FSM: A Fine-grained Splitting and Merging Framework for Dual-balanced Graph Partition

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ABSTRACT

As the initial and critical step for distributed graph processing tasks, partitioning a large graph into smaller subgraphs by minimizing the number of cutting vertices and edges, namely cut size or replication factor, plays a crucial role in reducing the elapsed time. However, many prior works have primarily focused on optimizing the cut size by considering only vertex balance or edge balance, leading to significant workload imbalance and consequently hindering the performance of downstream tasks. Therefore, in this paper, our focus is on addressing the dual-balanced graph partition problem, that is minimizing the cut size while simultaneously guaranteeing both vertex and edge balance. We propose a lightweight effective twophase framework, namely fine-grained splitting and merging (FSM), which decomposes the graph into more and smaller partitions and then merges them together. FSM offers the flexibility of integrating with various state-of-the-art single-balanced techniques. We develop two efficient algorithms Fast Merging and Precise Merging to enable trade-offs between computational efficiency and partitioning quality. Experimental results on large real-world graphs demonstrate that FSM achieves state-of-the-art cut size while maintaining dual balance. The runtime for downstream tasks PageRank, connected component, and diameter estimation, can be reduced by a large proportion, up to 9.43%, 11.35%, and 17.94%, respectively.

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1 INTRODUCTION

Graphs have been widely used across many fields ranging from social networks and recommendation systems to transportation networks. Numerous graph algorithms, such as PageRank [29], diameter estimation [18], and maximal clique enumeration [6, 8], have been proposed. However, the increasing scale of graphs poses challenges in terms of both time and memory when applying these

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Table 1: Vertex size imbalance of SOTA partitioners.

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Graph	\mathcal{B}_V	σ_V	\mathcal{B}_E	R	\mathcal{B}_V	σ_V	\mathcal{B}_E	\mathcal{R}	\mathcal{B}_V	σ_V	\mathcal{B}_E	R		
NE	1.99	41.23%	1.00	1.53	3.12	73.29%	8.26	1.02	2.31	40.81%	1.00	1.04		
HEP-100	1.94	42.61%	1.00	1.55	2.21	36.10%	1.00	1.06	1.90	30.57%	1.00	1.04		
METIS	1.77	39.78%	1.03	4.59	2.52	71.93%	1.03	1.09	1.85	44.65%	1.03	1.14		

graph algorithms. To tackle these challenges, several graph processing frameworks have been developed, including Pregel [25], GraphX [12], and PowerGraph [11]. To enable parallel graph computations, these frameworks demand a partitioning strategy that divides the huge graph into smaller partitions and allocates them across the machines within a cluster.

1.1 Motivation

Graph partitioning is the task of dividing the graph G into a family of groups of vertices and edges, leading to two kinds of partitioning paradigms, i.e., vertex partitioning and edge partitioning. Vertex partitioning (also known as edge cutting) delivers the vertices of G into pairwise disjoint subsets by cutting the edges. In contrast, edge partitioning (also known as vertex cutting) divides the edges of G into pairwise disjoint subsets by replicating (i.e., cutting) crossboundary vertices. The number of cutting edges (in vertex partitioning) and vertices (in edge partitioning) can be also called *cut* size. Existing studies [5, 15, 37, 39, 40] have demonstrated that the communication cost of downstream distributed tasks is positively correlated with the cut size of the partitions. Additionally, the deviation between partitioned sets of vertices (known as vertex balance) and the deviation between partitioned sets of edges (known as edge balance) are essential for achieving workload balance [5]. Thus, lots of efforts have been made to reduce the cut size while ensuring vertex balance [19, 30, 33, 36] or edge balance [26, 31, 38, 39]. However, even the state-of-the-art (SOTA) methods struggle to simultaneously optimize these three metrics. Table 1 presents the performance of three representative partitioners including NE [39], HEP ($\tau = 100$) [26], and METIS [19] over real-world graphs downloaded from WebGraph [2-4], where each graph is partitioned into p = 32 parts. Let \mathcal{B}_V (resp. \mathcal{B}_E) denote the vertex balance (resp. edge balance), i.e., the fraction of the largest vertex (resp. edge) size over the average vertex (resp. edge) partition size, σ_V denote the coefficient variation of vertex sizes across partitions, and R denote the replication factor (i.e., the cut size).

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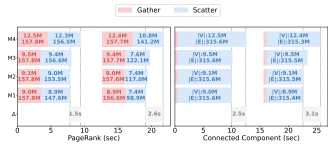


Figure 1: PageRank and Connected Component task on arabic graph using NE partitioner.

As shown in Table 1, we can observe that all three partitioners achieve promising replication factors and edge balances, but (I) exhibit very poor vertex balances, which suggests that they generate excessively large partitions; (II) the sizes of these partitions are highly skewed, indicating the existence of partitions with a significantly large number of vertices, as well as partitions with very few vertices. Focusing on only one single balance while minimizing the replication factor can lead to two troublesome problems as follows. (1) Terrible Workload Imbalance. In Bulk Synchronous Parallel (BSP) based systems, a global synchronization checkpoint is set between two iterations. The next iteration begins only when all parallel computations in the current iteration are completed. Therefore, the machine with the largest workload becomes the performance bottleneck. Figure 1 illustrates the corresponding elapsed time of gather and scatter operations on PowerGraph [11] for the top two iterations of the PageRank and Connected Component tasks on four machines (M1, M2, M3, and M4), along with the involved numbers of vertices and edges. While the number of involved edges across the four machines is roughly consistent in the first iteration, there exists a notable discrepancy in elapsed time among them. Discrepancies in the number of vertices can lead to differences in cache hit rates, as well as disparities in the numbers of involved edges in the subsequent iteration. For example, in the second iteration of PageRank, the discrepancies on the involved edges become significant in the scatter phase, thereby resulting in variations in machine elapsed time.

(2) Excessive Memory Consumption. In distributed graph computing frameworks deployed over homogeneous and memory-constrained clusters, each machine needs to maintain necessary information, such as neighbor sets and PageRank values for vertices. The excessively large partitions due to vertex imbalance will significantly increase memory consumption. As for PowerGraph, the partition with the largest number of vertices becomes the memory bottleneck. As depicted in Table 1, for the graph hollywood, the largest partition delivered by METIS has reached 8 times the average size. The machine hosting such a large partition is highly susceptible to encountering memory bottleneck, which can result in downstream task failures. As confirmed in the experiments (see Section 5.4), we find that SOTA partitioners like NE and HDRF [31] run out of memory on large graphs due to their vertex imbalance.

Therefore, it is crucial to minimize the replication factor while guaranteeing both vertex and edge balance. However, dual-balanced graph partitioning has not yet received sufficient attention and research despite its importance.

1.2 Challenges and Contributions

Challenge 1: Intractable Complexity. Previous study [39] has established the graph partitioning problem, minimizing the replication factor such that the edge balance is bounded by a constant, which is an NP-hard problem. The inherent complexity coupled with large graphs with billions of vertices and edges already poses significant challenges in terms of time and memory. Dual-balanced graph partitioning problem generalizes the single-balanced graph partition by introducing an additional constraint, making it more difficult to find an optimal solution that satisfies both balances simultaneously. Therefore, the partitioning algorithm itself is expected to be lightweight and efficient. These requirements force us to avoid using overly complex algorithms and redundant data structures.

Challenge 2: Skewed Degrees. Most real-world graphs follow the power-law distribution, revealing a skewed degree distribution that the majority of vertices have low degrees, while only a small subset of vertices are highly connected, known as hub vertices. Hub vertices play a crucial role in influencing the density of partitions, posing a significant challenge in achieving dual balance. For instance, the inclusion of hub vertices in a partition results in a substantial increase in the number of edges. This, in turn, is likely to result in poor edge balance.

Existing dual-balanced partitioners [1, 24, 40] have made efforts to address these challenges. EBV achieves dual balance by incorporating a scoring function that considers both vertex and edge loads. BPart modifies FENNEL [36] and requires simultaneous optimization of both vertex and edge balance. MDBGP [1] transforms the multi-dimensional balance partitioning problem into a mathematical optimization problem and solves it using gradient descent, but it constrains the number of partitions to powers of two. Beyond that, the optimization process of MDBGP involves multiple rounds of $O(n^2)$ intersection point calculations, which is not feasible for partitioning large graphs. To summarize, these partitioners (I) suffer from poor replication factors, resulting in a substantial communication cost that becomes a bottleneck again, and (II) do not support constraints on vertex-and-edge balance parameters or fail to achieve the desired constraints. There is a notable imbalance in both the vertex and edge dimensions for many graphs.

Contributions. To realize dual-balanced graph partitioning, in this paper, we propose the <u>F</u>ine-grained <u>S</u>plitting and <u>M</u>erging (FSM) framework. The underlying principle behind the FSM framework is to deal with vertex balance and edge balance incrementally, ensuring them one by one. The FSM framework consists of two phases, i.e., fine-grained splitting and subgraph merging.

Specifically, in the fine-grained splitting phase, FSM performs primary exploration of the graph by decomposing it into small-size subgraphs. The aim is to group the highly correlated vertices and edges together, producing a family of fundamental subgraphs. Moreover, in this phase, we can just concentrate on one balance at a time while minimizing replication factors. As a result, the framework becomes highly versatile and powerful, making it possible to integrate with various state-of-the-art single-balance techniques.

In the subgraph merging phase, we assemble the subgraphs produced above into larger partitions. Thus, we can consider the partitioning problem from a broader perspective, enabling us to

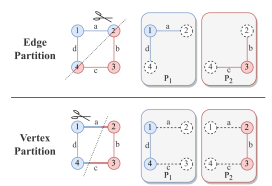


Figure 2: Edge partition and vertex partition.

neutralize the imbalances effectively and enhance the overall performance of the partitioning process. In particular, we formulate a subgraph allocation problem and develop two lightweight and effective merging algorithms, *Fast Merging* and *Precise Merging*.

