A Memory-Efficient Algorithm and Its Implementation of Variable-Size All-to-All Communication

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This paper proposes a memory-efficient algorithm of variable-size all-to-all communication based on bitonic sort and in-place merge. The algorithm takes $O(N \log^2 P)$ computation and communication time and requires merely $O(P)$ extra space for the communication among $P$ processes having $N$ elements of data to be exchanged for each. We also discuss another algorithm which takes $O(MP \log P)$ time where $M$ is the size of the largest chunk which a process must send to another process and thus is expected to be $O(N/P)$ in most practical applications to make the time complexity $O(N \log P)$. We implemented the first sort-based algorithm to show it works with reasonable efficiency.

1. Introduction

One of important motivations of parallel high-performance computing is, besides obvious expectation of parallel speedup, to enlarge the size of problems to be solved. This means that parallel algorithms must be aware of memory efficiency as well as timing efficiency. In a sense, memory efficiency is more critical than timing efficiency because we may allow 10% execution time extension while 10% memory size excess cannot be acceptable or we have to pay a huge memory/disk swap cost.

Memory efficiency of collective communications sometimes governs that of application programs using them. For example, if a program has a large distributed array which dominates the memory requirement of the program and we need to perform FFT on the array, all-to-all communication for transposition must be performed with memory awareness. Bad news to MPI users is that MPI_Alltoall() does not have in-place option\(^1\) to replace the data in send-buffer with those received. Therefore we need implement a hand-made version of all-to-all to restrict additional space to $O(1)$ or waste memory space as large as the array.

The variable-size variance of all-to-all communication, e.g., MPI_Alltoallv(), is also useful for parallel applications especially for load distribution/exchange among parallel processes. For instance, our particle-in-cell simulator\(^1\) needs an all-to-all distribution of particle subsets to initialize the simulation and/or to correct unacceptable load imbalance. Unfortunately, devising and implementing a memory efficient variable-size all-to-all is not trivial work. That is, in a fixed-size all-to-all, a process $p$ can exchange a fixed-size chunks with another process $q$ to place the chunks to their destinations, or can shift chunks along a rings connecting processes. However, these simple techniques are not applicable for the variable-size version because the pair of the chunks or those on the circular shift path may have arbitrary variable sizes.

The work presented paper aims at devising memory efficient and reasonably time efficient all-to-all algorithms applicable to memory intensive parallel applications including our particle-in-cell simulator by which we are motivated to pursue the work. In the rest of the paper, after defining the problem to be solved, we discuss two algorithms, one based on bitonic sort and the other with pairwise exchange. We also show our experiment results on the first algorithms to evidence its reasonable timing efficiency.

2. Problem Definition and Assumptions

For the sake of explanation simplicity\(^2\), we define our variable-size all-to-all communication problem as follows. Let $s(i,j)$ be a sequence of data elements $a_{i1}^j, \ldots, a_{in}^j$ which a parallel process $i \in [1,P]$ accommodates at initial and a process $j \in [1,P]$ finally stows in its memory. The initial layout of $s(i,1), \ldots, s(i,P)$ in the process $i$ is the concatenation of them in this order denoted by $s(i,1) \bullet \cdots \bullet s(i,P)$. The stable solution for the process $j$ is to have the

\(^1\) Good news is that in-place option is expected to be incorporated a (near) future version of MPI specifications.
\(^2\) Also for the easiness of our implementation, so far.
concatenation $s(1, j) \bullet \cdots \bullet s(P, j) \equiv S(j)$, while unstable one is a permutation of data elements in $S(j)$.

Let $|s(i, j)|$ be the length (or number of data elements) of the sequence $s(i, j)$ and $N$ be that of the initial and final concatenation of the sequences regardless the process accommodating it. That is, for all $i \in [1, P]$, the following is satisfied.

$$
\sum_{k=1}^{P} |s(i, k)| = \sum_{k=1}^{P} |s(k, i)| = N.
$$

The memory overhead of an algorithm is the maximum space required in addition to that for $N$ elements throughout the procedure to solve the problem. It is obviously desirable that the overhead is $O(1)$ but $O(P)$ overhead is acceptable. In fact, since a data element may be anything and thus may tell us neither its initial housing process, its initial location in the process, nor its final destination, it is almost inevitable that an array containing $|s(i, 1)|, \ldots, |s(i, P)|$ is given as the input for the process $i$ besides the sequences $s(i, 1) \bullet \cdots \bullet s(i, P)$, as MPI_Alltoallv() requires an input argument $sendcounts[P]^{*1}$.

