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Network-based correlated correspondence for unsupervised domain adaptation of hyperspectral satellite images

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Abstract—Adapting a model to changes in the data distribution is a relevant problem in machine learning and pattern recognition since such changes degrade the performances of classifiers trained on undistorted samples. This paper tackles the problem of domain adaptation in the context of hyperspectral satellite image analysis. We propose a new correlated correspondence algorithm based on network analysis. The algorithm finds a matching between two distributions, which preserves the geometrical and topological information of the corresponding graphs. We evaluate the performance of the algorithm on a shadow compensation problem in hyperspectral image analysis: the land use classification obtained with the compensated data is improved.

I. INTRODUCTION

Domain adaptation problems occur naturally in many applications of machine learning to real-world datasets [1]. In remote sensing image analysis this problem arises frequently, since the acquisition conditions of the images (cloud cover, acquisition angle, seasonal variations) are most often different. As a consequence, even if the images contain the same type of objects, the observed data distribution undergoes a d-dimensional and often nonlinear spectral distortion, i.e. a distortion that is local, class-specific and that impacts differently each region of the electromagnetic spectrum [2], [3].

One way to solve this problem is to perform an adaptation between the two d-dimensional image domains, in order to achieve a relative compensation of the shift by matching the data clouds to each other. Provided that the data are expressed as graphs and embed a topological structure, this problem can be seen as a graph matching problem [4]. In hyperspectral remote sensing, the adaptation problem has been tackled equivalently in [5], where the authors use the distance between nodes to constrain the possible assignments and enforce structure preservation of the graph.

Recent alternative methodologies consider the domains as manifold to be aligned [6] and use labeled examples [7] or local geometric matching [8] to perform the alignment. If the first option has been successfully applied to remote sensing data in a multimodal setting [9], local geometric matching must be handled with care, as the datasets to be matched are of large size and comparing all possible arrangements of local patterns is computationally really expensive.

To find the best matching, one can resort to energy minimization on Markov Random Fields (MRF). In the last years, this topic has received a lot of attention in computer vision (e.g. for dense stereo matching, segmentation and image stitching) [10] and efficient algorithms can be used to solve the optimization, especially if it consists solely of unary and pairwise constraints. In hyperspectral remote sensing, MRF are widely used to enforce spatial constraints in the classifiers [11], but, to our knowledge, the only attempt to use them for domain adaptation is found in [12]. In that study, the two domains are considered as observations of a single hidden MRF. Nodes in the source and target graphs are matched by finding a hidden node that is likely to have generated both. To compute unary and pairwise potentials, a Gaussian distribution of the distances between nodes and of the edges lengths is assumed. In this paper, we propose a MRF formulation of the graph matching problem. As a first contribution, we extend the algorithm in [13], originally developed to register a pair of 3D meshes, to the d-dimensional case by using descriptors issued from network theory [14] and secondly propose an alternative optimization to reduce the number of candidate matches and allow working on larger graphs. Contrarily to [12], we rely on node descriptors that are more robust to deformations between source and target domains. We also avoid the requirement of having the same number of nodes in both domains. Experiments on relative shadowing compensation show important benefits for land use pixel classification of a dataset comprising a 144-channels hyperspectral image and a LiDAR digital surface model.

II. THE CORRELATED CORRESPONDENCE ALGORITHM (CC)

Consider two graphs, one for the source data, $S$, and one for the target data, $T$. Each graph is composed of a set of nodes, $V$, connected by a set of edges, $E$. The total number of nodes, $K$, can differ from one graph to another. We will refer to the source graph as $G^S = (V^S, E^S)$ and to the target graph as $G^T = (V^T, E^T)$, with $K^S$ and $K^T$ nodes, respectively.

The correlated correspondence algorithm (CC, [13]) solves a three-dimensional graph matching problem that consists in
finding correspondences between the nodes of both graphs. Each node in the source graph \( \{ V_k^S \}_{k=1}^K \) is associated with a correspondence variable, \( c_k \), whose value represents the index of the node in \( V^T \) to which \( V_k^S \) has been matched. The CC solution to the graph matching problem is given by the \((K^S \times 1)\) vector \( c \) containing all the nodes assignments.