In summary, our contributions can be summarized as follows:

- (1) We formulate the problem of dual-balanced graph partitioning that minimizes the replication factor while both vertex balance and edge balance are guaranteed.
- (2) We develop a lightweight effective two-phase framework, namely fine-grained splitting and merging (FSM), for dual-balanced graph partitioning, that chunks the graph into smaller subgraphs and then merges them to form larger subgraphs. The FSM framework can incorporate any existing graph partitioning algorithm as well as any merge algorithm.
- (3) To enable trade-offs between computational efficiency and partitioning quality, we propose two efficient algorithms *Fast Merging and Precise Merging*. We provide a theoretical optimality analysis of *Fast Merging* and provide the approximation ratio.
- (4) We conduct extensive distributed experiments on 11 large graphs against 11 competitors. Compared to the state-of-the-art graph partitioners, FSM exhibits notable efficiency improvements across three distributed graph processing tasks: PageRank, connected component, and approximate diameter. FSM achieves efficiency improvements up to 29.7%, 25.3%, and 57.02% for these tasks, respectively, compared to without using FSM.

2 PRELIMINARIES AND OVERVIEW

2.1 Dual-balanced Graph Partitioning

Let G=(V,E) denote an undirected graph, where V and E represent the sets of vertices and edges, respectively. For simplicity, the number of vertices and edges are denoted by n and m, respectively. A p-partitioning is to divide G into p partitions, where $p \geq 2$ and $p \in \mathbb{N}$. Generally, there are two different partitioning paradigms as follows.

DEFINITION 1. (p-Edge Partitioning, also called Vertex Cutting). An edge partitioner divides the edge set E into p pairwise disjoint subsets such that $\bigcup_{i=1}^{p} E_i = E$ and $E_i \cap E_j = \emptyset$ for $i \neq j$.

In edge partitioning, each edge is assigned exactly into one subset. We say a vertex u is a boundary vertex if it is contained in more than one subset, which indicates that u is replicated in these subsets.

EXAMPLE 1. As shown at the top of Figure 2, vertices 2 and 4 are boundary vertices. The adjacent edges a and b of vertex 2 are assigned into two different subsets, resulting in a copy of vertex 2.

DEFINITION 2. (p-Vertex Partitioning, also called Edge Cutting). A vertex partitioner divides the vertex set V into p pairwise disjoint subsets, i.e. $\bigcup_{i=1}^{p} V_i = V$ and $V_i \cap V_j = \emptyset$ for $i \neq j$.

In contrast to the edge partitioner, each vertex is assigned exactly into one subset. As shown at the bottom of Figure 2, the vertices $\{1,4\}$ and $\{2,3\}$ are assigned to two different subsets. Notice that, to guarantee the completeness of the partitions (i.e., without losing any edges), the cutting edges a and c have to be replicated in many applications, leading to vertex cuts as well. Moreover, Bourse et al. [5] have proved that vertex cuts are smaller than edge cuts on power-law graphs. Therefore, we focus on the widely-used edge partitioning in this paper.

DEFINITION 3. (Vertex Balance & α -Vertex Balanced Partitioning). The vertex balance of a p-partitioning, denoted by \mathcal{B}_V , is defined as $\mathcal{B}_V = \frac{\max_{i=1}^p |V_i|}{\sum_{i=1}^p |V_i|/p}. \ A \ p$ -partitioning is α -vertex balanced if $\mathcal{B}_V \leq \alpha$.

Definition 4. (Edge Balance & β -Edge Balanced Partitioning). The edge balance of a p-partitioning, denoted by \mathcal{B}_E , is defined as $\mathcal{B}_E = \frac{\max_{i=1}^p |E_i|}{\sum_{i=1}^p |E_i|/p}. \ A \ p$ -partitioning is β -edge balanced if $\mathcal{B}_E \leq \beta$.

In the task of edge partitioning, \mathcal{B}_E can be rewritten as $\mathcal{B}_E = \frac{\max_{j=1}^p |E_i|}{|E|/p}$ since each edge is just contained in one subset E_i .

Definition 5. (Replication Factor). The replication factor of a p-partitioning, denoted by \mathcal{R} , is defined as $\mathcal{R} = \frac{1}{|V|} \sum_{i=1}^{p} |V_i|$.

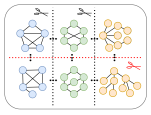
EXAMPLE 2. As shown at the top of Figure 2, both partitions contain two edges. Thus, we have the edge balance $\mathcal{B}_E=1.0$. Vertex 2 and vertex 4 are replicated once each, resulting in a replication factor of $\mathcal{R}=\frac{4+2}{4}=1.5$.

Problem Statement. (Dual-balanced Graph Partitioning). Given a graph G and three parameters p, α , and β , the task of dual-balanced graph partitioning, denoted by MIN- $\mathcal{R}(p,\alpha,\beta)$, is to return a p-edge partitioning of G with the minimum replication factor such that $\mathcal{B}_V \leq \alpha$ and $\mathcal{B}_E \leq \beta$.

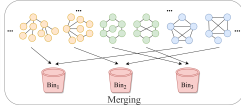
2.2 Overview of the Approach

To reach the objective of minimizing cut size while guaranteeing dual balance, we propose the Fine-grained Splitting and Merging (FSM) framework. FSM adopts an incremental algorithmic approach, by dividing the whole task into two phases as illustrated in Figure 3.

Fine-grained Splitting Phase. In the first *Fine-grained Split* phase, our objective is to obtain more and smaller partitions that have a low cut size and satisfy one single balance constraint at least. Any state-of-the-art edge partitioner or vertex partitioner that achieves a single balance can be used for fine-grained splitting. To maintain allocation information (the subgraph to which a vertex or an edge belongs) and cutting information (the replication status of a vertex or an edge), we develop a succinct data structure called *subgraph information* (denoted as gInfo), which can reduce storage costs and accelerate the subsequent merging process.







(a) Fine-grained Split Phase

(b) Subgraph Merging Phase

Figure 3: Overview of the FSM framework.

Subgraph Merging Phase. The core problem addressed in the *Subgraph Merging* phase is to merge the subgraphs obtained in the *Fine-grained Splitting* phase to achieve the final dual-balanced partition. Here, we draw inspiration from the Longest Processing Time (LPT [13]) algorithm. As shown in Figure 3(b), we can abstract the final partitions as bins and shuffle the fine-grained subgraphs. The problem then becomes assigning each subgraph to the appropriate bin. We adopt the *attempt merging* approach for each subgraph to determine its candidate bin. We further design the *attempt merging* process based on efficiency and quality requirements.

Novelty of the approach. For distributed downstream tasks, several systems have been developed, such as PowerLyra and TopoX. These systems adopt different partitioning strategies for high and low-degree vertices. In addition, there are many standalone partitioning algorithms, including state-of-the-art single-balanced algorithms like NE, HEP, and 2PS, as well as dual-balanced algorithms like BPart, MDBGP, and EBV. Single-balanced algorithms only optimize a single dimension, while existing dual-balanced algorithms and systems demonstrate suboptimal performance in terms of replication factor. FSM addresses these shortcomings by applying an effective two-phase method. The initial phase produces a set of smaller subgraphs with a minimized cut size, optimizing at least one single balance. The second phase merges these subgraphs to enable the dual-balanced guarantee. The two-phase framework is flexible to accommodate powerful splitting or merging strategies. Thus, FSM allows an excellent replication factor while implementing adaptive methods to achieve bounds on two dimensions.

3 FINE-GRAINED SPLITTING

Instead of directly partitioning the graph G into p partitions, we decompose G into more and smaller p' = k * p subgraphs. It is like assembling building blocks - with many small building blocks, we can always create toys of similar sizes. Although the sizes of these small building blocks may vary, we can flexibly combine them to avoid creating toys that are too big or too small.

Let us recap the problem and its challenges. We need to find a graph partition that satisfies a certain level of dual-balance in terms of vertices and edges while minimizing the replication factor \mathcal{R} . Solving this problem is inherently difficult. Most existing graph partitioners can achieve good replication factors under a single balance constraint, and we can consider them as small building blocks. These building blocks may have different sizes initially, but the abundance of blocks and their small individual sizes provide us with flexibility and room for optimization in subsequent steps.

The existing partitioners can achieve desirable cut sizes under a single balance, but only fulfilling two out of the three objectives. However, we can leverage the small subgraphs (as building blocks) obtained from these partitioners to flexibly assemble them in order to achieve dual balance. Specifically, if edge balance is prioritized, we then utilize small subgraphs with both edge balance and low $\mathcal R$ as building blocks. This logic applies equally to prioritizing vertex balance as well, thus providing small subgraphs with vertex balance and low $\mathcal R$ as building blocks.

3.1 Subgraph Information

To maintain the vertex and edge information of the k*p partitions, we develop a succinct data structure called *subgraph information* (denoted as gInfo), which can reduce storage costs and accelerate the subsequent merging process.

Specifically, gInfo consists of p' dense bitsets (shorted as bitset) named rep. When prioritizing edge balance, the rep bitset is used to track whether a vertex belongs to a partition. There is also an array named pvec, with a length of m for edge partitioners and n for vertex partitioners, which is used to record the initial partition assignment of each vertex or edge.

The rep (replication) is typically implemented with a bitset (dense binary string). If the v-th position of rep[b] is set to 1, it indicates that vertex v has a replica in P_b . Using a bitset has several advantages. (1) Memory efficiency: Compared to hash-based sets, it can save memory especially when p' is not large (2) Accelerating merging: Using a dense bitset can reduce the constant factor during set merging operations such as bitset OR and bitset popcount (to calculate the number of 1s).

3.2 Prioritizing Edge Balance

To obtain the edge-balance building blocks, we can use SOTA edge partitioners such as NE and HEP. (1) NE [39] is a state-of-the-art edge partitioner known for achieving edge balance and small \mathcal{R} , by dividing the partitions into core and boundary vertex sets. It iteratively expands the partitions and prioritizes moving vertices from the boundary set to the core set, particularly those with fewer external connections. Then, it assigns the corresponding edges to the current partition. (2) HEP [26] is a hybrid partitioner that combines NE and HDRF [31]. It partitions high-degree vertices with HDRF, alleviating the skewness of vertices across partitions. This approach also reduces the memory overhead of the adjacency list.

