Teaching MPI from Mental Models

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Abstract—The Message Passing Interface (MPI) is the de facto standard for programming large scale parallelism, with up to millions of individual processes. Its dominant paradigm of Single Program Multiple Data (SPMD) programming is different from threaded and multicore parallelism, to an extent that students have a hard time switching models. In contrast to threaded programming, which allows for a view of the execution with central control and a central repository of data, SPMD programming has a symmetric model where all processes are active all the time, and none is privileged in any sense, and where data is distributed.

This model is counterintuitive to the novice parallel programmer, so care needs to be taken how to install the proper ‘mental model’.

We identify problems with the currently common way of teaching MPI, and propose a way that is geared to explicit reinforcing the symmetric model. Additionally, we teach starting from realistic scenarios, rather than writing artificial code just to exercise a newly-learned routine.

This motivation implies that we reverse the commonly used order of presenting MPI routines, starting with collectives, and later introducing point-to-point routines only as support for certain symmetric operations, avoiding the process-to-process model.

I. INTRODUCTION

The Message Passing Interface (MPI) library \cite{1,2} is the de facto tool for large scale parallelism as it is used in engineering sciences. Its main model for parallelism is described as Single Program Multiple Data (SPMD): multiple instances of a single program run on the processing elements, each operating on its own data. The synchronization between the MPI processes is done through explicit send and receive calls.

The main motivation for MPI is the fact that it can be scaled to more or less arbitrary scales, currently up to a million cores \cite{3}. Contrast this with threaded programming, which is limited more or less by the core count on a single node, currently about 70.

While MPI programs can solve many or all of the same problems that can be solved in a multicore context, the programming approach is different, and requires an adjustment in the programmer’s ‘mental model’ \cite{4,5} of the parallel execution. This paper addresses the question of how to teach MPI to best effect this shift in mindset.

A. The traditional view of parallelism

The problem with mastering the MPI library is that beginning programmers take a while to overcome a certain mental model for parallelism. In this model, which we can call ‘sequential semantics’, there is only a single strand of execution\cite{6} where simple operations (such as between scalars) are done traditionally in a sequential manner, but certain operations (typically on arrays) are magically done in parallel.

Interestingly, work by Ben-David Kolikant \cite{6} shows that students with no prior knowledge of concurrency, when invited to consider parallel activities, will still think in terms of centralized solutions.

This mental model corresponds closely to the way algorithms are described in the literature, and it is actually correct to an extent in the context of threaded libraries such as OpenMP, where there is indeed initially a single thread of execution, which in some places spawns a team of threads to execute certain sections of code in parallel. However, in MPI this model is factually incorrect, since there are always multiple processes active, with none essentially privileged over others, and no shared or central data store.

B. Misconceptions in programming MPI

Then sequential semantics mental model invites the student to adopt certain programming techniques, such as the master-worker approach to parallel programming. While this is often the right approach with thread-based coding, where we indeed have a master thread and spawned threads, it is usually incorrect for MPI.

The strands of execution in an MPI run are all long-living processes (as opposed to dynamically spawned threads), and are symmetric in their capabilities and execution. Lack of recognition of this symmetry also induces students to solve problems by having a form of ‘central data store’ on one process, rather than adopting a symmetric, distributed, storage model.

For instance, we have seen a student solve a data transposition problem by collecting all data on process 0, and subsequently distributing it again in transposed form. While this may be reasonable in shared memory with OpenMP, with MPI it is unrealistic in that no process is likely to have enough storage for the full problem. Also, this introduces a sequential bottleneck in the execution.

In conclusion, we posit that beginning MPI programmers may suffer from a mental model that makes them insufficiently realize the symmetry of MPI processes, and thereby arrive at inefficient and nonscalable solutions.

We now consider the way MPI is usually taught, and offer an alternative that is less likely to lead to an incorrect mental model.

\footnote{We carefully avoid the word ‘thread’ which carries many connotations in the context of parallel programming.}
II. Teaching MPI, the Usual Way

The MPI library is typically taught as follows. After an introduction about parallelism (covering speedup and such), and shared versus distributed memory parallelism, students learn about the initialization and finalization routines, and the MPI_Comm_size and MPI_Comm_rank calls for querying the number of processes and the rank of the current process.

