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# Dense community detection in multi-valued attributed networks

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#### ABSTRACT

The proliferation of rich information available for real world entities and their relationships gives rise to a general type of graph, namely *multi-valued attributed graph*, where graph vertices are associated with a number of attributes and a vertex may have multiple values on an attribute. It would be useful if we can cluster such graphs into densely connected components with homogeneous attribute values. Recent work has studied graph clustering in attributed graphs considering the full attribute space. However, full space clustering often leads to poor results due to irrelevant attributes.

In this paper, we study subspace clustering in multi-valued attributed graph and propose an algorithm SCMAG for community detection. Our algorithm uses a cell-based subspace clustering approach and identifies cells with dense connectivity in the subspaces. Random walk with restart is used to measure the structural connectivity and attribute similarity. An indexing scheme is designed to support efficiently calculating cell connectivity from random walk scores. We also propose a new cell combining strategy on dimensions of categorical attributes and a novel mechanism to handle multi-valued attributes. Experimental results on IMDB data and bibliographic data demonstrate that SCMAG significantly outperforms the state-of-the-art subspace clustering algorithm and attributed graph clustering algorithm. Case studies show SCMAG can find dense communities with homogeneous properties under different subspaces.

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#### 46 **1. Introduction**

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47 Graph clustering has attracted a lot of attention recently in the literature. The goal of graph clustering is to group densely connected vertices into a cluster. Graph clustering has found broad applications in community detection, image seg-48 mentation, identification of functional modules in large protein-protein interaction networks, etc. Nowadays with rich infor-49 50 mation available for real world entities and their relationships, graphs can be built in which vertices are associated with a set of attributes describing the properties of the vertices. Such attributes can be numerical or categorical. For some attribute, an 51 52 entity can have more than one value on that attribute. For example, the genre of a movie can be both "drama" and "crime"; 53 the research topics of a researcher can be "database", "data mining" and "bioinformatics". Multi-valued attributes are very 54 common in many real world data. This gives rise to a new and more general type of graph, called *multi-valued attributed* 55 graph.

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Traditional graph clustering methods [29,21,34,26,27] mainly focus on the connections in a graph and try to achieve a dense subgraph within a cluster. But these methods do not consider attribute information associated with graph vertices. On the other hand, a graph summarization method [31] partitions a graph according to attribute values, but does not enforce dense connections within a partition.

60 There have been some recent studies [6,37] on clustering attributed graph, i.e., SA-Cluster and its fast version Inc-Cluster, 61 which partition the graph into several densely connected components with similar attribute values. SA-Cluster finds non-62 overlapping clusters in the full attribute space. Although SA-Cluster differentiates the importance of attributes with an attri-63 bute weighting strategy, it cannot get rid of irrelevant attributes completely. As graph vertices may have many attributes, the 64 high-dimensional clusters are hard to interpret, or there is even no significant cluster with homogeneous attribute values in 65 the full attribute space. If an attributed graph is projected to different attribute subspaces, various interesting clusters embedded in subspaces can be discovered which, however, may not exhibit in the full attribute space. In this paper, we study 66 67 subspace clustering on multi-valued attributed graph, and discover clusters embedded in subspaces. Such subspace clusters should not only have homogeneous attribute values but also have dense connections, i.e., correspond to communities with 68 69 homogeneous properties. The detected clusters in multi-valued attributed graphs can overlap, as the nodes may belong to multiple clusters in different subspaces. In contrast, non-overlapping community detection partitions the network into sev-70 eral disjoint clusters. Thus, overlapping communities are more common and natural than non-overlapping communities in 71 attributed graphs. For example, an individual in a social network belongs to many social circles corresponding to different 72 73 relationships, such as friends, schoolmates, family, research community and so on. Let us look at an example to illustrate the 74 motivation for this subspace clustering problem.

Fig. 1(a) shows a coauthor network with three attributes {*Topic, Affiliation, Country*}. A vertex represents an author and an edge represents the coauthor relationship between two authors. The attributes and their possible values are listed in Table 1. *Topic* is a multi-valued attribute and an author can have one or more topics, e.g., author  $v_4$ . The problem is how to partition the coauthor network into clusters with close collaborations and homogeneous attribute values.

If we apply SA-Cluster [6], the coauthor network is partitioned into 4 clusters, as shown in Fig. 1(b). But this clustering result is not satisfactory, because (1) the attribute values in the same cluster are still quite different, if considering the full attribute space; and (2)  $v_9$  is disconnected from his coauthors and forms a single-node cluster. This is not reasonable. In a high-dimensional attributed graph this problem may become even worse, as it is hard to find clusters with dense connectivity and homogeneous attribute values in the full space.

If we consider subspace clustering, then we can find two clusters under the subspace {*Affiliation, Country*} in Fig. 2(a), and another two clusters under the subspace {*Topic, Affiliation*} in Fig. 2(b). These subspace clusters make much more sense because they not only have homogeneous attribute values in the respective subspace, but also have a cohesive structure within a cluster. Note that, if traditional subspace clustering methods, e.g., CLIQUE [2] and ENCLUS [5] are applied, under the subspace {*Affiliation, Country*} we will have two more clusters { $v_1$ ,  $v_4$ ,  $v_6$ }, { $v_2$ ,  $v_3$ ,  $v_5$ ,  $v_7$ } respectively with attribute values [*Univ., AU*] and [*Univ., CN*]. However, we can easily find these two clusters have very sparse connections, thus not closely collaborating groups.

This example shows neither the recent attributed graph clustering algorithms [6,37] which consider the full attribute space, nor the traditional subspace clustering algorithms [2,5] which completely ignore the structural connectivity are suitable to solve the problem of multi-valued attributed graph clustering. The unique challenges in this problem include the following:

how to discover subspaces under which densely connected clusters with homogeneous attribute values are embedded?
 For example, there are meaningful clusters under the two subspaces shown in Fig. 2(a) and (b), but there is no cluster
 with dense structural connectivity under the subspace {*Topic, Country*};

98 2. how to properly handle categorical attributes and multi-valued attributes. Traditional subspace clustering algorithms,
99 e.g., CLIQUE [2] and ENCLUS [5], only handle numerical attributes. But in real data categorical and multi-valued attributes
100 are very common, for example, v<sub>4</sub> has two topics as *DB* and *DM*. Which cluster should v<sub>4</sub> belong to in Fig. 2(b)?

3. how to enforce both structural connectivity and attribute similarity requirements, like in the clusters in Fig. 2 (a) and (b)?

There have been some recent studies which combine attribute subspace clustering and dense subgraph mining on a graph 103 with feature vectors, e.g., CoPaM [19], GAMer [12,13], DB-CSC [11]. CoPaM and GAMer follow a cell-based subspace clustering 104 approach to find clusters that show high similarity in a feature subspace and are densely connected in the given graph, while 105 DB-CSC takes a density-based approach to find subspace clusters. They have the different limitations: (1) CoPaM and GAMer 106 do not have a cell merging strategy for adjacent cells, and the formation of a cell depends on the initial vertex to start with; 107 108 (2) CoPaM strictly requires all nodes in a cluster have the same value on each attribute in the subspace. GAMer and DB-CSC 109 use a single parameter maximal width w to control the value difference on all the attributes in the concerned subspace within 110 a cell. But for different attributes, which can be categorical or numerical ones, it is hard to set a uniform threshold to control 111 the attribute value differences; (3) The time complexity of GAMer and DB-CSC increases exponentially with the number of 112 vertices in the input graph, which can hardly scale with large graphs. Even though adjusting by various parameter settings, 113 GAMer is proven to be too slow to generate results in our experiments; and (4) According to the experimental results 114 reported in [12,13,11], only clusters with small size are found by GAMer and DB-CSC on real datasets, i.e., the average size 115 of the found clusters is only around 10 in a graph with 10k vertices. Similarly, according to [19], the largest found subgraph

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by CoPaM has 18 vertices in a graph with 1942 vertices. Overall, the efficiency challenge of detecting large communities by
 CoPaM, GAMer and DB-CSC have not been verified yet. Large communities naturally exist in many real-world networks, e.g.
 in terms of research areas, the research community of data mining in the coauthor network would contain far more than
 1000 individuals.

Different from these methods, in this paper we propose a novel subspace clustering algorithm on attributed graphs. Our clustering framework is similar to that in ENCLUS [5]. We first find interesting subspaces with good clustering using an entropy-based method. Then we identify cells with dense structural connectivity in the grid under the subspaces. Clusters are formed by merging adjacent cells. Random walk with restart on an attribute augmented graph is used to measure the structural connectivity between vertices as well as the attribute similarity. Our main contributions include:

We study the problem of graph subspace clustering on attributed graphs. We give a cell-based subspace clustering def inition, and propose a novel algorithm, called SCMAG (Subspace Clustering on Multi-valued Attributed Graph). With our
 formulation, we can discover more meaningful clusters which are closely connected and exhibit high attribute similarity
 under some subspaces. Subspace clusters may overlap and they provide multiple angles to exhibit cluster patterns under
 different subspaces.

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- We use random walk with restart to unify the structural closeness and attribute similarity into a single measure. Based on
   the random walk score, we design a novel cell combining strategy on dimensions of categorical attributes and an effective
   mechanism to handle multi-valued attributes.
- We consider both structural connectivity and attribute similarity in the subspace clustering process. We design an index
   scheme to speedup calculating the connectivity of a cluster. An upper bound is also derived for efficiently pruning clusters
   with low connectivity.
- We conducted extensive experiments on real datasets to evaluate the performance of our method in terms of structural density and attribute value similarity, and demonstrate its superiority over existing methods. We also show some interesting clusters we discovered in the case study.

The rest of this paper is organized as follows. Section 2 discusses related work. Section 3 introduces the preliminary concepts and describes our random walk based closeness measure. Our subspace clustering algorithm SCMAG is described in Section 4. We present extensive experimental results in Section 5. Finally Section 6 concludes our paper.

#### 144 **2. Related work**

Subspace clustering [2,22,18,23] has been studied extensively since late 1990s. CLIQUE [2] was one of the first subspace 145 146 clustering algorithms which uses an APRIORI search framework to find dense subspace clusters. ENCLUS [5] is an extension 147 of CLIQUE which proposes to use entropy to find subspaces with good clustering. DOC [25] is a density-based projective clus-148 tering algorithm which uses a Monte Carlo procedure to approximate with high probability an optimal projective cluster. PROCLUS [1] focuses on the clustering results to find appropriate subspace such that the inner-cluster points are close. 149 These subspace clustering algorithms only consider numerical attributes but completely ignore the relationships between 150 objects. Thus they are not suitable to cluster attributed graphs. A comprehensive survey of subspace clustering algorithms 151 152 proposed in recent years can be found in [30].

