# GSAP: A GPU-Accelerated Stochastic Graph Partitioner

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# ABSTRACT

Graph partitioning is essential for understanding the structure of a dataset, such as social networks and web pages. Among various graph partitioners, stochastic block partitioning (SBP) has shown promise in handling complex graphs with varying community sizes or strong intra-community connections. However, the sequential nature of the Monte Carlo Markov Chain iterations and the stochastic proposal generation process limit the efficiency and scalability of SBP. To overcome this limitation, this paper introduces GSAP, a GPU-accelerated stochastic graph partitioner, to enhance the runtime performance of SBP. We propose a parallel algorithm to speed up the generation process of stochastic proposals on GPU. Additionally, we accelerate the calculation of the minimal description length by dividing the formulation into several independent computations. To achieve better performance, we introduce an efficient blockmodel update algorithm to dynamically manage the blockmodel matrix on GPU. Our experimental results on the 2022 HPEC GraphChallenge dataset demonstrate that GSAP can achieve up to 12.3× and 60.9× runtime speedup on a single A4000 GPU compared to two CPU-parallel state-of-the-art SBP algorithms.

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#### 1 INTRODUCTION

Many real-world datasets [\[50\]](#page-10-1), such as social networks and web pages, use graphs to represent complex structures and relationships. Graph partitioning [\[49,](#page-10-2) [75\]](#page-10-3), which divides a graph into several parts, is a crucial technique for understanding these complex datasets. For example, the spectral algorithm [\[64\]](#page-10-4) utilizes the eigenspectrum of the modularity matrix to partition graphs based on connectivity between vertices. The Girvan–Newman algorithm [\[65\]](#page-10-5) partitions a graph by iteratively removing edges with the highest betweenness centrality.

SBP [\[66](#page-10-6)[–68\]](#page-10-7) is one of the graph partitioning algorithms based on iterative inference over stochastic blockmodels [\[45\]](#page-10-8) to identify the optimal partition of a graph. Stochastic blockmodels represent a graph's structure by a two-dimensional sparse matrix. The elements in the matrix record edge count between blocks. The SBP algorithm consists of three phases: (1) the block-merge phase, (2)



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the vertex-move phase, and (3) the golden-section search. In the block-merge phase, selected blocks are merged. During the subsequent vertex-move phase, Monte Carlo Markov Chain (MCMC) iterations [\[18\]](#page-9-0) use the Metropolis-Hastings algorithm to move vertices between blocks, refining the current partition. Next, the goldensection search phase identifies the optimal partition that achieves the minimum description length (MDL) of the blockmodel based on the current partition. Compared to conventional graph partitioning approaches, SBP performs well on complex graphs which have varied community sizes or strong intra-community connections. However, the serial nature of MCMC iterations and the stochastic proposal generation process limit the runtime efficiency and scalability to large graphs.

As a result, the recent IEEE High Performance Extreme Computing Conference (HPEC) introduced the SBP Challenge (SBPC) [\[44\]](#page-10-9) to seek novel acceleration techniques from the the high-performance computing (HPC) community. F-SBP [\[70\]](#page-10-10), the 2019 award winner, leverages existing sampling algorithms to enhance SBP runtime. Faster-SPB [\[69\]](#page-10-11), the 2021 champion, implements an aggressive initial merging strategy to significantly reduce the number of iterations required. H-SBP [\[73\]](#page-10-12) proposes a hybrid approach by combining MCMC iterations with the asynchronous Gibbs algorithm for parallelization. uSAP [\[2\]](#page-9-1), the 2023 award winner, introduces a novel initial block-merge strategy based on strongly connected components and employs task graph parallelism to enhance the runtime of SBP. However, these methods are still limited to handling only small to medium-sized graphs. I-SBP [\[72\]](#page-10-13), 2023 GraphChallenge winner, presents a distributed SBP algorithm to partition larger graphs, while their single-node version show limited performance.

Despite improved runtime performance by existing works, their scalability is largely constrained by CPU parallelism. Compared to CPU, modern GPU offers order-of-magnitude more parallelism and memory bandwidth, which is particularly suitable for handling large graphs. This advantage has inspired us to leverage the power of GPU computing to accelerate SBP. To this end, we propose a GPUaccelerated stochastic graph partitioner called GSAP. To the best of our knowledge, this is an early attempt that successfully accesses SBP using GPU computing. We summarize our contributions as follows:

- We propose a parallel algorithm to accelerate the generation of stochastic proposals on GPU, significantly speeding up the process compared to the CPU-parallel approach.
- We decompose the formulation of MDL into several independent calculations, enabling the parallel computation for each proposal during the block-merge and vertex-move phases. This approach significantly speeds up the heavy workload of MDL computation.
- We introduce an efficient blockmodel update algorithm for dynamically updating the blockmodel matrix during the blockmerge and vertex-move phases on GPU. This significantly enhances the runtime performance compared to the CPU-parallel approach.

