Predicting the Output Structure of Sparse Matrix Multiplication with Sampled Compression Ratio

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Abstract—Sparse general matrix multiplication (SpGEMM) is a fundamental building block in numerous scientific applications. One critical task of SpGEMM is to compute or predict the structure of the output matrix (i.e., the number of nonzero elements per output row) for efficient memory allocation and load balance, which impact the overall performance of SpGEMM. Existing work either precisely calculates the output structure or adopts upper-bound or sampling-based methods to predict the output structure. However, these methods either take much execution time or are not accurate enough. In this paper, we propose a novel sampling-based method with better accuracy and low costs compared to the existing sampling-based method. The proposed method first predicts the compression ratio of SpGEMM by leveraging the number of intermediate products (denoted as FLOP) and the number of nonzero elements (denoted as NNZ) of the same sampled result matrix. And then, the predicted output structure is obtained by dividing the FLOP per output row by the predicted compression ratio. We also propose a reference design of the existing sampling-based method targeting high prediction accuracy and low costs compared to the existing sampling-based method. We construct 623 test cases with various matrix dimensions and sparse structures to evaluate the prediction accuracy. Experimental results show that the proposed method achieves an average absolute relative error of 1.30% and 7.93%, respectively, on average, and 25% and 158%, respectively, in the worst case.

Index Terms—Sparse matrix multiplication, SpGEMM, predicting output structure, nonzero structure, size estimation

I. INTRODUCTION

Sparse general matrix multiplication (SpGEMM) is a fundamental building block in numerous scientific and machining learning applications such as Markov clustering [1], algebraic multigrid solvers [2], [3], molecular dynamics simulations [4], multi-source breadth first search [5], and finite element simulations based on domain decomposition [6].

Given two sparse matrices $A$ and $B$, SpGEMM computes the matrix multiplication $C = AB$, where $C$ is the sparse output matrix. To reduce the memory footprint and computation complexity when performing SpGEMM, all the input and output sparse matrices should be stored in a sparse matrix format. However, processing the sparse matrices with a sparse matrix format causes many performance issues due to the irregularities of the three involved matrices and the unknown structure of the output matrix.

Two essential aspects critical to SpGEMM’s performance are the memory allocation method [7], [8] for the output matrix and the load balance method [9]–[11] when performing SpGEMM. Both performance issues are highly related to the output structure (i.e., the number of nonzero elements of each output row) of SpGEMM and how the output structure is computed [7]–[11].

Existing methods to compute the output structure include the precise method [7], [10]–[12], the upper-bound method [7], [8], [13], [14], and the sampling-based method [9], [15]. The precise method computes the exact output structure, which is called the symbolic phase, before performing the actual numeric matrix multiplication [10]. The benefit is that the SpGEMM library adopting the precise method does not need to allocate the intermediate result matrix. However, the major problem with the precise method is that the computation complexity of the symbolic phase is similar to that of the numeric phase [7], [10], [11]. As a result, the symbolic phase (precise method) takes non-trivial computing overheads.

In contrast, the upper-bound method computes the number of necessary intermediate products (denoted as FLOP) per output row as the output structure, which is a low-cost computation step [7], [8]. However, the upper-bound method may allocate much memory space for the intermediate result matrix. For example, for a SpGEMM task with a compression ratio of 10, the memory footprint of the intermediate result matrix is 10× that of the actual result matrix. The compression ratio of a SpGEMM task (or simply of a result matrix) is defined by dividing the total FLOP to perform SpGEMM by the total number of nonzero elements (denoted as NNZ) of the result matrix.

Due to these limitations of the aforementioned methods, in this paper, we focus on developing a novel sampling-based method targeting high prediction accuracy and low computing cost. The existing sampling-based method first randomly samples two sub-matrices of the two input matrices and computes the NNZ of the sampled result matrix (denoted as sampled NNZ). It then divides the sampled NNZ by $p$ to predict the NNZ of the result matrix, where $p$ represents the matrix size proportion of the sampled result matrix to the entire result matrix.

