z_N^{s,m}]$ and $Z_m^{t} = [z_1^{t,m}, \ldots, z_N^{t,m}]$ are the domain-invariant features of the $m$th source domain data $X^{s,m}$ and the target domain data $X^t$ generated from the $m$th geodesic path, $N_m$ and $N_t$ are the numbers of source and target domain samples. Note that the features from the same pair of source and target domains (e.g., $Z_l^{s,m}$ and $Z_l^{t,m}$ for any $m \neq \tilde{m}$) are homogenous because they are obtained by using the same projection matrix. However, the features generated from different pathes (e.g. $Z_l^{s,m}$ and $Z_l^{t,m}$ for any $m \neq \tilde{m}$) are heterogeneous because different projection matrices are used. A straightforward way for multiple source domain adaptation is to independently learn one classifier for each pair of source and target domains and then fuse multiple classifiers in a late-fusion fashion. However, some source domains might be more relevant to the target domain, so the corresponding classifiers should be more important for the final classification. To this end, we propose to jointly learn multiple classifiers as well as the optimal weights of source domains.
B. Formulation

To simplify the notation, we denote \( \left[ \mathbf{z}^m \right]_{m=1}^{M} \) as the samples using the \( m \)th type of domain-invariant features, where \( N^m \) is defined as \( N^m = N^m_l + N^m_u \) with \( N^m_l \) being the number of labeled samples and \( N^m_u \) being the number of unlabeled samples. In the task of unsupervised DA, the labeled samples come from source domains and unlabeled samples are the target domain data.

Given each type of domain-invariant features, we propose to train one binary SVM classifier by using the labeled samples from the source domain as well as the unlabeled samples from the target domain. Formally, let us denote the classifier for the \( m \)th type of features as \( f^m(x^m) = \mathbf{w}^m_m \phi_m(x^m) + b_m \) (i = 1, …, \( N^m \)), where \( \phi_m(x^m) \) is the nonlinear feature mapping function, \( \mathbf{w}_m \) and \( b_m \) are the weight vector and the bias of the SVM classifier, respectively. When learning those \( M \) classifiers, we also propose to learn the weight coefficient \( \gamma_m \) for each classifier \( f^m(x^m) \). To this end, we formulate the learning task as follows:

\[
\begin{align*}
\min_{\mathbf{w}^m_m, \alpha^m_m, \beta^m_m} & \quad \sum_{m=1}^{M} \left( \frac{1}{2} \| \mathbf{w}^m_m \|^2 + C \sum_{i=1}^{N^m} (\xi^m_i + \xi^m_m)^+ \right) \\
+ & \frac{\theta}{2} \left( \| f^m_m - y^m \|^2 + \gamma_m \| \mathbf{f}^m_m - \mathbf{v}_u \|^2 \right) \\
\text{s.t.} & \quad \mathbf{w}^m_m \phi_m(x^m) + b_m - f^m_m \leq \varepsilon + \xi^m_i, \xi^m_i \geq 0, \\
& \quad f^m_m - \mathbf{w}^m_m \phi_m(x^m) + b_m - \varepsilon \leq \xi^m_i, \xi^m_i \leq 0, \\
& \quad 1 \geq \gamma \geq 0
\end{align*}
\] (4)

where these terms are described as follows:

1) \( \gamma = [\gamma_1, \ldots, \gamma_M] \): the vector of weight coefficients for \( M \) classifiers;
2) \( y^m_i = [y^m_i, \ldots, y^m_{N^m_i}] \): the label vector of labeled samples from the \( m \)th pair of source and target domains, with \( y^m_i \in \{+1, -1\} \) for \( i = 1, \ldots, N^m \) ;
3) \( \mathbf{v}_u = [v_1, \ldots, v_{N_u}] \): the virtual labels of unlabeled samples in the target domain;
4) \( f^m = [f^m_1, \ldots, f^m_{N^m_i}] \): the expected decision value vector for the samples when using the \( m \)th type of features, where \( f^m_1 = [f^m_{N^m_i+1}, \ldots, f^m_{N^m_i}] \)
5) \( \xi^m_i \) and \( \xi^m_{i+m} \): slack variables for \( \epsilon \)-insensitive loss;
6) \( C \) and \( \theta \): two regularization parameters.

