Multi-class Semi-supervised Learning based Upon Kernel Spectral Clustering

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Abstract—This paper proposes a multi-class semi-supervised learning algorithm using kernel spectral clustering (KSC) as a core model. A regularized KSC is formulated to estimate the class memberships of data points in a semi-supervised setting using the one-vs-all strategy while both labeled and unlabeled data points are present in the learning process. The propagation of the labels to a large amount of unlabeled data points is achieved by adding the regularization terms to the cost function of the KSC formulation. In other words, imposing the regularization term enforces certain desired memberships. The model is then obtained by solving a linear system in the dual. Furthermore, the optimal embedding dimension is designed for semi-supervised clustering. This plays a key role when one deals with a large number of clusters.

Index Terms—Semi-supervised learning, kernel spectral clustering, low embedding dimension for clustering, multi-class problem.

I. INTRODUCTION

The incorporation of some form of prior knowledge of the problem at hand into the learning process is a key element that allows an increase of performance in many applications.

In many contexts, ranging from data mining to machine perception, obtaining the labels of input data is often difficult and expensive. Therefore in many cases one deals with a huge amount of unlabeled data, while the fraction of labeled data points will typically be small.

Semi-supervised algorithms aim at learning from both labeled and unlabeled data points. In fact in semi-supervised learning one tries to incorporate the labels (prior knowledge) in the learning process to enhance the clustering/classification performance. The semi-supervised learning can be classified into two categories, i.e. transductive and inductive learning. Transductive learning aims at predicting the labels for a specified set of test data by taking both labeled and unlabeled data together into account in the learning process. In contrast, in inductive learning the goal is to learn a decision function from a training set consisting of labeled and unlabeled data for future unseen test data points. Throughout this paper we refer to semi-supervised inductive learning as semi-supervised learning.

The semi-supervised inductive learning itself can be categorized into semi-supervised clustering and classification. The former addresses the problem of exploiting additional labeled data to adjust the cluster memberships of the unlabeled data. The latter aims at utilizing both unlabeled and labeled data to obtain a better classification model, and higher quality predictions on unseen test data points.

In some classical semi-supervised techniques, a classifier is first trained using the available labeled data points and then the labels for the unlabeled data points are predicted using out-of-extension. In the second step, unlabeled data that are classified with the highest confidence score are added incrementally to the training set and the process is repeated until the convergence is satisfactory [11]–[13]. Several semi-supervised algorithms have been proposed in the literature, see [4]–[10]. For instance, the Laplacian support vector machine (LapSVM) [6], is one of the graph-based methods with a data-dependent geometric regularization which provides a natural out-of-sample extension. The authors in [7] used local spline regression for semi-supervised classification by introducing splines developed in Sobolev space to map the data points to class labels. A transductive semi-supervised algorithm called ranking with Local Regression and Global Alignment (LRGA) to learn a robust Laplacian matrix for data ranking is proposed in [8]. In this approach, for each data point, the ranking scores of neighboring points are estimated using a local linear regression model. A label propagation approach in graph-based semi-supervised learning has been introduced in [9]. The authors in [10] developed a semi-supervised classification method based on class membership, motivated by the fact that similar instances should share similar label memberships.

Spectral clustering methods belong to a family of unsupervised learning algorithms that make use of the eigenspectrum of the Laplacian matrix of the data to divide a dataset into natural groups such that points within the same group are similar and points in different groups are dissimilar to each other [11]–[13].

Kernel spectral clustering (KSC) is an unsupervised algorithm that represents a spectral clustering formulation as a weighted kernel PCA problem, cast in the LS-SVM framework [14]. In contrast to classical spectral clustering, there is a systematic model selection scheme for tuning the parameters and also the extension of the clustering model to out-of-sample points is possible.

In [15], for the sake of dimensionality reduction, kernel maps with a reference point are generated from a least squares support vector machine core model via an additional regularization term for preserving local mutual distances together with reference point constraints. In contrast with the class of kernel eigenmap methods, the solution (coordinates in the low dimensional space) is characterized by a linear system instead of an eigenvalue problem.

Recently the authors in [16] have extended the kernel spectral clustering to binary semi-supervised learning (semi-KSC) by in-
corporating the information of labeled data points in the learning process. Therefore the problem formulation is a combination of unsupervised and binary classification approaches. Contrary to the approach described in [16], a non-parallel semi-supervised classification (NP-Semi-KSC) is introduced in [17]. It generates two non-parallel hyperplanes which are then used for out-of-sample extension.

It is the purpose of this paper to develop a new Multi-class Semi-Supervised KSC-based algorithm (MSS-KSC) using a one-versus-all strategy. In contrast to the methods described in [1]–[3], [6]–[9], in the proposed approach we start with a purely unsupervised algorithm as a core model and the available side information is incorporated via a regularization term. Given $Q$ labels, the approach is not restricted to find just $Q$ classes (semi-supervised classification) and instead it is able to uncover up to $2^Q$ hidden clusters (semi-supervised clustering). In addition, it uses low embedding dimension to reveal the existing number of clusters which is important when one deals with large number of clusters. There is a systematic model selection scheme for tuning the parameters and it is provided with the out-of-sample extension property. Furthermore the formulation is constructed for the multi-class semi-supervised classification and clustering.

Here KSC [14] is used as the core model. In this case thanks to the discriminative property of KSC one can benefit from unlabeled data points. Unlike the KSC approach that projects the data to a $k-1$ dimensional space for being able to group the data into $k$ clusters, in this paper the embedding dimension is rather equal to the number of available class-labels in the semi-supervised learning framework. Therefore the highlights of this manuscript can be summarized as follows:

- Using an unsupervised model as the core model and incorporating the available side-information (labels) through a regularization term.
- Addressing both multi-class semi-supervised classification and semi-supervised clustering.
- Extension of the binary case to multi-class case and addressing the encoding schemes.
- Realizing low embedding dimension to reveal the existing number of clusters.