Traditional graph clustering methods [28] aim to have a dense subgraph within a cluster. Many graph clustering 153 algorithms have been proposed based on various criteria including normalized cuts [29], modularity [21], structural 154 density [34], weighted kernel k-means [10] or stochastic flows [26]. Satuluri et al. [27] study how to sparsify a graph 155 156 for scalable clustering without sacrificing quality. Recently, several new community model based different types of 157 dense subgraph are proposed for community search, such as quasi-clique community model [8] and k-truss community model [14]. But these methods do not consider attribute information associated with graph vertices. Some 158 159 recent studies consider summarizing or clustering attributed graphs in the full attribute space. Tian et al. [31] proposed OLAP-style aggregation approaches to summarize large graphs by grouping nodes based on user-selected attri-160 butes. This method achieves homogeneous attribute values within clusters, but ignores the structure connectivity. 161 Recently, Zhou et al. have proposed a graph clustering algorithm, SA-Cluster [6] and its fast version Inc-Cluster 162 [37], based on both structural and attribute similarities. [4] proposed a spectral method for clustering on attributed 163 graphs. A parameter-free method [15] is proposed to identify dense clusters and outliers in attributed graphs. Long 164 et al. [17] proposed a collective factorization model for multi-type relational data clustering. Yang et al. [35] studied 165 the problem of detecting overlapping communities in networks with node attributes. However, these algorithms 166 consider clustering in the full attribute space, thus it may be hard to find good clusters in high-dimensional attrib-167 168 uted graphs.

Random walk with restart (RWR) [33] provides a good relevance measure between two nodes in weighted graphs, and it 169 170 has wide applications in many domains, such as automatic captioning of images, generalizations to the connection sub-171 graphs, personalized PageRank. Jeh and Widom [16] defined a measure called SimRank, which measures the similarity between two vertices in a graph by their neighborhood similarity. Pons and Latapy [24] proposed a method to use short ran-172 dom walks of length l to calculate the similarity between two vertices for community detection in a graph. Desikan et al. [9] 173 174 proposed an incremental algorithm to effciently compute PageRank for the dynamic graph by partitioning the graph into a 175 changed part and an unchanged part. [36] studied a new problem of reverse top-k proximity search in graphs based on ran-176 dom walk with restart.

There have been some recent studies which mine cohesive subgraph patterns from a graph with feature vectors, and 177 combine the concepts of dense subgraph mining and subspace clustering. These methods include CoPaM [19], GAMer 178 [12,13], and DB-CSC [11]. Following a cell-based subspace clustering approach, CoPaM and GAMer mine clusters contain-179 ing all objects falling within a cell, i.e., the attribute values differ by at most a given threshold, with some density con-180 181 straint. However, neither methods have a cell merging strategy to merge adjacent cells, which is different from our work. The formation of a cell also depends on the initial vertex to start with and its feature vector. DB-CSC takes a density-182 183 based approach for finding subspace clusters. Similar to GAMer, DB-CSC uses a single parameter maximal width w to control the value difference on all the attributes in the concerned subspace within a cell. But for different attributes, which 184 can be categorical or numerical ones, it is hard to set a uniform threshold to control the attribute value differences. 185 186 Another issue is that the time complexity of GAMer and DB-CSC increases exponentially with the number of vertices

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in the graph, which can hardly scale with large graphs. In addition, according to the experimental results in [12,13,11], only clusters with small size are found and shown by GAMer and DB-CSC on real datasets, i.e., the average size of the found clusters is only around 10 in a graph with 10k nodes.

#### 190 **3. Preliminary concepts**

In this section, we firstly give our problem definition and a brief introduction of ENCLUS, a subspace clustering method for numerical data, on which our graph clustering method is built. Then we describe how to measure the structural connectivity and attribute similarity of graph vertices. We also propose a novel strategy to handle the categorical and multi-valued attributes in graph subspace clustering.

195 3.1. Problem definition

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**Definition 3.1** (*Multi-valued Attributed Graph*). A multi-valued attributed graph is denoted as G = (V, E, A, w), where V is the vertex set, E is the edge set,  $A = \{A_1, ..., A_d\}$  is a set of categorial or totally ordered numerical attributes. Each vertex  $v \in V$  is associated with an attribute vector  $A(v) = [A_1(v), ..., A_d(v)]$ , where  $A_i(v)$  is the attribute value that vertex v has on attribute  $A_i$ . The domain of a categorical attribute  $A_i \in A$  contains a finite set of values, denoted as  $Dom(A_i) = \{a_{i1}, ..., a_{in_i}\}$ ,  $|Dom(A_i)| =$  $n_i$ . If  $A_i$  is a multi-valued categorical attribute,  $A_i(v)$  contains a set of attribute values from  $Dom(A_i)$ , i.e.,  $A_i(v) \subseteq Dom(A_i)$ . The domain of a numerical attribute  $A_j \in A$  is  $\mathbb{R}$ .  $w : E \mapsto \mathbb{R}^+$  is an edge weight function. For each  $(u, v) \in E$ , the edge weight is denoted as w(u, v). We denote the size of vertex set as |V| = N.

Many real-world networks can be represented as multi-valued attributed graphs with weighted edges, such as co-author network and IMDB network. Continue with our example in Fig. 1(a), the weight of edge (u, v) in co-author networks can be assigned with the number of co-authored papers published by authors u and v, which represents the collaboration strength. The larger weight of w(u, v) is, the stronger relationship between u and v is.

**Remark 3.1.** In the paper, our proposed subspace clustering method can also handle graphs with multi-valued edge labels, in which the edges can be associated with multiples values in attributes. To handle such a graph, denoted as *G*, with multivalued edge labels, we need to construct a new attributed graph *G'* as follows. For each edge (u, v) in *G*, a new vertex  $e_{uv}$  is inserted into *G'*. We assign  $e_{uv}$  with the same attribute vector of edge (u, v). For two edges (u, v) and (u, w) adjacent to *u* in *G*, a new edge  $(e_{uv}, e_{uw})$  is added between two vertices  $e_{uv}, e_{uw}$  in *G'*. Then, *G'* is a multi-valued attributed graph by Definition 3.1, our subspace clustering methods can be applied on *G'*.

If an attribute set  $S \subseteq A$ , we say S is a subspace of A. The problem of subspace clustering on a multi-valued attributed graph is to find a set of subspaces  $\{S_1, \ldots, S_m\}$  with good clustering, where  $S_i \subseteq A, 1 \le i \le m$ . Under each subspace  $S_i$ , discover embedded clusters which have both high structural connectivity and high attribute similarity in the concerned subspace. We will define the formal requirements of "good subspace" and "embedded clusters" in Section 4.1.

#### 219 3.2. ENCLUS: a brief introduction

Our proposed graph subspace clustering algorithm is built based on ENCLUS [5], an entropy-based subspace clustering method on data objects with numerical features. We briefly describe ENCLUS here. It first divides each numerical attribute into intervals like [l, u), where l and u denote the interval lower bound and upper bound, respectively. Then the space is partitioned to form a grid. A cell has the form  $\{c_1, c_2, ..., c_d\}$  where  $c_i = [l_i, u_i)$  is an interval of attribute  $A_i$ . The subspace entropy is defined as follows.

**Definition 3.2** (*Subspace Entropy*). Given a set of attributes  $S = \{A_1, \ldots, A_k\} \subseteq A$ , the subspace entropy of S is defined as

$$H(A_1,\ldots,A_k) = -\sum_{a_1 \in Dom(A_1)} \cdots \sum_{a_k \in Dom(A_k)} p(a_1,\ldots,a_k) \log p(a_1,\ldots,a_k)$$
(1)

where  $p(a_1, ..., a_k)$  is the percentage of graph vertices whose attribute value vector is  $[a_1, ..., a_k]$ .

The framework of ENCLUS consists of the following three steps.

1. Find interesting subspaces with good clustering by an entropy-based method;

- 232 2. Find clusters in the identified subspace by merging adjacent dense cells;
- 233 3. Output the clusters.

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The term "good clustering" means that a subspace contains a good set of clusters. Formally the goodness of a sub-235 236 space is measured by entropy. Given an entropy threshold, a subspace whose entropy is below the threshold is consid-237 ered to have good clustering. ENCLUS uses a bottom-up approach similar to APRIORI to find good subspaces. It starts 238 with finding one-dimensional subspaces with good clustering. Then it generates the candidate two-dimensional sub-239 spaces from the one-dimensional ones and checks the candidates against the raw data to identify those that actually 240 have good clustering. This process is repeated with increasing dimensionalities until no more subspaces with good clus-241 tering are found. The correctness of this bottom-up approach is based on the *downward closure property* of entropy, as proved in [5]. 242

Under each good subspace, adjacent dense cells are combined to form a cluster. Two *k*-dimensional cells are called "*adjacent*" if they have the same attribute value intervals on k - 1 dimensions, and on the remaining dimension, the interval upper bound of a cell is equal to the interval lower bound of the other cell. Finally the clusters are returned to users.

#### 246 3.3. A unified closeness measure

247 ENCLUS only considers the attribute values of data objects, but ignores the connections between them. In our graph subspace clustering problem, to calculate the structural connectivity of vertices, we propose to use random 248 walk with restart to measure the closeness between two vertices. According to the properties of small-world net-249 works, the average shortest distance between two nodes would be significantly small. For example, in the Facebook 250 social network, the average shortest distance was 4.74 in 2011 [3]. This follows from the fact that two "distant" 251 252 can have a short path flow through hubs. In this case, the number of paths between two nodes would be more effec-253 tive for measuring their connectivity than the accurate shortest distance. As a result, it is better to use random walk with restart to measure the connectivity of vertices, than simple distance measures, e.g., short distance and nearest 254 neighbors. 255

**Definition 3.3** (*Neighborhood Random Walk*). Let *P* be the  $N \times N$  transition probability matrix of a graph. Given *L* as the maximum length a random walk can go,  $c \in (0, 1)$  as the restart probability, the neighborhood random walk matrix is

$$R = \sum_{l=1}^{L} c(1-c)^{l} P^{l}$$
(2)

The random walk score of two vertices  $v_i, v_j \in V$  is  $R(v_i, v_j)$ , measuring the probability that  $v_i$  can reach  $v_j$  within *L* steps.

