Synergistic Partitioning in Multiple Large Scale Social Networks

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Abstract—Social networks have been part of people’s daily life and plenty of users have registered accounts in multiple social networks. Interconnection among multiple social networks adds a multiplier effect to social applications when fully used. With the sharp expansion of network size, traditional stand-alone algorithms can no longer support computing on large scale networks while alternatively, distributed and parallel computing become a solution to utilizing the data-intensive information hidden in multiple social networks. As such, synergistic partitioning, which takes the relationships among different networks into consideration and focuses on partitioning the same nodes of different networks into same partitions. With that, the partitions containing the same nodes can be assigned to the same server to improve the data locality and reduce communication overhead among servers, which are very important for distributed applications. To date, there have been limited studies on multiple large scale network partitioning due to three major challenges: 1) the need to consider relationships across multiple networks given the existence of intricate interactions, 2) the difficulty for stand-alone programs to utilize traditional partitioning methods, 3) the fact that to generate balanced partitions is NP-complete. In this paper, we propose a novel framework to partition multiple social networks synergistically. In particular, we apply a distributed multilevel k-way partitioning method to divide the first network into k partitions. Based on the given anchor nodes which exist in all the social networks and the partition results of the first network, using MapReduce, we then develop a modified distributed multilevel partitioning method to divide other networks. Extensive experiments on two real data sets demonstrate that our method can significantly outperform baseline independent-partitioning method in accuracy and scalability.

I. INTRODUCTION

With the rapid development of computer science and technology, social network service has been thriving for several years, and has been playing increasingly important role in people’s daily life. Social network allows individuals to create public profiles, to establish connections with a list of users with whom to share photo/video/music/blogging, and to send/receive messages across the connections within the system, etc [1, 2]. Plenty of users have signed up for several social networks focusing on different kinds of content, and are maintaining different relationships in different networks simultaneously.

In recent years, social network size drastically increases. A recent report from BI Intelligence indicates that, among the largest social networks, Facebook still has the largest user population at 1.16 billion monthly active users (MAUs), which is followed by YouTube with 1 billion, and China’s social media network, Qzone, takes the third position with 712 million MAUs [3].

Co-processing refers to the information integration among multiple social networks. For example, current friend recommendation function derives information from a single network environment. However, through co-processing, information from multiple networks can be integrated and the efficiency and accuracy of two related friends can be greatly enhanced. The ever-expanding user base creates more challenges for co-processing the large amount of data stored in the multiple social networks environment. In our previous work, we have developed a novel community detection method, MCD (Mutual Community Detector) [4] which uses the heterogeneous information in multiple networks, but this method is limited to small networks. How to generalize MCD to large scale heterogeneous networks has been a burning question for us.

Users maintaining accounts in multiple networks are defined as anchor users. Making full use of the interactions of anchor users in different social networks will create multiplier effect for social applications. However, it is troublesome for large sized networks (notably the number of links) to be stored in main memory. Distributed and parallel computing on distributed file system will be more efficient than stand-alone computing on local disk.

As of now, there have been intensive research on partitioning within single network environment [5]. But problems about partitioning within multiple relevant network environment, especially within multiple large scale relevant networks, has less been studied. In this paper, we take into account data locality while partitioning multiple social networks so that the same partition on different social networks can be assigned to the same server. This will decrease the communication overhead among servers.

However, synergistic partitioning across multiple large scale social networks is very difficult for the following chal-
In this section, we will briefly review some classical network partition algorithms.

Researchers have developed many sequential algorithms for network partition, which can be referred to local improvement method and global method. Local improvement method takes a partition of a network as input, and tries to decrease the cut size with some local search methods. To perform a local improvement method, initial partitions should be generated by some other methods beforehand, and the performance is strongly affected by the quality of initial partitions.