The detailed algorithm procedure is outlined in Algorithm 1. In lines 3-7, we iterate through each edge e(u, v). We retrieve the

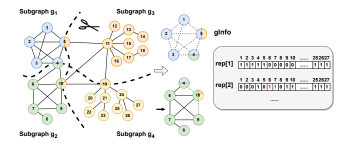


Figure 4: An example of performing splitting.

subgraph ID b to which e belongs through pvec. We set the corresponding bits for vertices u and v in the bitset rep[b] to 1 to indicate their presence in g_b .

EXAMPLE 3. As illustrated in Figure 4, we begin by obtaining the partition information for each edge through the edge partitioner and storing it in the pvec array. When processing the edge e(6,7), we first retrieve its assigned subgraph ID, i.e., 2, from pvec. Then, we set the bits at positions 6 and 7 in the bitset rep[2] to 1.

3.3 Overhead Analysis

The time and memory overheads of the Fine-grained Splitting phase mainly depend on the partitioner involved. We denote the time and memory overheads of partitioning p subgraphs using the chosen partitioner $T_1(p)$ and $M_1(p)$, respectively. When the splitting factor k is selected, the time and memory overhead of the splitting stage are $T_k(k * p)$ and $M_k(k * p)$, respectively. For each of the k * ppartitions, we need to maintain a bitset. When applied to edge partitioners, we track the replication status of vertices, so the length of the bitset is n. The memory overhead of the bitset part is proportional to O(k * p * n/C) and O(m) for pvec. C represents the word width which is determined by the underlying hardware and compiler implementation. Similarly, for vertex partitioners, the overall memory overhead is O(k * p * m/C + n). It should be noted that Algorithm 1 is usually synchronized during partitioning because many partitioners [26, 31, 39, 40] need to maintain vertex or edge allocation to facilitate partitioning. Thus, the time and memory overheads of these recording steps have already been included in $T_k(k * p)$ and $M_k(k * p)$.

Algorithm 1: Splitting by Edge Partitioner

Input : target partition count p, splitting factor k and graph G

Output: subgraph information gInfo (rep and pvec)

- $p' \leftarrow p * k$
- $_2$ pvec ← EdgePart(G, p')
- g foreach g(u,v) ∈ E do
- $b \leftarrow \text{pvec [e]}$

- return glnfo

4 SUBGRAPH MERGING

In the *Merging* phase of the FSM framework, the focus shifts from the cutting size and single-balance to the optimization of the other balance metric 'incrementally'. Through the *fine-grained splitting* phase, we obtain p' = k * p small subgraphs (denoted as g). In this phase, the task is to merge these subgraphs into p target partitions $\{P_i\}_{i=1}^p$ while achieving a balanced distribution along the other dimension as much as possible. Taking edge partitioners as an example, where the edge balance of the small subgraphs has already been ensured, our goal is to merge exactly k of $\{g_i\}_{i=1}^{p'}$ into one P_j . Thus, we can obtain p partitions with approximately equal numbers of edges. Next, we need to determine which k subgraphs to choose for each bin.

4.1 Merging Outline

4.1.1 Problem Formulation. We have a set of p' subgraphs $\{g_i\}_{i=1}^{p'}$ along with their information set $\{gInfo_i\}_{i=1}^{p'}$, where each gInfo has a bitset rep. We want to allocate this subgraph set to p bins to form the final partition $\{P_i\}_{i=1}^{p}$. To maintain the single balance achieved from the Fine-grained Splitting phase, the load of each bin is set to k, meaning each bin can accommodate exactly k subgraphs.

Let $\lambda(bin_i)$ denote the popcount of rep in the *i*-th bin, which is resulted from merging k subgraphs. The objective of subgraph allocation is to distribute $\{g\}$ among the bins in a way that ensures approximately equal popcounts across the bins, while minimizing the maximum popcount among the bins. Thus, we consider the following minimization problem:

minimize
$$\lambda(bin_M) = \max\{\lambda(bin_1), \lambda(bin_2), ..., \lambda(bin_n)\}$$

4.1.2 Algorithm Outline. The merging algorithm can be divided into three main steps.

(1) Subgraph selection. We sort the p' subgraphs in non-ascending order based on their λ . Now subgraph set $\{g\}$ is sorted, so we have $\lambda(g_1) \geq \lambda(g_2) \geq ...\lambda(g_{p'})$. In each iteration, we allocate the first subgraph in the current subgraph set, that is, the subgraph with the largest λ at the current stage (denoted as g_c).

(2) Bin selection. In the second step, we select a candidate bin (bin_{cand}) as the destination for the subgraph g_c .

We maintain a set of currently open bins, called openbin, which represents bins that still have available capacity for additional subgraphs. Initially, openbin includes all p empty bins and gradually shrinks as subgraphs are assigned to bins during the merge process.

Within openbin, we use the *attempt merging* approach to select the candidate bin. This involves iterative merging the pending subgraph with each bin in openbin and calculating λ of the resulting bin after the merge. The bin with the smallest λ is chosen as the candidate bin.

(3) **Merging.** During the merge processing, subgraph g_c is merged into bin_{cand} , leading to a new subgraph $bin_{cand} \cup g_c$. The bin_{cand} is then updated to $bin_{cand} \cup g_c$. We then remove g_c from the subgraph set $\{g\}$ and set the newid of g_c to the id of its candidate bin. If bin_{cand} reaches the maximum capacity of k subgraphs, it is closed and removed from openbin. We continue this allocation process for each subgraph until all subgraphs are allocated to a bin. Finally, we retrieve the final partitions through pvec and newid.

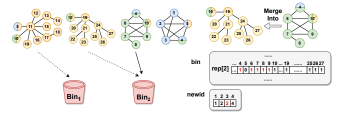


Figure 5: An example of performing merging.

```
Algorithm 2: Fast Merge

Input : subgraph information glnfo, target partition number p and splitting factor k

Output: partition result

1 Initialize min heap Q with p empty nodes
```

```
2 Sort(q, rep, non increasing)
```

```
3 foreach b \in [p'] do
4 | candidate ← Q.Pop
5 | candidate ← MergeBin(candidate, g_b)
6 | newid [b] = candidate.id
7 | if candidate.size < k then
8 | Q.Push(candidate)
9 | else
10 | result ← result \bigcup candidate
```

11 return result

Example 4. As shown in Figure 5, we sort the four subgraphs in non-ascending order based on their λ , resulting in the sorted subgraph set with $\lambda=11,10,6,5$. We simplify the attempt merging step and directly add λ to determine the size of the new bin. When considering subgraph g_3 , its candidate bin is bin_2 because $\lambda(bin_2)$ is currently the smallest. Then we merge bin_2 and g_3 by performing a bitwise OR operation between their corresponding rep. Finally, we update the entry in newid for g_3 to indicate its final partition is 2.

4.2 Merging Algorithm

Based on whether overlap is computed during the bin selection step, we propose two algorithms: Fast Merge and Precise Merge. By offering these two merging options, we provide flexibility to accommodate different requirements and trade-offs between computational efficiency and partitioning quality.

Fast Merge. The Fast Merge algorithm prioritizes speed by simply removing the *attempt merging*, as we assume that there is no overlap in rep between the g_c and all the bins. As a result, the *attempt merging* operation, which previously involved bitwise OR operations, can now be simplified to a direct summation of the λ s. In this case, we can simplify the bin selection step by selecting the bin with the smallest λ within openbin. To further improve speed, we utilize a min-heap to maintain the bin with the smallest λ within openbin. The Pop operation on the heap (Algorithm 2, lines 4-5) directly produces the candidate bin for merging. If the number of small subgraphs in the newly merged bin does not reach the threshold of k, the bin is placed back into the heap. Otherwise, the merged partition is directly considered as one of the final partitions (Algorithm 2, lines 7-10).

Algorithm 3: Precise Merge

12 return result

```
Input : subgraph information glnfo, target partition count
            p and splitting factor k
  Output: partition result
1 Initialize openbins with p empty bins
2 Sort(g, rep, non increasing)
b foreach b ∈ [p'] do
      foreach bin \in openbins do
          newbin \leftarrow MergeBin(bin, g_b)
5
          Check and update candidate
6
      candidate \leftarrow MergeBin(candidate, q_b)
7
      newid [b] \leftarrow \text{candidate.id}
      if candidate.size = k then
          openbins ← openbins \ candidate
10
11
          result ← result ( ) candidate
```

Precise Merge. The *Precise Merge* algorithm is a further refinement of the greedy approach, as it carefully evaluates the overlap when conducting the *attempt merging* step. Specifically, during the allocation of subgraph g_c , it is merged with each bin, and λ of the resulting merged bin is calculated (Algorithm 3, lines 4-6). This approach takes into account the potential reduction in λ due to the overlap between g_c and the bins. Once a bin reaches the threshold of k subgraphs, it will no longer be considered in the subsequent merge process and will be included as one of the final partition results (Algorithm 3, lines 9-11).

Overhead Analysis. Both *Fast Merge* and *Precise Merge* require maintaining p bins. Similar to gInfo, each bin consists of a bitset (O(L/C)). Moreover, we need an array named newid of length k*p to record the final partition assignment of each subgraph g. L represents the number of bits in the bitset. If prioritizing edge balance, it is equal to |V|, otherwise, it is equal to |E|. The bins in *Fast Merge* are maintained using a min heap, and the space complexity is O(k*p). During the entire *Subgraph Merging* process, we first sort the k*p subgraphs, resulting in time complexity of $O(k*p*\log(k*p))$. Then each of the k*p subgraphs needs to merge with its candidate bin once, resulting in a time complexity of O(k*p*L/C) for the entire operation.

Precise Merge performs attempt merging on each of the k * p subgraphs. Each attempt merging for each subgraph requires a bitset OR operation with all p buckets, resulting in a time complexity of O(p*L/C). Hence, the overall time complexity for Precise Merge is $O(k*p*\log(k*p)+k*p*(1+p)*L/C)$. Fast Merge efficiently selects candidate bins using a min heap, which can be completed in $O(\log p)$ for each subgraph. Hence, the overall time complexity for Fast Merge is $O(k*p*\log(k*p)+k*p*(\log p+L/C))$.