For the attributes on graph vertices, we do the following preprocessing. If  $A_i$  is a numerical attribute, it is discretized into 263 n<sub>i</sub> intervals first. There are some studies, e.g., MAFIA [20], which consider adaptive interval partitioning of numerical attri-264 265 butes in subspace clustering. But this is not the focus of our work. In our more general problem setting, we also consider categorical and multi-valued attributes. We have to address two problems: (1) how to measure the similarity between 266 two categorical values and determine which categorical values are "adjacent" for merging cells with categorical attributes? 267 and (2) if a vertex has multiple values on one attribute, which cell should the vertex be assigned to in the formed grid? To 268 solve these problems, we consider the neighborhood of a vertex to exploit the connections with its neighbors as well as the 269 attribute similarities with its neighbors. We construct the attribute augmented graph similarly as in [6] [32]. The definition is 270 271 as follows.

**Definition 3.4** (*Attribute Augmented Graph*). Given a multi-valued attributed graph denoted as G = (V, E, A, w) where  $A = \{A_1, ..., A_d\}$ . The domain of an attribute  $A_i$  is denoted as  $Dom(A_i) = \{a_{i1}, ..., a_{in_i}\}$ ,  $|Dom(A_i)| = n_i$ . An attribute augmented graph is denoted as  $G_a = (V \cup V_a, E \cup E_a)$ , where  $V_a = \{v_{ij} | 1 \le i \le d, 1 \le j \le n_i\}$  is the set of attribute vertices. An attribute vertex  $v_{jk} \in V_a$  represents the value  $a_{jk}$  on attribute  $A_j$ . An edge  $(v_i, v_{jk}) \in E_a$  iff  $a_{jk} \in A_j(v_i)$ , i.e., vertex  $v_i$  has value  $a_{jk}$  on attribute  $A_j$ . An edge  $(v_i, v_j) \in E$  is called a structure edge and an edge  $(v_i, v_{jk}) \in E_a$  is called an attribute edge.

For instance, we construct an attribute augmented graph shown in Fig. 3(b) for the co-author network in Fig. 3(a). In this attribute augmented graph, we add a set of dummy vertices  $V_a\{v_8, v_9, v_10, v_{11}, v_{12}\}$ , e.g.  $v_8$ , which represents the attribute value "DB" in "Topic". Authors with corresponding attribute values are connected to the two vertices respectively in dashed lines, e.g. the edge between  $v_1$  and  $v_8$ , indicating that author  $v_1$  has the value "DB" in the attribute of "Topic". With the newly inserted attribute edges, authors who are originally connected become more densely conncetted if they share common attribute values, e.g.,  $v_5$  and  $v_7$ .

With such a graph augmentation, we express the attribute similarity as vertex closeness in the graph: two vertices which share an attribute value are connected by a common attribute vertex. In the attribute augmented graph, two structure vertices  $v_i$  and  $v_j$  are close either if they are connected through many other structure vertices, or if they share many common attribute vertices as neighbors, or both. We simulate a length-*L* neighborhood random walk on the attribute augmented graph, which goes through both structure edges and attribute edges. The random walk score effectively combines structural closeness and attribute similarity into a unified closeness measure between graph vertices. The transition matrix  $P_A$  of the attribute augmented graph is a  $|V \cup V_a| \times |V \cup V_a|$  matrix. We denote  $|V_a| = N_A$ .

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Fig. 3. An example of attribute augmented graph with topics, affiliation and country.

We use  $N(v_i)$  to denote the set of structure vertices in  $v_i$ 's neighborhood, i.e.,  $N(v_i) = \{v_j | v_j \in V, (v_i, v_j) \in E\}$ , and use  $N_A(v_i)$  to denote the set of attribute vertices in  $v_i$ 's neighborhood, i.e.,  $N_A(v_i) = \{v_{jk} | v_{jk} \in V_a, (v_i, v_{jk}) \in E_a\}$ . For any  $v_i \in V$ , we assume  $\sum_{v_j \in N(v_i)} P_A(v_i, v_j) = \sum_{v_{jk} \in N_A(v_i)} P_A(v_i, v_{jk}) = 0.5$ , i.e., structure edges take 0.5 transition probability and attribute edges take 0.5 transition probability. Thus the transition matrix  $P_A$  is defined as follows. The transition probability from vertex  $v_i$  to  $v_j$  is

$$p_{\nu_{i},\nu_{j}} = \begin{cases} \frac{w(\nu_{i},\nu_{j})}{2\sum_{(\nu_{i},\nu_{k})\in E}w(\nu_{i},\nu_{k})}, & \text{if } (\nu_{i},\nu_{j})\in E\\ 0, & \text{otherwise} \end{cases}$$
(3)

Similarly, we define the transition probability from vertex  $v_i$  to attribute vertex  $v_{jk}$  as

$$p_{v_i,v_{jk}} = \begin{cases} \frac{1}{2|\mathcal{A}||A_j(v_i)|}, & \text{if } (v_i, v_{jk}) \in E_a \\ 0, & \text{otherwise} \end{cases}$$
(4)

The transition probability from vertex  $v_{ik}$  to vertex  $v_i$  is

$$p_{v_{jk},v_i} = \begin{cases} \frac{1}{|N(v_{jk})|}, & \text{if } (v_{jk}, v_i) \in E_a \\ 0, & \text{otherwise} \end{cases}$$
(5)

where  $N(v_{jk})$  denotes the set of vertices in  $v_{jk}$ 's neighborhood. The transition probability between two attribute vertices  $v_{ik}$ and  $v_{jl}$  is 0 since there is no edge between them.

$$p_{\nu_{ik},\nu_{il}} = 0, \forall \nu_{ik}, \nu_{jl} \in V_a \tag{6}$$

Integrating Eqs. (3)–(6) together, we can get the transition probability matrix  $P_A$  on the attribute augmented graph. Following Eq. (2) we can compute the random walk matrix  $R_A$  from  $P_A$ .  $R_A(v_i, v_j)$  is the closeness between vertices  $v_i$  and  $v_j$  in the attribute augmented graph.

314 We denote  $R_A$  as four submatrices

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$$R_A = \begin{bmatrix} Q_{VV} & Q_{VA} \\ Q_{AV} & Q_{AA} \end{bmatrix}$$

where  $Q_{VV}$  is an  $N \times N$  matrix representing the random walk scores between structure vertices,  $Q_{VA}$  is an  $N \times N_A$  matrix representing the random walk scores from structure vertices to attribute vertices,  $Q_{AV}$  is an  $N_A \times N$  matrix representing the random walk scores from attribute vertices to structure vertices, and  $Q_{AA}$  is an  $N_A \times N_A$  matrix representing the random walk scores between attribute vertices.

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We further perform a row normalization of Q<sub>W</sub> as

$$\widetilde{Q}_{VV}(\boldsymbol{v}_i, \boldsymbol{v}_j) = \frac{Q_{VV}(\boldsymbol{v}_i, \boldsymbol{v}_j)}{\sum_{\boldsymbol{v}_k \in V} Q_{VV}(\boldsymbol{v}_i, \boldsymbol{v}_k)}$$
(7)

With the normalized scores, it is easier for users to determine the cluster connectivity threshold, i.e., not depending on the absolute scores which may vary from graph to graph.

#### 328 3.4. Combining cells with categorical attributes

329 Let us get back to the first problem: how to measure the similarity between two categorical values and determine which categorical values are "adjacent" for merging cells with categorical attributes? As an example, consider attribute Conference in a 330 331 bibliographic data, with its domain {VLDB, SIGMOD, ICDE, ICCV, CVPR, ...}. How can we tell which conferences are more similar and which are less similar? Intuitively, if many authors publish in both VLDB and SIGMOD, then VLDB and SIGMOD are 332 333 similar. Consider the random walk submatrix  $Q_{AA}$ . For two different values  $a_{ii}$ ,  $a_{ir}$  of the same categorical attribute  $A_i$ , the 334 entry  $Q_{AA}(v_{il}, v_{ir})$  measures the random walk closeness between these two values. The rationale is, if  $a_{il}$  and  $a_{ir}$  of attribute  $A_i$  are shared by many vertices, they will have a high random walk score, indicating their closeness in the attribute aug-335 336 mented graph. Thus we define the categorical value adjacency based on random walk score. For an attribute value  $a_{ij}$ , the 337 adjacent values are defined as 338

$$Adj(a_{il}) = \{a_{ir}|TopK_{r=1\ r\neq l}^{n_i}(Q_{AA}(v_{il}, v_{ir}))\}$$
(8)

The adjacent values of a categorical attribute value  $a_{il}$  is the set  $\{a_{ir}\}$  with the top-*K* largest random walk score  $Q_{AA}(v_{il}, v_{ir})$ . Then in the cell-based subspace clustering approach, we can combine two *k*-dimensional cells, if they share common values in (k-1) dimensions, and on the remaining dimension they have either adjacent numerical values or adjacent categorical values as defined above.

#### 345 3.5. Handling multi-valued attributes

346 Let us consider the second problem we listed: if a vertex has multiple values on one attribute, which cell should the vertex be 347 assigned to in the formed grid? Consider vertex  $v_4$  in Fig. 1(a), which has values DB and DM on attribute Topic. Should  $v_4$  be assigned to the "DB" cluster or the "DM" cluster? If we consider  $v_4$ 's neighborhood, we can find  $v_4$  only has one collaborator 348  $v_3$  working on DB, but has three collaborators  $v_5$ ,  $v_6$ ,  $v_7$  working on DM. This indicates DM is more important to  $v_4$  than DB 349 350 (in this example we assume  $v_4$  has the same collaboration strength with each of his collaborators, as concrete edge weights 351 are not given in Fig. 1(a). In our algorithm, we actually consider the edge weights). To distinguish the multi-values, we consider the random walk submatrix  $Q_{VA}$ . An entry  $Q_{VA}(v_i, v_{jk})$  measures the closeness between vertex  $v_i$  and attribute vertex 352  $v_{ik}$ . The rationale is, besides vertex  $v_i$  itself has attribute value  $a_{ik}$ , if many of  $v_i$ 's neighbors also have attribute value  $a_{ik}$ ,  $v_i$ 353 and  $v_{jk}$  will have a high random walk score, indicating their closeness in the attribute augmented graph. Thus we will reas-354 sign the attribute value  $a_{ik}$  to  $v_i$  which has the maximum closeness to  $v_i$ , i.e., 355

$$A_{j}(v_{i}) = a_{jk}, k = \arg \max_{r=1}^{n_{j}} Q_{VA}(v_{i}, v_{jr})$$
(9)

Finally, each vertex is associated with an attribute vector containing a single value in each attribute.