One of the ground-breaking network partition methods is Kernighan-Lin (KL) method [7], which aims to divide a network into 2 balanced partitions. In it, an initial balanced bisection should be given, and then iterations will be performed to try to find a sequence of node pair exchanges that can lead to a reduction in the cut size. Time expenditure of each iteration in KL method is \(O(|n|^3)\). KL method is so important that plenty of local improvement methods are variations of it. For example, Dutt optimized the KL method and made it possible to complete KL process in \(O(|E| \max \{\log |n|, d_{\text{max}}\})\) time, where \(d_{\text{max}}\) means the maximum of node degree [8]. Fiduccia-Mattheyses (FM) [9] is another KL inspired method. For FM method, it does not choose node pairs as in KL method but single nodes to improve edge-cut in each iteration which may result in unbalanced partitions, and each iteration can be finished in \(O(|E|)\) time.

Unlike the KL and FM methods, Simulated annealing (SA) method [10] do not use greedy search but is based on statistical mechanics. It is a general purpose local search method, which tries to randomly select a neighbor node to estimate if it can improve the result of solution. But related experiments show that SA is not suitable for networks that are sparse or has some local structure [11].

Global method takes the whole graph and number of partitions \(k\) as input, and generates a \(k\)-way partition. Most of these methods first bisect the graphs, and then apply bisection step recursively until \(k\) node subsets generated. Global methods can be divide into 2 categories according to whether the spatial location information of nodes are included in the networks, geometric methods and coordinate-free methods [5].

Recursive coordinate bisection (RCB) [12] is the prime example of geometric algorithms. The algorithmic thought of the RCB method is very simple, a coordinate axis should be chosen at the beginning, then it tries to find an orthogonal plane of the axis that can divide the node set \(V\) into 2 equal sized partitions. Inertial method [13] proposed by Farhat and Lesoinne elaborates the RCB method. In the inertial method, the axis with minimum angular momentum [14] of the node set is chosen instead of coordinate axis in the RCB method. Leland and Hendrickson then combined the KL method with inertial method together to improve the bisection generated by the inertial method.

In some networks, space location information of nodes is not embedded or can not get high-quality partitions even if given. In this situation, some algorithms only considering the combinational network structure have been proposed. The recursive graph bisection (RGB) method [15] begins by finding a pseudo peripheral node in the network, i.e., one of a pair of nodes that are approximately at the greatest network distance.
from each other in the network. Finally, the nodes are sorted with respect to these distances and then the RGB method divides the nodes into two equal sized sets.

Pothen, Simon and Liou developed an eigenvector based method - recursive spectral bisection (RSB) method [16]. The RSB method sorts its nodes with respect to eigenvector coordinates and then divides the sorted nodes into two partitions. Based on RSB, Barnard and Simon first proposed a multilevel network graph partition method - multilevel-RSB to speed up the RSB method. The multilevel method consists of three phases: coarsening phase, partitioning phase and uncoarsening phase, which we will dilate in Section IV. The multilevel idea in the multilevel-RSB method has a profound influence on subsequent network partition methods, and lots of multilevel algorithms have been proposed by researchers, such as multilevel-KL methods [17, 18], multilevel k-way partition [19], multilevel graclus method [20], et al.

The KL method and SA method have been proved to be P-complete [21]. However, several parallel approximations of KL have been developed based on multi-processors (not multi-servers). The earliest KL based parallel network method which also is the earliest parallel method for network bisection is proposed by Gilbert and Zmijewski [22]. Savage and Wloka developed a Mob heuristic method [21]. Özturan et al proposed a new network bisection method based on iterative local node exchange between processors. Based on local improvement and a parallel multilevel evolutionary algorithm, Sanders and Schulz developed a balanced network partition method - KaFFPaE [23]. In 2013, Rahimian et al proposed a distributed graph partition method - ja-be-ja based on local search and SA techniques [24]. Some other parallel network partition methods can be found in [5] and most of parallel methods are essentially parallelization of well-known sequential algorithms, which usually are developed on multi-processor but stand-alone servers.

III. PROBLEM FORMULATION

The problem we are trying to address in this paper is synergistic partitioning in multiple large scale social networks. In this section, we formulate the problem and describe the assumptions used during the work.