We have evaluated GSAP on a single A4000 GPU using the datasets provided by the HPEC SBP Challenge [\[44\]](#page-10-9) and compared the runtime performance over two state-of-the-art SBP algorithms, uSAP [\[2\]](#page-9-1) and I-SBP, [\[72\]](#page-10-13) on a 14-core CPU. The experimental results shows that GSAP achieves 12.3× speedup over uSAP and 60.9× speedup over I-SBP when partitioning graphs of 50K–200K vertices. For the largest graphs of 1M vertices, GSAP can finish SBP in just 15 minutes, whereas uSAP and I-SBP both fail to complete in two hours.

#### 2 BACKGROUND

Graph partitioning aims to identify unique community structures within a graph by determining the partition of each vertex. This partition provides insights into the interactions among vertices and helps identify vertices that belong to specific communities of interest. Some well-known methods adopt the spectral approach [\[64\]](#page-10-4) to identify partitions in a graph. Some algorithms depend on graph modularity, which measures the density of connections within partitions. High modularity in graphs indicates more intra-partition connections and fewer inter-partition links. Nevertheless, this approach faces challenges, including resolution limits that affect the detectable size of partitions and difficulties in determining the optimal number of communities. To tackle these issues, recent studies have chosen generative statistical models for graph partitioning. Specifically, [\[66](#page-10-6)[–68\]](#page-10-7) utilize degree-corrected stochastic blockmodel [\[45\]](#page-10-8) for partition estimation, where each partition is defined as a block.

## 2.1 Stochastic Block Partitioning

Stochastic Block Partitioning (SBP) utilizes the stochastic blockmodel to identify the optimal partition of a graph, as depicted in Figure [1.](#page-1-0) This approach models the block structure with a sparse matrix  $M_{B\times B}$ , where B is the number of identified blocks within the graph. Each matrix element,  $M_{a,b}$ , records the edge count starting from block  $a$  to block  $b$ . The log posterior probability function, denoted by  $P(G|B)$  in Equation [1,](#page-1-1) is used to describe the current partition of graph  $G$ , given the blockmodel B.  $M_{i,j}$  is edges count between blocks *i* and *j*.  $D_i^{in}$  and  $D_j^{out}$  represent the total in-degree and out-degree for blocks  $i$  and  $j$ , respectively.

<span id="page-1-1"></span>
$$
P(G|B) = \sum_{i,j} M_{i,j} \cdot \log \left( \frac{M_{i,j}}{D_i^{in} D_j^{out}} \right)
$$
 (1)

SBP is divided into three distinct phases: block-merge, vertexmove, and golden-section search. SBP iterates among these phases to discover the ideal partition with the optimal block count,  $B^*$ , that minimizes the total description length  $(MDL)$ , as defined in Equation [2.](#page-1-2) To overcome the issue of local minima, the algorithm starts with each node assigned to its own block. Then, blocks are progressively merged, followed by Monte Carlo Markov Chain (MCMC) updates on individual vertices to determine the optimal partition for the current number of blocks.

<span id="page-1-2"></span>
$$
MDL = Eh\left(\frac{B^2}{E}\right) + V\log B - P(G|B)
$$
  
where  $h(x) = (1+x)\log(1+x) - x\log x$  (2)

In the block-merge phase, a merge is proposed for every block, followed by calculating the resulting change in MDL. The merges resulting in smaller MDL changes are performed according to the reduction rate parameter. In the vertex move phase, a move is proposed for each vertex, followed by the calculation of Metropolis-Hastings acceptance ratio based on MDL to determine whether the move should be accepted or rejected. Subsequently, the blockmodel matrix  $M$  is updated. The vertex-move phase is the major bottleneck in SBP due to the sequential MCMC updates that iterate over every vertex until convergence is achieved. The profiling results presented in [\[73\]](#page-10-12) reveal that the vertex-move phase can account for up to 98% of the total runtime. This high percentage is due to the timeconsuming nature of the proposal generation process and the MDL computation for each vertex. Furthermore, the blockmodel matrix must be updated after proposals are accepted during the vertexmove phase. Iterating through these three steps makes the vertexmove phase the bottleneck of the SBP algorithm.

<span id="page-1-0"></span>

Figure 1: Overview of the SBP algorithm. Initially, each vertex in the graph is assigned to a block. The graph then undergoes iterative block-merge and vertex-move phases until the partitioning process achieves the minimum description length (MDL) through a golden-section search algorithm.

# 2.2 HPEC SBP Challenge

Despite SBP's ability to handle complex graphs, its major limitation is the long runtime, caused by the sequential nature of the Monte Carlo Markov Chain iterations and the stochastic proposal generation process. To overcome the limitation, the recent IEEE HPEC launched the SBP Challenge (SBPC) to seek novel acceleration approaches for the SBP algorithm. To evaluate the performance of a graph partitioner, SBPC provides a comprehensive evaluation dataset containing synthetic graphs based on the stochastic model [\[45\]](#page-10-8). These graphs are generated by sampling the connection between vertices from a Poisson distribution with a correction term to simulate real-world graph characteristics. Table [1](#page-2-0) details the attributes of the dataset, which includes four graph categories ranging from 1K to 1M vertices. Each category represents varying degrees of partitioning challenges and graph complexity with the difficulty level of the categories increasing from easiest to hardest.