**Remark 3.2.** Assume that for a co-author network in Fig. 1(a), one author  $v_4$  has the same number of collaborators respectively working on *DB* and *DM* in its neighborhood. But, for  $v_4$ , its closeness value to attributes "DB" and "DM" could still be different. This is because even with the same number of "DB" and "DM" authors, the neighborhood structure of "DB" and "DM" clusters can still differ. Then, in Eq. (1), the closeness values are calculated based on the graph structure and attributes vectors of  $v_4$ 's neighbors, which would be different. Actually, in real applications, the tie closeness will seldomly happen, which helps to choose a unque attribute value with the highest closeness score.

#### 366 4. Our algorithm SCMAG

Our graph subspace clustering framework is similar to that of ENCLUS described in Section 3.2. In the following, we will first define the criteria for interesting subspace with good clustering tendency and find such subspaces. As each attribute  $A_i$ has  $n_i$  values, for a subspace  $\{A_1, \ldots, A_k\}$ , the *k*-dimensional space is partitioned to form a grid. We will identify cells in the grid with high coverage and high connectivity. Adjacent qualified cells will be merged to form a maximal cluster in the subspace.

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372 4.1. Criteria of subspace clustering

We follow the entropy-based subspace clustering method ENCLUS [5] to find interesting subspaces. The subspace entropy is defined in the Definition 3.2.

Remark 4.1. Note that the definition of subspace entropy assumes that each attribute of an object only has one value. Thus,
 ENCLUS cannot be directly applied to compute the subspace entropy due to the multi-values in a single attribute. On the
 other hand, our method SCMAG can handle such graphs. The variant methods of ENCLUS for multi-valued attribute graphs
 will be discussed in Section 5.1.

In addition, we want the attributes of a subspace to be correlated. If the attributes are independent of each other, the subspace does not give more information than looking at each attribute independently. We measure the correlation of a subspace S using mutual information between all individual dimensions of the subspace as below.

$$I(\{A_1,...,A_k\}) = \sum_{i=1}^k H(A_i) - H(A_1,...,A_k)$$

We consider a subspace  $S = \{A_1, \dots, A_k\}$  as an interesting subspace, if S is more strongly correlated than any of its subsets  $S' \subseteq S$ . To measure the increase in correlation of a subspace, we define the *interest* of a subspace.

**Definition 4.1** (*Subspace Interest*). Given a set of attributes  $S = \{A_1, \dots, A_k\} \subseteq A$ , the subspace interest of S is defined as the minimum increase in correlation of S over its (k - 1)-dimensional subsets.

*interest*
$$(A_1, \ldots, A_k) = I(\{A_1, \ldots, A_k\}) - \max_{i \in I} I(\{A_1, \ldots, A_k\} - \{A_i\})$$

Given a maximum entropy threshold  $e_{max}$  and a minimum interest threshold  $i_{min}$ , we want to find all subspaces  $S \subseteq A$  such that  $H(S) \leq e_{max}$  and *interest* $(S) \geq i_{min}$ .  $e_{max}$  guarantees the low entropy of generated subspaces, and  $i_{min}$  contributes to throw out subspaces with low increased correlation.

<sup>395</sup> Under a subspace  $S = \{A_1, \dots, A_k\}$ , the space is partitioned to form a grid. We identify cells with high coverage and con-<sup>396</sup> nectivity, according to the following definition.

**Definition 4.2** (*Coverage and Connectivity*). Given a cell *u* in a subspace, the coverage of *u* is measured by the number of vertices in *u*, i.e., V(u) = |u|. The connectivity of *u* is measured by the sum of random walk scores of all pairs of vertices, divided by the cell size

$$D(u) = \frac{\sum_{v_i, v_j \in u} Q_{W}(v_i, v_j)}{|u|}$$
(10)

Given a minimum coverage threshold  $v_{min}$  and a minimum connectivity threshold  $d_{min}$ , we only consider cells such that  $V(u) \ge v_{min}$  and  $D(u) \ge d_{min}$ . We will discuss the parameter setting in Section 4.5.

405 4.2. Properties

406 From the above definitions of subspace clustering criteria, we can derive two lemmas as below.

407 **Lemma 4.1.** If a k-dimensional subspace  $\{A_1, \ldots, A_k\}$  has entropy below the threshold  $e_{max}$  and there exists at least one high 408 coverage cluster C, then the property is established in any (k - 1)-dimensional projections of this space.

This property has been proved in [2,5]. From Lemma 4.1, we can use the bottom-up search strategy to find interesting subspaces level by level.

411 **Lemma 4.2.** Suppose two cells  $u_1, u_2$  both satisfy the  $d_{min}$  and  $v_{min}$  thresholds. When merging  $u_1$  and  $u_2$  to form a cluster 412  $C = u_1 \cup u_2, C$  also satisfies  $V(C) \ge v_{min}$  and  $D(C) \ge d_{min}$ .

413 **Proof 4.1.** 
$$V(C) = V(u_1) + V(u_2) \ge v_{min}$$

$$D(C) = \frac{\sum_{v_i, v_j \in u_1} \tilde{Q}_W(v_i, v_j) + \sum_{v_i, v_j \in u_2} \tilde{Q}_W(v_i, v_j)}{|u_1| + |u_2|} + \frac{\sum_{v_i \in u_1, v_j \in u_2} \tilde{Q}_W(v_i, v_j) + \sum_{v_i \in u_2, v_j \in u_1} \tilde{Q}_W(v_i, v_j)}{|u_1| + |u_2|} \\ \ge \frac{\sum_{v_i, v_j \in u_1} \tilde{Q}_W(v_i, v_j) + \sum_{v_i, v_j \in u_2} \tilde{Q}_W(v_i, v_j)}{|u_1| + |u_2|} \ge \frac{|u_1| \times d_{\min} + |u_2| \times d_{\min}}{|u_1| + |u_2|} = d_{\min} \quad \Box$$

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#### 417 4.3. *Mining subspace clusters*

#### 419 Algorithm 1. SCMAG

### **Input:** $G, e_{max}, i_{min}, v_{min}$ , and $d_{min}$ .

Output: all clusters satisfying coverage and connectivity requirements in interesting subspaces.

1: 2:	Calculate the unified random walk matrix $R_A$
3:	Assign each vertex $v_i$ with new attribute value in multi-valued attributes according to Eq. (9)
4:	Let $k = 1$ and $S_k$ be all one-dimensional subspaces
5:	for each subspace $s \in S_k$ do
6:	<b>if</b> $H(s) \leq e_{max}$ , and exists a cell $u$ in $s$ s.t. $V(u) \geq v_{min}$ <b>then</b>
7:	$CS_k = CS_k \cup \{s\}$
8:	if $interest(s) \ge i_{min}$ , and exists a cell u in s s.t. $D(u) \ge d_{min}$ then
9:	$Result = Result \cup Find - Clusters(s)$
10:	$S_{k+1} = \text{Generate} - \text{Candidate}(CS_k)$
11:	<b>if</b> $S_{k+1} = \emptyset$ , <b>then</b> go to step 13
12:	k = k + 1, go to step 5
13:	Output clusters in Result

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According to Lemma 4.1, we will use a bottom-up search strategy, similar to the APRIORI style approach, to find subspaces satisfying  $e_{max}$  and  $i_{min}$  thresholds. We will start with 1-dimensional subspaces and join two *k*-dimensional subspaces which share a common prior (k - 1)-dimensional subspace to form a (k + 1)-dimensional subspace candidate by a procedure Generate – Candidate. If a subspace violates  $e_{max}$  requirement, we do not need to further grow it. Once interesting subspaces are found, a procedure Find – Clusters is invoked to find subspace clusters. Algorithm 1 shows the subspace clustering algorithm SCMAG.

Algorithm 2 lists the procedure Find – Clusters for finding subspace clusters. We first identify cells with high coverage and connectivity in a subspace *s*. Then we perform a depth-first search to merge adjacent cells to form a maximal cluster. DFS – Find – Clusters is a recursive algorithm which considers each attribute value  $a_{jr_j}$  of a cell *u* in the *k*-dimensional space and tries to combine *u* with other cells having adjacent values to  $a_{jr_j}$  on attribute  $A_j$  and identical values on the remaining attributes. Finally all cells with the same label *cluster\_num* forms a maximal cluster in the subspace.

#### 463 Algorithm 2. Find-Clusters

**Input:** All cells in subspace  $s = \{A_1, ..., A_k\}$ ,  $v_{min}$ , and  $d_{min}$ . **Output:** all clusters in *s*.

1:	Initialize $cluster\_num = 1$
2:	for each cell <i>u</i> in <i>s</i> do
3:	<b>if</b> $V(u) < v_{min}$ or $D(u) < d_{min}$ <b>then</b>
4:	Remove <i>u</i> from <i>s</i>
5:	for each cell <i>u</i> in <i>s</i> do
6:	if u.clu_id is undefined then
7:	$DFS - Find - Clusters(u, cluster\_num)$
8:	<i>cluster_num</i> = <i>cluster_num</i> + 1
9:	<b>Procedure</b> DFS – Find – Clusters
	<b>Input:</b> starting cell $u = [a_{1r_1}, \ldots, a_{kr_k}]$ , cluter_num.
	Output: a maximal cluster.
10:	Set u.clu_id = cluter_num
11:	for $j = 1:k$ do
12:	for each $a_{jt} \in Adj(a_{jr_i})$ do
13:	$u_t = [a_{1r_1}, \ldots, a_{jt}, \ldots, a_{kr_k}]$
14:	if <i>u<sub>t</sub>.clu_id</i> is undefined <b>then</b>
15:	$DFS - Find - Clusters(u_t, cluster\_num)$

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Fig. 4. Random walk scores.

#### 504 4.4. Computing cell connectivity efficiently

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We impose a connectivity requirement on a cell u, i.e.,  $D(u) = \frac{\sum_{v_i, v_j \in u} \widetilde{Q}_{VV}(v_i, v_j)}{|u|} \ge d_{min}$ . It takes  $O(|u|^2)$  to calculate D(u) for a cell u. To speed up the connectivity calculation, we design an index and a connectivity upper bound to prune cells with low connectivity.