A. Terms and Definitions

Definition 1: (Social network) A social network can be mathematically described as \( G = (V,E) \), where \( V \) and \( E \) represent the node set and the edge set of \( G \), respectively. An integer \( n = |V| \) is used to record the size of \( V \) and \( m = |E| \) represents the number of edges. \( v(i) \) represents node \( i \) and corresponding degree is \( d(v(i)) \). \( e_{i,j} \in E \) means the link between \( v(i) \) and \( v(j) \).

Definition 2: (Anchor node and Non-anchor node) Users involved in multiple social networks simultaneously are called as anchor users, and named as anchor nodes in network set \( GS = \{G^1, G^2, \ldots, G^t\} \). All the anchor nodes constitute the anchor node set \( A \), and \( A = \cap V^t, 1 \leq i \leq t \). The left nodes are called non-anchor nodes, constituting the non-anchor node set - \( NA \). So, \( A \cap NA = \emptyset \) and \( A \cup NA = \cup V^t, 1 \leq i \leq t \).

Definition 3: (Input space) Input space \( InS \) means the network set and the anchor node set in this paper, so \( InS = (GS,A) \). From \( InS \), we are able to get all the structural information about the multiple social networks.

Definition 4: (Balanced network partition) For an undirected network \( G = (V,E) \), balanced (or uniform) network partition refers to the problem of partitioning \( V \) into equal sized components and the number of links across boundaries of different components is minimized. All the nodes and links within a component constitute a partition. If the number of components is given as \( k \), then the problem is called as balanced \( k \)-way network partitioning problem.

B. Problem Formulation

To take full advantage of users’ relationships in multiple social networks, cooperative analysis is required. However, in view of the facts that even a single social network is so large that the network data can not be stored in the main memory, but may need to placed onto local disk or remote distributed filesystem [25]. In the distributed data placement case, for the existence of anchor nodes connecting multiple social networks together closely, network partitioning methods have to take the relationships among the social networks into account.

Links in social networks can reflect people’s interactions in the real world to a great extent. So, we can assume that anchor users’ interpersonal relationships in different social networks should be similar. If we take these anchor nodes as samples, and assuming these samples are evenly distributed, it would be reasonable to believe that there are a lot of similar structures revolving around the samples hidden in the networks. Furthermore, if we expand the similar structures to the whole networks, we will be able to get potential similar partition pairs. Finally, we can align the partitions of all the networks and put similar partitions onto same servers. As a result, data locality will increase and communication volume between servers will reduce remarkably, which are very important in distributed applications. For example, in the ideal case of cooperative partitioning dividing two social networks into \( k \) partitions, applications only need searching local subnetworks and do not need searching the subnetworks stored in other servers, and the communication volume may decrease from \( O(m^2) \) to \( O(m) \). Fig.2 shows a simple diagram of result comparison between synergistic partitioning method and independent partitioning method.

Mathematically, synergistic partition problem can be described as: For an input space \( InS = (GS, A) \), try to find a partition set \( PS = \{P^1, P^2, \ldots, P^t\} \), where the amount of anchor nodes divided into same partitions in different networks - \( SumAN \) is maximized and the edge-cut size \( EC \) is minimized. Let \( pidx(G,v(i)) \) and \( P(G,j) \) be the partition ID of \( v(i) \) and partition \( j \) in network \( G \), respectively. \( w_{i,j} \) represents the weight of link \( e_{i,j} \). In the networks studied in this paper, \( w_{i,j} = 1 \) if \( e_{i,j} \in E \), and else \( w_{i,j} = 0 \).

\[
\text{count}(v(i)) = \begin{cases} 
1 & \text{if all } pid(G^j, v(i)) \text{ are the same, where } 1 \leq j \leq t. \\
0 & \text{others.}
\end{cases} 
\]

\[
SumAN = \sum_{v(i) \in A} \text{count}(v(i)) 
\]
B. Distributed Multilevel k-way partitioning

Algorithms guaranteed to find out near-optimal partitions in a single network have been studied for a long period, which have been discussed in Section II. But most of the methods are stand-alone, and performance is limited by the server’s capacity. Inspired by the multilevel k-way partitioning (MKP) method proposed by Karypis and Kumar [19, 26] and based on our previous work [25], we try to use MapReduce [27] to speedup the MKP method. As the same with other multilevel methods, MapReduce based MKP also includes three phases: coarsening, init partitioning and un-coarsening. Fig.3 gives a vivid description about MKP method.

