Self-weighted Multiview Clustering with Multiple Graphs

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Abstract
In multiview learning, it is essential to assign a reasonable weight to each view according to the view importance. Thus, for multiview clustering task, a wise and elegant method should achieve clustering multiview data while learning the view weights. In this paper, we propose to explore a Laplacian rank constrained graph, which can be approximately as the centroid of the built graph for each view with different confidences. We start our work with a natural thought that the weights can be learned by introducing a hyperparameter. By analyzing the weakness of this way, we further propose a new multiview clustering method which is totally self-weighted. More importantly, once the target graph is obtained in our models, we can directly assign the cluster label to each data point and do not need any postprocessing such as $K$-means in standard spectral clustering. Evaluations on two synthetic datasets indicate the effectiveness of our methods. Compared with several representative graph-based multiview clustering approaches on four real-world datasets, the proposed methods achieve the better performances and our new clustering method is more practical to use.

1 Introduction

Many computer vision objects, such as image and video, involve instances represented with multiple views. It is essential to study how to efficiently cluster such kind of data. Suppose we use a graph to capture entities and their relation in each view, we will naturally obtain multiple graphs. Therefore, from this perspective, graph-based multiview clustering can be transformed into a multiple graph learning problem.

Prior to the multiple graph learning method, a direct way to handle multiview data is to concatenate all the features into a new one and then utilize it to construct a single graph. However, this seemingly simple method neglects the relation among all the views and doesn’t make sense theoretically. Recently, in practice many works would prefer to construct a graph for each view and then jointly utilize them to learn a unified one. Inspired by co-training [Blum and Mitchell, 1998], which is widely used in multiview semisupervised learning, [Kumar and Daumé, 2011] searched for the clusters that agree across all the views. Later, to minimize the disagreement between every pair of views, [Kumar et al., 2011] applied the co-regularization technique to spectral clustering model and reformulated multiview clustering as a jointly maximization problem. Similarly, [Cai et al., 2011] designed a graph-based multiview spectral learning model to integrate heterogeneous image features. These methods extend pairwise concept to multiple views condition and look for the most consistent representation. Following a more comprehensive thought is that each individual view may contain specific information as well as common information, [Tang et al., 2009] presented the linked matrix factorization model to extract the structure information shared by all the graphs.

Almost every aforementioned method works on the assumption that all the views are reliable. However, in real-world clustering problems, the views of data are inherently strong or weak, and meanwhile some may have been corrupted by noises. This means the final results will be degraded if we fail to distinguish different views. [Xia et al., 2014] addressed this problem by separating noise from each graph and learned a shared low-rank transition probability matrix, which will be the input to the standard Markov chain method for clustering. [Kumar et al., 2011; Cheng and Zhao, 2009] proposed to roughly compute the weights by combining with the prior knowledge. This kind of approaches are manually intervening or shallow. In view of this point, some other methods [Li et al., 2015; Xie and Sun, 2013] looked for a new weight learning strategy which has been widely used in multiview learning. It explicitly defines the view weights in the objective and then learns them as variables, which is parallel with the thought of our first method. Nevertheless, this so-called adaptive or automatic weight learning strategy has to resort to an additional hyperparameter (For simplicity, we replace with $\gamma$ when referring this parameter in the following text), which is usually difficult to set in a specific task.

The first motivation of this work is: it is very common that graph-based clustering methods appear in the context of spectral learning, such as [Kumar and Daumé, 2011; Kumar et al., 2011; Cai et al., 2011; Tang et al., 2009; Xia et al., 2014; Cheng and Zhao, 2009; Li et al., 2015; Xie and Sun, 2013]. This means that in the last step of these methods, it often needs to apply a simple clustering method...
such as $K$-means to the learned indicator matrix. However, the uncertainty of this postprocessing operation will increase the instability of the original performance. We avert this trouble by extending an existing single graph learning approach [Nie et al., 2016b] to multiview domain. Inspired by the recent works, we propose the Parameter-weighted Multiview Clustering (PwMC) method to weightedly combine each single graph learning model. The second motivation is that, as many previous works, PwMC involves an undesired parameter $\gamma$. There are two reasons to argue. 1) In context of unsupervised learning where no instance is labeled, $\gamma$ can not be obtained by traditional supervised hyperparameter tuning techniques, such as cross validation. 2) It is observed that the final experimental performance is sensitive to $\gamma$ and the optimum of $\gamma$ varies on different datasets (see Experiments part of Section 3.3), which causes PwMC (and the related multiview clustering methods with the similar weight learning strategy) not practical to use. Thus, to remove $\gamma$ while without loss of too much precision, we further propose a new Self-weighted Multiview Clustering (SwMC) method. These two motivations are exactly corresponding to our contributions of this paper. The toy examples and experiments on real-world datasets demonstrate the effectiveness of our proposed methods.