The index is based on the following observation: for a vertex  $v \in V$ , there are only a small number of vertices with high closeness to v, but a majority of vertices have very low closeness to v. Fig. 4 shows the random walk scores of a vertex to all other vertices in a graph based on a bibliographic data we used in experiments. We can observe that most of the vertices have a very low random walk score close to 0. Therefore, we maintain two vertex sets for each vertex – one set of vertices with high closeness, denoted as *HC*, and the other set of vertices with low closeness, denoted as *LC*. We will start computing D(u) from the *HC* sets of vertices in u and see if we can stop early and tell whether  $D(u) \ge d_{min}$  or not.

We build the *HC* and *LC* sets for each vertex in *V* as follows. Consider the random walk submatrix  $\tilde{Q}_W$ . For a vertex  $v_i \in V$ , we sort all entries  $\tilde{Q}_W(v_i, v_j), \forall v_j \in V$  in the decreasing order. We insert the sorted entries  $\tilde{Q}_W(v_i, v_j)$  into  $v_i$ .*HC* one by one until  $\sum_{v_j \in v_i, HC} \tilde{Q}_W(v_i, v_j) \ge \delta$ , where  $\delta \in [0, 1)$  is a fixed threshold for all vertices. The remaining entries are inserted into  $v_i$ .*LC*. Table 2 shows the *HC* and *LC* sets for vertices in a graph under the setting  $\delta = 0.75$ . As we can see, the *HC* set usually contains a small number of vertices with high closeness, and *LC* contains a large number of vertices with low closeness.

For a cell u, we calculate D(u) in two stages. In the first stage we calculate the closeness entries in the *HC* sets only, as an estimation of the true D(u) value:

$$D_{HC}(u) = \frac{\sum_{v_i \in u} \sum_{v_j \in u, v_j \in v_i, HC} Q_{VV}(v_i, v_j)}{|u|}$$
(11)

 $If D_{HC}(u) \ge d_{min}, we can stop early and conclude D(u) \ge d_{min}. If D_{HC}(u) < d_{min}, we move to the second stage to compute D_{LC}(u) as$ 

$$D_{LC}(\boldsymbol{u}) = \frac{\sum_{\boldsymbol{v}_i \in \boldsymbol{u}} \sum_{\boldsymbol{v}_j \in \boldsymbol{v}_i, LC} Q_W(\boldsymbol{v}_i, \boldsymbol{v}_j)}{|\boldsymbol{u}|} \tag{12}$$

Finally we have  $D(u) = D_{HC}(u) + D_{LC}(u)$ . In addition, we can also derive an upper bound of D(u) to prune those cells with  $D(u) < d_{min}$ .

Lemma 4.3. Consider a cell  $u = \{v_1, ..., v_k\}$ . We compute the cell connectivity D(u) in two stages. In the first stage, we calculate  $D_{HC}(u)$  according to Eq. (11) by iterating over each vertex in u. During this stage, if  $D_{HC}(u) \ge d_{min}$ , we can stop and conclude  $D(u) \ge D_{HC}(u) \ge d_{min}$ . In the second stage, we calculate  $D_{LC}(u)$  according to Eq. (12) by iterating over each vertex in u.

In the first stage, assume we have iterated over vertices  $\{v_1, \ldots, v_l\}$ . Let  $w = \sum_{i=1}^l \sum_{v_j \in u, v_j \in v_i, HC} \tilde{Q}_{VV}(v_i, v_j)$ , then we can have an upper bound as  $\hat{D}_1(u) = \frac{w+k-\partial l}{|v||}$ .

In the second stage, assume we have iterated over vertices  $\{v_1, \dots, v_l\}$ . Let  $w' = \sum_{i=1}^k \sum_{v_j \in u, v_j \in v_i, HC} \widetilde{Q}_{VV}(v_i, v_j) + \sum_{i=1}^l \sum_{v_j \in u, v_j \in v_i, LC} \widetilde{Q}_{VV}(v_i, v_j)$ , then we can have an upper bound as  $\widehat{D}_2(u) = \frac{w' + (1-\delta)(k-l)}{|u|}$ .

If either  $\hat{D}_1(u) < d_{min}$  or  $\hat{D}_2(u) < d_{min}$ , then we can immediately stop and conclude  $D(u) < d_{min}$ .

**Proof 4.2.** The foundation to derive the upper bounds is that,  $\tilde{Q}_{W}$  is a row normalized matrix. That is,  $\forall v_i \in V$ , the following holds:

$$\sum_{\nu_j \in \nu_i, HC} \widetilde{Q}_{WV}(\nu_i, \nu_j) + \sum_{\nu_{j'} \in \nu_i, LC} \widetilde{Q}_{WV}(\nu_i, \nu_{j'}) = 1$$
(13)

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Table 2	Table	2	
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Vertices sorted in *HC* and *LC*,  $\delta = 0.75$ .

Vertex	HC set	LC set
$v_1 \\ v_2$	$\begin{array}{l}(\textit{v}_5,0.5),(\textit{v}_2,0.28)\\(\textit{v}_1,0.3),(\textit{v}_3,0.25),(\textit{v}_9,0.2)\end{array}$	$(v_3, 0.01), \dots, (v_n, 0.001)$ $(v_6, 0.03), \dots, (v_n, 0.003)$
$v_n$	$(v_{10}, 0.5), (v_8, 0.35)$	$(v_7, 0.01), \dots, (v_1, 0.002)$

545 According to the definitions of *HC* and *LC* sets, we have:

$$\sum_{v_j \in v_i, \text{HC}} \widetilde{Q}_W(v_i, v_j) \ge \delta$$

$$\sum_{v_j \in v_i, \text{LC}} \widetilde{Q}_W(v_i, v_j) \le 1 - \delta$$
(14)
(15)

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549 Based on these properties we can derive:

$$D(u) = \frac{\sum_{i=1}^{k} \sum_{j=1}^{k} \widetilde{Q}_{W}(v_{i}, v_{j})}{|u|} = \frac{\sum_{i=1}^{l} \sum_{v_{j} \in u, v_{j} \in v_{i}, HC} \widetilde{Q}_{W}(v_{i}, v_{j})}{|u|} + \frac{\sum_{i=1}^{l} \sum_{v_{j} \in u, v_{j} \in v_{i}, LC} \widetilde{Q}_{W}(v_{i}, v_{j})}{|u|} + \frac{\sum_{i=l+1}^{k} \sum_{j=1}^{k} \widetilde{Q}_{W}(v_{i}, v_{j})}{|u|} \\ \leq \frac{w + (1 - \delta)l + (k - l)}{|u|} = \frac{w + k - \delta l}{|u|} = \widehat{D}_{1}(u)$$

The inequality holds because  $\sum_{i=1}^{l} \sum_{v_j \in u, v_j \in v_i, LC} \widetilde{Q}_{VV}(v_i, v_j) \leq (1 - \delta)l$  according to Eq. (15), and  $\sum_{i=l+1}^{k} \widetilde{Q}_{VV}(v_i, v_j) \leq k - l$ according to Eq. (13). Thus we can derive the upper bound  $\widehat{D}_1(u)$ .

555 Similarly, to prove  $\widehat{D}_2(u)$  we have

$$D(u) = \frac{\sum_{i=1}^{k} \sum_{j=1}^{k} \widetilde{Q}_{W}(v_{i}, v_{j})}{|u|} = \frac{\sum_{i=1}^{l} \sum_{j=1}^{k} \widetilde{Q}_{W}(v_{i}, v_{j})}{|u|} + \frac{\sum_{i=l+1}^{k} \sum_{v_{j} \in u, v_{j} \in v_{i}, HC} \widetilde{Q}_{W}(v_{i}, v_{j})}{|u|} + \frac{\sum_{i=l+1}^{k} \sum_{v_{j} \in u, v_{j} \in v_{i}, LC} \widetilde{Q}_{W}(v_{i}, v_{j})}{|u|} \\ \leq \frac{w' + (1 - \delta)(k - l)}{|u|} = \widehat{D}_{2}(u)$$

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The inequality holds as  $\sum_{i=l+1}^{k} \sum_{v_j \in u, v_j \in v_i, LC} \tilde{Q}_W(v_i, v_j) \leq (1 - \delta)(k - l)$  according to Eq. (15). Thus we can derive the upper bound  $\hat{D}_2(u)$ .  $\Box$ 

561 Algorithm 3. Calculate-Connectivity

**Input:** A cell  $u = \{v_1, ..., v_k\}$  which contains k vertices, minimum cell connectivity  $d_{min}$ , a fixed threshold  $\delta$ . **Output:**  $D(u) \ge d_{min}$  or not.

1: initialize 
$$w = 0$$
  
/\* first stage \*/  
2: for  $l = 1 : k$  do  
3: for  $v_j \in u, v_j \in v_l.HC$  do  
4:  $w = w + \tilde{Q}_{VV}(v_l, v_j)$   
5: if  $w/|u| \ge d_{min}$  then  
6: return  $D(u) \ge d_{min}$   
7: if  $(w + k - \delta l)/|u| < d_{min}$  then  
8: return  $D(u) < d_{min}$   
/\* second stage \*/  
9: for  $l = 1 : k$  do  
10: for  $v_j \in u, v_j \in v_l.LC$  do  
11:  $w = w + \tilde{Q}_{VV}(v_l, v_j)$   
12: if  $w/|u| \ge d_{min}$  then  
13: return  $D(u) \ge d_{min}$   
14: if  $(w + (1 - \delta)(k - l))/|u| < d_{min}$  then  
15: return  $D(u) < d_{min}$ 

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We design an efficient algorithm Calculate-Connectivity in Algorithm 3 for calculating connectivity based on Lemma 4.3.

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#### 605 4.5. Parameters setting

SCMAG has four parameters  $e_{max}$ ,  $i_{min}$ ,  $v_{min}$  and  $d_{min}$ . For a *k*-dimensional subspace  $S = \{A_1, \ldots, A_k\} \subseteq A$ , the subspace entropy  $H(A_1, \ldots, A_k) \in [0, k \log_2 p]$  holds, if we assume  $|Dom(A_i)| = p$  for each attribute  $A_i \in S$ .  $k \log_2 p$  is reached when the *k* attributes are independent of each other. Thus the maximum entropy threshold  $e_{max}$  can be determined within the range  $[0, k \log_2 p]$ . In practice, for detecting good clusters in  $k(2 \leq k \leq 6)$ -dimensional subspace with p = 20, we find that  $e_{max} = 10$ always yields a good performance. Similarly, for a *k*-dimensional subspace  $S = \{A_1, \ldots, A_k\} \subseteq A$ , we can derive *interest* $(A_1, \ldots, A_k) \in [0, \log_2 p]$ . Thus the minimum interest threshold  $i_{min}$  can be determined within  $[0, \log_2 p]$ .  $i_{min} \leq 1$  is a good choice, which can avoid producing too few clusters.