Coarsening phase is a multilevel process and a sequence of smaller approximate networks $G_i = (V_i, E_i)$ are constructed from the original network $G_0 = (V, E)$, where $|V_i| < |V_{i-1}|$, $i = 1, 2, ..., s$. To construct coarser networks, node combination and edge collapsing should be performed. The task can be formally defined in terms of matchings [28]. A matching is a set of node pairs $M = \{i, j\}, i \neq j$ and $e_{i,j} \in E$, and in which each node can only appear for no more than once. For a network $G_i$ with a matching $M_i$, if $(j, k) \in M_i$ then $v(j)$ and $v(k)$ will form a new node $v(q) \in V_{i+1}$. The weight of $v(q)$ equals to the sum of weight $v(j)$ and $v(k)$, besides, all the links connected to $v(j)$ or $v(k)$ in $G_i$ will be connected to $v(q)$ in $G_{i+1}$. The total weight of nodes will remain unchanged during the coarsening phase but the total weight of edges and number of nodes will be reduced. Define $W(T)$ to be the sum of edge weight in $T$ and $N(T)$ to be the number of components in $T$. It will be that:

\[ W(E_{i+1}) = W(E_i) - W(M_i) \]
\[ N(V_{i+1}) = N(V_i) - N(M_i) \]

Analysis in [29] shows that for the same coarser network, smaller edge-weight corresponds to smaller edge-cut. With the help of MapReduce framework, we use a local search method to implement an edge-weight based matching (EWM) scheme to collect larger edge weight during the coarsening phase. For the convenience of MapReduce, we design a new network representation format: each line contains essential information about a node and all its neighbors (NN), such as node ID, node weight (VW), edge weight (w), et al. The whole network data are distributed in distributed filesystem, such as HDFS [30], and each data block only contains part of node set and corresponding connection information. Function $map()$ takes a data block as input and search locally to find node pairs to match according to the edge weight, $reduce()$ is in charge of node combination, renaming and sorting. With the new node IDs and matching, a simple MapReduce job will be able to update the edge information and write the coarser network back onto HDFS. The complexity of EWM is $O(|E|)$ in each iteration and pseudo code about EWM is shown as follow.

**Input:**
- A network, $G_h$;
- Maximum weight of a node, $maxVW = n/k$;

**Output:**
- A coarser network, $G_{h+1}$;

1. $map()$ function:
   2. for all node $i$ in current data block do

Fig. 2. Diagrammatic sketch of synergistic partitioning method and independent partitioning method on two social networks - network I (datum network) and network II (synergistic network). Anchor users and corresponding node IDs are shown on the right. (d) shows the result of synergistic partitioning method taking the partition result of anchor nodes in datum network into consideration. (c) is the result of an independent partitioning method. Compared the partition results of network II shown in (c) and (d) with the result of network I in (a), similarity of partitions between (a) and (d) is higher than that between (a) and (c).

\[ EC(G) = \sum_{v(i), v(j) \in P(G, u)} w_{i,j} \]  
Where $v(i) \in P(G, u)$, $v(j) \in P(G, w)$ and $u \neq w$.  

\[ EC = \sum_{i=1}^{t} EC(G^i) \]  

IV. PROPOSED FRAMEWORK

In this section, we describe the heuristic framework for synergistic partitioning among multiple large scale social networks, and we call the framework - SPMN.

A. The Basic Idea

Data processing in SPMN can be roughly divided into two stages: datum generation stage and network alignment stage.

