Acceleration of the KINETICS Integrated Dynamical/Chemical Computational Model using MPI

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Abstract

Understanding the evolution of a planet's atmosphere not only provides a better theoretical understanding of planetary physics and the formation of planets, but also grants useful insight into Earth's own atmosphere. One of the tools used at JPL for the modeling of planetary atmospheres and protostellar disks is KINETICS. KINETICS can simulate years of complex dynamics and chemistry. At the moment, KINETICS is in use by or planned to be used for:

2. Modeling of the composition of the protostellar disks which lead to star formation.
3. Examination of pollutants in Earth's atmosphere.
4. Modeling of Titan's atmosphere

Because of the complexity of KINETICS, it requires large amounts of computation to run simulations at a meaningful level of detail. Using MPI (Message Passing Interface), a popular multi-core programming model, will permit more useful experiments to be run in less time, accelerating the discovery process. The advantages of MPI lie in its exposure of communication, allowing programmers to manually tune the communication pattern of an application. MPI will be used with the existing FORTRAN code to parallelize and accelerate performance-critical sections of code in KINETICS.

I. Introduction

The understanding of the formation of a planetary body's atmosphere is important for both practical applications and pure science. By studying the composition of the atmosphere of Mars (for example), it is possible to not only gain a better theoretical understanding of planetary physics and the formation of planets but also gain insight into Earth's own atmosphere. However, the study of a body of particles as large as a planet or moon's atmosphere is not a simple task, and must be accomplished primarily using computer simulations.

To be able to trust the results of a simulation, it is necessary to represent the state of a massive body of molecules as accurately as possible. Doing this is extremely demanding in terms of both communication and computation. Therefore, it is not feasible to solve perfect chemical and dynamic models for an atmosphere. Rather, approximations must be used. This leaves us with a computationally difficult task rather than a computationally impossible task.

To run atmospheric models and other financial, medical, and scientific applications as demanding as it, multi-core programming is a necessity. Multi-core programming, as opposed to sequential programming, uses many processing units executing independent instruction streams concurrently in order to complete a task quicker than a single, faster processing unit. This ability to perform simultaneous tasks is extremely useful in scientific computation, and allows applications to be significantly accelerated. One of the tools used at JPL for the modeling of planetary atmospheres and protostellar disks is KINETICS. KINETICS allows the user to provide an initial input file containing information on the body being simulated and take the body of interest through years of complex dynamics and chemistry. At the moment, KINETICS is in use by or planned to be used by:

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2. Modeling of the composition of the protostellar disks which lead to star formation.
3. Examination of pollutants in Earth's atmosphere.

While KINETICS has years of development behind it (or because of the work that has been invested in it) it requires huge amounts of computation time to run simulations at a meaningful level of detail. Modifying the FORTRAN code base to take advantage of multiple cores will allow more useful experiments to be run in less time, accelerating the discovery process. To do this, MPI (Message Passing Interface) will be used, a common parallel programming model. The advantages of MPI lie with its exposure of communication to the user. Communication of data between physically separate computation units is often the most significant source of performance degradation in multi-core applications. MPI ensures that no data is shared between processors which is not explicitly communicated, and in this way allows the programmer to tune the communication patterns and reduce total communication. As a result, MPI also places considerably more burden on the programmer to optimize the application than other parallel programming models, such as OpenMP or Java Threads. MPI will be used with the existing Fortran code to parallelize critical sections of code in KINETICS which consume the largest amount of processor time.

II. Methods

MPI is a parallel programming model commonly used in cluster computing where several machines with multiple processor are connected by high latency, low bandwidth interconnect. The advantage of MPI is that it exposes any
and all communication to the programmer, allowing them to manually manage any data transmitted. This comes at
the cost of more programming burden being placed on the programmer, in that any shared data must be explicitly
communicated, but provides the parallel programmer with more power to optimize the communication patterns of
their application.

An MPI program is just a collection of processes, each running identical code and only differentiated by their
process IDs. These processes can communicate with each other whether using TCP. Because the TCP protocol is
used for communication, from the programmers point of view there is no difference between transmitting data to
another process on the same rack or in another room.

The code transformations necessary to move from sequential FORTRAN code to parallel MPI/FORTRAN code are
often extensive and included:

1. Removal of COMMON blocks. COMMON blocks provide variables which can be accessed from anywhere
   in an application. While they can be useful, they tend to make code less readable and finding inputs and
   outputs of subroutines a greater challenge. In order to aid in the analysis of the code base, any sections
   which were to be parallelized had references to COMMON blocks removed.
2. Loop transformations. The most common form of parallelization is across iterations in a loop. So long as
   the work done in each iteration is independent of the others, this can be a simple and very beneficial
   parallelization technique. However, most code (particularly legacy code) was not designed with
   parallelization in mind, but rather expressed how the programmer originally understood the concepts. It was
   necessary to transform many of the loops in KINETICS to 1) create loops which would contained sufficient
   work as to be beneficially parallelizable, and 2) ensure each iteration of these loops were independent and
   could be parallelized.
3. Insertion of MPI calls. For this program to be parallelized, many calls to communication subroutines had to
   be inserted in the original FORTRAN code. These can be split into two categories. First, MPI_BCAST calls
   were inserted in order to transmit to all MPI processes the current state of the program and ensure that they
   all had the correct inputs when entering a parallel section of code. Second, MPI_RECV and MPI_SEND
   calls were inserted after those parallel sections of code in order to gather the results back to a master
   process, which would then go on to use those results for additional, sequential computation.