Discussion. As analyzed above, *Fast Merge* runs faster as it has a smaller time complexity. Moreover, it boasts a theoretical guarantee on its approximation rate (see Section 4.3). By utilizing the potential reduction in λ due to the overlap between subgraphs and bins, the merging performance of *Precise Merge* and the downstream tasks based on *Precise Merge* are better than *Fast Merge* in most instances,

while with a marginal increase in the merging time. Thus, *Precise Merge* generally emerges as the preferable option, particularly when confronted with intricate bucket merging scenarios, especially as the parameter *k* increases. Nevertheless, we notice that there are cases where *Fast Merge* outperforms *Precise Merge*. The theoretical analysis of *Fast Merge* also provides some hints to *Precise Merge*.

4.3 Theoretical Analysis

We analyze the upper bound of the *Merge* algorithm, that is, the approximation rate compared to the optimal case. To simplify the problem, we do not consider the overlap of bitsets during merging, i.e., we discuss the *Fast Merging* algorithm (which is comparable to *Precise Merging* in terms of merging quality) where we use the λ summation instead of bitset OR. We define $\eta_i = \frac{\lambda(g_i)}{\lambda(g_{p'})}$. Since $\{g\}$ is sorted non-increasingly, we have $\eta = \{\eta_1, \eta_2, ..., \eta_{p'}\}, \ \eta_i \geq 1$ for $\forall i \in [p']$. This optimization objective is also known as the k-partitioning problem (NP-complete when k = 3 [10]).

THEOREM 1. When k=2, i.e., when p'=2*p subgraphs are merged into p bins, Fast Merge can find the optimal solution.

PROOF. According to the *Fast Merge* algorithm, we place each subgraph g into the current smallest bin. In the case of k=2, the process is simplified to sequentially placing p subgraphs into p bins, followed by placing the remaining p subgraphs in reverse order. Now, consider the scenario where the maximum bin size obtained from the *Fast Merge* algorithm is denoted as ω_0 . We will argue by contradiction that there is no other combination that can yield a smaller maximum bin size ω' than ω_0 .

Assume that $\omega' = \eta_m + \eta_n$, where m and n are two positive integers representing the indexes of two bins in the new combination. Consider three possible cases for the values of m and n:

Case 1: m, n > p. In this case, g_m and g_n belong to the smallest p subgraphs and are placed in the same bin. The remaining (p-1) bins need to accommodate the largest p subgraphs. According to the pigeonhole principle, there must be at least two of the largest p subgraphs placed in the same bin. However, this would result in a bin that is larger than ω' , which contradicts the assumption. Therefore, this case is not possible.

Case 2: $m, n \leq p$. Let $\omega_0 = \eta_a + \eta_{2p+1-a}$. Now, we consider the new combination where g_a and g_{i_1} are in the same bin. In this case, i_1 satisfies $i_1 > 2p + 1 - a$. If $i_1 \leq 2p + 1 - a$, then $\eta_{i_1} \geq \eta_{2p+1-a}$. However, $\eta_m + \eta_n \geq \eta_a + \eta_{i_1} \geq \eta_a + \eta_{2p+1-a} = \omega_0$, which contradicts the assumption $\eta_m + \eta_n = \omega' < \omega_0$. Similarly, for $\eta_{a-1} + \eta_{i_2}$, i_2 also needs to satisfy $i_2 > 2p + 1 - a$.

In summary, there are a subgraphs $g_a,g_{a-1},...,g_1$. But there are only a-1 subgraphs g_i corresponding to them, because these g_i need to satisfy $i \in (2p+1-a,2p]$. This contradicts the assumption and is not possible.

Case 3: $m \le p < n \text{ and } m + n \ne 2p + 1.$

Let $\omega_0=\eta_a+\eta_{2p+1-a}$. Now, we consider the new combination where g_a and g_{j_1} are in the same bin. We can conclude that j_1 satisfies $j_1>2p+1-a$. If $j_1\leq 2p+1-a$, then $\eta_{j_1}\geq \eta_{2p+1-a}$. However, $\eta_m+\eta_n\geq \eta_a+\eta_{j_1}\geq \eta_a+\eta_{2p+1-a}\geq \omega_0$, which contradicts the assumption $\eta_m+\eta_n<\omega_0$. Furthermore, for the subgraph g_{j_2} combined with g_{a-1},η_{j_2} must satisfy $j_2>j_1>2p+1-a$, which implies that $\eta_{j_2}\leq \eta_{j_1}\leq \eta_{2p+1-a}$.

If $j_2 < j_1 < 2p+1-a$, according to $\eta_{a-1} \ge \eta_a$ and $\eta_{j_2} \ge \eta_{j_1} \ge \eta_{2p+1-a}$, then we have $\eta_m + \eta_n \ge \eta_{a-1} + \eta_{j_2} \ge \eta_a + \eta_{j_1} \ge \omega_0$, which is a contradiction. Based on this reasoning, we can deduce that $2p \ge j_a > ... > j_2 > j_1 > 2p+1-a$. Hence, it is not possible to satisfy the conditions. In conclusion, there is no such new combination that allows $\omega' < \omega_0$.

Approximation Ratio. Let ω_0 denote the optimal size of the largest bins. The proposed *Fast Merging* algorithm achieves the following approximation ratio:

$$\frac{\omega'}{\omega_0} < 1 + \frac{k-1}{max(p-1+k,p'-(p-1)\eta_1)}$$

Theorem 2. Suppose subgraph g_l is the last subgraph placed in the largest bin, with the size of η_l . Let us assume that there are $l(0 \le l \le p'-1)$ subgraphs in total whose vertex sets have sizes greater than or equal to η_l , while the remaining p'-l subgraphs have vertex sets with sizes less than η_l . It holds that $\frac{\omega'}{\omega_0} < (1 + \frac{p-1}{l})$.

PROOF. First, we define $\varphi_j = \eta_m - \eta_j$, which represents the difference between the total size of the merged subgraph in bin j and the total size of the merged subgraph in the largest bin. We have:

$$\omega_0 = \frac{1}{p} \left(\sum_{i=1}^{p'} \eta_i + \sum_{j=1}^{p} \varphi_j \right) \ge \frac{1}{p} \sum_{i=1}^{p'} \eta_i \tag{1}$$

(The equality holds only when $\sum_{j=1}^p \varphi_j = 0$, i.e., when the vertex sizes are equal in all bins.) From (1) and $\eta_i \geq 1$, we have:

$$\omega_{0} \geq \frac{1}{p} \sum_{i=1}^{p'} \eta_{i} = \frac{1}{p} \left(\sum_{\eta_{i} \geq \eta_{l}} \eta_{i} + \sum_{\eta_{i} < \eta_{l}} \eta_{i} \right) \geq \frac{1}{p} \left(l * \eta_{l} + (p' - l) \right).$$

$$\eta_{l} \leq \frac{p * \omega - (p' - l)}{l} = \frac{p}{l} * \omega_{0} - \left(\frac{p'}{l} - 1 \right) \tag{2}$$

The equality holds only when there exist l subgraphs with sizes of η_l and p'-l subgraphs with sizes of 1. When allocating g_l to a target bin, there are three cases: (1) No bin has reached k subgraphs and closed. (2) Some bins have reached k subgraphs and closed, but the target bin is still the bin with the smallest total size among all bins. (3) Some bins have reached k subgraphs and closed, but the target bin is not the bin with the smallest total size among all bins.

Now we consider the first two cases because the sizes of the subgraphs to be merged are not significantly different. In these two cases, for $\forall j$, we have:

$$\varphi_j \le \eta_l$$
(3)

Equation (3) holds only when g_l is the last subgraph to be allocated, and before its allocation, the vertex sizes of all subgraphs are evenly distributed, i.e., l=p'-1. Therefore, we can conclude that Equation (1) and Equation (3) cannot hold simultaneously. From Equation (1), Equation (2), and Equation (3), we have:

$$\omega' = \frac{1}{p} \left(\sum_{i=1}^{p} \eta_i + \sum_{j=1}^{p} \varphi_j \right) \le \frac{1}{p} \left(\sum_{i=1}^{p'} \eta_i + (p-1)\eta_l + 0 \right)$$

$$< \left(1 + \frac{p-1}{l} \right) \omega_0 - \frac{p-1}{p} \left(\frac{p'}{l} - 1 \right)$$

$$< \left(1 + \frac{p-1}{l} \right) \omega_0.$$

THEOREM 3. For the index l of the subgraph g_l placed last into the largest bin, there is a lower bound: $l \ge p - 1 + k$.

PROOF. Now let us assume that $l . Let <math>bin_M$ be the largest bin, and let $|bin_M|$ denote the number of subgraphs in bin_M . Since initially all p bins are empty and our greedy strategy prioritizes filling smaller bins first, after distributing the first p subgraphs, each bin contains exactly one subgraph. We have:

$$|bin_M| \le (l) - p + 1 < (p - 1 + k) - p + 1 < k.$$

At this point, bin_M is already the largest bin. Therefore, the subgraphs with indices l+1 and beyond will be allocated to the remaining bins to balance their sizes with bin_M , and bin_M will not receive any more subgraphs. In other words, the final number of subgraphs in bin_M is less than k, which contradicts the definition of the k-partitioning problem.

THEOREM 4. For the index l of the subgraph placed last into the largest bin, there is another lower bound: $l \ge p' - (p-1)\eta_1$.

PROOF. Considering the subgraph set sorted in non-increasing order of their sizes, all subgraphs allocated after g_l are used to fill the size gaps between bins. Therefore, we can derive that:

$$\sum_{\eta_i < \eta_l} \eta_i \le (p-1) * \eta_l$$

Additionally, since $\sum_{\eta_i < \eta_l} \eta_i \ge p' - l$, we have: $p' - l \le (p-1)\eta_1$. \square

Combining Theorems 2, 3, 4, we conclude that:

$$\frac{\omega'}{\omega_0} < 1 + \frac{p-1}{max(p-1+k,p'-(p-1)*\eta_1)}$$

4.4 Adaptive Selection of Splitting Factor

In this section, we discuss how to address the constraints α and β for vertex balance and edge balance, respectively.