When got the anchor node set $A$, the framework will apply a distributed multilevel $k$-way partitioning method onto the datum network to generate $k$ balanced partitions. During this process, the anchor nodes are ignored and all the nodes are given the same treatment. We call this process datum generation stage. When finished, partition result of anchor nodes will be generated, we store them in a set - Map(anidx, pidx), where anidx is anchor node ID and pidx represents the partition ID the anchor node belongs to.

After the datum generation stage, synergistic networks will be partitioned into $k$ partitions according to the Map(anidx, pidx) to make the synergistic networks to align to the datum network, and during this process $max\{SumAN\}$ and $min\{EC\}$ are the objectives. We call this process network alignment stage.
3: if match[i] == -1 then
4: maxIdx = -1;
5: sortByEdgeWeight(NN(i));
6: for all \( v_j \in NN(v_i) \) do
7: if (match[j]) == -1 & \( VW(i) + VW(j) < \)
8: maxVW then
9: maxIdx = j;
10: end if
11: match[i] = maxIdx;
12: match[maxIdx] = i;
13: end for
14: end if
15: end for
16: reduce() function:
17: new newNodeID[n + 1];
18: new newVW[n + 1];
19: set idx = 1;
20: for all \( i \in [1, n] \) do
21: if \( i < \) match[i] then
22: set newNodeID[match[i]] = idx;
23: set newNodeID[i] = idx;
24: set newVW[i] = newVW[match[i]] = VW(i) +
25: VW(match[i]);
26: idx + +;
27: end if
28: end for
29: end for

After several iterations, a coarsest undirected weighted
network \( G_s \), consisting of only hundreds of nodes will be
generated. For the network size of \( G_s \), stand-alone algorithms
with high computing complexity will be acceptable for init
partitioning. Meanwhile, the weights of nodes and edges of
closer networks are set to directly reflect the weights of
the finer network during the coarsening phase, so \( G_s \) contains
sufficient information to intelligently satisfy the balanced
partition and the minimum edge-cut requirements. Plenty of
traditional bisection methods are quite qualified for the task.
In this paper, we adopt the KL method with an \( O(|E|^2) \)
computing complexity to divide \( G_s \) into two partitions and
then take recursive invocations of KL method on the partitions
to generate balanced \( k \) partitions.

Un-coarsening phase is inverse processing of coarsening
phase. With the initial partitions and the matchings of the
coarsening phase, it is easy to run the un-coarsening process
on the MapReduce cluster.

C. Synergistic Partitioning Process

In this section we talk about the synergistic partitioning
process on the synergistic networks with the knowledge of
partition results of anchor nodes from datum network. The
synergistic partitioning is also a MKP process but quit different
from general MKP methods.

In the coarsening phase, anchor nodes are endowed with
higher priority than non-anchor nodes. When choosing nodes
to pair, we assume that anchor nodes and non-anchor nodes
have different tendencies. Let \( G_s \) be the datum network
and \( G^s = \{G^{s1}, G^{s2}, \ldots, G^{s(t-1)}\} \) be the synergistic network set.

For an anchor node \( v(i) \), it would prefer to combine with
another anchor node \( v(j) \) which has the same partition ID
in the datum network, i.e., \( pidx(G^{s}, v(i)) = pidx(G^{s}, v(j)) \)
where \( v(i) \in A, v(j) \in A \) and \( i \neq j \). Second, if there is
no appropriate anchor node, it would try to find a non-anchor
node to pair. When planning to find a non-anchor node to pair,
the anchor node, assuming to be \( v(i) \), would like to find a
correct direction, and it would prefer to match with the non-
anchor node \( v(j) \), which has lots of anchor nodes as neighbors
with the same \( pidx \) with \( v(i) \). When combined together,
the new node will be given the same \( pidx \) as the anchor node.
To improve the accuracy of synergistic partitioning among
multiple social networks, an anchor node will never try to
combine with another anchor node with different \( pidx \).