This paper is organized as follows. In Section II a brief review about kernel spectral clustering is given. In Section III we formulate our multi-class semi-supervised classification algorithm using a one-vs-all strategy. In Section IV the semi-supervised clustering algorithm is discussed. The model selection of the proposed method is discussed in Section V. In Section VI numerical experiments are carried out to demonstrate the viability of the proposed method. Both synthetic and real-life data sets in different application domains such as in image segmentations and community detection in networks are considered.

II. BRIEF OVERVIEW OF KSC

The KSC method corresponds to a weighted kernel PCA formulation providing a natural extension to out-of-sample data i.e. the possibility to apply the trained clustering model to out-of-sample points. Given training data $D = \{x_i\}_{i=1}^M$, $x_i \in \mathbb{R}^d$, the primal problem of kernel spectral clustering is formulated as follows [14]:

$$\min_{w_1, b, \epsilon} \frac{1}{2} \sum_{\ell=1}^{k-1} w^{(\ell)} T w^{(\ell)} - \frac{1}{2M} \sum_{\ell=1}^{k-1} \gamma \epsilon^{(\ell)} T V \epsilon^{(\ell)}$$

subject to \( \epsilon^{(\ell)} = \Phi w^{(\ell)} + b^{(\ell)} 1_M, \ell = 1, \ldots, k - 1 \)

where \( \epsilon^{(\ell)} = [\epsilon_1^{(\ell)}, \ldots, \epsilon_M^{(\ell)}]^T \) are the projected variables and \( \ell = 1, \ldots, k - 1 \) indicates the number of score variables required to encode the \( k \) clusters. \( \gamma \epsilon \in \mathbb{R}^+ \) are the regularization constants. \( b^{(\ell)} \) is a bias term which is a scalar.

Here \( \Phi = [\varphi(x_1), \ldots, \varphi(x_M)]^T \) and a vector of all ones with size \( M \) is denoted by \( 1_M. \varphi(\cdot) : \mathbb{R}^d \rightarrow \mathbb{R}^h \) is the feature map and \( h \) is the dimension of the feature space which can be infinite dimensional. \( w^{(\ell)} \) is the model parameters vector in the primal. \( V = \text{diag}(v_1, \ldots, v_M) \) with \( v_i \in \mathbb{R}^+ \) is a user defined weighting matrix.

Applying the Karush-Kuhn-Tucker (KKT) optimality conditions one can show that the solution in the dual can be obtained by solving an eigenvalue problem of the following form:

$$V P_\ell \Omega \alpha^{(\ell)} = \lambda \alpha^{(\ell)}$$

where \( \lambda = M/\gamma \), \( \alpha^{(\ell)} \) are the Lagrange multipliers and \( P_\ell \) is the weighted centering matrix:

$$P_\ell = I_M - \frac{1}{1_M^T V 1_M} 1_M 1_M^T V$$

where \( I_M \) is the \( M \times M \) identity matrix and \( \Omega \) is the kernel matrix corresponding to \( i-j \)th entry \( \Omega_{ij} = (\varphi(x_i))^T \varphi(x_j) \). In the ideal case of \( k \) well separated clusters, for a properly chosen kernel parameter, the matrix \( V P_\ell \Omega \) has \( k-1 \) piecewise constant eigenvectors with eigenvalue 1.

It should be noted that no assumption about the data is made upon applying the KSC algorithm. Thanks to the bias term \( b^{(\ell)} \), as follows from one of the KKT conditions associated with the primal optimization (1), the kernel matrix gets automatically centered and it will be pre-multiplied by the centering matrix \( P_\ell \), in the dual (for more details see [14]).

The eigenvalue problem (2) is related to spectral clustering with random walk Laplacian. In this case, the clustering problem can be interpreted as finding a partition of the graph in such a way that the random walker remains most of the time in the same cluster with few jumps to other clusters, minimizing the probability of transitions between clusters. It is shown that if

$$V = D^{-1} = \text{diag}(\frac{1}{d_1}, \ldots, \frac{1}{d_M})$$

where \( d_i = \sum_{j=1}^{M} K(x_i, x_j) \) is the degree of the \( i \)-th data point, the dual problem is related to the random walk algorithm for spectral clustering.

From the KKT optimality conditions one can show that the score variables can be written as follows:

$$e^{(\ell)} = \Phi w^{(\ell)} + b^{(\ell)}$$

$$= \Phi \Phi^T \alpha^{(\ell)} + b^{(\ell)} = \Omega \alpha^{(\ell)} + b^{(\ell)}, \ell = 1, \ldots, k - 1$$

For the model selection, i.e. selection of the number of clusters \( k \) and kernel parameter, several criteria have been proposed in literature including a Fisher [18], Balanced Line
Fit (BLF) [14] or Silhouette criterion [19]. These criteria utilize the special structure of the projected out-of-sample points to estimate the out-of-sample eigenvectors for selecting the model parameters. When the clusters are well separated the out-of-sample eigenvectors show a localized structure in the eigenspace.

The out-of-sample extensions to test points \( \{x_i\}_{i=1}^{N_{\text{test}}} \) is done by an Error-Correcting Output Coding (ECOC) decoding scheme. First the cluster indicators are obtained by binarizing the score variables for test data points as follows:

\[
q^t_{\text{test}} = \text{sign}(\epsilon^t_{\text{test}}) = \text{sign}(\Phi^T_{\text{test}} w^{(t)} + b^{(t)})
\]

where \( \Phi^T_{\text{test}} = [\varphi(x_1), \ldots, \varphi(x_{N_{\text{test}}})]^T \) and \( \Omega_{\text{test}} = \Phi^T \).

The decoding scheme consists of comparing the cluster indicators obtained in the test stage with the codebook (which is obtained in the training stage) and selecting the nearest codeword in terms of Hamming distance.

In what follows we study two scenarios:

- In the first case the number of available class labels is equal to the actual number of existing classes.
- The second case corresponds to the situation where the number of available class labels is less than both the number of existing classes and the number of existing clusters.

In this paper, the terminology semi-supervised classification is used for referring to the first case and the problem of the second case is referred to as semi-supervised clustering.