Edge balance guarantee. For a specified parameter β , the FSM method initially sets the upper limit of edge capacity for small subgraphs to β times the average number of edges. It then uses a single-balanced partitioner to generate smaller subgraphs.

Vertex balance guarantee. If the vertex balance across the subgraphs generated by the initial fine-grained splitting does not satisfy the demand, it is difficult to achieve the balance effect by using the merge method. FSM achieves the vertex balance by adjusting the splitting factor k. Notice that increasing k improves vertex balance but may also lead to a higher replication factor \mathcal{R} . Therefore, for the given parameter α representing vertex balance, we can find the smallest k such that $\mathcal{B}_V \leq \alpha$. This objective can be accomplished through a binary search over the splitting factor. Specifically, if the partitioning result for the current k adheres to the requirement $\mathcal{B}_V \leq \alpha$, the value of k is decreased. Conversely, if the vertex balance requirement is not satisfied, the value of k is increased.

5 EXPERIMENTAL STUDY

5.1 Experimental Setting

Dataset. Nine real-world graphs from SNAP [22], WebGraph [2–4], and network repository [32] are used in the experiments. Table 2 lists their statistics.

Table 2: Statistics of Graphs: |V|, |E|, \overline{d} , size, and type denote the number of vertices and edges, the average degree, file size in binary edge-list form, and graph type, respectively.

Name	Graph	V	E	\overline{d}	Size	Туре
tw	twitter-2010	42 M	1.5 B	70.5	11 G	Soc. Net.
id	indochina-2004	7.4 M	192 M	40.7	1.5 G	Mas. Net.
it	it-2004	41 M	1.1 B	55.0	8.5 G	Mas. Net.
wk	wikipedia	26 M	599 M	46.2	4.5 G	Mas. Net.
uk	uk-2005	39 M	921 M	46.7	6.9 G	Mas. Net.
<i>u</i> 7	uk-2007-05	105 M	3.7 B	70.7	28 G	Mas. Net.
uu	uk-union	132 M	5.5 B	83.2	41 G	Mas. Net.
sk	sk-2005	51 M	1.9 B	76.2	15 G	Mas. Net.
wb	webbase-2001	116 M	993 M	17.2	7.4 G	Mas. Net.
hw	hollywood-2011	2.0 M	229 M	230.7	1.8 G	Misc. Net.
ar	arabic-2005	23 M	631 M	55.5	4.8 G	Misc. Net.

Methods. We compare the proposed method FSM with 11 partitioners, including streaming, hybrid, and in-memory algorithms. Degree information is taken as input for streaming algorithms.

- DBH [38]: DBH combines random partitioning with degree information by assigning each edge to the hash partition corresponding to the adjacent vertex with the lower degree.
- MDBGP [1]: MDBGP transforms the problem of partitioning multi-dimensional balanced graphs into an optimization problem and solves it using gradient descent.
- BPart [24]: BPart relaxes the vertex balance constraint of FENNEL and then optimizes the dual balance based on this relaxation.
- PowerLyra [7]: PowerLyra proposes a Hybrid algorithm that combines random hashing to allocate edges of low-degree and high-degree vertices using different strategies.
- Hybrid-BL (TopoX) [23]: On the basis of Hybrid algorithm, Topox incorporates fusion for low-degree vertices (multi-hop neighbors) and fission for high-degree vertexs, introducing the Hybrid-BL algorithm.
- HDRF [31]: HDRF allocates each edge based on a scoring function that takes into account both vertex degrees and the distribution of vertices and edges in partitions.
- EBV [40]: EBV also utilizes a similar scoring function to allocate edges, but it focuses more on dual balance.
- CLUGP [21]: CLUGP is a restreaming edge partitioning algorithm.
 It is pipelined into three steps: streaming clustering, cluster partitioning, and partition transformation.
- 2PS [27]: 2PS algorithm consists of two stages: streaming clustering stage and re-streaming stage. 2PS-HDRF is selected as the competitor as it exhibits better partitioning performance.
- NE [39]: NE applies a neighbor heuristic that prioritizes expanding vertices with fewer external connections to generate partitions incrementally.
- HEP-1, HEP-10, HEP-100 [26]: HEP partitions high-degree edges by using HDRF and partitions low-degree edges by using NE.
- FSM-NE: FSM framework utilizes NE as the *Fine-grained Split* partitioner and precise merge, shorted as FSM-N.
- FSM-HEP: FSM framework utilizes HEP-100 as the Fine-grained Split partitioner and precise merge, shorted as FSM-H.

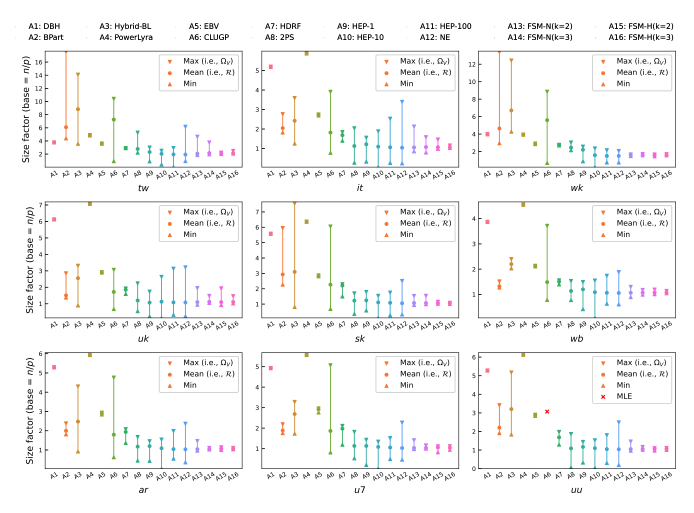


Figure 6: Maximum, mean, and minimum vertex size by 11 partitioners with p = 32.

Evaluation Metric. As existing dual-balanced partitioners do not support constraints on α and β or fail to achieve the desired constraints, we follow their settings and evaluate the performance following three metrics: the replication factor \mathcal{R} , vertex size coefficient variation σ_V , and max vertex size factor Ω_V ($\Omega_V = \frac{\max_{l=1}^P |V_l|}{n/p}$). We do not report edge balance \mathcal{B}_E since all partitioners can ensure $\mathcal{B}_E \leq 1.05$. The vertex balance \mathcal{B}_V is not reported as it can be represented as Ω_V/\mathcal{R} . We also investigate the effect of different partitions over downstream tasks, such as PageRank [29], connected component computation, and approximate diameter [18], by reporting the elapsed time.

We set up a distributed cluster with 8 machines, each having Intel(R) Xeon(R) W-2135 CPU @ 3.70GHz and 64 GB of RAM. They are set up with PowerGraph, running at full thread capacity, and connected via Gigabit Ethernet.

Reproducibility: The datasets used in the paper are available at: http://snap.stanford.edu/, https://law.di.unimi.it/ and https://networkrepository.com/. We employ the implementation offered by [1, 7, 19, 21, 26, 27, 39] and re-implement EBV, FENNEL, and

Hybrid-BL using C++. Furthermore, we develop standalone versions of BPart and TopoX using C++. The source codes of FSM are available at: https://github.com/lcj2021/split-merge-partitioner/.

5.2 Overall Partitioning Performance

Figure 6 illustrates the size factor of each partition that is generated by 11 partitioners in detail, where the *size factor* of a partition is defined as the ratio of its vertex size to the average vertex size. Here, we present the data for 9 graphs, excluding id and hw. For complete details, please refer to our technical report. We evaluate the performance of these partitioners based on the following three metrics. The number of partitions p is set to 32 by default.

5.2.1 Max Vertex Size Factor. The max vertex size factor, i.e., Ω_V , can directly reflect the **maximum memory overhead and computation workload** incurred by all machines in the cluster when performing distributed tasks. A good edge partitioner should minimize Ω_V to ensure that large-scale distributed tasks can be executed smoothly. The large $\mathcal R$ of streaming partitioners leads to their Ω_V being inevitably large, as $\Omega_V \geq \mathcal R$. As shown in Figure 6, NE, HEP, and 2PS perform similarly, with a low $\mathcal R$ but a high Ω_V . They

Table 3: Partitioning results on 11 graphs with p = 32, where the lowest is highlighted in bold and '-' indicates partition failure due to memory or time limits.