After that, the typical sequence is
1) two-sided communication, with first blocking and later non-blocking variants;
2) collectives; and
3) any number of advanced topics such as derived data types, one-sided communication, subcommunicators, MPI I/O et cetera, in no particular order.

This sequence is defensible from a point of the underlying implementation: the two-sided communication calls are a close map to hardware behaviour, and communications are both conceptually equivalent to, and can be implemented as, a sequence of point-to-point communication calls. However, this is not a sufficient justification for teaching this sequence of topics.

A. Criticism

We offer three points of criticism against this traditional approach to teaching MPI.

First of all, there is no real reason for teaching collectives after two-sided routines. They are not harder, nor require the latter as prerequisite. In fact, their interface is simpler for a beginner, requiring one line for a collective, as opposed to the 'master thread' and 'parallel regions' they will find it much harder to come to associate parallelism with a model that has a 'master thread' and 'parallel regions' they will find it much harder to make idiomatic use of the symmetric model of MPI.

Our second point of criticism is regarding the blocking and non-blocking two-sided communication routines. The blocking routines are typically taught first, with a discussion of how blocking behaviour can lead to load unbalance and therefore inefficiency. The non-blocking routines are then motivated from a point of latency hiding and solving the problems inherent in blocking. In our view such performance considerations should be secondary. Non-blocking routines should instead be taught as the natural solution to a conceptual problem, as explained below.

Thirdly, starting with point-to-point routines stems from a 'Communicating Sequential Processes' (CSP) view of a program: each process stands on its own, any global behaviour is an emergent property of the run. This may make sense for the teacher who know how concepts are realized 'under the hood', but it does not lead to additional insight with the students. We believe that a more fruitful approach to MPI programming starts from the global behaviour, and then derives the MPI process in a top-down manner.

We will now outline our proposed order for teaching the MPI concepts.

B. Teaching OpenMP

In scientific computing, the other dominant parallel programming system is OpenMP [8]. Often this is taught earlier because it is supposedly easier, or because its parallelism would be easier to grasp. Regardless our opinion on the first estimate, we argue that OpenMP should be taught after MPI because of its more intuitive parallelism model. If students come to associate parallelism with a model that has a 'master thread' and 'parallel regions' they will find it much harder to make idiomatic use of the symmetric model of MPI.

III. Teaching MPI, Our Proposal

As alternative to the above sequence of introducing MPI concepts, we propose a sequence that focuses on practical scenarios, and that actively reinforces the mental model of SPMD execution.

A. Process symmetry

Paradoxically, the first way to get students to appreciate the notion of process symmetry in MPI is to run a non-MPI program. Thus, students are asked to write a ‘hello world’ program, and execute this with mpiexec, as if it were an MPI program. Every process executes the print statement identically, bearing out the total symmetry between the processes.

Next, students are asked to insert the initialize and finalize statements, with three different ‘hello world’ statements before, between, and after them. This will prevent any notion of the code between initialization and finalization being considered as an OpenMP style ‘parallel region’.

A simple test to show that while processes are symmetric they are not identical is offered by the exercise of using the MPI_Get_processor_name function, which will have different output for some or all of the processes, depending on how the hostfile was arranged.

B. Functional parallelism

The MPI_Comm_rank function is introduced as a way of distinguishing between the MPI processes. Students are asked to write a program where only one rank prints the output of MPI_Comm_size.

Having different execution without necessarily different data is a case of ‘functional parallelism’. At this point there are few examples that we can assign. For instance, in order to code the evaluation of an integral by Riemann sums \( \pi/4 = \int_0^1 \sqrt{1-x^2} \, dx \) is a popular one) would need a final sum collective, which has not been taught at this point.

A possible example would be primality testing, where each process tries to find a factor of some large integer \( N \) by traversing a subrange of \([2, \sqrt{N}]\), and printing a message if a factor is found. Boolean satisfiability problems form another example, where again a search space is partitioned without involving any data space; a process finding a satisfying input can simply print this fact.
C. Introducing collectives

At this point we can introduce collectives, for instance to find the maximum of a random value that is computed locally on each process. This requires teaching the code for random number generation and, importantly, setting a process-dependent random number seed. Generating random 2D or 3D coordinates and finding the center of mass is an examples that requires a send and receive buffer of length greater than 1, and illustrates that reductions are then done pointwise.