### III. SEMI-SUPERVISED CLASSIFICATION

In this section we assume that there is a total number of \( Q \) classes \( (C_j, j = 1, \ldots, Q) \). The corresponding number of available class labels is also equal to \( Q \). Suppose the training data set \( \mathcal{D} \) consists of \( M \) data points and is defined as follows:

\[
\mathcal{D} = \{ \text{Unlabeled data} \}_{D_U} \cup \{ \text{Labeled data} \}_{D_L}
\]

where \( \{x_i\}_{i=1}^{M} \in \mathbb{R}^d \). The labels are available for the last \( N_L = M - N \) data points in \( \mathcal{D}_L \) and are denoted by

\[
Z = [z_1^T, \ldots, z_M^T]^T \in \mathbb{R}^{(M-N) \times Q},
\]

with \( z_i \in \{+1, -1\}^Q \) is the encoding vector for the training point \( x_i \).

In the proposed method we start with an unsupervised algorithm as a core model. Then by introducing a regularization term, we incorporate the available side information, which in this case are the labels, to the core model. Here the kernel spectral clustering is used as the core model. Because as it has been shown in [14] in contrast to classical spectral clustering, KSC has a systematic model selection scheme for tuning the parameters and it is provided with the out-of-sample extension property.

The one-vs-all strategy is utilized to build the codebook, i.e., the training points belonging to the \( i \)-th class are labeled by \(+1\) and all the remaining data from the rest of the classes are considered to have negative labels. Both the labeled and unlabeled data points are arranged such that the top \( N \) data points are the unlabeled ones and the rest, i.e., \( N_L \), are the labeled data points. We consider the labels of the unlabeled data points to be zero as in [16]. In our formulation unlabeled data points are only regularized using the KSC core model.

### A. PRIMAL-DUAL FORMULATION OF THE METHOD

We formulate the multi-class semi-supervised learning in the primal as the following optimization problem:

\[
\min_{w^{(t)}, b^{(t)}, \epsilon^{(t)}} \frac{1}{2} \sum_{\ell=1}^{Q} u^{(t)}_\ell w^{(t)}_\ell - \gamma_1 \sum_{\ell=1}^{Q} \epsilon^{(t)}_\ell V \epsilon^{(t)}_\ell + \gamma_2 \sum_{\ell=1}^{Q} \left( \epsilon^{(t)}_\ell - c^{(t)}_\ell \right)^T A \left( \epsilon^{(t)}_\ell - c^{(t)}_\ell \right)
\]

subject to \( \epsilon^{(t)} = \Phi w^{(t)} + b^{(t)}_\ell, \ell = 1, \ldots, Q \),

where \( c^{(t)}_\ell \) is the \( \ell \)-th column of the matrix \( C \) defined as

\[
C = \left[ \epsilon^{(1)}, \ldots, \epsilon^{(Q)} \right] M \times Q = \begin{bmatrix} 0_{N \times Q} & 0_{N_L \times N} \end{bmatrix} \frac{Z}{M \times Q},
\]

where 0 is a zero matrix of size \( N \times Q \) and \( Z \) is defined as previously. \( b^{(t)}_\ell \) is a bias term which is a scalar. The matrix \( A \) is defined as follows:

\[
A = \begin{bmatrix} 0_{N \times N} & 0_{N_L \times N} \end{bmatrix} \frac{I_{N \times N} \times L_{N \times N}}{L_{N \times N} \times L_{N \times N}}
\]

where \( I_{N \times N} \) is the identity matrix of size \( N_L \times N_L \).

The available prior knowledge, i.e., the labels, is added to the KSC model through the third term in the objective function of (3). This term aims at minimizing the difference between the score variables of the labeled data points, i.e., \( e_i \) for \( i \in D_L \), and the actual labels provided by the user. Therefore it enforces the \( e_i \) values for the labeled data points to be close enough to the actual labels in the projection space. Furthermore, since we do not intend to prejudge about the memberships of the unlabeled data points, the matrix \( A \) is taking place in the third term in the objective function.

**Lemma III.1.** Given a positive definite kernel function \( K : \mathbb{R}^d \times \mathbb{R}^d \rightarrow \mathbb{R} \) with \( K(x,z) = \varphi(x)^T \varphi(z) \) and regularization constants \( \gamma_1, \gamma_2 \in \mathbb{R}^+ \), the solution to (3) is obtained by solving the following dual problem:

\[
(I_M - RS \Omega) \alpha^{(t)} = \gamma_2 S \tau \epsilon^{(t)}, \ell = 1, \ldots, Q,
\]

where \( R = \gamma_1 V - \gamma_2 A \), \( \alpha^{(t)} = [\alpha_1^{(t)}, \ldots, \alpha_M^{(t)}]^T \) are the Lagrange multipliers and \( S = I_M - (1/\lambda_m^2) R M \), \( \lambda_m \) and \( I_M \) are defined as previously.

**Proof:** The Lagrangian of the constrained optimization problem (3) becomes

\[
\mathcal{L}(w^{(t)}, b^{(t)}, \epsilon^{(t)}, \alpha^{(t)}) = \frac{1}{2} \sum_{\ell=1}^{Q} u^{(t)}_\ell w^{(t)}_\ell - \gamma_1 \sum_{\ell=1}^{Q} \epsilon^{(t)}_\ell V \epsilon^{(t)}_\ell + \gamma_2 \sum_{\ell=1}^{Q} \left( \epsilon^{(t)}_\ell - c^{(t)}_\ell \right)^T A \left( \epsilon^{(t)}_\ell - c^{(t)}_\ell \right) + \sum_{\ell=1}^{Q} c^{(t)}_\ell \left( \epsilon^{(t)}_\ell - \Phi w^{(t)}_\ell - b^{(t)}_\ell \right)_1 M,
\]
where \( \alpha^{(\ell)} \) is the vector of Lagrange multipliers. Then the Karush-Kuhn-Tucker (KKT) optimality conditions are as follows,

\[
\begin{align*}
\frac{\partial L}{\partial w^{(\ell)}} &= 0 \rightarrow w^{(\ell)} = \Phi^T \alpha^{(\ell)}, \quad \ell = 1, \ldots, Q, \\
\frac{\partial L}{\partial \alpha^{(\ell)}} &= 0 \rightarrow 1^T_M \alpha^{(\ell)} = 0, \quad \ell = 1, \ldots, Q, \\
\frac{\partial L}{\partial c^{(\ell)}} &= 0 \rightarrow \alpha^{(\ell)} = (\gamma_1 V - \gamma_2 A)c^{(\ell)} + \gamma_2 e^{(\ell)}, \quad \ell = 1, \ldots, Q, \\
\frac{\partial L}{\partial b^{(\ell)}} &= 0 \rightarrow c^{(\ell)} = \Phi w^{(\ell)} + b^{(\ell)}1_M, \quad \ell = 1, \ldots, Q.
\end{align*}
\]