		DBH	MDBGP	BPart	Hybrid-BL	PowerLyra	EBV	CLUGP	HDRF	2PS	NE	HEP-1	HEP-10	HEP-100	FSM-N(k=2)	FSM-N(k=3)	FSM-H(k=2)	FSM-H(k=3)
	Ω_{V}	5.30		2.38	4.30	5.95	2.94	4.76	2.08	1.68	2.36	1.46	1.54	1.98	1.47	1.10	1.14	1.11
ar	R	5.29	-	2.00	2.48	5.94	2.90	1.80	1.94	1.18	1.03	1.20	1.09	1.04	1.04	1.05	1.07	1.07
hw	Ω_{V}	14.79		15.69	13.08	17.87	4.66	11.91	3.71	7.32	2.93	3.50	3.14	3.01	1.98	1.95	1.97	1.96
IIW	R	14.74		5.67	9.14	17.85	4.64	9.01	3.55	2.86	1.55	2.56	2.02	1.55	1.64	1.68	1.62	1.67
id	Ω_{V}	4.99		3.96	3.71	5.62	2.52	3.03	2.16	1.91	3.02	1.79	1.87	2.34	1.50	1.39	1.38	1.11
Iu	R	4.98		1.85	2.55	5.61	2.49	1.47	1.60	1.10	1.02	1.13	1.08	1.06	1.03	1.04	1.05	1.05
it	Ω_{V}	5.20		2.76	3.59	5.90	2.75	3.91	1.84	2.04	3.39	1.55	1.88	2.53	2.13	1.58	1.46	1.16
п	R	5.19		2.05	2.42	5.88	2.73	1.82	1.69	1.13	1.04	1.22	1.10	1.06	1.06	1.08	1.07	1.08
sk	Ω_V	5.58		5.95	7.55	6.37	2.87	6.05	2.29	1.70	2.50	1.79	1.54	1.75	1.53	1.54	1.15	1.09
SK	R	5.58		2.93	3.10	6.36	2.83	2.26	2.21	1.23	1.06	1.25	1.12	1.08	1.07	1.07	1.06	1.06
tw	Ω_{V}	3.78		17.57	14.08	4.89	3.61	10.41	2.91	5.26	6.16	3.00	2.50	2.92	4.64	3.77	2.22	2.47
tw	R	3.78		6.09	8.83	4.87	3.58	7.25	2.90	2.78	1.93	2.31	2.04	1.95	2.06	2.15	2.08	2.14
u7	Ω_{V}	4.93		2.19	3.28	5.58	2.96	5.07	2.11	1.82	2.27	1.43	1.34	1.51	1.41	1.16	1.11	1.12
u/	R	4.93		1.89	2.69	5.58	2.92	1.86	1.98	1.13	1.03	1.13	1.08	1.06	1.03	1.04	1.06	1.05
uk	Ω_V	6.13		2.85	3.31	7.09	2.92	3.06	1.93	2.24	3.21	1.74	2.63	3.13	1.95	1.48	1.94	1.46
uĸ	R	6.12		1.51	2.55	7.07	2.90	1.72	1.85	1.20	1.08	1.07	1.13	1.08	1.10	1.12	1.10	1.11
uu	Ω_V	5.28		3.41	5.17	6.14	2.90		1.98	1.86	2.49	1.44	1.53	1.80	1.46	1.17	1.10	1.11
uu	R	5.28		2.21	3.21	6.13	2.87	- 1	1.68	1.09	1.04	1.17	1.10	1.05	1.05	1.06	1.06	1.06
wb	Ω_{V}	3.87	_	1.52	2.40	4.56	2.13	3.71	1.56	1.53	1.88	1.49	1.55	1.74	1.30	1.18	1.19	1.14
WD	R	3.87		1.32	2.20	4.55	2.12	1.49	1.51	1.14	1.06	1.20	1.09	1.07	1.07	1.07	1.07	1.08
wk	Ω_V	4.00		13.40	12.43	3.95	2.92	8.85	2.77	3.04	2.06	2.56	2.36	2.15	1.67	1.72	1.67	1.70
-WK	R	3.99		4.64	6.71	3.94	2.88	5.60	2.75	2.47	1.51	2.20	1.59	1.50	1.58	1.63	1.58	1.61

Table 4: Time (second) and memory (GB) overhead over graphs uu, u7, and sk with p = 32.

		DBH	MDBGP	BPart	Hybrid-BL	PowerLyra	EBV	CLUGP	HDRF	2PS	HEP-1	HEP-10	HEP-100	NE	FSM-H(k=2)	FSM-N(k=2)
	Time	395.447	TIE	1669.37	1144.15	470.985	2758.66	MIE	1021.64	1856.99	684.97	365.45	325.41	7777.95	644.09	10429.5
uu	Memory	0.99	TLE	44.84	107.32	0.99	41.78	IVILE	0.99	10.3	17.06	27.56	39.52	98.79	53.22	110.84
7	Time	513.042	TIE	964.16	647.43	403.585	1746.23	5277.93	748.69	1284.61	420.09	238.64	247.48	3629.46	463.33	5971.39
u/	Memory	0.79	ILE	30.93	72.29	0.79	28.48	31.50	0.79	8.27	12.41	20.01	26.80	67.91	36.11	76.32
ale	Time	172.645	TIE	484.36	307.08	227.753	801.82	2438.81	407.79	698.26	190.98	72.21	222.75	1869.25	203.66	1965.72
SK	Memory	0.38	ILE	15.94	42.85	0.38	14.76	16.42	0.38	4.07	6.46	11.40	12.91	35.06	17.49	39.37

perform even worse than HDRF on graphs such as it. Streaming partitioners such as DBH and EBV are limited to high \mathcal{R} , and their Ω_V are also high. We also find that HDRF performs better than other streaming partitioners. Hybrid-BL tends to outperform PowerLyra on most graphs except graphs like wk and tw. FSMs can achieve the lowest Ω_V among the 11 graphs, especially for FSM-HEP. FSM-HEPs outperform FSM-NE in most cases because HEP outperforms NE in terms of Ω_V in the Fine-grained Split phase initially.

5.2.2 Replication Factor. The replication factor, i.e., \mathcal{R} , is positively correlated with the *communication volume* between clusters. We can observe that NE has the lowest R on these graphs. The \mathcal{R} of HEP depends on the value of τ . The larger the τ values, the closer HEP is to NE, resulting in a smaller R. Conversely, if the au is smaller, HEP becomes closer to HDRF, resulting in a larger \mathcal{R} . Among re-streaming partitioners, 2PS produces low \mathcal{R} , while CLUGP suffers from high replication factors. The $\mathcal R$ of Hybrid-BL is lower than PowerLyra on most graphs except tw and wk. The Rof FSM depends on the choice of the splitting factor k. The smaller the value of k, the fewer subgraphs are split, which usually leads to a smaller $\mathcal R$ (See Section 5.3.2). In most graphs, FSM achieves a \mathcal{R} close to NE's under p = 32. Streaming partitioners have a shortcoming in terms of \mathcal{R} compared to others, with DBH having the largest $\mathcal R$ followed by EBV and HDRF having the smallest $\mathcal R$. They also have a significant gap in $\mathcal R$ compared to other in-memory and hybrid partitioners.

5.2.3 Coefficient Variation of Vertex and Edge. The coefficient variation of vertex and edge are denoted as σ_V and σ_E , respectively. They indicate the discreteness of vertex and edge number among partitions, reflecting the degree of heterogeneity in memory overhead and computation workload during distributed tasks. In a homogeneous cluster, where the computing capabilities and communication capabilities between nodes are identical, the computing loads and memory overheads between machines should be as close as possible during distributed tasks. As indicated in Table 5, streaming partitioners usually have an advantage in terms of both σ_V and σ_E . DBH and PowerLyra exhibit the most outstanding σ_V because they both use hash for partitioning. EBV and HDRF follow closely behind. In-memory partitioners such as NE and HEP can only guarantee a low σ_E , but not σ_V . Hybrid-BL considers the fusion of multi-hop neighbors, which makes it impossible to obtain low σ on graphs with large degree differences. The dual-balanced partitioner BPart needs to optimize the balance of vertices and edges simultaneously, but it exhibits inferior performance in dual balance. MDBGP fails to complete the partitioning within 72 hours on all 11 graphs. FSMs achieve rather low σ_V on most graphs and the effect becomes more significant as k increases.

5.2.4 Time and Memory Overhead. Due to the space limit, we present the time and memory overhead on the three largest graphs uu, u7, and sk. The results are shown in Table 4, where 'MLE' and 'TLE' represent 'Memory Limit Exceeded' and 'Time Limit Exceeded' (the maximum partitioning time is set to 72 hours), respectively. Among the partitioners, MDBGP (40-thread partitioning)

Table 5: Average σ_V and σ_E of partitioned graphs achieved by different methods.

	DBH	MDBGP	BPart	Hybrid-BL	PowerLyra	EBV	CLUGP	HDRF	2PS	NE	HEP-1	HEP-10	HEP-100	FSM-N(k=2)	FSM-N(k=3)	FSM-H(k=2)	FSM-H(k=2)
σ_V	0.0%		20.01%	22.66%	0.1%	0.5%	32.02%	6.7%	29.08%	43.95%	19.88%	27.9%	33.31%	14.24%	6.95%	6.52%	3.67%
σ_E	0.0%		62.04%	32.29%	0.3%	4.46%	21.25%	0.0%	4.94%	12.43%	0.0%	0.0%	0.0%	0.08%	0.15%	0.0%	0.0%

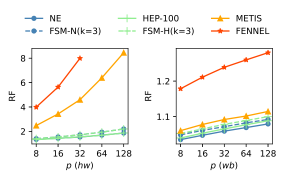


Figure 7: The trend of RF for different partitioners as p increases. FSM-N(k=3) and FSM-H(k=3) represent the FSM using NE and HEP, respectively, for partitioning with k=3.

takes more time than other methods due to its involvement in a $O(n^2)$ intersection point operations. In contrast, the streaming algorithms demonstrate the fastest partitioning time. In terms of memory overhead, the streaming algorithms PowerLyra, DBH and HDRF exhibit significant advantages, while the adjacency list-based algorithms such as BPart, NE, and Hybrid-BL require more memory. EBV sorts the edges according to the degree, which results in higher memory and time overhead compared to DBH and HDRF. The hybrid algorithms (HEPs) show intermediate performance in both time and memory overhead. The memory and time overheads of FSMs depend on the algorithms chosen for the Split Phase. From Table 4, it can be observed that FSMs roughly approximate the employed algorithms in terms of both time and memory.

5.3 Effect of Parameters

5.3.1 Effect of the Number of Partitions p. In this study, we apply four state-of-the-art partitioners (NE, HEP, METIS and FENNEL) to partition two graphs, wb and hw, with p ranging from 8 to 128. As shown in Figure 7, it is evident that all the values of $\mathcal R$ increase with the increase in p. This can be attributed to the fact that as the number of partitions p increases, there is a corresponding increase in the number of vertices at the partition boundaries. Consequently, this leads to unavoidable increases in the value of $\mathcal R$.

When analyzing the trends of the two graphs separately, we observe that increasing the value of p results in a significant rise in $\mathcal R$ for the hw graph. Particularly, for METIS, at p=128, its $\mathcal R$ has already reached around four times that of p=8. In contrast, the $\mathcal R$ for the wb graph remains consistently close to 1.0 across the entire range of p. For two FSM partitioners, we can observe that as p increases, FSM-N(k=3) and FSM-H(k=3) consistently achieve $\mathcal R$ that are similar to NE and HEP.