These examples evince both process symmetry and a first form of local data. However, a thorough treatment of distributed parallel data will come in the discussion of point-to-point routines.

It is an interesting question whether we should dispense with ‘rooted’ collectives such as MPI_Reduce at first, and start with MPI_Allreduce. The latter is more symmetric in nature, and has a buffer treatment that is easier to explain; it certainly reinforces the symmetric mindset. To this author, there are few reasons to prefer

MPI_Reduce( indata,outdata, /* root= */ 0 );

there are few reasons to prefer it certainly reinforces the symmetric mindset. To this author, in nature, and has a buffer treatment that is easier to explain; the students initially do exercises with the non-rooted variants.

This has the added advantage of not bothering the students initially with the asymmetric treatment of the receive buffer between the root and all other processes.

D. Distributed data

As motivation for the following discussion of point-to-point routines, we now introduce the notion of distributed data. In its simplest form, a parallel program operates on a linear array the dimensions of which exceed the memory of any single process.

The lecturer stresses that the global structure of the distributed array is only ‘in the programmer’s mind’: each MPI process sees an array with indexing starting at zero. The following snippet of code is given for the students to use in subsequent exercises:

```c
int myfirst = ......;
for (int ilocal=0; ilocal<nlocal; ilocal++) {
    int iglobal = myfirst+ilocal;
    array[ilocal] = f(iglobal);
}
```

At this point, the students can code a second variant of the primality testing exercise above, but with an array allocated to store the integer range. Since collectives are now known, it becomes possible to have a single summary statement from one rank, rather than a partial result statement from each.

The following exercise is done in the classroom:

Each student holds a piece of paper in the right hand – keep your left hand behind your back – and executes the following program:

1) If you are not the rightmost student, turn to the right and give the paper to your right neighbour.

2) If you are not the leftmost student, turn to your left and accept the paper from your left neighbour.

This introduces students to some subtleties in the concept of parallel correctness: a program may give the right result,
but not with the proper parallel efficiency. Asking a class to solve this conundrum will usually lead to at least one student suggesting splitting processes in odd and even subsets.

3) Back to data exchange: The foregoing detours into the behaviour of two-sided send and receive calls were necessary, but they introduced asymmetric behaviour in the processes. We return to the averaging operation given above, and with it to a code that treats all processes symmetrically. In particular, we argue that, except for the first and last, each process exchanges information with its left and right neighbour.

This could be implemented with blocking sends and receive calls, but students recognize how this could be somewhere between tedious and error-prone. Instead, to prevent deadlock and serialization as described above, we now offer the MPI_Sendrecv routine. Students are asked to implement the classroom exercise above with the sendrecv routine. Ideally, they use timing or tracing to gather evidence that no serialization is happening.

As a non-trivial example (in fact, this takes enough programming that one might assign it as an exam question, rather than an exercise during a workshop) students can now implement a swap-sort algorithm using MPI_Sendrecv as the main tool. For simplicity they can use a single array element per process. (If each process has a subarray one has to make sure their solution has the right parallel complexity. It is easy to make errors here and implement a correct algorithm that, however, performs too slowly.)

Note that students have at this point not done any serious exercises with the blocking communication calls, other than the ping-pong. No such exercises will in fact be done.

F. Non-blocking sends

Non-blocking sends are now introduced as the solution to a specific problem: the above schemes required paired-up processes, or careful orchestration of send and receive sequences. In the case of irregular communications this is no longer possible or feasible. Life would be easy if we could declare ‘this data needs to be sent’ or ‘these messages are expected’, and then wait for these messages collectively. Given this motivation, it is immediately clear that multiple send or receive buffers are needed, and that requests need to be collected.

Implementing the three-point averaging with non-blocking calls is at this point an excellent exercise.

Note that we have here motivated the non-blocking routines to solve a symmetric problem. Doing this should teach the students the essential point that each non-blocking call needs its own buffer and generates its own request. Viewing non-blocking routines as a performance alternative to blocking routines is likely to lead to students re-using buffers or failing to save the request objects. Doing so is a correctness bug that is very hard to find, and at large scale it induces a memory leak since many requests objects are lost.

G. Taking it from here

At this point various advanced topics can be discussed. For instance, Cartesian topologies can be introduced, extending the linear averaging operation to a higher dimensional one. Subcommunicators can be introduced to apply collectives to rows and columns of a matrix. The recursive matrix transposition algorithm is also an excellent application of subcommunicators.