(1) Low-Low: Low block overlap and low block size variation

(2) Low-High: Low block overlap and high block size variation (3) High-Low: High block overlap and low block size variation

(4) High-High: High block overlap and high block size variation

<span id="page-2-0"></span>

Table 1: The four categories of the SBPC dataset. The Low-Low category represents low complexity, making it the easiest benchmark, while the High-High category represents high complexity, making it the hardest benchmark. Each category contains graphs with vertex counts ranging from 1K to 1M.  $|V|$  represents the number of vertices in the graph,  $|E|$  denotes the number of edges, and  $B$  is the number of partitions.

#### 2.3 Related Works and Their Limitations

Many novel approaches have been proposed to accelerate the SBP algorithm. For example, F-SBP [\[70\]](#page-10-10) reduces the SBP runtime by utilizing an existing sampling algorithm for subgraph selection and performing partitioning across multi-core clusters. This approach preserves the graph's partition structure and accelerates the algorithm without significantly losing the accuracy. Nevertheless, this method does not achieve a significant reduction in runtime to effectively address the challenges posed by large graphs.

Faster-SPB [\[69\]](#page-10-11) introduces an aggressive initial merging strategy to considerably decrease the initial block count at the first iteration, thereby reducing the overall number of partitioning iterations needed. However, the aggressive initial merging strategy may merge blocks that cause negative effects on the partition quality. Its parallelism control strategy carefully manages the amount of parallelism during different phases of the algorithm to improve the performance while it does not perform well due to the synchronization overhead.

H-SBP [\[73\]](#page-10-12) implements the asynchronous Gibbs algorithm to parallelize the MCMC iterations. This innovative approach enables the parallelization of computations that are typically sequential within each MCMC in SBP. By serially processing a select portion of the most influential vertices and parallelizing the remainder, H-SBP maintains accuracy while significantly enhancing efficiency. The method demonstrates strong scaling capabilities, particularly in making the vertex move phase parallelizable. Despite these advances, H-SBP may encounter convergence issues with larger graphs.

uSAP [\[2\]](#page-9-1) introduces a new strategy based on strongly connected components for initial block merging, decreasing the number of iterations needed for partitioning. To speed up runtime, uSAP implements a dynamic, batch-oriented task graph parallel algorithm for vertex moves, leveraging Taskflow [\[6,](#page-9-2) [27,](#page-9-3) [32,](#page-10-14) [33,](#page-10-15) [35\]](#page-10-16) to accelerate SBP.

EDiSt [\[71\]](#page-10-17) introduces a distributed version of SBP, allowing the algorithm to efficiently scale across a larger number of compute nodes without encountering convergence problems. They also develope a version of SBP optimized for shared memory parallelism on larger clusters. However, the all-to-all communication pattern in EDiSt becomes a significant bottleneck as the number of nodes increases.

I-SBP [\[72\]](#page-10-13) combines three distinct heuristics [\[70,](#page-10-10) [71,](#page-10-17) [73\]](#page-10-12) to accelerate SBP. However, it still encounters limitations similar to those observed in other approaches.

Despite various approaches developed to accelerate the SBP algorithm, several significant challenges remain unresolved. We highlight three major challenges below:

2.3.1 Limitation to CPU Parallelism. Nearly all existing works leverage multi-core CPU parallelism to accelerate the SBP algorithm. However, the effectiveness of CPU-based parallelism diminishes as graph sizes increase. Therefore, the limitations of CPU parallelism for large graphs remain a critical challenge.

2.3.2 Lack of GPU Methods for SBP. To the best of our knowledge, no existing works focus on developing a GPU-accelerated SBP algorithm. Utilizing GPUs to handle large graphs presents a promising solution due to their ability to massively parallelize computations in SBP, such as generating stochastic proposals and calculating MDL.

2.3.3 Challenges with GPU Implementation. Although GPU offers substantial speedup and efficient handling of large volumes of data, designing a GPU-accelerated SBP presents several challenges. For instance, an efficient GPU data structure and kernel algorithm are necessary to dynamically represent partitioned graphs during the SBP process. The irregular data patterns pose challenges in developing GPU kernel algorithms for generating stochastic proposals and computing the MDL on GPU. Furthermore, it is critical to update the blockmodel on GPU to minimize the runtime of SBP. Effectively addressing these challenges is key to optimizing the algorithm's performance.