The minimum cell coverage threshold  $v_{min}$  should not be set too large, since only gigantic clusters will be generated in that case.  $v_{min}$  depends on the graph size and user interests. In our experiments, for a graph with  $|V| \ge 10,000$ , we usually choose  $v_{min}$  no less than 100. For a cell  $u, D(u) \in [0, 1]$  holds. Thus  $d_{min}$  can be determined within [0, 1].  $d_{min}$  should not be set too low, and  $d_{min} \in [0.2, 0.6]$  is highly recommended to find densely connected clusters.

#### 617 4.6. Time complexity analysis

618 Our algorithm has two major components: random walk and subspace clustering. The time complexity of calculating random walk is  $O(L(|V| + |V_a|)^{\omega})$ , where  $\omega < 2.376$  is the fast matrix product exponent [7], and |V| is the number of structure 619 vertices and  $|V_a| = \sum_{i=1}^{d} n_i$  is the number of attribute vertices. Under an attribute subspace, the time complexity of calculating 620 subspace entropy is O(|V|d), where d is the attribute number, and the time complexity of calculating cell connectivity is 621  $O(|V|^2)$ . The total number of qualified subspaces in worst would be  $2^d$ , then the time complexity of subspace clustering is 622  $O((|V|d + |V|^2)2^d)$ . As a result, the total time complexity of our algorithm is  $O(L(|V| + |V_a|)^{2.376} + (|V|d + |V|^2)2^d)$ . In real appli-623 cation, the number of qualified interesting subspaces can be controlled by our parameters e.g. e<sub>max</sub> and d<sub>min</sub>, and would be far 624 less than the worst case. We would test the number of qualified subspaces in our experiments. 625

#### 626 5. Experimental study

In this section, we conducted extensive experiments to evaluate the performance of our algorithm on the IMDB movie dataset and a bibliographic dataset. Moreover, we presented some interesting case studies. Our experiments were run on a Dell PowerEdge R900 server with 2.67 GHz six-core CPU and 128 GB main memory running Windows Server 2008. All algorithms were implemented in C++, except that random walk was implemented in MATLAB.

#### 631 5.1. Comparison methods

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We evaluate the clustering quality and efficiency of our method. We compare with ENCLUS [5], which does not consider
 structural connectivity between instances in clustering. We develop two variations of ENCLUS, which only differ in handling
 multi-valued attributes as described below.

- Single-ENCLUS For instances having multi-values in an attribute, Single-ENCLUS randomly selects one of the values and ignores the rest.
- Multi-ENCLUS For instances having multi-values in an attribute, Multi-ENCLUS creates multiple single-valued instances
   for each of the multi-values.
- We also compare with a closely related method GAMer [12,13] which combines subspace clustering with dense subgraph 640 mining. Since GAMer outperforms CoPaM [19] in terms of cluster quality and efficiency, we do not compare with CoPaM in 641 642 the experiment. The GAMer program is publicly available at http://dme.rwth-aachen.de/en/gamer. We first transform multivalued attributed data to binary vectors. GAMer has several parameters and we follow the recommended parameter settings 643 in its program.  $n_{min}$  is the minimum cluster size (the same as our parameter  $v_{min}$ ), and  $\gamma_{min}$  is the minimum density of a clus-644 ter, which is similar as our parameter  $d_{min}$  but different in definition. A non-empty cluster C fulfills the density requirement if 645  $\frac{\min_{v \in C}\{\{(v,u)|(v,u) \in E, u \in C\}\}}{|C|-1} \ge \gamma_{min}$  is the minimum subspace dimension, which should be at least 2. Another parameter maxi-646 mal width w should be set to 0 to handle categorical attributes, i.e., only vertices with the same categorical values will fall 647 into the same cell. The ranges of parameters  $n_{min}$  and  $\gamma_{min}$  are listed in Table 3, and other parameters are set to the default 648 values in Table 4. We test different parameter value combinations in the experiments. 649

<b>Table 3</b> Parameter range	es of $n_{min}$ and $\gamma_{min}$ in GAMer.
Parameter	Range
n <sub>min</sub> γ <sub>min</sub>	10, 20, 50, 100, 200, 300, 400, 500, 600, 700 0.10, 0.20, 0.30, 0.40, 0.50, 0.60

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## Table 4 Default parameter settings in GAMer

F							
Parameter	s <sub>min</sub>	w	а	b	С	r <sub>obj</sub>	r <sub>dim</sub>
Default value	2	0	0	1	0	0.5	0.5

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We also compare our method with the full space attributed graph clustering SA-Cluster [6] in Section 5.7.

#### 651 *5.2. Evaluation measures*

We use density and entropy defined below to evaluate the quality of clusters. Density measures the intra-cluster edge weights normalized by cluster size, thus the larger the better. Entropy measures the attribute value distribution within a subspace cluster, thus the lower the better.

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$$density(\{C_i\}_{i=1}^{R}) = \frac{1}{R} \sum_{i=1}^{R} \frac{\sum_{u,v \in C_i} w(u,v)}{|\{v|v \in C_i\}|}$$
(16)  
$$entropy(\{C_i\}_{i=1}^{R}) = \frac{1}{R} \sum_{i=1}^{R} \frac{1}{|\mathcal{A}_i|} \sum_{A \in \mathcal{A}_i} entropy(A, C_i)$$
(17)

where  $entropy(A, C_i) = -\sum_{j=1}^{|Dom(A)|} p_{ij} \log_2 p_{ij}$ , and  $p_{ij} = \frac{\sum_{v \in C_i, a_j \in A(v)} \frac{1}{|V|v \in C_i|}}{|V|v \in C_i|}$ , i.e., the percentage of vertices in cluster  $C_i$  having value  $a_j$ on attribute A. In the above definitions, density only depends on edge connectivity in clusters, which is not relevant with attribute space. Entory is defined in subspace. For a cluster  $C_i$ ,  $A_i$  is the attribute subspace of  $C_i$ , where  $A_i \subset A$ .

#### 661 5.3. IMDB

#### 662 5.3.1. Dataset description

We use IMDB data<sup>1</sup> to build a multi-valued attributed graph, where a vertex represents a movie and an edge exists between two movies if they share one or more actors. The edge weight is the number of common actors. The multi-valued attributed graph contains 17,506 movies and 609,996 edges. A vertex is associated with 8 attributes which are described in Table 5. The numerical attributes *Actor Number* and *Voter Number* are discretized using equal-frequency binning, and *Average Rating* and *Variance of Rating* are discretized using equal-width binning. The release year of all movies is between 1990 and 2000. The other three attributes are categorical. A movie may have multi-values in *Genres*, e.g., the genres of movie "The Shawshank Redemption" are *crime* and *drama*.

Unless otherwise specified, we use the default parameter values shown in Table 6, and set the indexing threshold  $\delta = 0.8$ . We test three different  $d_{min}$  values of 0.30, 0.15, and 0.0.

#### 5.3.2. Cluster quality comparison

In the first experiment, we compare the cluster quality of our method SCMAG and ENCLUS based methods by varying the parameter  $e_{max}$  – maximum allowed subspace entropy.

Fig. 5(a) shows the entropy. The entropy by our method SCMAG with  $d_{min} = 0.30$  and 0.15 consistently outperforms the other methods. This shows when a connectivity requirement is enforced, high-quality clusters are discovered from subspaces which have not only a dense structure but also similar attribute values. This phenomenon can be interpreted as follows: if a set of movies share many actors, i.e., dense connectivity, they also have similar attribute values. In contrast, the entropy of Single-ENCLUS and Multi-ENCLUS is much higher. When  $d_{min} = 0.0$ , the entropy of SCMAG increases and is close to that of Single-ENCLUS. Moveover, a general trend we observe is that entropy decreases for all methods when the parameter  $e_{max}$  increases.

Fig. 5(b) shows the density. SCMAG with  $d_{min} = 0.30$  achieves the highest density and outperforms the other methods by a large margin. This is natural as we enforce a stricter connectivity requirement with a larger  $d_{min}$  value, the resultant clusters have a larger density. In addition, our method with  $d_{min} = 0.30$  and 0.15 shows a stable density when  $e_{max}$  increases. In contrast, the other three methods have a decreasingly small density when  $e_{max}$  increases.

In the second experiment, we compare the cluster quality of different methods by varying the parameter  $v_{min}$  – minimum cell coverage. As we can see from Fig. 6(a), SCMAG with  $d_{min} = 0.30$  has the smallest entropy around 0.5–0.7. It also shows a decreasing trend when  $v_{min}$  increases. The other baseline methods have a much larger entropy. In Fig. 6(b), SCMAG with  $d_{min} = 0.30$  achieves the largest density and outperforms the other methods by a large margin.

From these two experiments, we can conclude SCMAG with  $d_{min} = 0.30$  performs the best on both entropy and density. Multi-ENCLUS performs the worst on both criteria.

<sup>1</sup> http://www.imdb.com.

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#### Table 5 IMDB attributes

ID	Name	Туре	Domain
1	Actor number	Numerical	10
2	Voter number	Numerical	20
3	Average rating	Numerical	20
4	Variance of rating	Numerical	20
5	Year	Numerical	11
6	Genres	Categorical, multi-value	29
7	Language	Categorical, single-value	135
8	Sound-mix	Categorical, single-value	33

#### Table 6

Default parameter values: IMDB.

Parameter	e <sub>max</sub>	i <sub>min</sub>	$v_{min}$	d <sub>min</sub>
Default value	10	0.10	150	0.30





#### 692 5.3.3. GAMer results

<sup>693</sup> Under the default parameter settings in Table 4, we test GAMer on IMDB data by varying different value combinations of <sup>694</sup>  $n_{min}$  and  $\gamma_{min}$  in Table 3. However, GAMer fails to return any meaningful results. The outcomes of all the runs can be summar-<sup>695</sup> ized as two cases: "timeout" and "no cluster found". "timeout" means GAMer cannot finish in 5 h, and "no cluster found" <sup>696</sup> means GAMer finishes with no cluster found. Fig. 7(a) shows the results under different  $n_{min}$  and  $\gamma_{min}$  values, where white <sup>697</sup> cells represent "no cluster found" and gray cells represent "timeout".