To predict the output structure of SpGEMM, the existing sampling-based method has to compute the precise FLOP per
output row and the total FLOP of the result matrix. And then, the predicted compression ratio of the result matrix is computed by dividing the total FLOP by the predicted total NNZ of the result matrix. At last, the output structure is predicted by dividing the FLOP per output row by the predicted compression ratio. Since the FLOP per output row and the total FLOP are precisely calculated, the accuracy of the predicted output structure, the predicted compression ratio, and the predicted NNZ of the result matrix can be seen as equivalent.

An intuition is that when the NNZ of a sampled result matrix is larger than its expectation, the FLOP of the same sampled result matrix (denoted as sampled FLOP) may also be larger than its expectation. Note that the expectations of the sampled NNZ or sampled FLOP are the precise NNZ($C$) or FLOP($C$) multiplied by $p$, where $p$ represents the proportion of the matrix size of the sampled result matrix to the entire result matrix. In other words, the relative errors of the sampled NNZ and sampled FLOP compared to their expectations may have a positive correlation.

Based on the aforementioned intuition, we propose a novel sampling-based method, which exploits the potential positive correlation of the sampled NNZ and the sampled FLOP of the same sampled result matrix. The proposed method divides the sampled FLOP by the sampled NNZ to obtain a sampled compression ratio as the predicted compression ratio of the result matrix. By doing so, the prediction of the compression ratio may achieve a certain degree of error neutralization between the sampled FLOP and sampled NNZ. For example, if the relative errors of the sampled FLOP and sampled NNZ are $25\%$ and $30\%$, respectively, the relative error of the predicted compression ratio will be $3.85\%$ (analyzed in Section IV-D). At last, the predicted output structure is easily computed by dividing the FLOP per output row by the predicted compression ratio.

As for the computing overheads, the existing sampling-based method uses the inner-product dataflow [9] to compute the sampled matrices, which is less efficient than the row-wise dataflow [16], [17] on both the computing and sampling of the two input matrices. Therefore, in this work, we implement the proposed method using the row-wise dataflow. To fairly compare the prediction accuracy of the proposed method and the existing method, we also implement a reference design of the existing sampling-based method.

We select 25 representative real-world sparse matrices from the SuiteSparse [18] dataset and compose 623 test cases by multiplying them with each other to evaluate the prediction accuracy of the predicted NNZ($C$). The relative prediction errors of the proposed method and the reference design are $1.30\%$ and $7.93\%$ (smaller is better), respectively, on average, and $25\%$ and $158\%$, respectively, in the worst case. We also conduct experiments to show that the parallel implementation of the proposed method only takes on average $0.72\%$ execution time of an entire state-of-the-art SpGEMM library (BRMerge-Precise) [16].

The main contributions of this work are listed as follows:

- We propose a novel sampling-based method that utilizes both the sampled FLOP and sampled NNZ to predict the output structure of SpGEMM.
- We propose a row-wise implementation of the proposed method and a reference design of the existing sampling-based method.
- We conduct comprehensive evaluations to show that the proposed method is much more accurate than the reference design of the existing method. We also conduct experiments to show that the parallel implementation of the proposed method only takes on average $0.72\%$ execution time of an entire state-of-the-art SpGEMM library (BRMerge-Precise) [16].

The rest of this paper is organized as follows. Section II introduces the notations and backgrounds. Section III discusses the related work. Section IV describes the proposed prediction method and the reference design of the existing prediction method. Section V details the efficient parallel implementation of the proposed method. Section VI shows the evaluation accuracy of the proposed method and the reference design. This section also shows the computing overheads of the proposed method. Section VII concludes this paper.

The source code of this paper is available at https://github.com/lorentzbf/Size-Prediction.git.

II. PRELIMINARIES AND BACKGROUNDS

A. Notations

Table I defines the notations used in this paper. The matrix dimensions of the three matrices $A$, $B$, and $C$ are $M \times K$, $K \times N$, and $M \times N$, respectively.

| Notation | Explanation |
|----------|-------------|
| $A$ | The first input matrix |
| $B$ | The second input matrix |
| $C$ | The result matrix |
| NNZ($C$) | Number of nonzero elements |
| FLOP($C$) | Number of intermediate products |
| CR | Compression ratio, defined as FLOP/NNZ |

TABLE I: Notations used in this paper.