For a non-anchor node, it would prefer to make a pair
with an anchor node neighbor which belongs to the dominant
partition in the non-anchor node’s neighbors. Here, dominant
partition in a node’s neighbors means the number of anchor
nodes with this partition ID is largest. Next, a non-anchor node
would choose a general non-anchor node to pair. At last, a non-
anchor node would not like to combine with an anchor node
being part of the partitions which are in subordinate status.
After combined together, the new node will be given the same
\( pidx \) as the anchor node. To ensure the balance among
the partitions, about 1/3 of the nodes in the coarsest network are
unlabeled.

Fig. 3 shows a diagrammatic sketch of synergistic parti-
tioning process. Take \( G_0 \) for example, nodes \( v_1 \sim v_7 \)
and the corresponding links information are stored on the same
server and the other nodes are stored on another server. Nodes
in pairs \( \{v_1, v_2\} \) and \( \{v_{10}, v_{11}\} \) are all with the same \( pidx \),
so they should be tackled first. \( v_9 \) and \( v_{15} \) choose a correct
direction to make a pair. Then, \( v_3 \) can not pair with \( v_1 \), so it
chooses \( v_4 \) to combine. Finally, after searching locally, \( v_7 \) can
not find a local neighbor to pair, but has to make a pair with
its remote neighbor \( v_8 \). The situation of \( v_8 \) is the same as \( v_7 \).
As discussed in part A of Section IV, max\{SumAN\}, \(\min\{EC\}\) and trying to balance the size of partitions are the objectives in synergistic partitioning process. However, when put together, it is impossible to achieve them simultaneously. So, we try to make a compromise among them and develop a heuristic method to tackle the problems. First, according to the conclusion - smaller edge-weight corresponds to smaller edge-cut and the pairing tendencies, we propose a modified EWM (MEWM) method to find a matching in the coarsening phase, of which the edge-weight is as large as possible. At the end of the coarsening phase, there is no impurity in any node, meaning that each node contains no more than one type of anchor nodes. Besides, a "purity" vector attribute and a "pidx" attribute are added to each node to represent the percentage of each kind of anchor nodes swallowed up by it and the "pidx" of the new node, respectively. Then, during the init partitioning phase, we treat the anchor nodes as labeled nodes and use a modified label propagation algorithm to deal with the non-anchor nodes in the coarsest network. At the end of the init partitioning phase, we will be able to generate balanced \(k\) partitions and to maximize the number of same kind of anchor nodes being divided into same partitions. Finally, we project the coarsest network back to the original network, which is the same as traditional MKP process. The pseudo code of coarsening phase in synergistic partitioning process is as follow.

**Input:**
- A network, \(G_h\);
- A map consists of anchor nodes with pidxs, \(Map(anidx, pidx)\);
- Maximum weight for a node, \(maxVW = n/k\);

**Output:**
- A coarser network, \(G_h+1\);

```java
1: map() function:
2: for all \(v_i\) in current data block do
3:   if match[i] == -1 then
4:     set flag = false;
5:     sortByEdgeWeight(NN(v_i));
6:   if \(v_i\) \in Map(anidx, pidx) then
7:     for all \(v_j \in NN(v_i)\) & match[j] == -1 do
8:       if \(v_j\) \in Map(anidx, pidx)
9:         & Map.get(v_i) == Map.get(v_j)
10:        & VW(i) + VW(j) < maxVW then
11:          match[i] = j; match[j] = i;
12:          flag = true; break;
13:       end if
14:     end for
15:     if flag == false, no suitable anchor node then
16:       for all \(v_k \in NN(v_i)\) & match[j] == -1 &\n17:         & VW(v_i) + VW(v_j) < maxVW do
18:         indirectNeighbor[i] = NN(v_i);
19:       sortByEdgeWeight(indirectNeighbor);
20:       for all \(v_k \in indirectNeighbor\) do
21:         if \(v_k\) \in Map(anidx, pidx)
22:           & Map.get(v_i) == Map.get(v_k)
23:             & match[j] == -1 then
24:             match[i] = j; match[j] = i;
25:             flag = true; break;
26:           end if
27:         end for
28:       if flag == true then
29:         break;
30:     end if
31:   else
32:     sortByEdgeWeight(NN(v_i));
33:     for all \(v_j \in NN(v_i)\) & \(v_j\) \notin Map(anidx, pidx)
34:       & VW(i) + VW(j) < maxVW &
35:       & match[j] == -1 do
36:         match[i] = j; match[j] = i; break;
37:       end for
38:     end if
39:   end if
40: end for
41: reduce() function:
42: new newNodeID[n + 1];
43: new newVW[n + 1];
44: set idx = 1;
45: for all \(i\) in newNodeID[] do
46:   if \(i < match[i]\) then
47:     newNodeID[idx] = i;
48:     newVW[idx] = maxVW
49:     idx ++;
50: end if
51: end for
52: new purity[i] = purity[i]*VW(i) + purity[j]*VW(j);
53: newPidx[i] = max\{pidx[i], pidx[j]\};
54: end for
```