We can observe, for example, when  $\gamma_{min} = 0.6$  and  $n_{min} \ge 200$ , no cluster can be found, as there is no cluster satisfying the  $\gamma_{min}$  and  $n_{min}$  requirements. When  $\gamma_{min} = 0.1$  and  $10 \le n_{min} \le 700$ , GAMer becomes timeout. This is because the search space is exponential in terms of the number of vertices in a graph, GAMer takes extremely long run time and a lot of memory, especially when the pruning is not effective.

702 5.3.4. Case study

We present some interesting subspace clusters found by SCMAG with  $d_{min} = 0.30$  on IMDB.

We first study a two-dimensional subspace {*Actor Number, Genres*}, under which we only found two clusters shown in Table 7. Cluster 1 contains 2380 drama movies with actor number in [25,4185], and cluster 2 contains 167 adult movies with actor number in [5,6]. These clusters quite make sense as drama movies usually have a big cast, while adult movies usually have a smaller cast.

We study another subspace {*Genres, Language*}, under which we found a total of 14 clusters. Table 8 shows the cluster list.
 We have the following interesting observations.

- Some typical clusters are from a combination of drama or comedy movies with one of the languages in English, German,
   Spanish, and French (clusters 1–4, 6–9).
- Interestingly, cluster No. 5 for Hindi drama movies is discovered with a high density value of 0.652, showing that these movies share many common actors. It is well known that India has the Bollywood just like America has the Hollywood, which has produced many popular movies.

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Fig. 7. GAMer results. White cell: no cluster found, gray cell: timeout, black cell: out of memory.

#### Table 7

Clusters in subspace {Actor Number, Genres} on IMDB graph.

Cluster	Actor number	Genres	Size	Density
1	[25, 4185]	Drama	2380	0.433
2	[5, 6]	Adult	167	0.316

#### Table 8

Clusters in subspace {Genres, Language} on IMDB graph.

Cluster	Genres	Language	Size	Density
1	Drama	English	4857	0.554
2	Drama	German	300	0.276
3	Drama	Spanish	387	0.523
4	Drama	French	459	0.409
5	Drama	Hindi	219	0.652
6	Comedy	English	2292	0.377
7	Comedy	German	298	0.303
8	Comedy	Spanish	160	0.416
9	Comedy	French	236	0.307
10	Comedy	Cantonese	189	0.497
11	Action	Cantonese	215	0.410
12	Music/documentary	English	970	0.281
13	Adult	English	525	0.940
14	Animation	English	272	0.396

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• Cantonese is a dialect spoken in southern China, Hong Kong, etc. Interestingly, we found a cluster of 189 Cantonese comedy movies (No. 10) with a density of 0.497. Another cluster of 215 Cantonese action movies (No. 11) corresponds to the world famous Chinese Kung Fu movies. These clusters reflect that the film industry of Hong Kong had brilliant achievements in 1990–2000, although Cantonese is only a local dialect.

Cluster No. 12 is formed by combining music and documentary English movies. According to our mechanism to compute the similarity between categorical values, we find "documentary" is close to "biography" and "music". Thus the cells of [Music, English] and [Documentary, English] can be merged. This cluster could not be found by Single-ENCLUS and

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Multi-ENCLUS. Five representative movies in this cluster include: *The Beatles Anthology (1995), Metallica: S&M (2000), The Wall: Live in Berlin (1990), The Filth and the Fury (2000), Metallica: Cunning Stunts (1998), all of which have genres of documentary and music.* 

• Interestingly, cluster No. 13 for adult English movies has a very high density value of 0.940, which reflects that most of these adult movies are starred by a common set of actors.

#### 5.4. Bibliographic data

#### 729 5.4.1. Dataset description

We use a bibliographic dataset collected from Arnetminer<sup>2</sup> to construct a multi-valued attributed graph, where a vertex 730 731 represents an author and an edge exists between two authors if they have co-authored one or more papers. The edge weight 732 is the number of co-authored papers by two authors. The multi-valued attributed graph contains 80,000 authors and 733 476,387 edges. A vertex is associated with 12 attributes which are described in Table 9. Topic is a categorical multi-valued attri-734 bute describing the research topics of authors, and an author can have multiple topics. We partition all venues in this bibliographic dataset into 45 groups according to area. Venue describes a list of venue groups an author has published in them. 735 736 For an author, we retain at most 3 values for attributes *Topic* and *Venue* respectively. The remaining attributes are numerical 737 and are discretized into 10 bins using equal-width binning. These attribute values are computed by Arnetminer algorithms.

<sup>738</sup> Unless otherwise specified, we use the default parameter values in Table 10, and set the indexing threshold  $\delta = 0.8$ . We <sup>739</sup> test three different  $d_{min}$  values of 0.40, 0.20, and 0.0.

#### 740 5.4.2. Cluster quality comparison

In the first experiment, we compare the cluster quality of different methods by varying  $e_{max}$ . Fig. 8(a) shows the entropy. The entropy by SCMAG with  $d_{min} = 0.40$  and 0.20 is very low when  $e_{max} = 7, 8, 9$ , but the entropy increases given a larger  $e_{max}$ . In contrast, the entropy of the other three methods shows a decreasing trend with  $e_{max}$ . Multi-ENCLUS has the largest entropy in all cases.

Fig. 8(b) shows the density. Again, our method SCMAG with  $d_{min} = 0.40$  achieves the highest density and outperforms the other methods by a large margin. This is due to the  $d_{min}$  constraint. In addition, SCMAG with  $d_{min} = 0.40$  and 0.20 has a stable density despite the increase of  $e_{max}$ , while the other three methods show a decreasing trend with the increase of  $e_{max}$ . Single-ENCLUS has the lowest density in all cases. These results demonstrate that our method can find dense clusters in subspaces with low entropy.

In the second experiment, we compare the cluster quality of different methods by varying the parameter  $v_{min}$ . Fig. 9(a) shows the entropy. We can observe that all methods have a lower entropy when  $v_{min}$  increases. This shows subspace clusters with a larger size have more similar attribute values in the bibliographic data. Our method SCMAG with  $d_{min} = 0.20$  and 0.0 and Single-ENCLUS score closely and achieve the lowest entropy. Multi-ENCLUS has the largest entropy in all cases. Fig. 9(b) shows the density. All methods have a stable density when  $v_{min}$  increases. SCMAG with  $d_{min} = 0.40$  achieves the largest density and outperforms the other methods by a large margin.

From these experiments, we can conclude SCMAG with  $d_{min} = 0.20$  achieves a good balance between entropy and density. Multi-ENCLUS performs poorly on both criteria.

#### 758 5.4.3. GAMer results

<sup>759</sup> Under the default parameter settings in Table 4, we also test GAMer on the bibliographic data by varying different value <sup>760</sup> combinations of  $n_{min}$  and  $\gamma_{min}$  in Table 3. Again GAMer fails to return any meaningful results. The outcomes of all the runs can <sup>761</sup> be summarized as three cases: "no cluster found", "timeout" and "out of memory" (GAMer throws a <sup>762</sup> *java.lang.OutOfMemoryError* exception). Fig. 7(b) shows the results under different  $n_{min}$  and  $\gamma_{min}$  values, where dark cells <sup>763</sup> represent "out of memory", and white and gray cells are similarly defined as in Fig. 7(a).

We can observe, for example, when  $n_{min} = 10$  and  $\gamma_{min} = 0.5$  or 0.6, "out of memory" happens. When  $\gamma_{min} = 0.6$  and  $n_{min} \ge 100$ , no cluster can be found, because there is no cluster satisfying the requirements. When  $\gamma_{min} = 0.1$  and  $10 \le n_{min} \le 300$ , GAMer becomes timeout; while for  $400 \le n_{min} \le 700$ , no cluster can be found. In summary, GAMer cannot find any meaningful clusters on the bibliographic data within a reasonable time.

#### 768 5.4.4. Case study

769 We present some interesting subspace clusters found by SCMAG with  $d_{min} = 0.40$  on the bibliographic data.

We first examine a two-dimensional subspace {*Topic, Venue*} under which we found 31 clusters. It is interesting to discover the combination of *Topic* and *Venue*, as research topics are closely correlated with published venues. This shows the subspace interest criterion is useful in finding correlated subspaces. We pick four clusters as examples and present their topics, representative venues, top 10 most cited authors in Table 11. In particular, cluster 3 combines several similar topics, e.g., *communication complexity, approximation algorithms, lower bounds, perfect graphs*, etc., related to algorithms and theory in the *Topic* dimension. Cluster 3 and cluster 4 could not be found by Single-ENCLUS and Multi-ENCLUS.

<sup>&</sup>lt;sup>2</sup> http://www.arnetminer.org.

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#### Table 9

Bibliographic data attributes.	
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ID	Name	Туре	Domain
1	Торіс	Categorical, multi-value	200
2	Venue	Categorical, multi-value	45
3	Publication	Numerical	10
4	Citation	Numerical	10
5	H-index	Numerical	10
6	G-index	Numerical	10
7	Activity	Numerical	10
8	Uptrend	Numerical	10
9	Newstar score	Numerical	10
10	Longevity	Numerical	10
11	Diversity	Numerical	10
12	Sociability	Numerical	10

#### Table 10

Default parameter values: bibliographic.



Fig. 8. Cluster quality vs. *e*<sub>max</sub>: bibliographic.





We study another subspace {*Activity, Uptrend, Venue*} under which we only found 1 cluster. The cluster has the highest
 attribute value intervals [1.09, 32.27), [0.75, 23.36) in *Activity* and *Uptrend* respectively. The authors in this cluster are very
 active, showing a strong uptrend in publication in the area of algorithms and theory (refer to venue information in cluster 3,
 Table 11). We list 5 authors with the highest uptrend values: *Pierre Fraigniaud, Flavio Chierichetti, David Woodruff, Devavrat Shah, and Phillip B. Gibbons.*

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#### Table 11

Clusters in subspace {Topic, Venue} on bibliographic graph.