<span id="page-3-0"></span>

Figure 2: Overview of GSAP. The task graph consists of three steps: (1) The block-merge phase, where each block proposes another block to merge with and calculates the corresponding change in MDL ( $\triangle MDL$ ). Subsequently, these proposals are transferred back to the CPU for merging. (2) The vertex-move phase, which performs parallel MCMC iterations by selecting batches of vertices to propose a move to another block and computing the acceptance probability of each proposal. (3) The golden-section search step, which determines when GSAP should stop by progressively narrowing the search to the number of blocks where the MDL is achieved.

# 3 GSAP

To overcome the above challenges, we introduce GSAP, a GPUaccelerated stochastic graph partitioner, to speed up the SBP algorithm using GPU. Figure [2](#page-3-0) shows the overall task graph of GSAP, where arrows indicate task dependencies. The task graph consists of three steps: the block-merge phase, the vertex-move phase, and the golden-section search. In the block-merge phase, each block stochastically proposes another block to merge with and calculates the corresponding change in  $MDL$  ( $\triangle MDL$ ). Subsequently, these proposals are transferred back to CPU, where the selected blocks perform merges. In the vertex-move phase, we select a batch of vertices to perform asynchronous Gibbs sampling for the MCMC iterations. Each selected vertex stochastically proposes a block to move to and computes the acceptance probability for this proposal. If the proposal is accepted, the vertex is moved to the proposed block. The proposals in both the block-merge phase and the vertexmove phase are generated in parallel on GPU. The golden-section search step determines when the algorithm will stop. When the MDL is achieved, we obtain the optimal partition of the graph. We highlight the GPU-accelerated steps in dark color, while the perform-merge step is executed on the CPU because its does not benefit from GPU acceleration. We aim to address the runtime bottleneck through GPU parallelism.

## 3.1 Data Representation

The blockmodel matrix is a crucial data structure used to represent the structure of a graph in the SBP algorithm. Elements within the matrix represent the weighted sum of edges between blocks, while

<span id="page-3-1"></span>

Figure 3: An example of data representation in blockmodels, illustrating both the matrix representation and the Compressed Sparse Row format (CSR) representation.

the diagonal elements denote the weighted sum of edges within each block, indicating self-connectivity. Typically, the blockmodel matrix appears as a sparse matrix, presenting challenges for efficient storage. To reduce the memory footprint and maximize the size of the graph that can be stored, we utilize the Compressed Sparse Row (CSR) format to store the blockmodel in GPU's DRAM. Figure [3](#page-3-1) illustrates a blockmodel represented in the CSR format, which consists of three arrays:

- adjNbr: The adjacency neighbor list array contains the blocks adjacent to each block.
- adjPtr: The adjacency pointer array indexes the start of each block's adjacency list in adjNbr.

• adjWgt: The adjacency weight array records the weight of each edge connecting each block to its neighboring blocks.

Given that the edges between blocks are directed, it is crucial to maintain information on both in-degree and out-degree neighbors. The CSR structure incorporates six arrays to store this information used by GSAP. For example, in Figure [3,](#page-3-1) consider the out-degree neighbors of block 0: an edge within the block has a weight of 3, representing self-connectivity. Additionally, block 0 has an out-degree neighbor, block 2, with the weighted sum of edges directed from block 0 to block 2 recorded as 5. Therefore, the adjacency list for block 0 includes entries for both blocks 0 and 2, with corresponding weights of 3 and 5 noted in the adjacency weight array.

To represent the partition of a graph, we use a block mapping array, denoted by Bmap, to record the block ID for each vertex. Whenever blocks merge or vertices move, we update this array to reflect the current partition. Additionally, to calculate the Δ for each proposal during both the block-merge and vertex-move phases, we maintain the in-degree and out-degree for each block in two separate arrays, BdegIn and BdegOut.

## 3.2 Generating Stochastic Proposals

The iterative proposal-generating process during the block-merge and vertex-move phases of the SBP algorithm is time-consuming. Each block (or vertex) initially selects an adjacent block based on the multinomial probability distribution derived from the weights of connecting edges. If a block (or vertex) has no neighbors, it randomly selects a block. Subsequently, a number  $x$  is drawn from a uniform distribution between 0 and 1 and compared to  $\frac{B}{\deg(u)+B}$ , where  $B$  represents the number of blocks in the current partition and deg[u] is the total degree of block  $u$ , calculated as the sum of BdegIn[u] and BdegOut[u]. If  $x$  is greater than the value, the proposal is generated from the adjoining block of  $u$ ; otherwise, a block is proposed randomly. This stochastic decision is implemented to prevent the proposal-generating process from being trapped in local optima of MDL. During the block-merge phase, this process is repeated for each block in the current partition, iterating num\_proposals times. In the vertex-move phase, each vertex in the graph iteratively generates a proposal until it reaches the MDL.