The label propagation algorithm (LPA) [31] is a graph-based semi-supervised learning algorithm. It can be used to predict the information of unlabeled nodes by a few of labeled nodes, and has many good natures such as simple, effective, speedy, favorable scalability and reliability. Inspired by the idea of local degree center based label propagation algorithm (LDC-LPA) [32], we develop a modified LPA method to deal with the weighted nodes and weighted links in the coarsest network. In each iteration in LPA, normalized probabilistic transfer matrix \(T_{n\times n}\) should be given, then we can update the probabilities matrix of each node \(Y_{n\times k}\) by \((y_{i,j}\) means the probability of \(v_i\) being labeled as \(j\)):

\[
Y \leftarrow TY
\]  

In consideration of weighted nodes and weighted links, we have to modify the transfer matrix \(T\). In the coarsest network, each node is a node set, but only labeled nodes are able to propagate their labels. Besides, each link contains plenty of links, but it maybe that parts of the links are from the labeled nodes. Take Fig.4 as example, \(v_c\) has four neighbors with different VWs and purities, and the links between \(v_c\) and its neighbors are with different EWs. We assume the percentage of links from the labeled nodes of a node, e.g. \(v_a\), to the unlabeled node, e.g. \(v_c\), equals to the purity of the node. Then, we can define \(T_{i,j}\) as:

\[
T_{i,j} = P(j \rightarrow i) = \frac{EW_{i,j} \times purity_i}{\sum_{v_i \in \text{neighbor}(j)} EW_{i,j} \times purity_i}
\]
To ensure the convergence of the modified LPA, we use the initialization method of LDC-LPA to initialize the probability vector for each labeled node and randomly assign probabilities to the unlabeled nodes. At the end of the iterations, a stable probability distribution matrix \( Y \) will be generated. We can sort \( Y \) by the first label and collect the top \( n/k \) nodes together to form the first partition. Then delete the collected nodes from \( Y \) and repeat the process for the other labels. Finally, we will get \( k \) approximately equal sized partitions.

\[ \text{V. EXPERIMENT AND ANALYSIS} \]

In this section, we conduct extensive experiments to evaluate the work proposed in this paper.

\[ \text{A. Experimental Environment and Data Set} \]

In this paper, all experiments are running on a Hadoop-1.1.1 cluster of Antivision Software Ltd., which consists of 20 PowerEdge R320 servers (Intel Xeon CPU E5-1410 @2.80GHz, memory 8GB) with 64-bit NeoKylin Linux OS connected by a Cisco 3750G-48TS-S switch.

Data sets used are parts of two real social networks: Foursquare and Twitter, which are crawled from May 7th to June 9th. In Foursquare, we crawl 1,064,011 users, among them about 413,182 have their accounts in Twitter as well. Numbers of followers and users of these 413,182 Twitter users follow in Twitter are 1,030,855,018 and 187,295,465 respectively. To compare with the baseline method, we sample a smaller sized subnetwork of Twitter, which has about the same size with Foursquare, to perform experiments. The details about data sets can be found in Tab.1.