Cluster	Торіс	Venue	Authors (top-10 citations)
1. Data mining	Data mining	TKDE, KDD, SDM, ICDM, PAKDD, KAIS	Jiawei Han, Sudhir Kumar, Philip S. Yu, Vipin Kumar, George Karypis, Heikki Mannila, Mohammed J. Zaki, Eamonn J. Keogh, Bing Liu, Michael J. Pazzani
2. Database	XML data	SIGMOD, VLDB, ICDE, EDBT, CIKM, Data Eng. Bulletin	Rakesh Agrawal, Jennifer Widom, Christos Faloutsos, Ramakrishnan Srikant, Serge Abiteboul, Miron Livny, Tomasz Imielinski, Johannes Gehrke, Rajeev Rastogi, Dan Suciu
3. Algorithms, theory	Communication complexity, approximation algorithms, finite sets, new designs, convex polygons, planar arrangements, lower bounds, perfect graphs	TCS, TIT, DAM, JCSS, SICOMP, Algorithmica, STOC	Rajeev Motwani, Christos Papadimitriou, Robert E. Tarjan, Prabhakar Raghavan, David R. Karger, Richard M. Karp, Jon M. Kleinberg, Leslie Valiant, Oded Goldreich, Moni Naor
4. Computer vision, pattern recognition	Character recognition, object recognition, two-view motion estimation, face recognition, image analysis	icip, icpr, icme, pr, cvpr, Tpami, prl, icdar	Anil K. Jain, Takeo Kanade, Jitendra Malik, Alex Pentland, Andrew Zisserman, Thomas Huang, Andrew Blake, Cordelia Schmid, Guillermo Sapiro, Azriel Rosenfeld

#### Table 12

Clusters in subspace {Citation, H-index, G-index, Venue} on bibliographic graph.

Cluster 1 database	Cluster 2 software engineering & scientific computing	Cluster 3 hardware & architecture	Cluster 4 algorithms & theory
Rakesh Agrawal	C.A.R. Hoare	A.L. Sangiovanni-Vincentelli	Rajeev Motwani
Hector Garcia-Molina	Leslie Lamport	Sharad Malik	Robert E. Tarjan
Jeffrey D. Ullman	Thomas A. Henzinger	Sartaj K. Sahni	Christos Papadimitriou
Jennifer Widom	Rajeev Alur	Lothar Thiele	Prabhakar Raghavan
Christos Faloutsos	David Harel	Sudhakar M. Reddy	David R. Karger
Jim Gray	Joseph Halpern	Jason Cong	Richard M. Karp
David J. DeWitt	Amir Pnueli	Robert Brayton	Jon M. Kleinberg
Michael Stonebraker	Moshe Vardi	Miodrag Potkonjak	Leslie Valiant
Ramakrishnan Srikant	Edmund Clarke	Massoud Pedram	Oded Goldreich
Serge Abiteboul	Robin Milner	Janak H. Patel	Moni Naor

In the subspace {*Citation, H-index, G-index, Venue*}, SCMAG found four clusters of authors from different research fields. Again the subspace combination of *Citation, H-index* and *G-index* is interesting, as these three attributes are positively correlated – H-index and G-index are computed from citations. Authors in these four clusters all have the highest values in *Citation, H-index* and *G-index* as [696, 51258), [12, 102) and [6, 25) respectively. We list 10 representative authors in each cluster in Table 12.

#### 5.5. Subspace number, cluster number and size

<sup>787</sup> We report the number of attribute subspaces (with valid clusters) found by different algorithms by varying  $e_{max}$  in Fig. 10. <sup>788</sup> With the increase of  $e_{max}$ , Single-ENCLUS, Multi-ENCLUS and SCMAG with  $d_{min} = 0.0$  output an increasingly large number of <sup>789</sup> subspaces. But with a minimum cell connectivity constraint  $d_{min} > 0$ , SCMAG does not output many subspaces when  $e_{max}$ <sup>790</sup> increases, as those subspace clusters which do not meet the connectivity requirement are not output. This result shows <sup>791</sup> our method can better control the subspace cluster quality and output only those clusters with both dense connectivity <sup>792</sup> and high attribute similarity.

793 We plot the cluster size (x axis) versus cluster number (y axis) in Fig. 11. A point (x, y) in a curve means there are 794 y clusters whose size is above x. On both IMDB and bibliographic graphs we can see the curves of Single-ENCLUS, Multi-ENCLUS and SCMAG with  $d_{min} = 0.0$  largely overlap, as they do not have any structural connectivity require-795 796 ment. We observe the following phenomena: (1) the size of clusters found by SCMAG with  $d_{min} > 0$  is significantly 797 smaller than the size of clusters found by the baseline methods. For example, the baseline methods find 12 gigantic clusters which contain more than 8753 (50%) movies on IMDB graph, and find 110 gigantic clusters which contain 798 799 more than 40,000 (50%) authors on bibliographic graph. Such gigantic clusters do not correspond to densely con-800 nected communities and are of little significance. (2) The number of clusters found by SCMAG with  $d_{min} > 0$  is orders 801 of magnitude smaller than that found by the baseline methods. Many clusters with a sparse connectivity are pruned 802 by our method. This result confirms that the connectivity consideration is necessary for subspace clustering on 803 attributed graphs.

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Fig. 10. Number of subspaces found.



Fig. 11. Clustering size vs. number.



804 5.6. Clustering efficiency comparison

We compare the efficiency of different subspace clustering algorithms by varying the parameter  $e_{max}$ . Fig. 12 shows the running time on IMDB and bibliographic graphs. Since SCMAG has an extra cost of random walk computation, it is 12.5 times slower on average than Single-ENCLUS on IMDB and 2.4 times slower on average on bibliographic graph.

#### 808 5.7. Comparison with SA-cluster

In this experiment, we compare our method with SA-Cluster [6] on IMDB (Table 13) and bibliographic graph (Table 14). Since Inc-Cluster [37] generates the same clustering results as SA-Cluster, we do not test Inc-Cluster. We set  $d_{min} = 0.30$  on IMDB and  $d_{min} = 0.40$  on bibliographic graph. SA-Cluster considers the full attribute space for clustering. We set the

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Table	. 12
IdDI	e 13

SA-Cluster	vs.	SCMAG:	IMDB.

	SA-Cluster	SCMAG
Cluster number Max cluster size Avg cluster size (except max one) Entropy Density	39 16,685(95%) 22 1.741 0.205	39 4857(28%) 732 0.598 0.452

#### Table 14

SA-Cluster vs. SCMAG: bibliographic.



Fig. 13. Scalability evaluation vs. vertex number |V|.

parameter of cluster number *K* in SA-Cluster to be the same as the number of clusters found by SCMAG. From both tables we can see SA-Cluster always outputs a gigantic cluster containing more than 90% nodes, which does not make much sense. The remaining clusters are extremely small, i.e., the average size of clusters except the gigantic one is 22 on IMDB and 18 on bibliographic graph. On the other hand, the clusters found by SCMAG are not extremely huge or small, as each cluster size has to exceed the minimum coverage  $v_{min}$ . In terms of cluster quality, SCMAG significantly outperforms SA-Cluster on both entropy and density. These results confirm that subspace clustering on attributed graphs generates high-quality clusters, i.e., with high structural connectivity and high attribute similarity.

819 5.8. Synthetic data

In this experiments, we use synthetic graph datasets to conduct scalability test and quality evaluation. Each graph has the number of edges as |E| = 4|V|, where |V| is the vertex number. A vertex is associated with 15 attributes, of which 10 attributes are categorical and multi-valued. The domain of each attribute is 20.

#### 823 5.8.1. Scalability test

To evaluate the scalability of SA-Cluster, we generate a series of synthetic graphs described above by increasing the number of vertices |V| from 10,000 to 100,000. We set the parameters  $e_{max} = 10$ ,  $i_{min} = 0.01$ ,  $v_{min} = 50$  and  $d_{min} = 0.3$ . We report the running time result in Fig. 13. As we can see, SA-Cluster scales very well with the increasing vertex number. The running time on the graph with |V| = 100,000 by SA-Cluster is only 22 times slower than the cost on the graph with |V| = 10,000, due to the effective pruning techniques.

#### 829 5.8.2. Quality evaluation

To evaluate clustering quality by all methods, we generate a series of synthetic graphs with ground-truth clusters. Each graph contains 5 clusters of 10 nodes by default. Each cluster has a density of 0.4, and 5 relevant dimensions out of 15

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#### Table 15

F1-score of clusters produced by different methods on ground-truth synthetic graphs.

Cluster size	GAMer	Single-ENCLUS	Multi-ENCLUS	SCMAG
10	0.97	0.60	0.60	0.98
15	0.86	0.53	0.52	0.98
20	-	0.45	0.45	0.97
25	-	0.41	0.40	0.97
30	-	0.36	0.36	0.98

832 dimensions. We also add the noise nodes into graphs, which has the same total number of nodes in clusters. Noise nodes do not belong to any clusters. We evaluate the quality of clusters on different graphs by increasing the number of clusters 833 size.The clustering quality is measured by F1-score criterion [13]. The results are reported in Table 15. For all graphs, the 834 parameters of each method are consistly set up. GAMer uses parameters  $n_{min} = 8$ ,  $\gamma_{min} = 0.3$ , and other parameters are set 835 to the default values in Table 4. SCMAG uses parameters  $e_{max} = 10$ ,  $i_{min} = 0.01$ ,  $v_{min} = 8$  and  $d_{min} = 0.3$ . Single-ENCLUS and 836 Multi-ENCLUS both use the same parameters  $e_{max} = 10$ ,  $i_{min} = 0.01$  and  $v_{min} = 8$ . 837

As we can see, our method SCMAG achieved the highest F1-scores among all methods on all graphs. For the graphs with 838 cluster size of 10 and 15, GAMer obtained higher scores than Single-ENCLUS and Multi-ENCLUS, due to the consideration of 839 840 structural density in clusters by GAMer. For the size of clusters exceeding 20 nodes, the program of GAMer failed to produce results by throwing an exception "java.lang.OutOfMemoryError", which may be resulted from expensive search space. 841

#### 6. Conclusions 842

We studied subspace clustering on multi-valued attributed graph for community detection. Different from existing attrib-843 844 uted graph clustering algorithms which consider the full space, we find interesting subspaces with good clustering and discover clusters with dense connectivity, homogeneous attribute values and good coverage. We use random walk with restart 845 on an attribute augmented graph to measure the structural closeness and attribute similarity. An indexing scheme is 846 designed to efficiently calculate cell connectivity. Moreover, we propose a new cell merging strategy and a mechanism to 847 848 handle multi-valued attributes based on graph neighborhood information.

849 Experimental results show that SCMAG significantly outperforms existing methods ENCLUS, GAMer and SA-Cluster, Some interesting discovered clusters are presented in case study to show the effectiveness of our method. 850

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