To leverage GPU parallelism for efficiently solving the iterative proposal-generating process in SBP, we implemented three generators using the NVIDIA cuRAND library. Initially, we launched the random number generator, uniform number Generator, and multinomial distribution generator. These are used to create three lookup tables, which is detailed in Figure [4.](#page-4-0) This setup significantly reduces the overhead of repeatedly generating random numbers. Specifically, The uniform number generator produces a large batch of  $x$  for each proposal. The random number generator is tasked with creating numerous random block IDs. The multinomial distribution generator determines an adjacent block for each block in the graph.

With the lookup table in place, we can initiate many GPU threads to generate proposals simultaneously. During the block-merge phase, we scale up the operation by launching threads equal to the product of the number of blocks in the current partition and the num\_proposals to handle proposals in parallel. For each block, the best proposal is selected based on the MDL. In the vertex-move

<span id="page-4-0"></span>

 $P_{00}$   $P_{01}$   $\ldots$   $P_{0k}$   $P_{10}$   $P_{11}$   $\ldots$   $P_{1k}$   $\ldots$ *num\_proposals*  $P_{10}$   $P_{11}$   $\ldots$   $P_{1k}$ *num\_proposals*  $P_{N0}$   $P_{N1}$   $\cdots$   $P_{Nk}$ *num\_proposals*

Figure 4: In both the block-merge and vertex-move phases, GSAP initially launches three concurrent kernels to build three lookup tables that store multinomial distributions, uniform numbers, and random numbers. Subsequently, GSAP launches many GPU threads to generate proposals in parallel using these lookup tables.

phase, we employ a parallel MCMC approach where a batch of vertices generates proposals simultaneously. This strategy considerably reduces the time required for sequential MCMC iterations. The generated proposals are stored in a proposal array, which is used to calculate the MDL for each proposal. The GPU kernel algorithm for generating proposals in the block-merge phase is described in [1.](#page-4-1) Each block considers its neighbor counts and then uses three pre-generated tables to propose candidates. The algorithm for the vertex-move phase is similar, but each vertex must identify the block ID of its adjacent vertices.

<span id="page-4-1"></span>

## 3.3 Computation of MDL

The MDL computation is another time-consuming step in SBP, as it is necessary to compute the MDL for each proposal. To speed up this heavy workload, we begin by decomposing the formulation into several independent calculations. Instead of calculating the entire MDL, we focus on the change in MDL caused by each proposal, denoted as  $\Delta MDL$  and defined in Equation [3.](#page-5-0) This equation quantifies the difference in the log posterior probability functions for the graph  $G$  by comparing the blockmodel  $B'$  of the new partition with the blockmodel  $B$  of the current partition.

<span id="page-5-0"></span>
$$
\Delta MDL = P(G|B') - P(G|B)
$$
\n(3)

During the block-merge phase, the primary change between the original and new partition occurs when a proposed block merges with another block. However, the connections among other blocks remain unchanged. To avoid unnecessary calculations of  $\triangle MDL$ , it is sufficient to focus only on the incoming and outgoing edges of the merged block  $b$  and the proposed block  $s$  before and after the merge. To simplify this process, we separate the calculations for incoming and outgoing edges, denoted as  $C^{in}$  and  $C^{out}$ , respectively. This ensures that each set is evaluated independently, as outlined in Equation [4.](#page-5-1)

<span id="page-5-1"></span>
$$
\Delta MDL = \left(C_{b'}^{out} + C_{b'}^{in}\right) - \left(C_b^{out} + C_b^{in}\right)
$$
  
where  $b' = b \cup s$  (4)

To calculate  $C^{in}b$  and  $C^{out}b$  for the original block  $b$ , we access the in-degrees  $D_i^{in}$  and out-degrees  $D_j^{out}$  of  $b$ 's adjacent blocks from adjNbr. We also retrieve the weights of the incoming  $M_{i,b}$ and outgoing  $M_{b,j}$  edges from adjWgt, as outlined in Equation [5.](#page-5-2) To leverage GPU parallelism, we employ a segmented reduction operation across the current blockmodel, allowing for the parallel computation of  $C_b^{in}$  and  $C_b^{out}$ . To avoid the duplicated computation of self-connectivity in Equation [5,](#page-5-2) we must subtract the result of this self-connectivity to maintain the correctness.

<span id="page-5-2"></span>
$$
C_b^{in} = \sum_{i,b} M_{i,b} \cdot \log \left( \frac{M_{i,b}}{D_i^{in} D_b^{out}} \right)
$$
  

$$
C_b^{out} = \sum_{b,j} M_{b,j} \cdot \log \left( \frac{M_{b,j}}{D_b^{in} D_j^{out}} \right)
$$
 (5)

To compute  $C_{b'}^{in}$  and  $C_{b'}^{out}$  for  $b'$ , a new block formed by merging block  $s$  into block  $b$ , it is necessary to verify if  $s$  and  $b$  share the same adjacent blocks. If they do, the connected edges must be integrated, and the total weight of these edges summed up, as detailed in Equation [6.](#page-5-3) We implement this operation on GPU, where each thread performs a serial merge operation and accumulates the result, as illustrated in Figure [5.](#page-5-4)