CC solves this graph matching problem as an MRF energy minimization problem. Each node on the source graph is associated with a unary potential and a number of pairwise potentials. Unary potentials \( \psi(c_k) \) encode the dissimilarity between a source node and each of the target nodes. Pairwise potentials \( \psi(c_k, c_l) \) encode the cost of assigning two neighboring source nodes to two specific target nodes, penalizing the assignments producing strong changes in the topology of \( G^S \).

There are two such potentials: \( \psi_{n}(c_k, c_l) \), which penalizes an increase of the distance between neighboring nodes in \( G^S \) and \( \psi_{f}(c_k, c_l) \), which penalizes a decrease of the distance between non-neighboring nodes in \( G^S \). CC minimizes the energy function

\[
E(c) = \sum_{k \in K^S} \psi(c_k) + \sum_{k,l \in N^S} \psi_{n}(c_k, c_l) + \sum_{k,l \in F^S} \psi_{f}(c_k, c_l),
\]

where \( N^S \) is the set of neighboring nodes on the source graph and \( F^S \) is the set of nodes which are far from each other. Although minimizing \( E(c) \) is in general NP-hard, efficient approximate algorithms exist such as the Tree-Reweighted Message Passing algorithm (TRWS [15]) used in this paper and Loopy Belief Propagation [16].

### III. THE NETWORK-BASED CORRELATED CORRESPONDENCE ALGORITHM (netCC)

In this section, we detail the proposed network-based correlated correspondence (netCC) algorithm.

#### A. Node descriptors

The first step necessary to compute all the potentials in Eq. (1) is to obtain a set of descriptors used to match the nodes. These descriptors must reflect proximity for nodes to be matched and dissimilarity for nodes not to be matched. If the graphs are relatively similar in shape and position in the input space (as in [5]), the Euclidean distance between the \( d \)-dimensional vectors can be used. However, when strong distortions (scaling, local transformations) are observed, like in Fig. 2, this distance is likely to produce too many mismatches, as the nodes in the source graph would be matched to their direct neighbors. Authors in [13] use local surface signatures based on local 2D histograms. These local histograms are obtained after projecting the neighboring points onto the tangent plane associated with the point. Although this works well for 3D meshes, it is not straightforward to extend this method to the \( d \)-dimensional case (where we do not have surface information) and we therefore propose an alternative method.

In the proposed netCC, we compute a vector of descriptors, which includes several node characteristics invariant to rotation and translation of the graph (Fig. 1):

- **Closeness centrality** [14], \( V^c \): how far (in term of shortest path) a node \( i \) is from all the other nodes.
- **Eccentricity**, \( V^c \): the maximum shortest path distance between node \( i \) and all other nodes.
- **Median distance to the manifold**, \( V^m \): the median Euclidean distance between node \( i \) and all the points in the domain.
- **Density**, \( V^d \): the average Euclidean distance between node \( i \) and its neighbors in the domain.
- **Distance to center**, \( V^{dc} \): the Euclidean distance between node \( i \) and the graph’s gravity center.

To obtain a descriptor vector for each node, these measures are stacked as \( x_i = [V^c_i \ V^c_i \ V^m_i \ V^d_i \ V^{dc}_i]^T \).

#### B. Unary potentials

Each node in the source graph \( \{ V_k^S \}_{k=1}^K \) yields an unary potential that encodes the cost of assigning it to each node in the target graph \( \{ V_i^T \}_{i=1}^I \):

\[
\psi(c_k = i) = d(x_k^S, x_i^T),
\]

where \( d \) is the Euclidean distance between the vectors of descriptors of the two nodes.

#### C. Pairwise potentials

For the pairwise potentials, we follow [13] and define potentials that favor assignments preserving distances between nodes of the source graph \( G^S \). Such distances are computed as the shortest path distances on the graph between the two nodes, \( d_{sp}(V_1, V_2) \). We also normalize all the distances on both graphs by the maximum distance on each graph in order to be able to compare the distances between pairs of nodes computed on \( G^S \) with those on \( G^T \). Such a normalization only deals with global scaling problems (as those considered in the experiments presented), but should be modified for non-uniform local scalings.