The baseline method used to compare with the SPMN framework is an independent network partitioning method - Metis, which is the state-of-the-art method for single network partitioning by now. With Metis, we first partition all the networks independently. Then for each partition \( P_i \), \( 1 \leq i \leq k \) in \( G^d \), we search in each synergistic network \( G^s \in G^s, 1 \leq s \leq d \) for a smaller sized subnetwork of Twitter, which has about the same size with Foursquare, to perform experiments. The details about data sets can be found in Tab.1.

<table>
<thead>
<tr>
<th>TABLE I. DATA SETS DESCRIPTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>Foursquare</td>
</tr>
<tr>
<td>n</td>
</tr>
<tr>
<td>1,064,011</td>
</tr>
</tbody>
</table>

To evaluate the accuracy and the balance performance of the framework SPMN, we compute the NMI (Normalized Mutual Information) [33] of anchor nodes in the datum network and synergistic networks. To measure the edge-cut, we edge-cut ratio to represent the percentage of edges cut off by the partition boundaries.

\[ \text{B. Experiments and Results} \]

In this section, first, we simply test the performance of SPMN on datum network (Foursquare network) and the result is shown in Fig.5. From Fig.5(a), we can see that SPMN consumes more time than Metis when the partition number \( k \) is small. But when \( k \) increases, growth of running time slows down. When \( k \) climbs to 32, Metis becomes slower than SPMN. By and large, the increments in both methods are very small. The reason is as follows. The first phase consumes the most time in MKP methods. For SPMN, time consumptions in the first and last phases have nothing to do with \( k \) and would be constants. Besides, they are running on MapReduce, so time consumption should be less than Metis. However, the coarsening phase depends two jobs (node pairing and edge renaming) and several iterations, and job initialization in MapReduce would take up great part. The init partitioning phase takes recursive iterations to generate initial partitions and the number of iterations is linear to \( k \) which is about the same with Metis. For Metis, only the first phase doesn’t concern partition number. If network size increases steadily, proportion of job initialization will decrease and the total time consumption in SPMN will be less than Metis. From Fig.5(b), we can see that the edge-cut size in SPMN is larger than in Metis, that is because Metis adds a refine process during the uncoarsening phase to further optimize the partitions.

Second, we verify the validity and practicability of SPMN on synergistic network (Twitter network). To increase the number of data sets and guarantee the connectedness in all networks, we deliberately and randomly ignore parts of anchor nodes and construct another three data sets which are only different in the number of anchor nodes. D1 has 413,182 anchor nodes, D2 has 328,014 anchor nodes, D3 has 203,491 anchor nodes and D4 has 109,842 anchor nodes. To test the scalability performance of SPMN, we enlarge the synergistic network by
TABLE II. SYNERGISTIC NETWORKS FOR SCALABILITY TEST (×10^{6})

<table>
<thead>
<tr>
<th>n</th>
<th>D1</th>
<th>D3</th>
<th>D6</th>
<th>D7</th>
<th>D8</th>
<th>D9</th>
</tr>
</thead>
<tbody>
<tr>
<td>m</td>
<td>1.18</td>
<td>1.5</td>
<td>1.8</td>
<td>2.1</td>
<td>2.4</td>
<td>2.7</td>
</tr>
</tbody>
</table>

Fig. 6. Performance of SPMN on Synergistic Network. (a) Edge-cut test; (b) NMI test on the partitions of anchor nodes in datum network and synergistic network; (c) Running time test; (d) Scalability test on synergistic networks.

VI. CONCLUSION

The synergistic partitioning we propose, combined with distributed computing, has proven to be an effective approach toward cospooling interconnected information in multiple social networks environment. We have seen significant improvement in NMI and scalability when comparing our approach with the state-of-the-art independent partitioning method. Currently, synergistic partitioning still has the pitfall of enlarging the size of edge-cut. Adding a refinement process to the uncoarsening phase could be a potential solution to this problem. Application using synergistic partitioning is able to partition multiple social networks considering the relationship among them, which is not taken into account by traditional methods. In this way, the cospooling in large scale social networks can be greatly advanced. For example, the accuracy and performance of production recommendation will be improved.

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