Elimination of the primal variables \( w^{(\ell)}, e^{(\ell)} \) and making use of Mercer’s Theorem [20], results in the following equation

\[
R\Omega \alpha^{(\ell)} + b^{(\ell)} R1_M = \alpha^{(\ell)} - \gamma_2 c^{(\ell)}, \quad \ell = 1, \ldots, Q.
\]

where \( R = \gamma_1 V - \gamma_2 A \). From the second KKT optimality condition and (7), the bias term becomes:

\[
b^{(\ell)} = (1/1^T_M R1_M)(-1^T_M \gamma_2 c^{(\ell)} - 1^T_M R\Omega \alpha^{(\ell)}), \quad \ell = 1, \ldots, Q.
\]

Substituting the obtained expression for the bias term \( b^{(\ell)} \) into (7) along with some algebraic manipulation one can obtain the solution in dual as the following linear system:

\[
\gamma_2 \left( I_M - \frac{R1_M 1^T_M}{1^T_M R1_M} \right) c^{(\ell)} = \alpha^{(\ell)} - R \left( I_M - \frac{1^T_M R}{1^T_M R1_M} \right) \Omega e^{(\ell)}. \tag{9}
\]

Remark III.1. It should be noted that since the optimization problem (3) does have equality constraints therefore the KKT conditions include the primal equality constraints and the gradient of the Lagrangian with respect to the primal variables [see [21, Chapter 5]]. In (6), the first three equations correspond to the derivative of the Lagrangian with respect to primal variables and the primal equality constraints are equivalently obtained by taking the derivative of the Lagrangian with respect to dual variables.

It should be noticed that one can also obtain the following linear system when the primal variables \( w^{(\ell)}, e^{(\ell)} \) are eliminated from (KKT) optimality conditions in (6):

\[
\begin{bmatrix}
\Omega - R^{-1} 1^T_M 1_M \\
1^T_M 0
\end{bmatrix}
\begin{bmatrix}
\alpha^{(\ell)} \\
b^{(\ell)}
\end{bmatrix}
= \begin{bmatrix}
-R^{-1} \gamma_2 e^{(\ell)} \\
0
\end{bmatrix}, \quad \ell = 1, \ldots, Q,
\]

where \( \alpha^{(\ell)} = [\alpha^{(1)}, \ldots, \alpha^{(M)}]^T \) and \( \Omega = \Phi \Phi^T \) is the kernel matrix. Matrix \( R \) is a diagonal matrix and it is invertible if and only if \( \gamma_1 v_i \neq \gamma_2 \) for \( i = 1, \ldots, M \).

The linear systems (5) and (9) have a unique solution when the associated coefficient matrix is full-rank which depends on the regularization parameters.

B. Encoding/Decoding scheme

In semi-supervised classification, the encoding scheme is chosen in advance since the number of existing classes is known beforehand. The codebook \( CB \) used for out-of-sample extension is defined based on the encoding vectors for the training points.

If \( Z = [z_{N+1}^T, \ldots, z_M^T]^T \) is the encoding matrix for the training points, the \( CB = [c_q]_{Q=1}^Q \), where \( c_q \in \{-1, 1\}^Q \), is defined by the unique rows of \( Z \) (i.e., from identical rows of \( Z \) one selects one row). Considering the test set \( D_{test} = \{x_{test,i}\}_{i=1}^{N_{test}} \) the score variables evaluated at the test points become:

\[
\begin{align*}
\epsilon_{test}^{(\ell)} &= \Phi_{test} w^{(\ell)} + b^{(\ell)}1_{N_{test}} \\
&= \Omega_{test} \alpha^{(\ell)} + b^{(\ell)}1_{N_{test}}, \quad \ell = 1, \ldots, Q,
\end{align*}
\]

where \( \Omega_{test} = \Phi_{test} \Phi^T \). The procedure for the multi-class semi-supervised classification is summarized in Algorithm 1.

**Algorithm 1: Multi-class semi-supervised classification**

**Input:** Training data set \( D \), labels \( Z \), tuning parameters \( \{\gamma_i\}_{i=1}^2 \) and kernel parameter (if any), test set \( D_{test} = \{x_{test,i}\}_{i=1}^{N_{test}} \) and codebook \( CB = [c_q]_{Q=1}^Q \)

**Output:** Class membership of test data points \( D_{test} \)

1. Solve the dual linear system (5) to obtain \( \alpha^{(Q)} \) and compute the bias term \( b^{(Q)} \) using (8).
2. Estimate the test data projections \( \{\epsilon_{test}^{(Q)}\}_{Q=1}^Q \) using (9).
3. Binarize the test projections and form the encoding matrix \( [\text{sign}(\epsilon_{test}^{(Q)}), \ldots, \text{sign}(\epsilon_{test}^{(Q)})]_{N_{test} \times Q} \) for the test points
4. \( \forall i, \) assign \( x_{test,i} \) to class \( q^* \), where \( q^* = \arg\min_q d_H (\epsilon_{test,i}, c_q) \) and \( d_H (\cdot, \cdot) \) is the Hamming distance.