B. CSR Storage Format

CSR (Compressed Sparse Row) storage format stores the sparse matrix in a compressed way. Fig. 1 illustrates the CSR storage format, which consists of three arrays named $rpt$, $col$, and $val$. The $val$ and $col$ arrays record the nonzero elements and their corresponding column indices in a sorted row-major and column-major order. The $rpt$ array records the start and end offsets for each row’s values and column indices in the $val$ and $col$ arrays.

C. Row-wise Dataflow

The row-wise dataflow is described by (1), which shows the computation of the $i_{th}$ output row ($C_{i\ast}$).

$$C_{i\ast} = \sum_{k} A_{ik} \cdot B_{k\ast}, \quad (1)$$
where \( C_{ix} \) and \( A_{ix} \) represent all the nonzero elements in the \( i_{th} \) row of \( C \) and \( A \), respectively, \( k \) belongs to the set of column indices of \( A_{ix} \), and \( B_{kx} \) represents all the nonzero elements in the \( k_{th} \) row of \( B \).

### III. Related Work

We first describe a highly related prediction method proposed in the database literature. And then, we describe two sampling-based methods that predict the output structure of SpGEMM. At last, we describe several SpGEMM libraries that utilize the output structure for memory allocation and load balance issues.

Bar-Yossef et al. [19] proposed a method to predict the number of distinct elements in a data stream. Let \( a = a_1, a_2, ..., a_n \) be an array with \( n \) elements, where \( a_i \in [0, m] \). Certain elements in \( a \) may have the same value. Let \( F = F(a) \), which represents the number of distinct elements in \( a \). To predict \( F \), this method first constructs a hash function \( h : [m] \rightarrow [0, 1] \).

And then, the method applies \( h(a) \) to all elements in \( a \) and maintain the smallest hashed value as \( v \). At last, the predicted number of distinct elements in \( a \) is computed by \( F^* = 1/v \).

The intuition here is that if the hashed values of all the elements in \( a \) are randomly distributed in \([0, 1]\), the smallest hashed value will be \( 1/F \). This prediction method is utilized in the following methods that predict the output structure of SpGEMM [9], [15]. The benefit of this method is minimum memory space usage when processing the data in \( a \) since it only maintains the smallest element. However, the time complexity is not necessarily low since all the elements in \( a \) are processed.

Amossen et al. [15] proposed a method to predict the total NNZ of the result matrix of SpGEMM, which is mainly based on the idea in [19]. This method constructs a hash function \( h : [m, n] \rightarrow [0, 1] \) and samples random subsets from the two input matrices as \( A' \) and \( B' \). It applies all the row and column indices of the intermediate products when computing \( C' = A'B' \) to the hash function \( h \). The \( k_{th} \) smallest element of the hashed values is maintained as \( v \). After this process, the predicted NNZ(\( C' \)) is computed by \( k/v \) and the predicted NNZ(\( C \)) is computed by \( k/v/p \), where \( p \) stands for the matrix size proportion of \( C' \) compared to \( C \).

Pham et al. [9] proposed a SpGEMM library BHash which implements Amossen’s prediction method. Specifically, Pham’s method randomly samples the rows from \( A \) as \( A' \) with the probability \( p = 1/10 \) and randomly samples columns from \( B \) as \( B' \) with the same probability. It then use the same method in [15] to predict NNZ(\( C' \)), where \( C' = A'B' \). At last, the predicted NNZ(\( C \)) is calculated by \( NNZ(C')/p^2 \).

The predicted NNZ(\( C \)) and the predicted output structure are used for the load balance of the computing tasks and memory allocation for the result matrix in BHash [9].

Most of the existing SpGEMM libraries only use the precise or upper-bound methods to compute/predict the output structure [7], [8], [10], [11], [13], [14], [20], [21]. For example, bhsparse [8] computes the FLOP per output row as the upper-bound output structure and classifies the rows with different FLOP to different bins. Rows in different bins are computed by different accumulation methods for better load balance. The memory allocation of bhsparse is also based on the upper-bound output structure. The nsparse [10] computes the upper-bound output structure and uses this information for the load balance of its symbolic phase, which computes the precise output structure.