Specifically, the first term in (4) is to regularize the complexity of the learned classifiers, which is similar to MKL. The second term is the \( \epsilon \)-insensitive loss which enforces the decision values of learned classifiers (i.e., \( \mathbf{w}_m \phi_m(x^m) + b_m \)) to be close to the expected decision values (i.e., \( f^m_m \)).

1) Data-dependent Regularizer: The data-dependent regularizer in the third term of (4) not only enforces the expected decision values of unlabeled data from different classifiers to be consistent with the virtual labels, but also encourages the classifiers from relevant source domains to have higher weights. Specifically, if the \( m \)th source domain is more relevant to the target domain, then the expected decision values (i.e., \( f^m_m \)) should be closer to the virtual labels of unlabeled data \( \mathbf{v}_u \). In this case, it will encourage the corresponding \( \gamma_m \) to be larger. On the other hand, a larger \( \gamma_m \) will also enforce the corresponding \( \mathbf{f}_u^m \) to be closer to \( \mathbf{v}_u \).

2) Virtual Labels: To construct the virtual label vector \( \mathbf{v}_u \), we first train an SVM classifier for each pair of domains separately, and use them to predict the decision values of the unlabeled samples. Given an unlabeled sample \( \mathbf{x} \), let us denote these \( M \) decision values by \( p_m |_{m=1}^{M} \), then the virtual label of \( \mathbf{x} \) is calculated by using the weighted sum of these \( M \) decision values, which is defined as \( \sum_{m=1}^{M} e^{-d_m^2} p_m \) where \( d_m \) is the distance between \( \mathbf{x} \) and the closest sample from the \( m \)th source domain, the parameter \( \delta \) is used to decide the weights of local neighbors.

C. Optimization

We alternatively optimize the weight coefficient vector \( \gamma \) and other variables related to the classifier.

1) Fix \( \gamma \): When fixing \( \gamma \), the optimization problem in (4) is separable with respect to \( m, m = 1, \ldots, M \). We therefore optimize these \( M \) subproblems one by one. To simplify the presentation, we omit the subscript \( m \) below unless necessary.

For each subproblem, we introduce the Lagrangian multipliers \( \alpha_l \)'s and \( \eta_i \)'s (resp. \( \alpha_u \)'s and \( \eta_i \)'s) for the constraints in (5) (resp. (6)), and set the derivatives of the Lagrangian with respect to the primal variables \( (\mathbf{f}^m, \mathbf{w}_m, b_m, \xi^m_i, \xi^m_{i+m}) \) to zeros, respectively. Then, we obtain the following equations:

\[
\begin{align*}
\mathbf{w}_m = & \gamma_m \Phi_m (\alpha^*_m - \alpha) , \\
\mathbf{f}^m = & \left( \gamma^m \right)_l \Phi_m \left( \mathbf{I}^{N^m_l}_{N^m_l} \mathbf{O}^{N^m_l}_{N^m_l} \right) \frac{1}{1 + \theta} \mathbf{v}_u
\end{align*}
\] (8)

as well as the constraints: \( \mathbf{1}' \alpha = 1 \alpha^* \), \( 0 \leq \alpha, \alpha^* \leq C \) where \( \alpha = [\alpha_1, \ldots, \alpha_N]' \), \( \alpha^* = [\alpha_1^*, \ldots, \alpha_N^*]' \), and \( \Phi_m = \Phi_m \left( \Phi_m ^T \right) \).