The time complexity of an algorithm is determined by two factors, computation cost and communication cost. Computation cost is mainly for memory operations to manipulate sequences and elements in them and, of course, is calculated taking parallel execution into account. As for communication cost, we simply assume that a transmission of $n$ data elements from a process $p$ to a process $q$ takes $O(n)$ time. That is, we neglect the constant overhead for the communication and the latency possibly depending on the distance between them in the process network. We also assume that processes are connected with a non-blocking network through which communications between any number of non-overlapping pairs can be performed simultaneously without any performance interference. Finally, if a time complexity has factors of $N$ and $P$ with the same degree, we neglect $P$ to reduce, for example, $N + P$ to $N$ because $N \gg P$.

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*1 The argument recvcounts is not necessary to be given explicitly. Since we assume the fixed order of initial and final sequences as the fixed-size MPI_Alltoall() does, sdispls and rdispls are also unnecessary. If it is required to specify the permutation of the sequences by the displacement arguments, pre- and post-processing can be implemented using a in-place sort with the in-place merger discussed later.

3. Algorithms

3.1 Fundamental Means

For designing of memory efficient algorithms, we have two fundamental means which requires merely $O(1)$ extra space regardless the size of data to operate on. The first is inter-process swap namely swap$(i, j, s_i, s_j)$ to exchange sequences $s_i$ and $s_j$ of size $n = |s_i| = |s_j|$ between the process pair $i$ and $j$. This operation can be straightforwardly performed by MPI_Sendrecv_replace() which is expected to use a fixed size data buffer to temporarily store sending data before directly receiving data into its sendbuf (or to store receiving data which is then copied to sendbuf). Since we have to repeat send/receive communications if the buffer is smaller than the size of $n$ elements, a large buffer of a few mega-bytes, for example, is desirable but the size is still $O(1)$ and is much smaller than $N$.

The second mean merge$(s_1, s_2)$, which operates on the concatenation of ordered sequences $s = s_1 \bullet s_2$ to merge the sequences replacing them, is more powerful and complicated than the swap. The algorithm, invented by Huang and Langston$^{2}$, takes $O(|s|)$ time while requiring $O(1)$ extra space for stable merge. The outline of the algorithm is as follows.

1. Let $n = |s|$ and split $s$ into $\sqrt{n}$ blocks $b_1, \ldots, b_{\sqrt{n}}$ so that each block has $\sqrt{n}$ elements.

2. Rotate a part of $s$ so that $b_{\sqrt{n}}$ has $\sqrt{n}$ largest elements, and then Rotate $s$ so that $b_{\sqrt{n}}$ is placed leftmost and thus becomes $b_1$. Then sort blocks $b_2, \ldots, b_{\sqrt{n}}$ using the last elements of block as keys by a stable sorting algorithm with $O(1)$ extra space, for example, by selection sort taking
That is, the sorting with ordering above results in the sequence $S(1) \cdots \cdot S(P)$. Since we assume $\sum_{i=1}^{P} |s(i,j)| = N$ for all $j$, each part of the sorted sequence $S(j)$ should reside in the process $j$ if we arrange that $j$ has the $j$-th subsequence $N$ long.

Bitonic sort\(^1\) is one of efficient and easily-parallelizable sorting algorithms. It operates on a bitonic sequence being the concatenation of an ascending sequence and a descending one, or a circular rotation of this fundamental sequence. The lower and upper halves of a bitonic sequence are easily found by scanning its first and second halves. More specifically, for a bitonic sequence $a_1, \ldots, a_{2n}$, the crossing index $i$ such that $a_i < a_{2i} \iff a_{i+1} \geq a_{2i+1}$ tells us the lower (or upper) half is the concatenation of $a_1, \ldots, a_i$ and $a_{n+1}, \ldots, a_{2n}$ while upper (or lower) half is $a_{i+1}, \ldots, a_{2n+i}$, if $a_i < a_{2i}$ (or $a_i > a_{2i}$). Since it is assured that the lower and upper halves are bitonic sequences again, iterative splitting of a bitonic sequence $a_1, \ldots, a_{2n}$ gives us subsequences $s_1, \ldots, s_{2^n}$ such that $|s_i| = n$ for all $i$, and $a_k \leq a_j$ for all $1 \geq i < j \leq 2^k$, $a_k \in s_i$ and $a_j \in s_j$.