<span id="page-5-3"></span>
$$
C_{b'}^{in} = \sum_{i,b} \left( M_{i,b} + M_{i,s} \right) \cdot \log \left( \frac{\left( M_{i,b} + M_{i,s} \right)}{D_i^{in} \left( D_b^{out} + D_s^{out} \right)} \right)
$$
  

$$
C_{b'}^{out} = \sum_{b,j} \left( M_{b,j} + M_{s,j} \right) \cdot \log \left( \frac{\left( M_{b,j} + M_{s,j} \right)}{\left( D_b^{in} + D_s^{in} \right) D_j^{out}} \right)
$$
  
where  $b' = b \cup s$  (6)

The computations of  $C^{in}b$ ,  $C^{out}b$ ,  $C^{in}_{b'}$  and  $C^{out}_{b'}$  are all independent and can be performed across different blocks without interference. This independence allows us to utilize GPU parallelism to significantly speed up these computations. Once the calculations are completed, the  $\triangle MDL$  for each block can be computed in parallel. Subsequently, we identify the best proposals for each block based on  $\Delta MDL$  and then sort them in ascending order. The results are transferred back to the CPU to perform the merge operation, wherein a portion of blocks to be merged is selected based on the num blocks reduction rate.

<span id="page-5-4"></span>

Figure 5: Illustration of how GSAP computes ΔMDL using segmented reduction and merge operations on GPU. (a) Calculating  $C_b^{out}$  for each block before merging. (b) Calculating  $C_{b'}^{out}$  for each block after merging.

In the vertex-move phase, the calculation procedures for  $C_b^{in}$ and  $C_b^{out}$  remain consistent with those in the block-merge phase. However, the calculation of ΔMDL needs to be modified due to changes in connections between block  $b$  and block  $s$  resulting from the movement of a vertex  $v$  from block  $b$  to block  $s$ . The blocks of  $v$ 's adjacent vertices must switch their connections from  $b$  to s. For block  $b$ , this involves removing edges that connected to the blocks of  $v$ 's adjacent vertices. For block  $s$ , it necessitates adding these edges or integrating them if they already exist in block s. This is accomplished by performing serial merge operations for the block

b and s. The revised  $\Delta MDL$  for the vertex-move phase is detailed in Equation [7.](#page-6-0)

<span id="page-6-0"></span>
$$
\Delta MDL = \left(C_{b'}^{out} + C_{b'}^{in} + C_{s'}^{out} + C_{s'}^{in}\right) - \left(C_{b}^{out} + C_{b}^{in} + C_{s}^{out} + C_{s}^{in}\right)
$$
  
where  $b' = b - \{v\}$  and  $s' = s + \{v\}$  (7)

## 3.4 Blockmodel Update

The blockmodel matrix is frequently updated during the blockmerge and vertex-move phases in the SBP algorithm. For example, as shown in Figure [6,](#page-6-1) if block 0 is merged into block 1 or vertex 0 moves to block 0, the blockmodel matrix must be updated. To update the blockmodel matrix, which is represented as a conventional twodimensional matrix, the random access method can be employed. In contrast, this method is not applicable to a blockmodel matrix in CSR format. In addition, the blockmodel matrix requires frequent updates due to the MCMC iterations in the vertex-move phase, which constitute the majority of the runtime in the SBP algorithm. This limitation highlights the critical need for dynamic management and updating of the blockmodel on GPU. Therefore, we propose a blockmodel update algorithm on GPU to enhance computational efficiency and address the challenges posed by the existing data structures.

<span id="page-6-1"></span>

Figure 6: An example of when should update the blockmodel matrix. (a) Block 0 is merged into Block 1. (b) Vertex 0 moves to Block 0. In both cases, the blockmodel matrix must be updated accordingly.

The process of blockmodel update in GSAP begins with the graph's adjacency lists in CSR format, as shown in Step 1 of Figure [7.](#page-6-2) The second step sorts vertices according to their block IDs, stored in Bmap, using the sort\_by\_key operation. Subsequently, we identify each vertex's neighbors and corresponding edges, which are available in Graph\_adjNbr and Graph\_adjWgt, and store them in the nbr and nbrWgt arrays. The next step is to map each neighbor in the nbr array to its corresponding block ID using Bmap, and then performing the segmented sort operations on the pairs of the nbr and nbrWgt. During this stage, we detect the starting position of

each subsegment for each block section. The results of these positions are stored in subseg\_hd, which is then used to perform the reduction operation and record the value of adjPtr for each block. This is accomplished by utilizing CUDA warp shuffle instructions to compare data across GPU threads. The final stage consists of a segment reduction operation to produce the adjNbr and adjWgt, which form the CSR-formatted blockmodel. We derive the adjPtr by applying an exclusive scan operation to the subseg\_hd array. The complete method is outlined in Algorithm [2.](#page-6-3)