Substituting those equations and constraints back into the Lagrangian, we arrive at the following dual form:

\[
\begin{align*}
\min_{\alpha, \alpha^*} & \quad \frac{1}{2} (\alpha - \alpha^*)^T \mathbf{K}_m (\alpha - \alpha^*) \\
+ & \frac{\gamma^m}{2} (\alpha - \alpha^*)^T + \frac{\eta}{2} (\alpha + \alpha^*) \\
\text{s.t.} & \quad \mathbf{1}' \alpha = 1 \alpha^*, \quad 0 \leq \alpha, \alpha^* \leq C
\end{align*}
\] (9)

where \( \gamma^m = \| \gamma^m \|_1 \), \( \mathbf{K}_m = \gamma_m \mathbf{K}_m + \frac{1}{2} \mathbf{I}^{N^m_l}_{N^m_l} \mathbf{O}^{N^m_l}_{N^m_l} \frac{1}{1 + \theta} \mathbf{I}_{N^m_l} \), \( \mathbf{K}_m = \Phi_m \Phi_m ^T \) is the kernel matrix using the \( m \)th type of features. Actually, the above dual formulation is in analogous to the support vector regression (SVR) formulation, and can be readily solved with LIBSVM [32].

2) Fix \( \gamma^m, \mathbf{w}_m \) and \( b_m \): When each classifier is known, \( \gamma^m |_{m=1}^{M} \) can be optimized as

\[
\begin{align*}
\min_{\gamma} & \quad \frac{1}{2} \sum_{m=1}^{M} \left( \frac{1}{\gamma_m} \| \mathbf{w}_m \|^2 + \theta \gamma_m \| \mathbf{f}^m - \mathbf{v}_u \|^2 \right) \\
\text{s.t.} & \quad 1 \geq \gamma \geq 0.
\end{align*}
\] (10)
Algorithm 1: Multiple Source Domain Adaptation

Input: Source domain data \(X^{s,m}(m = 1, \ldots, M)\) with the corresponding label vector \(y^{m}_T\), and unlabeled target domain data \(X^t\).

1. Obtain \(M\) domain-invariant features \([z^{m}_i, i = 1, \ldots, N^m_i, \ldots, N^m], m = 1, \ldots, M\) by using our DASC method as described in Section III.
2. Generate the virtual label vector \(v_u\) (Section IV-B);
3. Initialize \(t \leftarrow 1\);
4. repeat
5. if \(t = 1\) then
6. \hspace{1em} Set the weight coefficient vector \(\gamma \leftarrow 1/M\).
7. else
8. \hspace{1em} Based on the learnt \((\alpha_m, \alpha^*_m)\), calculate \(w_m\) and \(f^m\) by using (8) and (9), respectively.
9. \hspace{1em} Solve for the weight coefficient vector \(\gamma\) as
10. \hspace{2em} \(\gamma_m = \min \{(\sqrt{\|f^m\|^2}/\theta \|f^m - v_u\|^2), 1\}\) and \(\gamma \leftarrow \gamma / (1/\gamma)\).
11. Obtain \((\alpha_m, \alpha^*_m)\) by solving \(M\) SVR problems one by one as in (10) using LIBSVM [32].
12. Set \(t \leftarrow t + 1\);
13. until The change of the objective in (4) is less than a predefined threshold.

Testing:

14. Predict the labels of target domain samples by using (13).

Output: The labels of target domain samples.

where \(w_m\) and \(f^m\) can be respectively calculated from (8) and (9) by using the learnt dual variables \((\alpha_m, \alpha^*_m)\).

The above problem is a convex optimization problem with a box constraint. By setting the derivative with respect to \(\gamma\) to zero and applying the box constraint, we reach the solution as
\[\gamma_m = \min \{(\sqrt{\|w_m\|^2}/\theta \|f^m - v_u\|^2), 1\}\]
and then normalize \(\gamma\) as \(\gamma \leftarrow \gamma / (1/\gamma)\).