### IV. Proposed Method

This section describes the proposed sampling-based method and the reference design of the existing sampling-based method. The proposed method contains three progressive optimizations compared to the existing method: 1) adopting the row-wise dataflow with the associated sampling method, 2) calculating the precise NNZ of the sampled result matrix, 3) predicting the NNZ of the result matrix by using the sampled FLOP and sampled NNZ. To fairly compare the prediction accuracy of the proposed method and the existing method, we implement the reference design of the existing method by adopting the first two optimizations.

#### A. Computing Dataflow and Sampling Method

The existing methods select both rows and columns from the two input matrices [9], [15]. However, the two input matrices are usually stored in homogeneous storage format [8], [10]. For example, both input matrices may be stored in the CSR format, which is hard to select the columns from the matrix. Furthermore, even if the rows and columns can be selected with a low cost, the following inner-product dataflow may still be a performance bottleneck [16], [17]. To tackle these two performance issues, we adopt the row-wise dataflow for the sampling method and the computation of the samples.

The associated sampling method randomly samples fraction of rows from the \( A \) matrix as \( A' \), where \( p \) is usually a small value for a low cost. The method does not need to sample the \( B \) matrix since \( B \) is accessed according to the selected rows of \( A \) (see Section II-C). This can also be seen as the entire \( B \) matrix is sampled.

#### B. Computing Method for the Samples

The existing method does not compute the precise NNZ of the samples, but predicts it by utilizing a hash function \( h : [m, n] \rightarrow [0, 1] \) [9], [15]. The accuracy of the existing method is highly related to the hash function, which is difficult.
to be constructed. Moreover, the computation complexity of the existing method is similar to that of precisely computing the NNZ of the samples. The reason is that the computing bottlenecks of both methods lie in processing all the intermediate products of the samples. Therefore, we propose to directly compute the precise NNZ of the samples, which is automatically more accurate in computing the NNZ of the samples than the existing sampling-based method. We describe the parallel implementation of computing the precise NNZ of the samples in section V-B.

C. Reference Design

The reference design adopts the two techniques mentioned above. We then describe how the reference design predicts the output structure in more detail. We also analyze the relative error of the reference design.

We denote the computed NNZ of the samples as $z^*$, the expected NNZ of the samples as $z$, the predicted NNZ(C) as $Z^*$, and the precise NNZ(C) as $Z$. Note that throughout this paper, we use the additional subscript $1$ to denote the variables computed by the reference design. The expected NNZ of the samples $z$ is defined as $z = pZ$, where $p$ represents the matrix size proportion of the sampled result matrix compared to the entire result matrix. For example, if the number of rows of the $A$ matrix is 1000 and the number of sampled rows from the $A$ matrix is 3, $p$ will be equal to 0.003. Similar to the previous work [9], [15], the reference design predicts the NNZ(C) by $z^*/p$. We define the relative error of the $z^*$ to the expected $z$ as $\epsilon_1$. Then the entire prediction method and the error analysis of the reference design are shown in (2).

\[
Z^* = \frac{1}{p}z^* = \frac{1}{p}(1 + \epsilon_1)z = (1 + \epsilon_1)Z. \tag{2}
\]

To predict the output structure of the result matrix, the reference design (as well as the existing methods [9], [15]) has to compute the FLOP per output row and the total FLOP of the result matrix (denoted as $F$). And then, the predicted compression ratio is computed by $F/Z^*$, and the output structure is computed by dividing the FLOP per output row by the predicted compression ratio.

D. Proposed Prediction Method

The proposed prediction method utilizes the same information computed by the reference design, which includes the computed FLOP per output row and the NNZ of the samples. In addition, the proposed method also computes the FLOP of the samples. The cost of computing the FLOP of the samples is negligible after the FLOP per output row has been computed.