<span id="page-6-2"></span>

Figure 7: A detailed illustration of the blockmodel update approach based on the example in Figure [6.](#page-6-1) The final blockmodel in CSR format matches the matrix in Figure [6.](#page-6-1)

Algorithm 2: Blockmodel Update

- Input: CSR arrays of the input graph G and Bmap Output: CSR arrays of the current partition
- 1 V\_map = sort\_by\_key({0, 1, ...,  $|V| 1$ }, Bmap);
- <sup>2</sup> nbr = Graph\_adjNbr[Graph\_adjPtr[Vmap]];
- <sup>3</sup> nbrWgt = Graph\_adjWgt[Graph\_adjPtr[Vmap]];
- 4 find(nbr, Bmap); // Find the block ID for nbr.
- <sup>5</sup> segsort(nbr, nbrWgt);
- 6 subseg\_hds =  $find$  subseg  $hds(hbr)$ ; // Compare adjacent elements to identify the starting index of each subsegment.
- $\tau$  adjPtr =  $\text{prefix\_scan}(\text{subseg\_hds});$
- <span id="page-6-3"></span> $s$  adjWgt, adjNbr =  $segreduce(nbr, nhrWgt);$

## 4 EXPERIMENTAL RESULTS

We evaluate the performance of GSAP on the SBPC dataset in Table [1.](#page-2-0) GSAP is implemented using C++17 and NVIDIA CUDA 12.2 and compiled with the nvcc compiler on a host compiler of GNU GCC-12.3.0 with the -O3 optimization flag enabled. All experiments were carried out on a Ubuntu Linux 6.1.0-1022-oem x86\_64 single-node machine equipped with a 14-core (20 CPU threads) 13th Gen Intel(R) Core(TM) i5-13500 operating at 2.5 GHz, 128 GB RAM, and one RTX A4000 16 GB GPU.

#### 4.1 Baseline

We consider uSAP [\[2\]](#page-9-1) and I-SBP [\[72\]](#page-10-13) as baselines, which were previous winners in the 2023 SBPC. uSAP is implemented in Taskflow [\[32\]](#page-10-14) to achieve CPU parallelism, while I-SBP is implemented using OpenMP. uSAP and I-SBP are compiled with GNU GCC-12.3.0 and executed using 20 CPU threads, which is the maximum hardware concurrency supported by our system. Table [2](#page-7-0) lists the parameters used for SBP in uSAP, I-SBP, and GSAP.

<span id="page-7-0"></span>

Parameters	Values
num blocks reduction rate	0.4
num_proposals	10
max_num_nodal_itr	100
delta_entropy_threshold1	0.0005
delta_entropy_threshold2	0.0001
delta_entropy_moving_avg_window	3
num_batches_for_MCMC	

Table 2: Partitioning parameters used by I-SBP, uSAP, and GSAP.

#### 4.2 Overall Runtime Comparison

Table [3](#page-8-0) compares the runtime among uSAP, I-SBP (using 14 CPU cores), and GSAP (using one A4000 GPU) on the SBPC dataset. The results show that GSAP significantly outperforms both uSAP and I-SBP on nearly all graphs, except for the smallest graph containing 1K vertices. For graphs with 1K vertices, GSAP is slower than uSAP due to the overhead associated with memory allocation on GPU and data transfer between CPU and GPU. Additionally, the GPU requires setup time to initiate kernels and configure the execution environment. These overheads dominate the runtime of GSAP for the smallest graphs, leading to slower runtime. However, as graph size increases, the advantages of GSAP become more remarkable. For the largest graphs containing 1M vertices, GSAP achieves optimal partitions in about 15 minutes, whereas I-SBP and uSAP could not complete within 2 hours.

<span id="page-7-1"></span>

#### Figure 8: Runtime speedup of GSAP compared to uSAP and I-SBP on the SBPC dataset.

Figure [8](#page-7-1) illustrates the runtime speedup of GSAP over uSAP and I-SBP on graphs ranging from 5K to 200K vertices. GSAP achieves a 12.3× speedup over uSAP under the 200K-vertex graph in the high-low category and a 60.9× speedup over I-SBP under the 50Kvertex graph in the high-high category (hardest). For the remaining graphs, GSAP is 4.5× and 14.2× faster than uSAP and I-SBP on average. The runtime advantage of GSAP comes from our stochastic proposal generation approach, parallel computation of MDL, and the blockmodel update algorithm. Figure [9](#page-7-2) compares the runtime among GSAP, uSAP, and I-SBP in the low-low category. The runtimes for uSAP and I-SBP when processing graphs with 1M vertices are not reported, as neither algorithm completed the task within two hours.

<span id="page-7-2"></span>

Figure 9: Runtime comparison on the low-low category.