Notation. Throughout the paper, all the matrices are written as uppercase. For a matrix $M$, the $i$-th row and $ij$-th element of $M$ are denoted as $m_i$ and $m_{ij}$ separately. The trace of $M$ is denoted as $Tr(M)$. The $v$-th view representation of $M$ is written as $M^{(v)}$. The Frobenius norm of matrix $M$ is denoted by $\|M\|_F$. In particular, we use $1_n$ to denote a $n$ dimensional column vector where each element is 1.

2 The Proposed Framework

2.1 Graph-based Clustering Revisit

Suppose we have $n$ samples and they can be partitioned into $c$ clusters, graph-based clustering methods firstly construct a Similarity Matrix (SM, and we will not differ SM and graph in this paper) to represent the affinities of all the samples. Many early works have studied how to design a SM with high quality, such as [Zelnik-Manor and Perona, 2005; Cai et al., 2005]. These well-designed SMs will be the inputs of graph-based clustering methods, e.g., spectral clustering. However, an ideal SM $S \in \mathbb{R}^{n \times n}$ is supposed to exactly have $c$ connected components, by which way, $S$ can be directly used for the clustering task. Recently, [Nie et al., 2014a; Feng et al., 2014; Chen and Dy, 2016; Nie et al., 2016b] have leveraged this property in different ways. We briefly introduce the Constrained Laplacian Rank (CLR) method [Nie et al., 2016b] therein, which is easier to understand and will be the base of our proposed framework. Given an arbitrary input SM $A \in \mathbb{R}^{n \times n}$, the target SM can be learned by minimizing the following problem

$$\min_{s, 1_n=1, s_{ij} \geq 0, S \in C} \|S - A\|_F^2,$$  \hspace{1cm} (1)

where $S$ is nonnegative, whose each row sums up to 1, and $C$ represents the set of $n$ by $n$ square matrices with $c$ connected components. According to the graph theory in [Mohar et al., 1991; Chung, 1997], the connectivity constraint can be replaced with a rank constraint, and thus we have

$$\min_{s, 1_n=1, s_{ij} \geq 0, \text{rank}(L_S) = n-c} \|S - A\|_F^2,$$  \hspace{1cm} (2)

where $\text{rank}(L_S)$ means the rank of $L_S$. The Laplacian matrix $L_S = D_S - (S^T + S)/2$, where the degree matrix $D_S \in \mathbb{R}^{n \times n}$ is defined as a diagonal matrix whose $i$-th diagonal element is $\sum_j (s_{ij} + s_{ji})/2$. In this way, the target SM can be solved and we directly use it for clustering.

2.2 Parameter-weighted Multiview Clustering (PwMC)

CLR is a single view graph-based clustering method. In this paper, we introduce this technique into multiview clustering domain. For multiview data, let $m$ be the number of views and $A^{(1)}, A^{(2)}, ..., A^{(m)}$ be the corresponding input SMs, where $A^{(v)} \in \mathbb{R}^{n \times n} (1 \leq v \leq m)$. Thus, our goal comes to find out the target SM $S$ which is constrained as in Eq. (2) but can approximate each original input SM $A^{(v)}$. An intuitive way to address this problem is to simply calculate an average SM $\bar{A} = \frac{1}{m} \sum_{v=1}^{m} A^{(v)}$ and input it into CLR. Actually, this thought is equivalent to assigning the equal weight to each graph. In practice, we need to employ a group of meaningful weights to measure the importance of each view. In other words, the target SM $S$ is supposed to approximate every input SM with different confidences. This idea can be naturally modeled by minimizing the linear combination of the reconstruction error $\|S - A^{(v)}\|_F^2$ for each view. Thus, the formulated objective can be written as

$$\min_{\alpha^{(v)}, S} \sum_{v=1}^{m} \alpha^{(v)} \|S - A^{(v)}\|_F^2 + \gamma \|\alpha\|_2^2,$$  \hspace{1cm} (3)

s.t. $\alpha^{(v)} \geq 0, \alpha^T 1_m = 1, s_{ij} \geq 0, s_1 1_n = 1$,

$$\text{rank}(L_S) = n-c,$$

where $\alpha = [\alpha^{(1)}, \alpha^{(2)}, ..., \alpha^{(m)}]^T$ and $\gamma > 0$. The second term in problem (3) is used to smoothen the weight distribution. Straightforward, without this regularization term (or $\gamma \to 0$), the trivial solution will be obtained, i.e., the weight of best view will be assigned to 1 and other weights will be 0s. On the contrary, when $\gamma \to \infty$, the equal weights will be obtained. Since the weights severely depend on the parameter $\gamma$, we name this method as Parameter-weighted Multiview Clustering (PwMC). The detailed procedure of optimize the problem (3) is presented in Section 2.4.