Although the FLOP per output row and the FLOP(C) are precisely computed, we can still predict the FLOP(C) in a symmetric way as how the reference design predicts the NNZ(C) and establish the error analysis of the predicted FLOP(C). Similar to the notations for the reference design, we denote the computed FLOP of the samples as $f^*$, the expected FLOP of the samples as $f$, the predicted FLOP(C) as $F^*$, and the actual FLOP(C) as $F$. The expected FLOP of the samples $f$ is defined as $f = pF$, where $p$ represents the matrix size proportion of the sampled result matrix compared to the entire result matrix. We define the relative error of the $f^*$ to the expected $f$ as $\epsilon_f$. Then the entire prediction method and the error analysis for $F$ are shown in (3).

\[
F^* = \frac{1}{p}f^* = \frac{1}{p}(1 + \epsilon_f)f = (1 + \epsilon_1)F. \tag{3}
\]

An intuition is that for the same sampled result matrix, if the computed $z^*$ is greater than the expected $z$, the computed $f^*$ will be more likely to be greater than the expected $f$ rather than smaller than the expected $f$. In other words, the relative error of $z^*$ may have a positive correlation to the relative error of $f^*$ for the same samples.

Based on this intuition, we propose a novel prediction method as (4). The proposed method first computes the precise FLOP and NNZ of the same samples (denoted as $f^*$ and $z^*$, respectively) so that the two variables may keep a certain degree of positive correlation. And then, the proposed method computes the predicted compression ratio as $r^* = f^*/z^*$ at last, the predicted NNZ(C) is computed by dividing the actual FLOP(C) by the predicted compression ratio. The output structure is computed by dividing the FLOP per output row by the predicted compression ratio.

\[
Z^*_2 = \frac{F}{F^*} = \frac{F}{f^*}z^*_1 = \frac{F}{(1 + \epsilon_f)f}(1 + \epsilon_1)z = \frac{1 + \epsilon_1}{1 + \epsilon_f}p = \frac{1 + \epsilon_1 - \epsilon_f}{1 + \epsilon_f}Z, \tag{4}
\]

where $Z^*_2$ is the predicted NNZ(C) by the proposed method.

The relative error of the proposed method for predicting NNZ(C) is:

\[
\epsilon_2 = \frac{\epsilon_1 - \epsilon_f}{1 + \epsilon_f}. \tag{5}
\]

Based on (5), the relative error of the proposed method is close to the difference between $\epsilon_1$ and $\epsilon_f$ since the denominator is usually close to 1. Therefore, the proposed method may achieve good error neutralization between $\epsilon_1$ and $\epsilon_f$ if the two relative errors are positively correlated and close to each other. Specifically, if $\epsilon_1$ and $\epsilon_f$ are both positive values and $\epsilon_f \in [0, 2\epsilon_1]$, the relative error of $\epsilon_2$ will be smaller than $\epsilon_1$, which is our expectation. Similarly, if $\epsilon_1$ and $\epsilon_f$ are both negative values and $\epsilon_f \in [2\epsilon_1, 0]$, the relative error of $\epsilon_2$ will also be “smaller” than $\epsilon_1$; the “smaller” here means more close to 0. In a best case, if $\epsilon_f$ approaches $\epsilon_1$, $\epsilon_2$ will approach 0.

V. PARALLEL IMPLEMENTATION

This section describes the implementations of two performance-critical tasks used by the proposed method and the reference design. The goal of the parallel implementations is to achieve low computing overheads. We target the implementation at the multi-core CPUs instead of GPUs since these computing tasks are relatively small and irregular, which may not efficiently utilize the massively parallel computing resources on GPUs. We implement the parallel algorithms with the OpenMP framework [22].
A. Computing the FLOP per Output Row

The first performance-critical task is to compute the FLOP per output row. Algorithm 1 shows the parallel implementation of this task, where the floprC denotes the actual FLOP per output row, and the total_flop denotes the actual FLOP(C). The FLOP per output row is the upper bounds of the number of nonzero elements per output row. The computation complexity of this algorithm is relatively small since only the row offsets and column indices of the A matrix and the row offsets of the B matrix are processed. We parallelize this algorithm by statically assigning the same number of rows to each CPU thread. The total_flop (Line 8) is computed within a critical section provided by OpenMP [22].

Algorithm 1 Compute FLOP(C)

Input: A.rpt, A.col, B.rpt, M.
Output: floprC, total_flop.
1: total_flop = 0
2: for i = 0 to M-1 in parallel do
3:    local_flop = 0
4:    for j = A.rpt[i] to A.rpt[i+1] do
5:        local_flop += B.rpt[A.col[j]+1] - B.rpt[A.col[j]]
6:    end for
7:    floprC[i] = local_flop
8:    total_flop += local_flop
9: end for

B. Computing the Predicted NNZ(C) by the Proposed Method

The second performance-critical computing task is to compute the NNZ of the sampled result matrix. Since the other computation steps are straightforward and with a low cost, Algorithm 2 directly shows how the proposed method computes the predicted NNZ(C), which includes how the sampled NNZ is computed.