**Nearness preservation potentials:** The nearness preservation potentials \( \psi_n \) favor assignments in which two neighboring nodes \( V_k^S \) and \( V_l^S \) on \( G^S \) will stay close after matching with \( G^T \). Two nodes \( V_i^T \) and \( V_j^T \) on \( G^T \) are considered close if their shortest path distance is lower than a threshold \( \alpha \).

\[
\psi_n(c_k = i, c_l = j) = \begin{cases} 0 & \text{if } d_{sp}(V_i^T, V_j^T) < \alpha \\ 1 & \text{otherwise} \end{cases}
\]

The threshold \( \alpha \) is to be chosen by the analyst. Since its value corresponds to the radius of a local neighborhood sphere, it should remain quite small (e.g. \( \alpha \in [0.15...0.35] \)), in order to avoid considering all the nodes as neighbors.

**Farness preservation potentials:** The farness preservation potential \( \psi_f \) encodes the constraint that two nodes, which are far on \( G^S \), should be assigned to nodes that are far on \( G^T \). Two nodes are considered far if their shortest path distance is greater than a threshold \( \beta \).

\[
\psi_f(c_k = i, c_l = j) = \begin{cases} 0 & \text{if } d_{sp}(V_i^T, V_j^T) > \beta \\ 1 & \text{otherwise} \end{cases}
\]
In principle, we admit that both farness and nearness potentials are considered for disjoint set of pairs, i.e. \( \beta > \alpha \).

### D. Computational complexity

We apply a number of optimizations to reduce the computational cost of the minimization of Eq. (1). One shortcoming of this formulation is that each node \( V^S_k \) of the source graph can be matched to any target node \( V^T_i \). This means that unary potentials will be stored as vectors of length \( K^T \) and pairwise potentials as matrices of size \( (K^T \times K^T) \). Since we have \( K^S \) such potentials, this leads to a \( O(K^S K^T^2) \) memory requirement for each pairwise potential.

To reduce the number of farness potentials, we follow the strategy in [13]. We first solve the energy minimization problem of Eq. (1) without the farness potentials and we then check which potentials are violated by the solution. These potentials are added to the objective function and the optimization is iterated until no additional potentials are required.

We also reduce the number of candidate matches among the target nodes. We do so by considering only the \( m \) closest target nodes in descriptor space. In this way, we empirically choose \( m = 30 \), which seems to be a good tradeoff between the number of missed matches and the computational gain. By using only \( m \) candidate target nodes, the memory requirement for pairwise potentials reduces to \( O(K^S m^2) \). In our tests, increasing this value did not yield any performance improvement, but this might depend on the dataset.

### E. Full algorithm

Algorithm 1 contains the detailed steps to perform domain adaptation using netCC. Creating a graph where each point in the domain is a node would result in a very large graph that would be impractical to work with. Instead, we use vector quantization and build a graph on the centroids found by the clustering algorithm for each domain. We use two levels of clustering:

- A fine level with \( C \) clusters. We use these graphs to approximate geodesic distance between two data points on the manifold [17].
- A coarse level where we subsample the fine level to \( K^S < C \), respectively \( K^T < C \) clusters. The resulting graphs are used in the netCC algorithm, but all distances are computed on the fine graph.

The algorithm defines a local transformation that can be used to transfer points between the two domains. Therefore, given a \( d \)-dimensional source dataset, netCC creates an adapted \( d \)-dimensional dataset composed of each source point transferred to the target domain. A classifier can then be trained on the target dataset and applied directly to the adapted source points.

### IV. DATA AND SETUP OF EXPERIMENTS

#### A. Dataset

We use the dataset of the 2013 IEEE GRSS Data Fusion Contest [18] which consists of an hyperspectral image and a Digital Surface Model (DSM) derived from airborne LiDAR at a 2.5m spatial resolution with an image size of 1903 \( \times \) 329. The hyperspectral image has 144 spectral bands. The goal is to classify each pixel into a land use class using its spectral features.