Parallel implementation of bitonic sort with $P = 2^p$ processes consists of $p$ phases (calls of procedure bitonic_sort()) as shown in Figure 2 and 3. The $k$-th phase starts from $2^{p-k}$ bitonic sequences each of which lies across $2^k$ processes, then each sequences is split $k$ times so that each process has a bitonic sequence whose ascending and descending subsequences are merged to have a ascending or descending sequence. Note that local_sort() to make ascending (if UP) or descending (if DOWN) is unnecessary for our application to all-to-all because the initial sequence $s(i,1) \cdots \cdot s(i,P)$ is an ascending sequence in our ordering definition\(^2\).

The procedure bitonic_split() is for splitting a bitonic sequence lying across processes $i, \ldots, i + 2^k - 1$ performed by swap$(j, j + 2^{k-1}, s'_1, s'_{j+2^{k-1}})$ for all $j$ such that $i \leq j \leq i + 2^k - 1$, where $s'_1$ is a leading or trailing subsequence of the sequence $s_j$ which accommodates and is defined follows. Let $j'$ such that $i \leq j' \leq i + 2^k - 1$ and the crossing index lies in the process $j'$, $A$ be true iff the crossing index on a ascending sequence, and $L$ be true iff $j$ needs lower half of

\[^{1}\] Therefore our implementation uses $O(\sqrt{n})$ buffer, so far.

\[^{2}\] In our implementation, we omit reversing $s(i,1) \cdots \cdot s(i,P)$ for even number processes but give the sequence an annotation that it should be considered as reversed.
procedure sort(i, p) begin  
if i mod 2 = 1 then local_sort(UP);  
else local_sort(DOWN);  
for k = 1 to p do begin  
  up ← (i - 1) mod 2^k < 2^{k-1};  
  bitonic_sort(i, k, up);  
end  
end

procedure bitonic_sort(i, k, up) begin  
for j = k + 1 to P do begin  
  k = k - 1;  
  up ← up;  
  if i ≤ 2^j then bitonic_split(i + 2^j, lower);  
  else bitonic_split(i - 2^j, ¬lower);  
end  
bitonic_merge(up);  
end

Fig. 2 Pseudo code of parallel bitonic sort.

the bitonic sequence. Then s'_j is defined as follows.
- s_j itself if j ≠ j' and (j' < j ↔ A) ↔ L.
- Leading (or trailing) subsequence of s_j preceding (or following) the crossing
  index on j (i.e., j = j') if ¬(A ↔ L) (or A ↔ L).
- Nothing if j ≠ j' and (j' > j ↔ A) ↔ L.

The procedure bitonic_merge() is for merging a bitonic sequence on a process
performed by merge() after rotating the sequence if necessary.

Since the computation and communication cost of swap() is O(N) and the
computation cost of merge() is also O(N), the total time complexity of the
algorithm is O(N log^2 P). As for the space complexity, we just need an array of
2P + 1 whose k-th element has the number of elements of k-th chunk in the
sequence which is a subsequence of S(i) for some i. That is, the sequence consists up to three subsequences each of which is represented by the destination of the first chunk, the number of chunks and the number of data elements in the
subsequence. Therefore, the space complexity is O(P).

3.3 Algorithm with Pairwise Exchange
The second algorithm is similar to pairwise exchange for fixed-size all-to-all,
but the exchanging is asymmetric. That is, a pair of processes i and j such that
i < j performs swap(i, j, s_i, s_j) so that i gets a part of S(i) which j has but i
gives j arbitrary data elements which is not necessary a part of S(j). With this
asymmetric exchange, a process i ∈ [1, P] will have S(i) or some permutation of
its member elements by the following P − 1 communications whose global view
for P = 8 is shown Fig. 4.

(1) Repeat the following for j = 1 to i − 1. Concatenate all subsequence of
S(j) residing in i to make s_i and do swap(i, j, s_i, s_j) to get a sequence s_j.
(2) Repeat the following for j = i + 1 to P. Assemble a sequence s_i not having
any elements in $S(i)$ to be swapped with $s_j$ which should be a sequence of elements in $S(i)$, and then do swap\((i, j, s_i, s_j)$.

In the algorithm, we have a free hand to compose $s_i$ in the step (2). One reasonable way is to compose it by elements in $S(i+1)$, then those $S(i+2)$ if not suffice, and repeat this process until $|s_i| = |s_j|$. The time complexity of this algorithm is hard to analyze. Bad news is the worst case complexity is $O(NP)$ in some extreme cases. For example, if the process $1$ initially has $S(P)$ and other processes $i \in [2, P]$ have $S(i-1)$, the execution is serialized with $P-1$ communications of swap\((i, i+1, S(P), S(i))$. In practice, however, it is expected that initial setting is more moderate and the time complexity is $O(MP \log P)$ where $M$ is the maximum size of swapped chunks in each row in Fig. 4 and is expected to be $O(N/P)$ to make the complexity $O(N \log P)$.