2.3 Self-weighted Multiview Clustering (SwMC)

Although problem (3) provides a feasible scheme to fuse multiple graphs on the model level, it involves the undesired parameter $\gamma$, which is confirmed to be dataset-related by our experiments. To remove the parameter $\gamma$ in problem (3), we propose a self-weighted multiview clustering method. It sounds unreasonable that the suitable weights could be generated out of thin air. But in this part, we present a new formulation
that actually induces a self-conducted weight learning. The proposed objective is
\[
\min_{s_{ij} \geq 0, s_i, 1_n = 1, rank(L_S) = n - c} \sum_{v=1}^{m} \left\| S - A^{(v)} \right\|_F.
\] (4)

This equation looks pretty simplified and compact, but no weight factor is explicitly defined therein. We consider it as a normal issue and firstly write its Lagrange function
\[
\min_{S} \sum_{v=1}^{m} \left\| S - A^{(v)} \right\|_F + G(\Lambda, S),
\] (5)
where \(\Lambda\) is the Lagrange multiplier, \(G(\Lambda, S)\) serves as a proxy for the constraints to \(S\). Taking the derivative of Eq. (5) w.r.t \(S\) and setting the derivative to zero, we have
\[
\sum_{v=1}^{m} w^{(v)} \frac{\partial}{\partial S} \left\| S - A^{(v)} \right\|_F^2 + \frac{\partial G(\Lambda, S)}{\partial S} = 0,
\] (6)
where \(w^{(v)}\) is given as the following form\(^1\)
\[
w^{(v)} = 1 / \left( 2 \left\| S - A^{(v)} \right\|_F \right).
\] (7)

Obviously, from Eq. (7) we know that \(w^{(v)}\) is dependent on \(S\), which means the two factors of the first term in Eq. (6) are coupled with each other. But if we set \(w^{(v)}\) stationary, Eq. (6) can be considered as the solution to the following problem
\[
\min_{s_{ij} \geq 0, s_i, 1_n = 1, rank(L_S) = n - c} \sum_{v=1}^{m} w^{(v)} \left\| S - A^{(v)} \right\|_F^2,
\] (8)
which is simpler to be solved. And then, the calculated \(S\) from problem (8) can be further used to update \(w^{(v)}\) by Eq. (7). This inspires us to solve the original problem (4) by alternately optimizing \(S\) and \(w^{(v)}\) iteratively. We summarize this process into Algorithm 1. Moreover, if this alternating optimization strategy converges (it will be proved later), from Eq. (6) we know that the finally learned \(S\) will converge to the KKT condition of the problem (4).

**Algorithm 1** The algorithm of Self-weighted Multiview Clustering (SmWC) in Eq. (4)

**Input:** SMs for \(m\) views \(\{A^{(1)}, A^{(2)}, \ldots, A^{(m)}\}\) and \(A^{(v)} \in \mathbb{R}^{n \times n}\), number of clusters \(c\).

**Initialize** the weight for each view (e.g., \(\alpha^{(v)} = \frac{1}{m}\)).

**Repeat**
1. Calculate \(S\) by solving the problem (8).
2. Update \(\alpha^{(v)}\) by using Eq. (7).

**Until** converge

**Output:** \(S \in \mathbb{R}^{n \times n}\) with exactly \(c\) connected components

\(^1\)To avoid dividing by zero, in practice we use the following formula \(w^{(v)} = 1 / \left( 2 \sqrt{\left\| S - A^{(v)} \right\|_F^2 + \delta} \right)\), where \(\delta\) is some very small value, like 0.0001.