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**Algorithm 1 Domain adaptation with network-based correlated correspondence**

**Inputs**

- Source dataset \( X^S \in \mathbb{R}^d \)
- Target dataset \( X^T \in \mathbb{R}^d \)

1: Eigendecompose \( X^T \)
2: Project both datasets on the \( \text{npc} \) top eigenvectors
3: Cluster each dataset using \( k \)-means with \( C \) clusters
4: Build a \( k \)-NN graph per dataset using the cluster centers as nodes
5: Compute geodesic distances \( d_{sp} \) within each graph
6: Subsample graph \( G^S \) to keep \( K^S \) nodes
7: Subsample graph \( G^T \) to keep \( K^T \) nodes
8: Apply the netCC algorithm to find the optimal correspondence vector \( c \)
9: for each node \( k \in K^S \): do
10: Compute the displacement \( D_k = V^T_k - V^S_k \)
11: Apply the displacement to all the points in \( X^S \) assigned to node \( V^S_k \)
12: end for
13: Reconstruction : invert the projection to retrieve the \( d \)-dimensional data.
reach a performance of 0.36-0.47, which is acceptable.

The labels provided for this dataset are divided into two groups: one for training and one for testing. The training set only contains pixels that are not affected by the shadow while the testing set contains both lit and shadowed pixels.

### B. Experimental setup

The labels provided for this dataset are divided into two groups: one for training and one for testing. The training set only contains pixels that are not affected by the shadow while the testing set contains both lit and shadowed pixels.

To evaluate the performance of our adaptation procedure, we use a one-vs-one SVM with a Gaussian kernel. After model selection by cross-validation in the ranges $\gamma = [10^{-3}, ..., 10^2]$ and $C_{SVM} = [10^0, ..., 10^4]$, we obtained the optimal SVM parameters. For each pixel, the following features are used by the SVM:

- 144 spectral bands
- 1 LiDAR band
- 6 bands obtained by applying opening by reconstruction and closing by reconstruction on the LiDAR band with three different square structuring elements of sizes: $\{7, 19, 31\}$. Morphological filters are often used in remote sensing pixel classification to enforce the smoothness of the decision function in the spatial domain [19].

The spectral bands are normalized by their overall maximal value and the lidar band is normalized by its maximal value. We set the nearness threshold $\alpha = 0.2$ and the farness threshold $\beta = 0.3$. We use $C = 200$ for the fine graphs and experiment with two different values for $K^S = K^T = [50, 100]$. The value of $C$ depends on the dataset and should be set large enough to yield a good representation of data. Since the fine graphs are used only for the computation of the geodesic distances, which are computed only once at the beginning of the process, $C$ can be increased without impacting the performances.

We compare the proposed netCC algorithm with histogram matching applied on all the spectral bands. Since we are interested in the classification improvement after adaptation, we consider two separate testing scenarios: testing pixels under the shadow only and all testing pixels. In the experiments, we study the variability with regard to 2 parameters: the dimensionality of the space where the matching is performed (npc) and the number of nodes in the coarse graphs ($K^S$, $K^T$). For each choice of parameters, we run the whole graph matching procedure (clustering, matching) 10 times and report the average performances.

### V. Results and Discussion

With the setting described above, the proposed netCC algorithm converges in a limited number of iteration, typically around 100. This corresponds to an average running time of a few minutes on a laptop (Intel i5).

Table II shows the results when using only pixels under the shadow as test samples (around 1400 points). Without adaptation, we get a $\kappa$ score of 0.131, which is very low. We can see that by using histogram matching or netCC, we respectively reach a performance of 0.36-0.47, which is acceptable considering that this is a problem involving 15 classes, and some of them are very similar (e.g. there are three types of grass, two types of parking lots, two types of buildings). We obtain an improvement of 0.01 to 0.11 $\kappa$ points by using netCC over histogram matching. This corresponds to an increase in accuracy of about 11%. The choice of the number of centroids, $K^S$ and $K^T$, has an important influence on the results. This is of course dependent on the problem, but on our testing dataset, it seems $K^S = K^T = 50$ is enough to get a good representation of the data manifold and increasing to 100 mostly increases the number of clusters near the center of the