5.3.2 Effect of the Splitting Factor k. Fast merge prioritizes time efficiency, while precise merge often achieves better partition quality. In this study, we evaluate the effects of these two merge algorithms,

Table 6: Effects of α and β on the partitioning quality.

α	1.5	1.10	1.05	1.03	β	1.35	1.1	1.05	1.03
k	3	5	7	8	k	5	5	6	6
\mathcal{B}_V	1.32	1.10	1.05	1.02	\mathcal{B}_V	1.02	1.02	1.01	1.02
\mathcal{B}_E	1.00	1.00	1.00	1.00	\mathcal{B}_E	1.35	1.10	1.05	1.03
$\mathcal R$	1.11	1.12	1.12	1.13	$\mathcal R$	1.11	1.11	1.12	1.12

using HEP-100¹ as the partitioner in the *Fine-grained Split* phase with p=32 and k=2,3,4,5,6. We evaluate them based on merging quality (Ω_V and \mathcal{R}) and merging time. The results on graphs sk and tw are presented in Figure 8.

When using the naive HEP-100 for partitioning, it achieves a value of Ω_V of 1.73 and 2.92 on sk and tw, respectively. With the FSM framework, the $\mathcal R$ and Ω_V obtained from the *Precise Merge* are usually lower than those from the *Fast Merge*. We observe that as k increases, FSM consistently maintains $\mathcal R$ and Ω_V at similar levels, resulting in a favorable $\mathcal B_V$. In particular, the *Precise Merge* algorithm achieves an average $\mathcal B_V$ of 1.02 and 1.12 on the graphs sk and tw, respectively. For the sk graph (other similar graphs like id, it, ar, and wb), we find that FSM achieves a triple one score, where all $\mathcal B_E$, $\mathcal B_V$, and $\mathcal R$ are close to 1. This is close to the theoretical optimum, as any partition must satisfy $\mathcal R > 1$ when p > 1. The time gap between the two merge algorithms increases with k. However, even for a graph like sk with 51M vertices and 1.9B edges, the *Precise Merge* with k = 5 only takes 60 seconds.

5.3.3 Effect of α and β . We discuss how the dual-balanced constraints, α and β , affect the quality of FSM partitioning. To study the effect of α , we vary α from 1.03 to 1.50 by fixing $\beta=1$. Table 6 presents the results on graph uk. It shows that stricter vertex balance requirements (i.e., smaller α) necessitate a larger k, at the same time it also results in higher replication factors $\mathcal R$. To study the effect of β , we vary β from 1.03 to 1.35 by fixing $\alpha=1.03$. Compared to the case of fixing $\beta=1$, relaxing the constraint on β makes it easier to achieve the target of $\alpha=1.03$. It is worth noting that FSM maintains a lower replication factor $\mathcal R$ as well.

5.4 Results of Distributed Tasks

5.4.1 Accelerating Distributed Tasks. To evaluate the performance of FSM on downstream tasks, we conduct PageRank (PR), connected components (CC), and approximate diameter (AD) on a cluster of p=8 machines. PR is fixed to 100 iterations to force all vertices active so that we can test the effect under heavy communication overhead. CC is implemented based on label propagation, with fewer active vertices in each iteration. AD estimates the diameter by computing reached vertex pairs for each hop [18], with all vertices being re-activated in each hop. We fix the number of hops to 10 due to its long elapsed time. We repeat each experiment 3 times, and take the average as the result, ensuring that the standard error of the

 $^{^{1}}$ NE cannot guarantee edge balance when p is large.

Table 7: Runtime of distributed PageRank (sec). The lowest runtime is highlighted in bold, second lowest is underlined.

Graph	DBH	MDBGP	BPart	Hybrid-BL	PowerLyra	EBV	CLUGP	HDRF	2PS	HEP-1	HEP-10	HEP-100	NE	FSM-N(k=2)	FSM-N(k=3)	FSM-H(k=2)	FSM-H(k=3)
ar	329.23	361.90	255.27	226.50	217.80	222.30	235.13	216.23	173.60	179.47	180.50	176.00	192.70	166.53	159.47	164.33	173.37
hw	124.37	111.43	108.53	99.67	87.80	88.30	102.13	90.27	85.93	79.37	78.50	73.63	74.87	72.07	71.40	68.17	70.43
id	167.77	139.90	121.40	105.57	94.27	104.67	104.70	105.30	99.03	97.77	98.53	95.43	101.40	92.83	97.87	93.27	93.60
it	622.07	-	476.33	390.33	383.30	381.67	371.17	290.50	267.77	309.73	333.17	291.90	376.77	282.47	264.97	249.97	265.77
sk	767.60	-	828.17	477.60	523.47	510.20	412.47	436.27	373.60	334.30	337.63	323.10	341.60	332.67	311.50	308.17	308.10
tw	535.63	-	1028.97	MLE	424.90	479.10	843.23	438.57	494.80	422.40	405.17	394.40	395.80	423.13	420.50	392.93	398.83
u7	1527.50	-	1173.73	893.07	871.33	990.80	759.27	732.53	496.13	527.70	567.93	595.07	739.93	550.30	534.70	462.67	449.37
uk	648.80	-	448.30	368.53	421.50	377.97	314.10	315.60	281.83	307.20	294.83	294.33	321.50	277.60	261.90	274.53	259.63
uu	1892.30	-	1080.83	1371.20	MLE	1398.83	-	829.63	649.93	691.83	781.97	782.20	895.90	685.37	664.90	631.90	606.83
wb	1609.17	-	914.70	1121.27	866.33	921.30	872.10	853.97	733.93	836.27	765.97	748.83	782.23	778.20	741.77	699.17	699.30
wk	305.43	522.57	416.97	405.00	196.97	261.53	352.53	247.13	240.40	227.47	192.60	193.37	200.67	199.60	198.60	198.97	197.53

Table 8: Runtime of connected components (sec). The lowest runtime is highlighted in bold, second lowest is underlined.

Graph	DBH	MDBGP	BPart	Hybrid-BL	PowerLyra	EBV	CLUGP	HDRF	2PS	HEP-1	HEP-10	HEP-100	NE	FSM-N(k=2)	FSM-N(k=3)	FSM-H(k=2)	FSM-H(k=3)
ar	46.77	58.30	44.23	40.30	38.53	41.93	43.27	35.97	34.00	32.07	33.60	33.23	37.33	31.13	29.53	28.67	33.03
hw	12.73	14.03	14.17	12.40	10.37	10.90	12.00	10.30	9.87	9.57	9.20	9.17	8.47	9.37	8.33	8.83	7.70
id	24.30	25.87	20.87	20.07	16.80	21.37	19.97	17.27	18.23	18.50	18.57	19.47	20.70	17.97	17.33	17.77	16.87
it	77.90	-	80.07	75.47	62.97	65.70	66.30	50.37	55.87	49.07	51.57	52.87	60.87	49.90	48.43	46.17	43.50
sk	76.80	-	90.00	71.33	81.63	70.90	77.03	53.53	60.70	47.33	44.00	45.17	53.10	46.90	47.37	45.03	45.00
tw	64.13	-	103.67	MLE	63.73	66.43	100.97	61.23	77.03	57.27	56.80	55.47	61.83	56.33	55.83	50.57	51.63
u7	173.50	-	179.20	154.43	157.73	169.50	147.77	125.87	114.40	113.27	118.87	113.73	123.10	110.47	110.90	110.20	108.77
uk	104.90	-	83.60	82.23	86.23	83.23	71.63	61.80	64.20	57.47	67.37	66.10	73.20	57.57	54.70	57.50	61.20
uu	294.13	-	232.13	269.50	MLE	277.43	-	190.90	197.20	186.60	192.93	182.23	199.00	180.97	180.27	174.67	174.93
wb	509.60	-	444.87	473.73	423.10	480.37	444.03	430.77	409.97	399.53	432.57	421.30	432.13	412.83	396.10	415.50	418.57
wk	49.53	74.57	65.87	64.47	34.43	45.67	50.47	44.07	42.60	43.13	40.47	42.03	41.13	37.40	36.80	37.17	41.53

Table 9: Runtime of approximate diameter (sec). The lowest runtime is highlighted in bold, second lowest is underlined.

Graph	DBH	MDBGP	BPart	Hybrid-BL	PowerLyra	EBV	CLUGP	HDRF	2PS	HEP-1	HEP-10	HEP-100	NE	FSM-N(k=2)	FSM-N(k=3)	FSM-H(k=2)	FSM-H(k=3)
ar	954.67	1189.33	779.67	759.00	910.82	725.00	902.00	651.67	575.67	535.67	528.67	531.67	586.00	470.33	439.00	452.67	457.00
hw	233.00	292.00	200.00	225.00	239.91	149.67	211.33	133.33	162.00	130.00	144.67	127.67	131.33	112.33	111.33	116.33	115.33
id	304.00	477.00	229.00	244.00	249.54	284.00	189.00	200.67	199.33	167.00	185.33	210.00	339.67	153.00	146.00	144.00	146.00
it	1909.33	-	1543.00	1589.00	1705.89	1417.33	1684	1050.00	1042.00	1006.33	1073.33	1065.67	1376.67	954.67	861.33	851.67	867.67
sk	3243.33	-	2792.67	2423.00	2713.50	2131.67	2550.33	1859.33	1540.00	1393.67	1316.67	1333.33	1717.00	1414.67	1261.33	1256.67	1281.33
tw	2027.00	-	4324.33	MLE	1963.02	1900.67	4165.67	1713.00	2546.67	1634.67	1633.33	1816.67	2077.67	1611.33	1713.33	1487.33	1569.67
u 7	MLE	-	MLE	MLE	MLE	MLE	MLE	MLE	MLE	MLE	MLE	MLE	MLE	MLE	MLE	TLE	4326.00
uk	1835.00	-	1185.67	1395.33	1940.11	1444.67	1145.67	946.67	980.67	918.00	1103.33	1157.33	1174.33	846.33	753.33	858.67	771.67
uu	MLE	-	MLE	MLE	MLE	MLE	-	MLE	MLE	MLE	MLE						
wb	MLE	-	MLE	MLE	MLE	MLE	MLE	MLE	MLE	MLE	MLE	MLE	MLE	MLE	4587.33	4542.33	2041.33
wk	964.67	2363.33	1759.33	2286.33	702.78	918.67	1495.33	878.00	859.00	792.00	757.67	734.33	754.00	700.00	714.33	710.67	727.33

experiment time is less than 5%. For tasks that run out of memory, we label them as "MLE". For tasks that cannot be finished within 12 hours, we abort them and label them as "TLE". Tables 7, 8, and 9 list the results, where "-" indicates that the partitioning cannot be finished within 72 hours or exceeds the memory limit during the partitioning process.