So far, we have presented the general process to solve the problem (4). But what is the relation between it and our self-weighted multiview learning? **Inspecting Eq. (8), when the optimization process converges, its form can be described as the linear combination of reconstruction errors of different views if we view the \(w^{(v)}\) as the weight factor, which is exactly what we want to learn.** Since Eq. (8) is derived from solving the problem (4), it is a totally self-weighted process. Furthermore, if view \(v\) is good, then \(\left\| S - A^{(v)} \right\|_F\) should be small, and thus the learnt \(w^{(v)}\) for view \(v\) is large according to Eq. (7). Accordingly, a weak view will be assigned a small weight. This indicates that our self-weighted learning model is meaningful. In fact, the above process of solving problem (4) can be seen as a special case of the iteratively re-weighted (IR) technique [Nie et al., 2010; Daubechies et al., 2010; Nie et al., 2014b]. In this perspective, the effectiveness of this method will reflect the practical significance of the weight formula in IR.

### 2.4 Optimization

#### Optimization of PwMC

Once given a proper value of parameter \(\gamma\), we can adopt the alternating iterative strategy to optimize the problem (3).

**When \(\alpha\) is fixed**, we need to solve the following subproblem
\[
\min_{s_{ij} \geq 0, s_i, 1_n = 1, rank(L_S) = n - c} \sum_{v=1}^{m} \alpha^{(v)} \left\| S - A^{(v)} \right\|_F^2.
\] (9)

Let \(\sigma_i(L_S)\) denote the \(i\)-th smallest eigenvalue of \(L_S\), \(\sigma_i(L_S) \geq 0\) since \(L_S\) is positive semi-definite. Given a large enough \(\lambda^2\), the rank constraint in Eq. (9) can be eliminated and the problem (9) is equivalent to the following form:
\[
\min_{s_{ij} \geq 0, s_i, 1_n = 1} \sum_{v=1}^{m} \alpha^{(v)} \left\| S - A^{(v)} \right\|_F^2 + 2\lambda \sum_{i=1}^{c} \sigma_i(L_S).
\] (10)

When \(\lambda\) is large enough, note that \(\sigma_i(L_S) \geq 0\) for each \(i\), thus the optimal solution \(S\) to the problem (10) will make the second term \(\sum_{i=1}^{c} \sigma_i(L_S)\) equal to zero and the constraint \(rank(L_S) = n - c\) will be satisfied. In addition, according to Ky Fan’s Theory [Fan, 1950], we have the following equation:
\[
\sum_{i=1}^{c} \sigma_i(L_S) = \min_{F \in \mathbb{R}^{n \times c}, Tr(F^T L_S F) = 1} Tr(F^T L_S F).
\] (11)

Thus, combining with Eq. (11), the problem (10) is further equivalent to the following problem:
\[
\min_{S, F} \sum_{v=1}^{m} \alpha^{(v)} \left\| S - A^{(v)} \right\|_F^2 + 2\lambda Tr(F^T L_S F)
\] (12)
\[
s.t. s_{ij} \geq 0, s_i, 1_n = 1, F \in \mathbb{R}^{n \times c}, F^T F = I.
\]

\(^2\)Unlike the regular hyperparameter, this introduced \(\lambda\) will not influence the final experimental performance. In our method, following the CLR method, we determine the value of \(\lambda\) in a heuristic way to accelerate the procedure.
We solve this problem by optimizing variables $F$ and $S$ iteratively as follows.

i. **Solving $F$ When $S$ is fixed**, the problem (12) becomes

$$
\min_{F \in \mathbb{R}^{n \times c}, F} \text{Tr} \left( F^T L_S F \right).
$$

(13)

It is known that the optimal solution of $F$ is formed by the $c$ eigenvectors of $L_S$ corresponding to the $c$ smallest eigenvalues.

ii. **Solving $S$ When $F$ is fixed**, the problem (12) becomes

$$
\min_{s_{ij} \geq 0, s_i, s_{1:n} = 1} \sum_{v=1}^{m} \alpha_v (s_{ij} - a_v) ^2 + \lambda \sum_{i,j=1}^{n} \|f_i - f_j\|_2^2 s_{ij}.
$$

(14)

Since the problem (14) is independent for different $i$, we can solve the following problem separately for each $i$:

$$
\min_{s_{ij} \geq 0, s_i, s_{1:n} = 1} \sum_{j=1}^{n} \alpha_v (s_{ij} - a_v) ^2 + \lambda \sum_{j=1}^{n} \|f_i - f_j\|_2^2 s_{ij}.
$$

(15)

For simplicity, we denote $v_{ij} = \|f_i - f_j\|_2^2$ and $v$ as a vector with $j$-th element equal to $v_{ij}$ (and similarly for $s_i$ and $a_i$), the problem (15) can be written in vector form as

$$
\min_{s_{ij} \geq 0, s_i, s_{1:n} = 1} \left\| s_i - \left( \sum_{v=1}^{m} \alpha_v a_v - \frac{\lambda}{2} v_i \right) \right\|_2^2.
$$

(16)

This problem can be solved by an efficient iterative algorithm proposed in [Duchi et al., 2008]. To accelerate the computing, we can choose to update $t$ (One can set $t$ as a const, like 10) neighbors of $i$-th data. Thus, $S$ is totally sparse and can be rapidly calculated.