The method to compute the NNZ per output row is the same as the existing work [7], which utilizes the hash-based method. Line 12 to Line 28 in Algorithm 2 show the hash-based method. The memory space of the hash table for each CPU thread is set as the largest floprC (Line 6). However, the used memory space of the hash table for each output row is the FLOP of that row (Line 10). Line 9 selects a random row from the A matrix, where the random data is pre-computed and stored in an array named rand (Line 3). Line 7 is parallelized so that each CPU thread computes the same number of sampled rows. As a result, Line 29 and Line 30 are computed in a critical section [22]. In the end, the predicted NNZ(C) is easily computed as shown in Line 32.

In the implementation of both the proposed method and the reference design, we empirically set the number of the sampled rows as sample_num = min(0.003 * M, 300), where M is the number of rows of A. We set the maximum sampled rows as 300 to reduce the computing overheads when M is large. The accuracy loss is often negligible since the prediction accuracy is often very high when M is relatively large.

Algorithm 2 Compute $Z_2^*$

Input: A.rpt, A.col, B.rpt, B.col, floprC, total_flop, M
Output: $Z_2^*$
1: sample_num = min(0.003 * M, 300)
2: rand = new float [sample_num]
3: Generate sample_num random data in the range [0,1] and store them to the rand array
4: Calculate the max floprC as the max hash table size: max_tsize
5: sample_flop = 0, sample_nnz = 0
6: ht = new int [max_tsize] in each CPU thread
7: for r = 0 to sample_num - 1 in parallel do
8:    local_nnz = 0
9:    rid = M * rand[r]
10:   tsize = floprC[rid]
11:   Initialize the front tsize elements in the ht array to -1
12:   for i = A.rpt[rid] to A.rpt[rid+1] do
13:     B_row = A.col[i]
14:     for j = B.rpt[B_row] to B.rpt[B_row + 1] do
15:       hash = (B.col[j] * HASH_SCALE) % tsize
16:       while true do
17:         if ht[hash] == B.col[j] then
18:            break
19:        else if ht[hash] == -1 then
20:            local_nnz += 1
21:            ht[hash] = B.col[j]
22:            break
23:        else
24:            hash = (hash + 1) % tsize
25:        end if
26:     end while
27:   end for
28:   sample_nnz += local_nnz
29: end for
30: sample_flop += floprC[rid]
31: end for
32: $Z_2^* = \text{total_flop} / \text{sample_flop} \times \text{sample_nnz}$

VI. EXPERIMENTS

In this section, we compare the accuracy of the predicted NNZ(C) of the proposed method and the reference design. We also show the prediction overheads of the proposed method compared with a state-of-the-art SpGEMM library BRMerge-Precise [16].

A. Prediction Accuracy

Recall that for both the proposed method and the reference design, the output structure is computed by dividing the FLOP per output row by the predicted compression ratio, where the predicted compression ratio is computed by dividing the total FLOP by the predicted total NNZ. Since the FLOP per output row and the total FLOP are precisely computed, the accuracy
of the predicted output structure, the predicted compression ratio, and the predicted $NNZ(C)$ can be seen as equivalent. For simplicity, we only compare the prediction accuracy of the predicted $NNZ(C)$.

For the diversity of the evaluation, we select 25 representative real-world sparse matrices from the SuiteSparse matrix collection [18]. Table II shows the detailed information of the 25 sparse matrices. We try to multiply these 25 matrices with each other to obtain more test cases with various sparse structures and matrix dimensions. One problem is that the two input matrices may not be multiplied due to mismatched matrix dimensions. To tackle this problem, we reshape either the first or the second input matrix. For example, if the dimensions of the two input matrices are $10 \times 10$ and $5 \times 5$, we reshape the first matrix to a $10 \times 5$ matrix by keeping its left 5 columns. If the dimensions of the two input matrices are $5 \times 5$ and $10 \times 10$, we reshape the second matrix to a $5 \times 10$ matrix by keeping its top 5 rows.