All of the above steps required considerable by-hand analysis of the FORTRAN code for data dependencies.

When parallelizing an application like KINETICS, it is crucial to maintain consistency with the original source
code. When working with scientific code which will be used to expand the understanding of Earth and other planets,
this becomes even more crucial. The primary output of KINETICS consists of a text file with information about
each time step in the simulation, including concentrations of certain elements at different altitudes, latitudes, and
longitudes. In order to validate the output of this code the Unix diff utility was used to compare the contents of a
control output file and the output of the altered code. When the same compiler was used, no differences in these files
were tolerated. Different compilers with identical code did cause differences in the output, so some tolerance for
changes had to be made in those situations.

When first analyzing the KINETICS code, performance profiling using gprof and visualization tools was done in
order to identify those sections of code which were taking up the most execution time, and therefore whose
acceleration could have the largest impact on total run time.

In addition to the work done on the actual code, it was also necessary to include the ability to use MPI in the build
process. KINETICS supports several different compilers which can be used to build KINETICS. This is useful and
necessary in order to distribute KINETICS, as not every researcher will have access to the same architectures or
compilers depending on licensing issues. Therefore, it was necessary to insert a new option which included the MPI
compiler and the arguments to that compiler which would be necessary to produce results compatible with the
others. This process was a lesson in the inconsistencies which different compilers can have, and the numerical effect
they can have on the output of an application. Simply switching from the Absoft Fortran compiler to the Intel
Fortran compiler resulted in significant changes in the output of KINETICS, as a result of some aggressive optimizations on behalf of Intel. However, the Intel compiler actually picked up on some compile time errors which Absoft did not. Adding floating-point accuracy options to the Intel compilation brought the two compilers into close agreement, but this entire task taught us something about how carefully different compilers should be used and the skepticism their results should be treated with.

III. Results

One result of this internship is the composition of a manual on the use of MPI. This manual includes:

1. An overview of MPI. Targeted at those with no experience with MPI or parallel programming, this overview tries to relate MPI to more commonly understood features of computing to make MPI a more approachable programming model.
2. Detailed information on useful MPI subroutines.
3. More advanced performance concerns which are important to keep in mind when building an MPI application, as well as how to resolve them.
4. Compilation and Execution of an MPI program, taking the reader through the steps to build and run a simple MPI application.
5. A walk-through of the parallelization of matrix-matrix multiply using MPI.

The goal of this manual was to provide scientists (non-programmers) with the ability to parallelize their code.
In order to test the performance of the parallelized code, a data set representing a 1D column of the Titan atmosphere was used. Timing was achieved using the Unix time utility.

In Figure 3, we see performance gain of the KINETICS code from 1 process to 16 processes, achieving an eventual speedup of 4.5x (parallel efficiency of 29%). While the acceleration is significant, the gains are not efficient or linear. There are two explanations for this:

1. Amdahl's Law. Amdahl's Law places an upper bound on the amount of speedup that can be expected from parallelizing sections of code within a program. Because only two functions, RAD and MARCH, are being parallelized in this execution of KINETICS, only a certain percentage of the total execution time is being done in parallel. Even given perfect speed up of those sections and infinite processors, there is a theoretical lower bound on the measured execution time, which would be the time which the sequential sections take. As more of the code is parallelized this lower bound on execution time and upper bound on speedup become more beneficial, but time constraints dictated that no additional work could be done.

2. Communication. As more processes are added, more data needs to be transmitted. Rather than sending an array from process 0 to process 1 with 2 processes, it would be necessary to transmit that array to 15 other processes when using 16 processes. This places more load on the available bandwidth and can affect the observed speed up. Again, due to time constraints more efficient data transfer patterns could not be implemented, which would limit the impact of this problem.

IV. Conclusion

This paper demonstrated that there are significant improvements to be gained from parallelizing the KINETICS code. Using MPI to limit communication and parallelize performance-critical sections of code produced speedup of 4.5x.

While promising results were obtained, the difficulty of working with legacy code which was never designed for parallelization represented a serious bottleneck in this project. The loop structure of the code did not lend itself to parallelization, and required some major code transformations to be done and validated. The use of Fortran common blocks made it difficult to analyze data dependencies between subroutines. Even the compile process presented a
challenge as we learned that different compilers treat errors in different ways, produce slight differences with completely identical code, and rely in different libraries or syntax. Overall, this added overhead constituted the bulk of the work done while the actual parallelization tasks were relatively simple, straightforward, and quick.

Given more time, considerably better performance could be achieved. The use of more complex and efficient data transfer patterns could be implemented, following a policy of copy early and overlapping communication with computation. More of the source code could be parallelized, increasing the upper bound on speed up while potentially eliminating the need to copy certain data. Additionally, more testing could be completed to further certify the robustness of the parallel code.

The manual written as part of this work will hopefully be useful for future parallelization efforts on the KINETICS code and other scientific codes which lend themselves to parallelization.

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