D. Algorithm

We describe the whole algorithm for multiple source domain adaptation in Algorithm 1. We first initialize the weight coefficient vector to \(1/M\). Then, we solve the \(M\) SVR problems one by one as in (10) to obtain the dual variables \((\alpha_m, \alpha^*_m)\). After that, we can calculate \(w_m\) and \(f^m\), respectively, using (8) and (9), and obtain \(\gamma\) by solving (12). The weight coefficient vector \(\gamma\) is further normalized as \(\gamma \leftarrow \gamma / (1/\gamma)\). We repeat the above two steps until the change of the objective value in (4) is less than a predefined threshold. Actually, this algorithm always converges in our experiments (see Section V-D for the details).

Finally, the prediction of a target unlabeled sample \(x\) can be obtained by fusing these \(M\) classifiers, that is
\[f(x) = \sum_{m=1}^{M} \gamma_m(x)(\Phi_m(z^m)\Phi_m(\alpha^*_m - \alpha_m) + b_m)\]
(13)
where \(z^m\) is the \(m\)th feature of \(x\), and \(\alpha_m\) and \(\alpha^*_m\) are the learnt dual variables from the \(m\)th subproblem in (10).

V. Experiments

In this section, we evaluate our methods on two unsupervised domain adaptation settings: single source domain adaptation and multiple source domain adaptation.

A. Experimental Setup

For single source domain adaptation, we strictly follow the experimental setting in [2] by using four datasets with ten common classes, i.e., Amazon, Webcam, and DSLR collected in [5], and Caltech-256 [6]. Similar to [1], [2], and [5], we extract SURF features [33] and encode the images with 800-bin token frequency (TF) feature by using a codebook trained from a subset of Amazon images. Then, the features are normalized and z-scored to have zero mean and unit standard deviation in each dimension.
We treat each dataset as one domain, and perform the unsupervised domain adaptation task using each pair of domains, so in total we have 12 cases. Twenty labeled samples (resp., eight labeled samples) per class are selected randomly as training data when using Amazon, Webcam, and Caltech (resp., DSLR), as the source domain. All the samples are used as unlabeled training data when the dataset is used as the target domain. The test data is as the same as the unlabeled training data as in [1] and [2].

To validate the proposed learning method for multiple source domain adaptation, we additionally use ImageNet [34] and collect another dataset called Google-Image which contains the top ranked 100 images returned from Google-Image search engine by using the class name as the query. We perform the unsupervised multiple source domain adaptation task by leaving one dataset out as the target domain and using the rest datasets as five source domains. Due to the lack of label information on Google-Image dataset, we only use it as the source domain. For ImageNet, 20 labeled images per class are selected as training samples when it is used as one of the source domains. For other datasets, we use the same settings as in the single source domain adaptation task.

For all the settings, we perform 20 rounds of experiments with different randomly selected samples. We empirically set the number of intermediate covariance matrices to \( n = 8 \) as suggested in [1], and fix the feature dimension as \( d = 30 \) after using PCA, which results in 300-dim domain-invariant features. For LDA, we set the feature dimension to the class number minus one as in [4]. RBF kernel is used in SVM by setting the bandwidth parameter as the mean distance and we use the default tradeoff parameter (i.e. \( C = 1 \)). In multiple source DA, the tradeoff parameter \( \theta \) is set to 1. We set the bandwidth parameter \( \delta \) to 100, when calculating the virtual labels, and we observer the performance is stable when \( \delta \) is in the ranges of \([10, 1000]\).