However, didactically these topics do not require the careful attention that the introduction of the basic concepts needs, so we will not go into further detail here.

IV. FURTHER COURSE SUMMARY

We have taught MPI based on the above ideas as both an academic class (typical enrollment: 40 students) and a two-day intensive workshop (attendance 10–40 students depending on circumstances) of 6–8 hours per day. Students of the academic class are typically graduate or upper level undergraduate students; the workshops get attendance from postdocs, academics, and industry too. The typical background is applied math, engineering, physical sciences.

We cover the following topics, with division over two days in the workshop format:

- Day 1: familiarity with SPMD, collectives, blocking and non-blocking two-sided communication.
- Day 2: exposure to: sub-communicators, derived datatypes. Two of the following: MPI-I/O, one-sided communication, process management, the profiling and tools interfaces, neighbourhood collectives.
A. Exercises

On day 1 the students do approximately 10 programming exercises, mostly finishing a skeleton code given by the instructor. For the day 2 material students do two exercises per topic, again starting with a given skeleton.

The design of these skeleton codes is an interesting problem. The skeletons are intended to take the grunt work away from the students, to both indicate a basic code structure and relieve them from making elementary coding errors that have no bearing on learning MPI. On the other hand, the skeletons should leave enough unspecified that multiple solutions are possible, including wrong ones: we want students to be confronted with conceptual errors in their thinking, and a too-far-finished skeleton would prevent them from doing that.

Example: the prime finding exercise mentioned above (which teaches the notion of functional parallelism) has the following skeleton:

```c
int myfactor;
// Specify the loop header:
// for ( ... myfactor ... )
for {
    /** your code here ****/
}
if (bignum%myfactor==0)
    printf("Processor %d found factor %d\n", procno,myfactor);
}
```

This leaves open the possibility of both a blockwise and a cyclic distribution of the search space.

B. Evaluation and supervision

In our academic class the programming exercises are done partly in class and partly as homework. They are very lightly graded: the majority of the grade comes from a programming project that is the student’s own initiative and coded from scratch.

In the workshop format there is no grading, but the instructor plus possible assistants are continuously monitoring students to help them over stumbling blocks and clarify concepts, both in MPI and elementary programming. We have found that one instructor or assistant per 8–10 students is a good ratio; more students per instructor will have students fall by the wayside. The typical exercise takes 15–30 minutes.

V. Evaluation and Discussion

At the moment, no rigorous evaluation of the efficacy of the above ideas has been done. We intend to perform a comparison between outcomes of the proposed way of teaching and the traditional way by comparing courses at two (or more) different institutions and from different syllabi. The evaluation will then be based on evaluating the independent programming project.

However, anecdotal evidence suggests that students are less likely to develop 'centralized' solutions as described in section I-B. This was especially the case in our semester-long course, where the students have to design and implement a parallel programming project of their own choosing. After teaching the ‘symmetric’ approach, no students wrote code based on a manager-worker model, or using centralized storage. In earlier semesters, we had seen students do this, even though this model was never taught as such.

Even with our current approach, however, we still see students writing idioms that are contrary to the symmetric model. For instance, they will write

```c
for (p=0; p<nprocs; p++)
    if (p==myrank)
        // do some function of p
```

This code computes the correct result, and with the correct performance behaviour, but it still shows a conceptual misunderstanding. Further investigation of the mental model that leads to such constructs, and how it can be prevented, is needed.

In general, it is the opinion of this author that it pays off to teach from the basis of instilling a mental model, rather than of presenting topics in some order of (perceived) complexity or sophistication.

VI. Summary

In this paper we have introduced a non-standard sequence for presenting the basic mechanisms in MPI. Rather than starting with sends and receives and building up from there, we start with mechanisms that emphasize the inherent symmetry between processes in the SPMD programming model. This symmetry requires a substantial shift in mindset of the programmer, and therefore we target it explicitly.

Comparing our presentation as outlined above to the standard presentation, we recognize the downplaying of the blocking send and receive calls. While students learn these, and in fact learn them before other send and receive mechanisms, they will recognize the dangers and difficulties in using them, and will have the combined sendrecv call as well as non-blocking routines as standard tools in their arsenal.

REFERENCES