When $S$ is fixed, the problem (3) is reduced to minimization

$$
\alpha_v \geq 0, \alpha_v \neq 1 \Rightarrow \sum_{v=1}^{m} \alpha_v e(v) + \gamma \|\alpha\|_2^2,
$$

where $e(v) = \|S - A(v)\|_F^2$. It is easy to rewrite the problem (17) into the following form

$$
\min_{\alpha_v \geq 0, \alpha_v \neq 1} \left\| e + \alpha \right\|_2^2,
$$

(18)

which is identical with the problem (16). Now, we summarize the solving process of problem (3) in Algorithm 2.

### Optimization of SwMC

According to Algorithm 1, solving SwMC mainly contains two alternative steps, in which updating Eq. (7) is quite simple while the subproblem (8) needs to be further calculated. Seeing that the only difference between problems (8) and (9) is whether weights sum to 1, thus we can solve the subproblem (8) with the same way.

#### 2.5 Converge Analysis

According to the alternating optimization steps described in Algorithm 2, since we can find the optimal solution of each subproblem, obviously Algorithm 2 will converge. Now, we focus on proving the convergence of Algorithm 1.

**Lemma 1** [Nie et al., 2010] For any positive number $u$ and $v$, the following inequality holds:

$$
u - \frac{u^2}{2v} \leq v - \frac{v^2}{2u}.
$$

(19)

**Theorem 1** In each iteration of Algorithm 1, the updated target $SM$ will monotonically decrease the objective of problem (4), which generally makes the solution converge to the local optimum of the problem (4).

**Proof:** Let $S$ denote the updated $S$ in each iteration. According to the first step of loop in Algorithm 1, we have

$$
S = \arg \min_{s_{ij} \geq 0, s_i, s_{1:n} = 1} \sum_{v=1}^{m} w(v) \|S - A(v)\|_F^2.
$$

(20)

Combining with $w(v) = 1/(2\|S - A(v)\|_F)$, we can derive

$$
\sum_{v=1}^{m} \frac{\|S - A(v)\|_F^2}{\|S - A(v)\|_F} \leq \sum_{v=1}^{m} \frac{\|S - A(v)\|_F^2}{\|S - A(v)\|_F}.
$$

(21)

According to Lemma 1, we have

$$
\sum_{v=1}^{m} \|S - A(v)\|_F^2 \leq \sum_{v=1}^{m} \|S - A(v)\|_F^2.
$$

(22)

Summing Eq. (21) and Eq. (22) in the two sides, we arrive at

$$
\sum_{v=1}^{m} \|S - A(v)\|_F^2 \leq \sum_{v=1}^{m} \|S - A(v)\|_F^2.
$$

(23)
Thus, the iterative optimization will monotonically decrease the objective of the problem (4) in each iteration until it converges. When the convergence reaches, the equality in Eq. (22) holds, thus $\tilde{S}$ will satisfy Eq. (6), the KKT condition of problem (4). Therefore, in most cases, the Algorithm 1 will at least converge to a local optimal solution of problem (4).

3 Experiments

In this paper, we follow CLR method to construct a graph for each view as the initialized input graph $A^{(v)}$. Different from some regular kernel-based methods [Zelnik-Manor and Perona, 2005; Cai et al., 2005], this graph construction approach only needs to set one parameter $k$ which represents the number neighbors. For all the compared methods, we validate their performances by fixing $k$ as 10. Another advantage is that this approach naturally obtains the neat normalized graph for each view. It is a hard-to-get property, and many previous graph construction methods have to normalize multiview data before utilization. The standard clustering Purity and Normalized Mutual Information (NMI) metrics are used to measure the clustering performance in our experiments.

3.1 Toy Example

In this part, we conduct two toy experiments to verify the effectiveness of the proposed algorithms.