We eliminate the test case of " delaunay_n24" × "poisson3Da" since the reshaped " delaunay_n24" matrix is a hypersparse matrix and the randomly sampled result matrix contains no nonzero elements. We also eliminate the test case of "poisson3Da" × "patents_main" since the reshaped "patents_main" matrix contains no nonzero elements. As a result, we construct 623 matrix multiplication test cases to evaluate the accuracy.

We compare the three relative errors: $\epsilon_1 = (Z_1 - Z)/Z$, $\epsilon_f = (F^* - F)/F$, and $\epsilon_2 = (Z_2 - Z)/Z$, which are described in section IV. Recall that $F^*$ can be precisely computed with low costs, which means we do not need to predict the total $FLOP(C)$ in real cases. We show the relative prediction error of $\epsilon_f$ compared to $\epsilon_1$ to intuitively show their positive correlation.

Experiments on the 623 test cases show that the average absolute relative errors of $\epsilon_1$, $\epsilon_f$, and $\epsilon_2$ are 7.93%, 8.39%, and 1.30%, respectively, the median absolute relative errors of $\epsilon_1$, $\epsilon_f$, and $\epsilon_2$ are 3.04%, 2.90%, and 0.45%, respectively, and the worst absolute relative errors of $\epsilon_1$, $\epsilon_f$, and $\epsilon_2$ are 158%, 155%, and 25%, respectively. Moreover, the proposed method is more accurate than the reference design on 83.0% of the 623 test cases.

The overall results show that the proposed method is much more accurate than the reference design. Considering the difference between the proposed method and the reference design, we can infer that the relative errors of the sampled FLOP and sampled NNZ are positively correlated and close to each other in most cases. Therefore, the proposed method can achieve good error neutralization by dividing the sampled FLOP by the sampled NNZ (see section IV-D).

Moreover, the correlation coefficient [23] of $\epsilon_1$ and $\epsilon_f$ on the 623 test cases is 98.31%, which statistically shows a strong positive correlation between the sampled FLOP and sampled NNZ of the same randomly selected samples.
TABLE III: The relative errors of 20 representative test cases.

| A       | B          | sample_num | CR | NNZ(C) | $\epsilon_1$(%) | $\epsilon_f$(%) | $\epsilon_2$(%) |
|---------|------------|------------|----|--------|-----------------|-----------------|-----------------|
| 1       | 2cubes_sphere | consph     | 300| 1.5    | 64800734       | -2.31           | -2.6            | 0.29            |
| 2       | cage12     | patents_main | 300| 1   | 4611949       | 3.61            | 3.62            | -0.01          |
| 3       | cage15     | majorbasis | 300| 1.1  | 15990225      | 16.38           | 15.73           | 0.56            |
| 4       | delaunay_n24 | maro002   | 300| 1    | 12553686      | -22.49          | -22.81          | 0.42            |
| 5       | delaunay_n24 | cop20k_A  | 300| 1.01  | 15604104      | -45.88          | -46.32          | 0.81            |
| 6       | m133-b3     | rma10      | 300| 1.14  | 8336596       | 4.41            | 5.57            | -1.11          |
| 7       | majorbasis  | 2cubes_sphere | 300| 1.08  | 22688054      | -0.49           | -0.5            | 0.01            |
| 8       | maro002     | webbase-1M  | 300| 1.21  | 6866846       | -11.98          | -16.68          | 5.65            |
| 9       | mc2dept     | poisson3Da | 300| 1.02  | 1366481       | 66.23           | 70.54           | -2.53          |
| 10      | pwtk        | consph     | 300| 5.99  | 54168970      | -11.32          | -9.81           | -1.68          |
| 11      | shipsec1    | rma10      | 300| 4.67  | 27713808      | -7.48           | -5.53           | -2.07          |
| 12      | scircuit    | poisson3Da | 300| 1.04  | 1848459       | 8.76            | 8.06            | 0.65            |
| 13      | scircuit    | mac_econ_fwd500 | 300| 1.11  | 5313337       | 1.44            | 1.36            | 0.08            |
| 14      | rma10       | pdb1HYS    | 140| 8.3   | 23240867      | -3.07           | -2.43           | -0.66          |
| 15      | pwtk        | shipsec1   | 300| 5.41  | 77530890      | -3.59           | -4.29           | 0.73            |
| 16      | cage12      | hood       | 300| 1.23  | 83406736      | -0.31           | 0.2             | -0.51          |
| 17      | 2cubes_sphere | cant       | 300| 1.62  | 40235181      | -0.19           | -3.47           | 3.4            |
| 18      | rma10       | offshore   | 140| 1.53  | 25255211      | -0.01           | 0.28            | -0.29          |
| 19      | filter3D    | filter3D   | 300| 4.26  | 20161619      | 1.74            | 4.47            | -2.62          |
| 20      | hood        | poisson3Da | 300| 1.12  | 17777942      | -0.39           | 0.98            | -1.35          |