The log $P$ factor of the time complexity is to merge received sequences. A process $i$ should perform merging when it does swap\()$ with $j$ such that $j < i$ if $i$ has already received sequences in $S(j)$. The process $i$ also needs to merge received sequences in $S(k)$ such that $k > i$ when it starts swap\() with the process $i+1$ if we adopt the reasonable way to compose $s_i$ as discussed above. The number of merging operations and the size of sequences to be merged are expected to be small enough to make the time complexity $O(N \log P)$ but they could be too large resulting in $O(NP)$ complexity. In addition, the size of data structures to maintain the received sequences is expected to be $O(P)$ but can be $O(P^2)$ especially when we need the stability.

To summarize, this algorithm is faster than that based on bitonic sort in usual cases, but could be slower and inefficient with respect to the memory overhead depending on the initial setting. Therefore, we have to switch two algorithms by examining the initial setting. The implementation of this pairwise exchange algorithm is left for our future work together with the combination of two algorithms and switching criteria.

### 4. Experiments

We implemented the algorithm based on bitonic sort on our T2K Open Supercomputer using Fujitsu’s C compiler version 3.0 and MPI library version 3.0. We measured the performance with $N = 2^4$ data elements of 16 byte, or 256 MB memory space, for each process. The number of processes is set to $2^8$ varying $p$ from 1 to 8 to have $P = 2$ to 256, and each process is allocated on a core of quad-core Opteron.

As for the initial setting of $s(i, j)$ for the process $i$, we examined the following four cases.

**case-1:** $|s(i, j)| = N/P$

**case-2:** $|s(i, j)| = \begin{cases} 0 & 0 \leq (j-i) \mod P < P/4 \\ N/P & P/4 \leq (j-i) \mod P < P-1 \\ N/4 & (j-i) \mod P = P-1 \end{cases}$

**case-3:** $|s(i, j)| = \begin{cases} 0 & 0 \leq (j-i) \mod P < P/2 \\ N/P & P/2 \leq (j-i) \mod P < P-1 \\ N/2 & (j-i) \mod P = P-1 \end{cases}$

**case-4:** $|s(i, j)| = \begin{cases} 0 & (j-i) \mod P \neq P-1 \\ N & (j-i) \mod P = P-1 \end{cases}$

Finally, we also measured the performance of MPI\(_{\text{Alltoallv}}\)$ for all cases as reference.
The measured execution times are shown in Table 1. Although execution times vary depending on the initial settings, they are fit well to the time complexity $O(N \log^2 P)$. In fact, if we can neglect the execution time of $merge()$, the time for one $swap()$ is in the range 1.4–1.5 s in case-1, 1.7–2.1 s in case-2, 1.7–2.2 s in case-3, and 0.7–1.5 s in case-4. These numbers mean that the throughput of one $swap()$ is 120–380 MB/s, which is not excellent but reasonable.

The absolute execution time up to 78.6 s for 256 process in case-3 is much larger than that of $MPI_{Alltoallv}()$ which takes up to 4.5 s, but is acceptable for many purposes including our motivated example of load (re-)distribution in particle-in-cell simulation. Note that the execution time of one minute or so is significantly shorter than a straightforward solution relying on (local) disk storage, which might take 6–7 minutes for our 16-core nodes providing the effective throughput for swap-in/out of the node disk is 20 MB/s. Finally, the execution time of 4096 processes is expected to be about 2.5 minutes which is still significantly better than the solution with disk storage.

5. Conclusion

In this paper, we discussed two memory-efficient algorithms of variable-size all-to-all communication. The first algorithm based on bitonic sort takes $O(N \log^2 P)$ computation and communication time with $O(P)$ extra memory space. The second algorithm with asymmetric pairwise exchange will be faster than the first in usual cases because its $O(MP \log P)$ time complexity is expected to be $O(N \log P)$ if the data size $M$ of each exchange is $O(N/P)$.

We implemented the first algorithm on our T2K Open Supercomputer and measured its performance using up to 256 CPU cores. This evaluation confirmed that the execution time is proportional to $N \log^2 P$ and the absolute value up to 78.6 s with 256 MB data for each of 256 processes is acceptably small.

Our urgent future work is to analyze the behavior of the second pairwise exchange algorithm in detail to find criteria to bound its time complexity to $O(N \log N)$ and memory overhead to $O(P)$. Then we will implement a combined algorithm to switch two algorithms according to the criteria.

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