<table>
<thead>
<tr>
<th>Class</th>
<th>Color</th>
<th>Train</th>
<th>Test shadow</th>
<th>Test lit</th>
</tr>
</thead>
<tbody>
<tr>
<td>Healthy grass</td>
<td>Green</td>
<td>178</td>
<td>178</td>
<td>875</td>
</tr>
<tr>
<td>Stressed grass</td>
<td>Yellow</td>
<td>172</td>
<td>158</td>
<td>866</td>
</tr>
<tr>
<td>Synthetic grass</td>
<td>Orange</td>
<td>192</td>
<td>0</td>
<td>505</td>
</tr>
<tr>
<td>Trees</td>
<td>Brown</td>
<td>188</td>
<td>70</td>
<td>981</td>
</tr>
<tr>
<td>Soil</td>
<td>Gray</td>
<td>180</td>
<td>0</td>
<td>966</td>
</tr>
<tr>
<td>Water</td>
<td>Blue</td>
<td>182</td>
<td>0</td>
<td>143</td>
</tr>
<tr>
<td>Residential</td>
<td>Pink</td>
<td>196</td>
<td>78</td>
<td>986</td>
</tr>
<tr>
<td>Commercial</td>
<td>Purple</td>
<td>187</td>
<td>431</td>
<td>601</td>
</tr>
<tr>
<td>Road</td>
<td>Red</td>
<td>187</td>
<td>1</td>
<td>1031</td>
</tr>
<tr>
<td>Highway</td>
<td>Pink</td>
<td>191</td>
<td>326</td>
<td>707</td>
</tr>
<tr>
<td>Railway</td>
<td>Brown</td>
<td>172</td>
<td>142</td>
<td>848</td>
</tr>
<tr>
<td>Parking Lot 1</td>
<td>Green</td>
<td>192</td>
<td>0</td>
<td>1041</td>
</tr>
<tr>
<td>Parking Lot 2</td>
<td>Yellow</td>
<td>184</td>
<td>20</td>
<td>254</td>
</tr>
<tr>
<td>Tennis Court</td>
<td>Blue</td>
<td>181</td>
<td>0</td>
<td>247</td>
</tr>
<tr>
<td>Running Track</td>
<td>Pink</td>
<td>187</td>
<td>0</td>
<td>473</td>
</tr>
<tr>
<td>Sum</td>
<td></td>
<td>-2775</td>
<td>1404</td>
<td>10524</td>
</tr>
</tbody>
</table>

**TABLE I: Labeled pixels in the dataset**

Fig. 3: The 2013 GRSS DFC dataset.
This makes it harder to find a set of correspondences that satisfy all nearness potentials. On the other hand, we see that increasing the number of principal components \((n_{pc})\) has a positive effect on the performances. This is because with less principal components the reconstruction will be of lower quality.

Table III reports the results obtained when using all of the testing labels (about 12000 points). We get a small increase in performance when using netCC over histogram matching. The effect of DA is less visible due to the large number of test points in the lit area that do not benefit from adaptation.

Figure 4 shows classification maps for the right part of the image with the tested methods. Visually, the benefits of domain adaptation are obvious. We can see that without domain adaptation, the classifier fails to produce a meaningful result and classifies everything as water. The problem disappears when adapting the spectra prior to classification.

Figure 5 provides a magnified view of two areas where histogram matching and netCC result in different classifications. In the second row, we can see that histogram matching causes a grass area (visible on the labels and the flat lidar) to be classified as trees while netCC correctly classify it as grass. In the third row, a zoom on an urban area is shown. From the RGB and lidar images, it is clear this area contains buildings, some of which are misclassified as grass after histogram matching. With netCC, they are correctly classified as buildings, albeit not of the correct class. The same observation can be made for some roads which are misclassified as soil after histogram matching.

**VI. Conclusion**

In this paper, we propose an extension of the correlated correspondence algorithm for graph matching to handle \(d\) dimensional datasets. We propose the use of descriptors used in network analysis to guide the matching process. To reduce the computational costs, we use a two-levels graph and only allow matches between neighbors in the descriptor space.

Experiments on a challenging real-world dataset show good performances and a significant improvement over histogram matching. Future research directions could focus on the choice of descriptors and on the use of different potentials. Deeper exploration and sensitivity analysis of the various parameters would also be interesting, as well as the application to other datasets.

Another area for improvement is to study better ways to
transport the points for a given transformation of the nodes. In our current implementation, the local transformation is a simple translation, but more generic classes of transformation (e.g., affine) could be considered and possibly improve the performance of the classifier trained in the adapted domain.

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