To show the better accuracy of the proposed method more intuitively, we show the relative errors of $\epsilon_1$, $\epsilon_f$, and $\epsilon_2$ on 20 representative test cases in Table III. For most test cases, the prediction accuracy of the proposed method is much more accurate than the reference design. For example, the fifth test case in Table III shows that the relative error of predicting the NNZ(C) and FLOP(C) by only using the NNZ or FLOP of the samples are -45.88 and -46.32, respectively. In contrast, the relative error of predicting NNZ(C) by the proposed method is only 0.81. This test case shows significant error neutralization between the sampled FLOP and sampled NNZ. Also note that the relative errors of $\epsilon_1$, $\epsilon_f$, and $\epsilon_2$ of all the test cases exactly meet the equation described by (5).

The last five test cases in Table III show that the signs of $\epsilon_1$ and $\epsilon_f$ are different. We observe that when the signs of $\epsilon_1$ and $\epsilon_f$ are different, the absolute values of $\epsilon_1$ and $\epsilon_f$ are usually close to zero. As a result, the relative error of predicting NNZ(C) by the proposed method is also relatively small in such cases.

B. Computing Overhead

Two performance-critical tasks in the proposed method and the reference design are computing the FLOP per output row (denoted as computing FLOP) and computing the NNZ of the samples (see section V-A and section V-B). Since the computation complexity of computing the sampled NNZ is similar to predicting the NNZ(C) by the proposed method after the FLOP per output row has been computed, we directly show the computing overheads of predicting NNZ(C) by the proposed method (denoted as predicting $Z_2^*$). We compare the relative execution time of the two tasks compared to a state-of-the-art SpGEMM library BRMerge-Precise [16]. The execution time is measured as the average execution time of ten runs after one warm-up run. Fig. 2 shows the relative execution time in percentage on the matrix square benchmark with the 25 sparse matrix in Table II. The average relative execution time of computing FLOP and predicting $Z_2^*$ compared to BRMerge-Precise are 1.68% (up to 4.13%) and 0.72% (up to 1.89%), respectively, which is a relatively small computing overhead.

In most SpGEMM algorithms, computing the FLOP per output row is a necessary task for either the upper-bound allocation method [7], [8], [13], [14] or the load balance of the symbolic phase in the precise method [10], [11]. Therefore, the actual computing overheads of the proposed prediction method only take an average of 0.72% (up to 1.89%) execution time compared to the state-of-the-art SpGEMM library BRMerge-Precise, which is a negligible cost.

VII. Conclusion

Computing or predicting the output structure of SpGEMM is an important task for efficient memory allocation and load balance of SpGEMM, which greatly impacts the overall performance [7]–[10]. In this paper, we propose a novel sampling-based prediction method that utilizes the positive correlation between the sampled FLOP and sampled NNZ of the same samples to achieve a certain degree of error neutralization. The proposed method achieves much more accurate prediction accuracy than the reference design of the existing sampling-based method. For low computing overheads, the proposed method adopts the row-wise dataflow and only samples up.
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Fig. 2: Relative execution time of the two performance-critical tasks compared to BRMerge-Precise.

to 0.003 of the total rows of the first input matrix. We also propose the parallel implementation of the proposed method targeting the multi-core CPUs. The computing overheads of the proposed method only take on average 0.72% execution time compared to the overall execution time of a state-of-the-art SpGEMM library BRMerge-Precise [16].

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