Figure [9](#page-7-2) demonstrates GSAP's superior runtime performance, which is attributed to the GPU's order-of-magnitude greater parallelism compared to the CPU-parallel approaches used by uSAP and I-SBP. The advantage of GSAP grows with larger edge counts, highlighting its enhanced scalability. The limitations of the CPUparallel approaches in uSAP and I-SBP are primarily due to the iterative proposal-making process and the MDL calculation, where each block or vertex must account for every edge connected to its neighbors. This iteration over the entire graph significantly worsens the runtime for uSAP and I-SBP. In contrast, GSAP leverages the GPU's capabilities to process these tasks in parallel, substantially reducing runtime.

# 4.3 Runtime Breakdown

Figure [10](#page-8-1) provides a runtime breakdown of uSAP, I-SBP, and GSAP across 50K- and 200K-vertex graphs. We observe that the runtimes for the block-merge phase in GSAP account for up to 2%, while uSAP and I-SBP reach 4.2% and 7.7%, respectively. For the 200kvertex graph in the high-low category, the vertex-move phase in GSAP constitutes only 86% of the runtime, compared to at least 95% for uSAP and 92.3% for I-SBP. These results demonstrate that GSAP significantly reduces the time spent in both phases. GSAP does not optimize the golden-section search phase because it achieves better runtime performance on CPU.

Figure [11](#page-8-2) shows the average runtimes for each proposal during the block-merge and vertex-move phases. This demonstrates the effectiveness of our parallel algorithm in generating stochastic proposals on GPU. On the Low-High 200K-vertex graph, the average runtime of GSAP is 19.6× and 210.3× faster than uSAP and I-SBP, respectively. This improvement is due to our parallel algorithm

<span id="page-8-0"></span>

<span id="page-8-3"></span>Table 3: Runtime comparison among uSAP, I-SBP (using 14 CPU cores), and GSAP (using one NVIDIA A4000 GPU) on the SBPC dataset. 'Failed' indicates the algorithm could not reach the MDL.





<span id="page-8-1"></span>

Figure 10: Runtime breakdown of uSAP, I-SBP, and GSAP.

with the pre-generated tables for stochastic proposals and GPU parallelism allows us to generate proposals concurrently. However, each proposal in uSAP and I-SBP requires examining the number of neighbors for each block or vertex and then proposing stochastic candidates, which is very time-consuming.

Figure [12](#page-9-4) demonstrates the effectiveness of GSAP's blockmodel matrix update approach by comparing it with CPU-based updates. We observe that the speedups correlate with the increase in edge count. Notably, GSAP achieves a 31.5× speedup on Low-Low 200Kvertex graph. Iterating over all edges in such large graphs can significantly slow down the runtime on CPU. In contrast, GSAP's proposed blockmodel matrix update approach leverages the GPU's capabilities to perform updates on the CSR-formatted blockmodel matrix in parallel, resulting in substantial runtime improvements.

<span id="page-8-2"></span>

Figure 11: Average runtime of each proposal in the blockmerge phase and the vertex-move phase.

<span id="page-9-4"></span>

Figure 12: Runtime speedup of the proposed GPU-based blockmodel update over CPU.

# 4.4 Partition Quality Comparison

The normalized mutual information (NMI) is commonly used to evaluate the quality of partitioning results by comparing them with the ground truth [\[44\]](#page-10-9). An NMI value closer to one indicates a perfect correlation between the result and the ground truth, while a value closer to zero indicates no correlation. As shown in Table [4,](#page-8-3) GSAP generally outperforms uSAP and I-SBP across all graph categories. Specifically, GSAP achieves the highest or nearly the highest NMI scores in most cases for both 1K- and 50K-vertex graphs. For the 50K-vertex graph in the high-low category, GSAP scores 0.78, which is significantly higher than the lowest score of 0.33 achieved by I-SBP in the high-high category for 50K graphs. This implies that despite a drop in performance in this scenario, GSAP still significantly outperforms I-SBP. GSAP's effective performance is attributed to maintaining the original structure of the SBP algorithm, while introducing a parallel version of SBP that adheres to the foundational concepts as proposed in [\[66](#page-10-6)[–68\]](#page-10-7).

#### 5 CONCLUSION

In this paper, we have introduced GSAP, a GPU-accelerated stochastic graph partitioner, that significantly enhances runtime performance of SBP. Our experimental evaluations on the 2022 GraphChallenge dataset show that GSAP achieves up to 12.3× and 60.9× runtime speedup on a single A4000 GPU compared to the stateof-the-art algorithms on 14 CPU cores. Inspired by the success of GPU computing in graph processing [\[1](#page-9-5)[–17,](#page-9-6) [19](#page-9-7)[–26,](#page-9-8) [28–](#page-10-18)[31,](#page-10-19) [34–](#page-10-20) [43,](#page-10-21) [46–](#page-10-22)[49,](#page-10-2) [51](#page-10-23)[–63,](#page-10-24) [74–](#page-10-25)[76\]](#page-10-26), our future work plans to incorporate GPU task parallelism using the CUDA Graph to reduce the overhead associated with launching CUDA kernels for larger graphs.

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