### B. Results for Single Source Domain Adaptation

We compare our DASC with four baselines. OrigFeat, SGF [1], GFK [2], and information theoretical learning (ITL) [10], where OrigFeat uses the original TF features, SGF and its extended version GFK are the most relevant methods in which the subspaces are exploited, and ITL is a recently proposed unsupervised DA method. Specifically, in SGF [1], PCA is used to compute the subspaces for source
and target domains and then partial least squares (PLS) [35] is used to generate the discriminative features after interpolating intermediate subspaces, which is referred as SGF(PLS) here. In GFK [2], PCA is used to generate the subspace for the target domain while PCA and PLS are employed for the source domain, which are referred as GFK (PCA, PCA) and GFK (PLS, PCA) here, respectively. For our DASC, we employ PCA on the intermediate covariance matrices to obtain the projection matrices and then apply LDA to obtain the discriminative features, therefore we refer to it as DASC (PCA+LDA). To fairly compare with SGF and GFK, we additionally report SGF (PCA+LDA) which follows our method to use LDA for extracting the discriminative features and GFK (LDA, PCA) in which LDA is used to obtain the subspace for the source domain. Moreover, we also report the results only using PCA for SGF and our DASC, which are referred as SGF (PCA) and DASC (PCA), respectively.

Following [1], [2], and [10], we report the performances using the NN classifier for the above settings. For ITL, we strictly follow the validation scheme in [10] to select the best parameters. For SGF, we also strictly follow their parameter settings and report their best results by selecting the best subspace dimension $d$ according to the average accuracy over all the 12 cases. For GFK, we also follow their parameter settings, and the subspace dimension $d$ is determined by using the RoD criterion as in [2]. Moreover, we also report the results using the SVM classifier. We do not apply LDA for our DASC and other methods since label information can be exploited in SVM. But we still report the results for SGF and GFK when using PLS for dimension reduction or computing the subspace. Tables I and II summarize the mean accuracies and standard errors for all the 12 cases, and we also report the average accuracy over all cases for each method in Table II (see the last column).

We first consider the NN classifier using PCA only in which we do not use any label information when generating the features. Compared with OrigFeat, SGF (PCA), and GFK (PCA, PCA), our DASC (PCA) achieves the best results in 10 out of 12 cases, which indicates our domain-invariant features can better handle domain difference when compared with them. It also implies that it is better to reduce the data distribution mismatch by directly coping with the covariance matrix rather than using the subspace as in SGF and GFK (see the detailed discussion in Section III-B). We also note that DASC (PCA) generally achieves better results than ITL, which indicates it is better to shift covariance for domain adaptation rather than to use the information theory criterion as in ITL [10].

In general, domain adaptation methods are better than OrigFeat by using either NN or SVM classifier. When compared with other methods, our proposed method DASC (PCA+LDA) (resp. DASC) is the best in 9 (resp. 10) out of the 12 cases when using the NN (resp. SVM) classifier, and is comparable with other methods in the rest cases. In term of the average accuracy, our method is better than the second best result by 2.14% (resp. 1.45%) when using the NN (resp. SVM) classifier.

C. Results for Multiple Source Domain Adaptation

For multiple source unsupervised domain adaptation, we also compare our method with SGF [1] and GFK [2] as well as several state-of-the-art multiple source DA methods including multi-KMM [24], conditional probability-based multi-source
domain adaptation (CP-MDA) [25], DAM [7], and DSM [15].

The original TF features are used for [7], [15], [24], and [25] since all these methods are designed for homogenous source domains. We additionally compare two commonly used SVM baselines, SVM (single best) which is the best among all the SVMs trained on each source domain and SVM (fusion) which equally fuses the decision values from all the SVMs trained on different source domains. For SGF and GFK, we strictly follow their approaches in [1] and [2] for multiple source domain adaptation except using the SVM classifier to replace the NN classifier. For these baseline methods, we either use the recommended parameters in their papers or select the best ones according to the test results.

We report the mean accuracies in Table III. SVM (single best) generally performs better than SVM(fusion), which indicates it is necessary to select relevant source domains for multiple source domain adaptation. The proposed method achieves the best performances in four out of five cases. Even for the remaining one our method still achieves the second best result and the performance is also comparable with the best result. These results demonstrate the effectiveness of our domain-invariant features similarly as in the single source DA task. Moreover, the results also demonstrate our proposed learning method can effectively integrate different heterogeneous domain-invariant features and thus it achieves much better performance compared with other multiple source domain adaptation methods.