IV. SEMI-SUPERVISED CLUSTERING

In what follows we assume that there is a total number of \( T \) clusters and a few labels from \( Q \) of the clusters are available \( (Q \leq T) \). Therefore we are dealing with the case that some of the clusters are partially labeled. The aim is to incorporate these labels in the learning process to guide the clustering algorithm to adjust the membership of the unlabeled data. Next we will show how one can use the approach described in section III in this setting.

A. From solution of linear systems to clusters: encoding

Since the number of existing clusters is not known a priori, we cannot use the predefined codebook as in semi-supervised classification. Therefore a new scheme is developed for generating a codebook to be used in the learning process.

It has been observed that the solution vector \( \alpha^{(\ell)}, \ell = 1, \ldots, Q \) of the dual linear system (5) has a piecewise constant property when there is an underlying cluster structure in the data (see Fig. 2(d)). Once the solution to (5) is found, the codebook \( CB \in \{-1, 1\}^{p \times Q} \) is formed by the unique rows of the binarized solution matrix (i.e., \( [\text{sign}(\alpha^{(1)}), \ldots, \text{sign}(\alpha^{(Q)})] \)). The maximum number of clusters that can be decoded is \( 2^Q \) since the maximum value that \( p \) can take is \( 2^Q \). In our approach the number of encodings, i.e., \( p \), is tuned along with the model selection procedure. Therefore a grid search on the interval \([Q, 2^Q]\) is conducted to determine the number of clusters.
It should be noted that in Algorithm 1, the static codebook $CB$ (static in a sense that the number of codewords is fixed and only depends on $Q$) is known beforehand and is of size $Q \times Q$. On the other hand in Algorithm 2, the codebook $CB$ is no longer a static codebook and is of size $p \times Q$, where $p$ can be maximally $2^Q$. Furthermore, it is obtained based on the solution matrix $S_\alpha$ (see steps 2 and 3 in Algorithm 2).

**B. Low dimensional spectral embedding**

One may notice that as opposed to kernel spectral clustering [14] where the score variables lie in a $T - 1$ (where $T$ is the actual number of clusters) dimensional space, in our formulation the embedding dimension is $Q$ which can be smaller than $T$. This can also be seen as the optimized embedding dimension for clustering which plays an important role when the number of existing clusters is large. In fact one only requires $Q = \log T$ solution vectors to uncover $T$ clusters. Therefore one is able to deal with a larger number of clusters in a more compact way. In contrast with the KSC approach where one needs to solve an eigenvalue problem, in our formulation we solve a linear eigenvalue problem, in our formulation we solve a linear system.

In contrast with the KSC approach where one needs to solve an eigenvalue problem, in our formulation we solve a linear system. It should be noted that although the two approaches share almost the same computational complexity, the quality of the solution vector obtained by the proposed algorithm is higher than that of KSC as shown in Fig. 5 and 6. This demonstrates the advantage of prior knowledge incorporation. The proposed semi-supervised clustering is summarized in Algorithm 2.

**Algorithm 2:** Semi-supervised clustering

**Input:** Training data set $D$, labels $Z$, the tuning parameters $\{\gamma_i\}_{i=1}^2$, the kernel parameter $\alpha$ (if any), number of clusters $k$, the test set $D_{\text{test}} = \{x_{\text{test}}^i\}_{i=1}^{Q_{\text{test}}}$ and number of available class labels i.e. $Q$.

**Output:** Cluster membership of test data points $D_{\text{test}}$.

1. Solve the dual linear system (5) to obtain $\{\alpha^T\}_{i=1}^Q$ and compute the bias term $b^T_{\text{test}}$ using (8).
2. Binarize the solution matrix $S_\alpha = [\text{sign}(\alpha^{(1)}), \ldots, \text{sign}(\alpha^{(Q)})]_{M \times Q}$, where $\alpha^T = [\alpha^T_1, \ldots, \alpha^T_M]^T$.
3. Form the codebook $CB = \{c_q\}_{q=1}^p$, where $c_q \in \{-1, 1\}^Q$, using the $k$ most frequently occurring encodings from unique rows of solution matrix $S_\alpha$.
4. Estimate the test data projections $\{e_{\text{test}}^{(1)}\}_{i=1}^Q$ using (10).
5. Binarize the test projections and form the encoding matrix $[\text{sign}(e_{\text{test}}^{(1)}), \ldots, \text{sign}(e_{\text{test}}^{Q})]_{N_{\text{test}} \times Q}$ for the test points (Here $e_{\text{test}}^{(1)} = [e^{(1)}_{\text{test}}, \ldots, e^{Q_{\text{test}}}_{\text{test}}]$).
6. \forall i, assign $x_{\text{test}}^i$ to class/cluster $q^*$, where $q^* = \arg\min_q d_H(e_{\text{test},i},c_q)$ and $d_H(\cdot , \cdot)$ is the Hamming distance.

**V. MODEL SELECTION**

The performance of the multi-class semi-supervised model depends on the choice of the tuning parameters. In the case of RBF kernel the optimal values of $\gamma_1, \gamma_2$ and the kernel parameters $\sigma$ can be obtained by evaluating the performance of the model (classification accuracy) on the validation set using a grid search over the parameters. One may also consider to utilize Coupled Simulated Annealing (CSA) in order to minimize the misclassification error in the cross-validation process. CSA leads to an improved optimization efficiency due to the fact that it reduces the sensitivity of the algorithm with respect to the initialization of the parameters while guiding the optimization process to quasi-optimal runs [22].

In our experiments, based on the analysis given in [16, Section III.C] we set $\gamma_1 = 1$. Then we tune $\gamma_2$ and $\sigma$ through a grid search. The range in which the search is made is discussed for each of the experiments in section VI. In general in our experiments we observed that a good value for $\gamma_2$, most of the times, is selected from the range $[0,1]$.

Since labeled and unlabeled data points are involved in the learning process, it is natural to have a model selection criterion that makes use of both. Therefore, for semi-supervised classification, one may combine two criteria where one of them evaluates the performance of the model on the unlabeled data points (evaluation of clustering results) and the other one maximizes the classification accuracy [16], [17].