Overall elapsed time. We conclude that using FSM-N(k=3) or FSM-H(k=2) generally results in the shortest elapsed time on most graph tasks. Although NE achieves the lowest \mathcal{R} , we find that its task elapsed time is generally not as good as HEP and even falls behind the streaming partitioner HDRF (on the it graph). HEP demonstrates excellent performance in PR and CC tasks, with overall performance second only to FSM. Among the streaming partitioners, HDRF exhibits the best performance, followed by EBV, and DBH performs the worst. The two re-streaming algorithms

show impressive performance on graphs like it. This is likely due to the presence of clustering in graphs, which allows these algorithms to maintain good balance while achieving low \mathcal{R} . Power-Lyra demonstrates good performance on graphs such as tw and wk, while Hybrid-BL performs better on graph wb. We note that the improvement of FSM-N over NE is impressive. The maximum improvement on PR and CC tasks is 29.7% and 25.3%, respectively. Similar improvement is also observed in FSM-H. This is because FSM preserves the original low \mathcal{R} of NE and HEP and optimizes their maximum load.

Paging Fault due to vertex imbalance. In distributed environments with limited memory, large graph processing tasks often experience frequent page faults. In some cases, although the graph processing task does not exceed memory capacity, the memory usage approaches 100%, increasing the likelihood of page faults. These

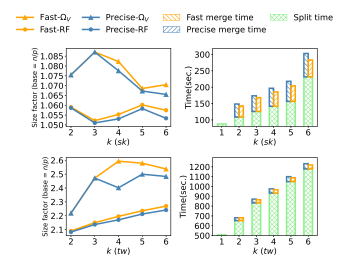


Figure 8: The merge effect and time of FSM-H with different k, where p=32 and k=1 indicates partitioned by naive HEP-

page faults contribute to higher CPU utilization by the kswapd0 process, which is used for swapping pages. For instance, the AD tasks on graph *wb* partitioned by FSM-N(k=3) and FSM-H(k=2) are adversely affected by increasing page faults and additional CPU utilization, resulting in a long elapsed time. Hence, in distributed tasks, the memory overhead on the largest partition should be minimized, even though it may not exceed memory.

5.4.2 Effectiveness on Different Cluster Size. To further explore the potential of FSM on small clusters (i.e., the number of machines utilized), we vary cluster sizes from 5 to 8 with 32GB memory per machine. Figure 9 shows the improvement of FSM compared to the fastest runner among baselines on three large graphs it, uk, and ar. As the cluster gets smaller, it becomes harder to process large graphs. On one hand, the reduction in the number of machines means that each machine needs to load larger partitions, which means a higher probability of memory exceeding. On the other hand, fewer machines mean fewer CPU cores, which means a longer elapsed time.

For the AD tasks, we mark the speedup as 100% since all other partitioners only get "TLE" (more than 12 hours) or "MLE", while FSMs can finish within 1.5 hours. Furthermore, we have observed that even with smaller clusters, FSM still provides significant improvements, particularly in the AD task. We can conclude that FSM, compared to other partitioners, not only achieves improvements in downstream task elapsed time but also *enables the processing of large graphs on small clusters*.

5.4.3 Guidance on Applicability. Based on extensive experimental results on various datasets, we have summarized the following scenarios where the FSM framework is applicable: (I) Graphs with rich communities and high-degree vertices, such as wb, it, uu, and id. These communities and high-degree vertices can facilitate the fine-grained splitting phase of FSM, providing small replication factors even for a large number of partitions. (II) Downstream tasks that require high memory usage. Tasks such as triangle counting

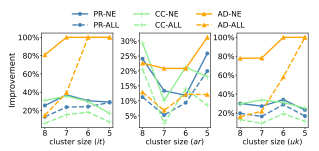


Figure 9: The speedup effect on different cluster sizes using FSM-N(k=3).

and approximate diameter often involve complex data structures that consume a significant amount of memory. This can lead to memory bottlenecks on individual machines. (III) When the cluster size is small, each machine needs to handle a larger computational load, and the issue of load imbalance in terms of computation and memory becomes more prominent. Therefore, using the FSM framework to balance the load can significantly enhance the performance of tasks on small clusters.

6 RELATED WORK

In-memory partitioners. One of the most established categories of in-memory partitioning algorithms is multilevel partitioning algorithms, which generally follow the three-step process of coarsening, partitioning, and refinement. Different algorithms are used in each step, and notable examples include METIS [19], KaFFPa [33], and Scotch [30]. These algorithms emphasize cut size and vertex balance but tend to perform poorly in terms of edge balance. NE [39] is a typical edge partitioning algorithm that achieves excellent $\mathcal R$ by prioritizing the expansion of vertices with fewer external connections. It currently provides the best quality among edge partitioners. Other algorithms based on the NE idea include HEP (along with NE++) [26] and DNE [15]. They also primarily focus on optimizing partition efficiency and scalability but often neglect the issue of vertex balance.

Streaming partitioners. Streaming algorithms [35] can be further classified into stateful and stateless categories. Stateful algorithms typically evaluate vertices or edges using specific scoring functions and allocate them to partitions with the highest scores. HDRF [31] prioritizes cutting high-degree vertices and performs well on power-law graphs. However, the scoring function does not consider vertex balance. FENNEL [36] is a vertex partitioning algorithm that takes into account both the number of adjacent and non-adjacent vertices. However, it does not consider edge balance. 2PS [27] and CLUGP [21] are two re-streaming algorithms. They utilize clustering algorithms to proactively gather global information, which aids in subsequent re-partitioning. The difference lies in the clustering strategy used. These re-streaming algorithms tend to yield better partitioning results but often sacrifice balance.

Stateless algorithms only use minimal graph information, such as degrees and vertex IDs. These algorithms often employ randomization or hashing techniques for partitioning. Grid [17] is a hashing-based edge partitioning algorithm that takes into account the load balancing of partitions. DBH [38] is a degree-based hashing edge partitioning algorithm, where the hashing function

incorporates the idea of "high-degree vertices are preferentially replicated", resulting in good performance on power-law graphs. Random partitioning [35] randomly assigns edges or vertices to partitions.

Dual-balanced partitioners. To the best of our knowledge, there are three works that consider dual-balanced partitions, but they often produce larger cut sizes (i.e., replication factor). Zhang et al. proposed the edge partitioning algorithm EBV [40], which incorporates both vertex and edge load as well as vertex replication into the scoring function and thus achieves a good dual-balance. However, like other streaming algorithms, it can only achieve suboptimal \mathcal{R} . Lin et al. extend the FENNEL algorithm and introduce a vertex partitioning scheme called BPart [24]. It achieves dualbalance through multiple rounds of re-partitioning and combining. However, it requires simultaneous optimization of both vertex and edge balance during the combining phase and only considers extreme merges (merging the current maximum with the minimum), without guaranteeing an approximation ratio. In terms of \mathcal{R} , BPart incurs significant losses compared to FENNEL, while FENNEL is inferior to METIS in \mathcal{R} . Different from the discrete algorithms, MD-BGP [1] uses Projected Gradient Descent to balance vertex-edge load. It exhibits high parallelism and emphasizes multi-dimension balance. However, it suffers from significant complexity and falls short in cut size, performing even worse than FENNEL.

Different from these methods, FSM employs an effective twophase method that can benefit from any SOTA single-balanced partitioners and achieve remarkable replication factor and promising scalability.

Postprocessing algorithms. Postprocessing algorithms improve the quality of partitions by making adjustments or refinements to them. ParE2H and ParV2H [9] are task-specific algorithms that refine given partitions to shorten the elapsed time for specific tasks. However, they usually require higher training costs and are limited to optimizing a particular task. In contrast, FSM achieves fast partitioning and provides acceleration benefits for a wide range of tasks. Additionally, FSM generates high-quality partitions with dual-balance, which can serve as pre-trained partitions for ParE2H and ParV2H. LS [14] adjusts edges and blocks of given partitions to achieve high-quality partitions under edge balance. However, similar to other edge partitioners, LS primarily focuses on reducing \mathcal{R} , which can potentially lead to a deterioration in vertex balance during the adjustment process. Therefore, LS only achieved a maximum improvement of 10.6% in downstream tasks [14].

Dynamic partitioners. Dynamic partitioners are used for partitioning dynamic graphs. They typically achieve dynamic optimization by migrating vertices and edges. Based on two optimization objectives: structural changes and computational load changes, dynamic partitioners can be divided into two categories. The former focuses on optimizing the dynamic graph structure, e.g., Leopard [16], Hermes [28], and Planar [41]. The latter considers the workload of the graph computation, e.g., Mizan [20] and CatchW [34].

FSM can be also extended to dynamic scenarios. For dynamically incoming graphs, a buffer can be set up to cache the incoming updates. The buffer can then be partitioned using a single-balanced partitioner, and the resulting partitions can be merged with the existing buckets using the Merge operation. This allows for the incorporation of dynamic updates while maintaining dual balance.

7 CONCLUSION

In this paper, we introduce a lightweight and efficient partitioning framework FSM that consists of two phases, namely *fine-grained splitting* and *subgraph merging*. The experimental results on large-scale graphs have demonstrated that the proposed framework FSM achieves a remarkable replication factor while achieving both vertex and edge balances, outperforming state-of-the-art partitioners. Furthermore, the performance of downstream tasks can be significantly improved benefiting from the partitioning result.

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