Toy_1 dataset. We design a two-view synthetic dataset where each view is a $90 \times 90$ matrix with three $30 \times 30$ block matrices diagonally arranged. Without loss of generality, the data within each block denotes the affinity of two corresponding points in one cluster, while the data outside all blocks denotes noise. Each element in all blocks is randomly generated in the range of 0 and 1, while the noise data is randomly generated in the range of 0 and $e$, where $e$ is set as 0.6 in the 1st matrix, and 1.0 in the 2nd matrix. Then they are normalized to be that the sum of each row is 1. The original input graphs are shown in Figure 1.

We have verified that CLR can directly recover the desired block diagonal matrix from view 1 but fails on view 2. Given some small value to the parameter $\gamma$, the learned weights of PwMC are close to 1/0 , which means in such a situation PwMC achieves the best performance (For the space limitation, we won’t present all the learned clean block diagonal matrix.) by selecting the best view. For SwMC, it also learns the clean block diagonal matrix but with the normalized weights 0.63/0.37. Seeing that the elements within diagonally blocks always contributes to clustering, it indicates that SwMC arrives at the optimal solution in a different style.

Toy_2 dataset. This synthetic dataset has the same scale with Toy_1 but with the different noise settings. The initial noise in view 1 and 2 is set as $e = 0.6$ and $e = 0.7$ respectively. By increasing the noise between the first and second block data to $e = 0.8$, we obtain the input matrix for view 1. Similarly, in view 2, we increase the noise between the second and third block data to $e = 1.0$. They are are normalized and then shown as Figure 2.

By performing CLR on each individual graph, their clustering performance are: Purity of view 1 and 2 are 0.66/0.63, NMI of view 1 and 2 are 0.58/0.58. The proposed methods can still integrate these two complementary graphs and recover the clean block diagonal matrix. The learned weights of both proposed methods are around 0.53/0.47. Noting that view 1 contains less noise than view 2, for SwMC method, according to Eq. (7), we find this phenomenon agrees with our prior inference in Section 2.3.

3.2 Performance Evaluation

Following [Li et al., 2015], we evaluate the performance of the compared methods on three multi-view datasets, MSRCv1 [Winn and Jojic, 2005], Caltech101 [Fei-Fei et al., 2007] (we use two regular subsets Caltech101-7 and Caltech101-20), Handwritten numerals (Digits) [Asuncion and Newman, 2007], which are briefly summarized in Table 1.

We compare the proposed PwMC and SwMC with following methods: Co-regularized spectral clustering [Kumar et al., 2011] (Co-reg), Multi-View Spectral Clustering [Cai et
Table 2: Clustering performance comparison, where SwMC is the unique one which has no parameter to tune.

<table>
<thead>
<tr>
<th></th>
<th>MSRCv1</th>
<th>Caltech101-7</th>
<th>Caltech101-20</th>
<th>Digits</th>
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<td>NMI</td>
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</table>

Figure 3: Clustering results comparison between SwMC and PwMC on different datasets.

3.3 Parameter-free Weight Learning

Figure 3 exhibits the clustering performance of the proposed SwMC comparing with PwMC on different datasets. Typically, $\gamma$ in PwMC is searched in logarithm form ($\log_{10}\gamma$ from 0 to 4 with step size 0.5). Although PwMC and SwMC are influenced by the different initializations, we start then with the equal weights in this experiment.

According to Figure 3, except for Digits, PwMC achieves better (or at least equal) performance than SwMC at some specific $\gamma$. Theoretically, a specific $\gamma$ corresponds to a weight distribution, which means the optimal weight combination will be obtained if one can search all the possible $\gamma$s. However, it is observed that the performance of PwMC varies dramatically when $\gamma$ changes, which makes that attempting to apply a fixed $\gamma$ throughout all the applications is not practical. Therefore, the proposed method SwMC is preferred. This new multiview clustering method does not depend on an additional parameter to learn the weights and without much precision loss, is interesting and can be acceptable.

4 Conclusion

In this paper, to recover the block diagonal matrix from multiple original input graphs, we firstly propose Parameter-weighted Multiview Clustering (PwMC) method. By analyzing its weaknesses, we further propose a new weight learning strategy in multiview clustering, named as Self-weighted Multiview Clustering (SwMC). The proposed methods are verified on two toy datasets and four benchmark datasets. Experimental results demonstrate the effectiveness of our methods. Especially, we note that SwMC achieves the close performance with PwMC but is practical to use. In future work, like [Nie et al., 2016a], we will consider extending this framework to semi-supervised context.
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