A common approach for evaluating the quality of the clustering results consists of using internal cluster validity indices [23] such as Silhouette, Fisher and Davies-Bouldin index (DB) criteria. In this paper the Silhouette index is used to assess the clustering results. The Silhouette technique assigns to the $i$th sample of $j$-th class, $C_j$, a quality measure $s(i)$ which is defined as:

$$s(i) = \frac{b(i) - a(i)}{\max\{a(i), b(i)\}}.$$  

Here $a(i)$ is the average distance between the $i$th sample and all of the samples included in $C_j$, $b_i$ is the minimum average distance from the $i$-th sample to points in different clusters. The silhouette value for each sample is a measure of how similar that sample is to samples in its own cluster versus samples in other clusters, and is in the range of $[-1, 1]$.

The proposed model selection criterion for semi-supervised learning, with kernel parameter $\sigma$, can be expressed as follows:

$$\max_{\gamma_1, \gamma_2, \sigma, k} \eta \text{Sil}(\gamma_1, \gamma_2, \sigma, k) + (1 - \eta) \text{Acc}(\gamma_1, \gamma_2, \sigma, k).$$  

(11)

It is a convex combination of Silhouette (Sil) and classification accuracy (Acc). $\eta \in [0,1]$ is a user-defined parameter that controls the trade off between the importance given to unlabeled and labeled instances. In case few labeled data points are available one may give more weight to Silhouette criterion and vice versa.

The silhouette criterion is evaluated on the unlabeled data points in the validation set. One can also consider to evaluate it on the out-of-sample solution vectors.

In equation (11), $k$ denotes the number of clusters that is unknown beforehand. In the case of semi-supervised classification where the number of classes is known a priori, one does not need to tune $k$ and thus it can be removed from the list of decision variables of the aforementioned model selection criterion.

In any unsupervised learning algorithm one has to find the right number of existing clusters over the specified range which...
is provided by the user. When there is a form of prior knowledge about the data under study, the search space is reduced. In our semi-supervised clustering the lower bound of the range in which the number of clusters are sought is $Q$ (assuming that $Q$ cluster labels are available). Therefore applying the proposed MSS-KSC algorithm will make it easier to reveal the lower level of the cluster hierarchy. In the proposed MSS-KSC approach one requires to solve a linear system. Therefore the complexity of the proposed MSS-KSC algorithm in the worst case scenario is $O(M^3)$ where $M$ is the number of training data points.

VI. EXPERIMENTAL RESULTS

In this section, some experimental results are presented to illustrate the applicability of the proposed semi-supervised classification and clustering approaches. We start with a toy problem and show the differences between the obtained results when semi-supervised classification and semi-supervised clustering are applied on the same data. (see Fig. 1 and 2).

The performance of the proposed algorithms is also tested on two moons and two spirals data sets which are standard benchmark for semi-supervised learning algorithms used in the literature [24].

Next we apply the proposed semi-supervised classification on some benchmark data sets taken from the UCI machine learning repository and the performance is compared with Laplacian SVM [6] and MeanS3VM [25]. Afterwards, we have tested the performance of the semi-supervised clustering on image segmentation tasks and the obtained results are compared with the kernel spectral clustering algorithm [14]. Finally the application of the semi-supervised classification is also shown in community detection of real-world networks.

A. Toy problems

The performance of the proposed semi-supervised classification and clustering algorithms are shown on a synthetic data set consisting of seven well separated Gaussians. Some labeled data points from three of them are available (see Fig. 1(a)). When the semi-supervised classification algorithm is used the data are grouped into three classes due the fact that the codebook used in semi-supervised classification is a static codebook and it consists of three codewords. On the other hand, in semi-supervised clustering algorithm the codebook is designed based on the solution vector of the associated linear system and is not static, i.e. the number of codewords is not fixed and is tuned. Therefore, by applying the semi-supervised clustering one is able to partition the data into seven clusters. As it can be seen from Fig. 1(d) and 2(b), the projected data points are embedded in 3 dimensional space and yet we are able to cluster them in contrast with kernel spectral clustering algorithm [14] which requires an embedding space with dimension 6 to be able to group the given data sets into 7 clusters.

We also conducted experiments on nonlinear toy problems such as two moons and two spirals and the obtained results are shown in Fig. 3. For two spirals data set two scenarios are tested corresponding to different positions of the labeled data point. A comparison is made with LRGAl algorithm\(^\dagger\) proposed in [8]. The LRGAl algorithm has two parameters $k$ and $\lambda$. In these experiments the parameter $k$ (size of the neighborhood) is set to 10 and $\lambda$ is searched within $[1, 10^{16}]$ using a logarithmic scale. As Fig. 3 shows, for the two moons data set the results of both method are comparable. However the results of the two spirals data set indicate that our proposed algorithm is less sensitive to the position of labeled data points\(^\dagger\) compared to LRGAl algorithm.

In these experiments, $\gamma_2$ and $\sigma$ are tuned through a grid search. The range in which the search (using a logarithmic scale) is made for $\gamma_2$ and $\sigma$ is shown in Fig. 2(c) and Fig. 3(g,h,i). From these figures, it is apparent that there exist a range of $\gamma_2$ and $\sigma$ for which the value of the utilized model selection criterion is quite high on the validation set.

B. Real-life benchmark data sets

Four benchmark data sets used in the following experiments are chosen from the UCI machine learning repository [26]. The benchmark consists of Wine, Iris, Zoo and Seeds data sets. In all cases, the data points are divided using proportion $80\%$ and $20\%$ into training and test counterparts respectively. Then one fourth of randomly selected data points in the training set are considered to be labeled and the remaining three fourths are unlabeled. The performance of the proposed semi-supervised classification approach (MSS-KSC), is compared with Laplacian SVM (LapSVMp)\(^\dagger\) [6] and MeanS3VM [25] using the one-vs-all strategy.

In this experiment, the procedure used for model selection is a two-step procedure which consists of Coupled Simulated Annealing [22] initialized with random sets of parameters for the first step and the simplex method [27] for the second step. After CSA converges to some local minima, the parameters that obtained the lowest misclassification error are used for initialization of the simplex procedure to refine our selection. At every iteration for CSA method a 10-fold cross-validation is utilized. In all the experiments the RBF kernel is used.