D. Convergence

We study the convergence of our multiple source domain adaptation algorithm as described in Algorithm 1. Fig. 4 shows the objective values of 20 rounds of experiments with different randomly selected training samples when using Caltech as the target domain. We can observe that our algorithm usually converges after 30−50 iterations. We also have similar observations in other cases.

E. Covariance Versus Subspace

We have discussed the advantages of using the covariance matrix on a Riemannian manifold when compared with using the subspace on a Grassmann manifold in Section III-B. SGF and GFK cannot bridge the data distribution mismatch in the common subspace of two domains. In contrast, our DASC does not suffer from such a problem because we directly cope with the covariance matrices for domain adaptation without the aid of subspaces. Fig. 5 shows the performance comparison of our DASC with SGF and GFK for the single source domain adaptation task using PCA and NN classifier. Our DASC is generally better than SGF and GFK in most cases when using different $d$’s (i.e., the dimension of subspaces for SGF and GFK and the feature dimension after using PCA for our DASC), which clearly demonstrates the effectiveness of our method.

VI. CONCLUSION

In this paper, we have proposed an effective method to generate domain-invariant features for unsupervised domain adaptation. Different from the existed methods such as SGF and GFK, which adopted the subspace assumption on a Grassmann manifold, we directly use the covariance matrix to represent a domain and construct a geodesic path between the source and target domains on a Riemannian manifold. Then, we extract domain-invariant features by projecting the samples onto the intermediate domains along the geodesic path. For the multiple source domain adaptation task, as the domain-invariant features from each pair of source and target domains may be different, we further propose a new SVM-based approach to simultaneously learn the target classifier as well as the optimal weights for multiple source domains. Extensive experimental results clearly demonstrate the effectiveness of our work.

REFERENCES


2Note that the results in Table I and II are obtained with the fixed parameter $d$, therefore the performances in this figure could be better by varying $d$. 

TABLE III

<table>
<thead>
<tr>
<th>Method</th>
<th>Caltech</th>
<th>Amazon</th>
<th>Webcam</th>
<th>DSLR</th>
<th>ImageNet</th>
</tr>
</thead>
<tbody>
<tr>
<td>SVM(single best)</td>
<td>40.14</td>
<td>53.04</td>
<td>76.59</td>
<td>69.68</td>
<td>31.21</td>
</tr>
<tr>
<td>SVM(fusion)</td>
<td>44.33</td>
<td>50.14</td>
<td>58.47</td>
<td>67.51</td>
<td>33.23</td>
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<tr>
<td>Multi-KMM [24]</td>
<td>47.07</td>
<td>54.22</td>
<td>64.54</td>
<td>70.44</td>
<td>35.69</td>
</tr>
<tr>
<td>CP-MDA [25]</td>
<td>42.90</td>
<td>47.81</td>
<td>53.35</td>
<td>46.88</td>
<td>30.73</td>
</tr>
<tr>
<td>DAM [7]</td>
<td>44.33</td>
<td>49.97</td>
<td>58.47</td>
<td>67.26</td>
<td>32.91</td>
</tr>
<tr>
<td>DSM [15]</td>
<td>33.68</td>
<td>42.59</td>
<td>45.25</td>
<td>66.30</td>
<td>27.50</td>
</tr>
<tr>
<td>SGF [1]</td>
<td>47.52</td>
<td>55.37</td>
<td>57.41</td>
<td>53.76</td>
<td>33.96</td>
</tr>
<tr>
<td>GFK [2]</td>
<td>39.79</td>
<td>50.26</td>
<td>45.00</td>
<td>42.42</td>
<td>26.57</td>
</tr>
<tr>
<td>Our method</td>
<td>49.17</td>
<td>57.90</td>
<td>84.63</td>
<td>80.13</td>
<td>35.48</td>
</tr>
</tbody>
</table>


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