For MeanS3VM method, the regularization parameters $C_1$ and $C_2$ are fixed to 1 and 0.1, respectively and the width parameter in RBF kernel is tuned with respect to the accuracy on the validation set. For the Laplacian SVMs, we tuned the kernel parameter and $\gamma$, with respect to the accuracy on the validation set. The remaining parameters, i.e. $\gamma_1$ and $NN$, are set to their default values ($\gamma_1 = 1$ and $NN = 6$).

The mean and standard deviation of the accuracy rates on test data points with respect to 10 random splits are reported in Table 1. Table 1 shows that the proposed MSS-KSC approach outperforms in most cases the other approaches on these tested problems. The effect of changing the value of the user defined parameter $\eta$, used for model selection, on the performance of the proposed algorithm with respect to 10 random splits can be seen in Fig. 4.

C. Image segmentation

In this section, the task is to segment the given image using the proposed semi-supervised clustering. Here the aim is to show

\(^\dagger\)Available at: http://www.cs.cmu.edu/~yiyang/LRGAl_ranking.m

\(^\dagger\)The equivalent of the query provided by the user.

\(^\dagger\)Available at: http://www.dii.unisi.it/~melacci/lapsvmp/
Fig. 1. Toy Problem: Seven well separated Gaussians. The labeled data points of only three classes are available and are depicted by the blue squares ( ■ ), green triangles ( ▲ ) and red circles ( • ). (a): Data points in the original space (b): Result of multi-class semi-supervised classification using LapSVMp with RBF kernel. (c): Result of the proposed multi-class semi-supervised classification with RBF kernel (Note that the algorithm detected three classes. The first class consists of one cluster whereas the second and third class consist of three clusters respectively). (d): The projections of the validation data points when the proposed semi-supervised classification algorithm is used (indicating the line structure in the projection-space).

Fig. 2. Toy Problem: Seven well separated Gaussians. The labeled data points of only three classes are available and are depicted by the blue squares ( ■ ), green triangles ( ▲ ) and red circles ( • ). (a): Result of the proposed multi-class semi-supervised clustering with RBF kernel. (b): The projections of the validation data points when semi-supervised clustering algorithm is used (indicating the line structure in the projection-space). (c): Model selection for semi-supervised clustering using Silhouette validity index corresponding to the best case $T = 7$. The asterisk (*) marks the optimal model. (d): Piecewise constant property of the solution vector.
Fig. 3. Toy Problems: two spiral and two moon data sets. The labeled data point is depicted by the red squares (■). First row (a,b,c): Data points in the original space. Second row: (d,e,f): Result of the proposed semi-supervised algorithm with RBF kernel. Third row: (g,h,i): Model selection of the proposed algorithm. (The asterisk (*) marks the optimal model for these examples.) Fourth row: (j,k,l): Result of the LRGA algorithm corresponding to the worst case when the parameter $k$ (size of the neighborhood) is set to 10 and $\lambda$ is searched within $[1, 10^{16}]$ using a logarithmic scale. Fifth row: (m,n,o): Result of the LRGA algorithm corresponding to the best case when the parameter $k$ (size of the neighborhood) is set to 10 and $\lambda$ is searched within $[1, 10^{16}]$ using a logarithmic scale.
F-measure, i.e. Variation of information (VI): it measures the distance to the unsupervised model, it is possible to improve the result that by incorporating the side-information (labels in this case) to the unsupervised model, it is possible to improve the result of the unsupervised algorithm.

Experimental results on two synthetic images and some color images from the Berkeley image data set [28] are shown in Fig. 5 and 6. For each image, a local color histogram with a $5 \times 5$ local window around each pixel is computed using minimum variance color quantization of eight levels. A subset of 500 unlabeled pixels together with some labeled pixels (see Table 2) are used for training and the whole image for testing.

For the synthetic images we provided a qualitative evaluation of both approaches, since the ground truth of these images were not available. For the Berkeley images data set for which the ground truth segmentations are known, the segmentations obtained by MSS-KSC and KSC are compared with the ground truth in Table 2. Two evaluation criteria are used:

- F-measure, i.e. $\frac{2 \times \text{Precision} \times \text{Recall}}{\text{Precision} + \text{Recall}}$ with respect to human ground-truth boundaries.
- Variation of information (VI): it measures the distance between two segmentations in terms of their average conditional entropy. Low values indicate good match between the segmentations [29].

In these experiments, the range in which the search (using a logarithmic scale) is made for tuning the parameters $\gamma_2$ and $\sigma$ are $[0, 1]$ and $[10^{-3}, 10^{1}]$ respectively. The length of the codebook $p$ is also tuned on the interval $[1, 25]$. The score variables obtained by the proposed MSS-KSC algorithm for two images are shown in Fig. 5 when Silhouette criterion is used. As it can be seen, the embedding dimension (spectral embedding) is three and yet we can detect more than four clusters from the given image. Unlike the toy example 1 for these images, due to the fact that clusters are not well separated, the line structure of the score variables is less clear. In Fig. 6, we plotted the maximum value of the Silhouette criterion for each $p$ (length of the codebook) while tuning $\gamma$ and $\sigma$. Therefore the predicted number of clusters is equal to $p$ for which the Silhouette value is maximum. The obtained results are shown in Fig. 5 and 7 which reveal that incorporating the prior knowledge (labels provided by human), can potentially increase the performance in the segmentation task with respect to a genuinely unsupervised
Table II

Comparison of KSC and MSS-KSC for image segmentations in terms of F-measure and variation of information indices

<table>
<thead>
<tr>
<th>Image ID</th>
<th>D^\text{val}</th>
<th>F-measure</th>
<th>Variation of information</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>D_u</td>
<td>D_L</td>
<td>D_u</td>
</tr>
<tr>
<td>100007</td>
<td>4</td>
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<td>3</td>
<td>500  6</td>
<td>3000  6</td>
</tr>
<tr>
<td>385039</td>
<td>5</td>
<td>500 14</td>
<td>3000 12</td>
</tr>
<tr>
<td>388067</td>
<td>3</td>
<td>500  6</td>
<td>3000  6</td>
</tr>
<tr>
<td>8049</td>
<td>3</td>
<td>500  6</td>
<td>3000  7</td>
</tr>
</tbody>
</table>

Note: For variation of information the lower the value the better, whereas for F-measure the higher value the better the segmentation is.

Fig. 5. (a),(f): Original image used for the KSC algorithm. (b),(g): Segmented image using the KSC approach. (c),(h): Labeled image used for the proposed semi-supervised clustering algorithm. (d),(i): Segmented image using the MSS-KSC approach. (e),(j): Score variables in the projection space.

D. Community detection

Community detection is an important topic related to complex networks [30]. It consists of finding clusters of strongly connected nodes such that nodes in the same community share more connections than nodes in different communities. Once properly identified, the community structure can help to shed light on the whole functioning of the network. Community detection is an unsupervised technique. However, if some form of prior knowledge of the community structure is present, semi-supervised techniques could in principle be used to improve the results [31], [32].

In this section the performance of the proposed method is analyzed in the community detection problems when there exist some form of prior knowledge about the existing communities. We conduct the experiments on two well-known real-world networks, i.e. Karate, Football data sets shown in Fig. 7, which are described briefly as follows:

Karate: The Zachary’s karate club network [33] consists of 34 member nodes, and splits into two smaller clubs after a dispute emerged during the course of Zachary’s study between the administrator and the instructor.

Football: This network [34] describes American college football games and is formed by 115 nodes (the teams) and 616 edges (the games). It can be divided into 12 communities according to athletic conferences.

Concerning the Karate network, a comparison with the methods described in [35] is performed. In [35] a percentage of node pairs, which then are determined whether they belong to must-link or cannot-link groups, is used in the learning process. The reported results in Table 1 of [36] for different percentages of node pairs are tabulated in Table III.

Since in the proposed approach we work with the labeled nodes, not pairs, we randomly select some nodes and labeled them according to the true community to which they belong. The averaged normalized mutual information (NMI) over 10 simulation runs for Karate network is reported in Table III. One can observe that the proposed method is able to achieve the
Fig. 6. Model selection curves corresponding to the obtained Silhouette value for different $p$ value. (a) Three labels were provided ($Q=3$). The maximum Silhouette value for the first syntactic image, while $\sigma$ and $\gamma_2$ are tuned, over a range of $p \in \{Q, \ldots, 2Q\}$. (b) Three labels were provided ($Q=3$). The maximum Silhouette value for the second syntactic image, while $\sigma$ and $\gamma_2$ are tuned, over a range of $p \in \{Q, \ldots, 2Q\}$.

Fig. 7. Image segmentation results using the proposed method and the KSC [14]. A subset of 500 and 3000 randomly chosen pixel histograms (unlabeled data points) together with some labeled data points are used for training and validation respectively. The whole image is used for testing. The original image is shown in the first row. The segmentation results obtained by KSC using the original images are shown in the second row. The third row shows the images labeled by human. The results of the proposed semi-supervised clustering algorithm applied on the labeled images are depicted in the fourth row.

maximum performance using less labeled nodes than the other algorithms. In particular with 10 labeled nodes the maximum value of NMI is achieved.

Concerning the Football network, we conducted the semi-supervised classification task. The training set consists of both labeled and unlabeled nodes. 40% of each class (community) is randomly selected and form the labeled training nodes and another 40% randomly selected nodes form the unlabeled nodes. The whole network is considered as the test set and the obtained result is compared with KSC approach. The partitions found by KSC and MSS-KSC are evaluated according to adjusted Rand index (ARI) [35]. The obtained ARI values on the test set after 10 runs is shown in Fig. 9 respectively. We can observe that, the prior knowledge incorporation helps to improve the performance with respect to KSC.

VII. CONCLUSIONS

In this paper, a multi-class semi-supervised formulation based on kernel spectral clustering has been proposed. The method is able to handle both semi-supervised classification and clustering. In the semi-supervised clustering case, an optimal embedding dimension is designed and utilized. The validity and
TABLE III


<table>
<thead>
<tr>
<th>pairs constraints</th>
<th>% pairs [c]</th>
<th>NMF-LSE</th>
<th>NMF-KL</th>
<th>SNMF</th>
<th>SP</th>
<th>The proposed method</th>
</tr>
</thead>
<tbody>
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<td></td>
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<td></td>
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<tr>
<td>2%</td>
<td>4 [4,8]</td>
<td>0.98</td>
<td>0.73</td>
<td>0.51</td>
<td>0.90</td>
<td>4</td>
</tr>
<tr>
<td>4%</td>
<td>6 [6,12]</td>
<td>0.99</td>
<td>0.85</td>
<td>0.60</td>
<td>0.96</td>
<td>6</td>
</tr>
<tr>
<td>5%</td>
<td>8 [8,16]</td>
<td>0.99</td>
<td>0.89</td>
<td>0.53</td>
<td>0.95</td>
<td>8</td>
</tr>
<tr>
<td>10%</td>
<td>16 [16,32]</td>
<td>1.00</td>
<td>0.89</td>
<td>0.57</td>
<td>1.00</td>
<td>10</td>
</tr>
<tr>
<td>20%</td>
<td>31 [31,34]</td>
<td>1.00</td>
<td>0.98</td>
<td>0.56</td>
<td>1.00</td>
<td>12</td>
</tr>
</tbody>
</table>

Note: The minimum and maximum number of nodes that could result in the given number of pairs are denoted by r and t.

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Fig. 8. Visualization of the networks when nodes are colored according to their degree value (a) American college football undirected graph. (b) Zachary’s karate club undirected graph.

Fig. 9. American college football network. (a) Obtained ARI value when KSC and MSS-KSC algorithm are used. (b) Kernel matrix showing the partitioning related to $\eta = 0.6$. A clear block structure revealing the presence of the 12 communities can be noticed.

REFERENCES


