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CLUSTEREASY

A Program for Simulating Scalar Field Evolution on Parallel Computers

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We describe a new, parallel programming version of the scalar field simulation program LATTICEEASY. The new C++ program, CLUSTEREASY, can simulate arbitrary scalar field models on distributed-memory clusters. The speed and memory requirements scale well with the number of processors. As with the serial version of LATTICEEASY, CLUSTEREASY can run simulations in one, two, or three dimensions, with or without expansion of the universe, with customizable parameters and output. The program and its full documentation are available on the LATTICEEASY website at <http://www.science.smith.edu/departments/Physics/fstaff/gfelder/latticeeasy/>. In this paper we provide a brief overview of what CLUSTEREASY does and the ways in which it does and doesn't differ from the serial version of LATTICEEASY.

I. INTRODUCTION

Studying the early universe requires describing the evolution of interacting fields in a dense, high-energy environment. The study of reheating after inflation and the subsequent thermalization of the fields produced in this process typically involves non-perturbative interactions of fields with exponentially large occupation numbers in states far from thermal equilibrium. Various approximation methods have been applied to these calculations, including linearized analysis and the Hartree approximation. These methods fail, however, as soon as the field fluctuations become large enough that they can no longer be considered small perturbations. In such a situation linear analysis no longer makes sense and the Hartree approximation neglects important rescattering terms. In many models of inflation preheating can amplify fluctuations to these large scales within a few oscillations of the inflaton field. Moreover, such large amplification appears to be a generic feature, arising via parametric resonance in single-field inflationary models and tachyonic instabilities in hybrid models.

The only way to fully treat the nonlinear dynamics of these systems is through lattice simulations. These simulations directly solve the classical equations of motion for the fields. Although this approach involves the approximation of neglecting quantum effects, these effects are exponentially small once preheating begins. So in any inflationary model in which preheating can occur lattice simulations provide the most accurate means of studying post-inflationary dynamics.

In 2000 G.F. and Igor Tkachev released LATTICEEASY [1], a C++ program for simulating scalar field evolution in an expanding universe. In the ensuing years LATTICEEASY has been used by us and other groups to study such topics as preheating, baryogenesis, gravity wave production, and more. These simulations have been extremely useful, but they have for the most part been confined to relatively simple toy models, primarily due to computational limitations. To study cosmology in more complex models such as the MSSM or GUT theories will require the use of large, parallel clusters. CLUSTEREASY is a version of LATTICEEASY that can be run in parallel on multiple processors.

Section II of this paper gives a brief overview of what LATTICEEASY does and how to use it, and notes the modifications that must be made in the LATTICEEASY files to run them in CLUSTEREASY. Section III describes the algorithms used to parallelize the simulations. For more detailed documentation the reader is referred to the LATTICEEASY website

<http://www.science.smith.edu/departments/Physics/fstaff/gfelder/latticeeasy/>

II. OVERVIEW

LATTICEEASY consists of several C++ files, but only two are designed to be modified by most users. Each particular scalar field potential that the program solves is encoded in a *model file* called `model.h`, in which the user enters equations for the potential and its various derivatives. The parameters that control individual runs are stored in a file called `parameters.h`. These parameters include physical quantities such as masses and couplings, numerical quantities such as the number of gridpoints and the size of the time step, and parameters to control what types of output are generated by the simulation.

To use CLUSTEREASY the user must replace all of the non-user-modifiable files from LATTICEEASY with the new, CLUSTEREASY versions. The parameter files from LATTICEEASY can be used with no changes, however, and the model files only need the addition of two lines, described in the online documentation.

The formats of the outputs created by CLUSTEREASY are the same as those from the serial version. One of the output options is to create a grid image that can be used to resume a run and continue it to later times. Grid images created by LATTICEEASY can be read in by CLUSTEREASY and vice-versa. The only way to distinguish CLUSTEREASY output from LATTICEEASY output is the file `info`, which contains basic information about the run such as the potential used and the physical and numerical parameters. In CLUSTEREASY this file has an additional line specifying the number of processors used for the run.

To run CLUSTEREASY you need a cluster with MPI. MPI is a standard set of libraries used for parallel programming in C, C++, and Fortran, and should be installed on any standard cluster. You also need the freely available Fourier Transform library FFTW. (See the online documentation for possible compatibility issues with the way FFTW is installed on different systems and how to resolve them.)

The makefile that comes with CLUSTEREASY assumes that the command for compiling a C++ MPI program is `mpiCC`. If this command is different on your system you will need to modify the makefile accordingly. Otherwise you should be able to compile the code simply by typing “make.” You should consult your system documentation for the correct syntax for running a parallel program, but on most clusters it is

```
mpirun -np <number of processors> latticeeasy
```

Note that the number of processors is determined at execution-time, not at compile-time.

III. PARALLEL ALGORITHMS

LATTICEEASY uses a staggered leapfrog algorithm with a fixed time step. This means that at each step the field values f and their derivatives \dot{f} are stored at two different times t and $t + dt/2$ respectively. The derivatives are used to advance the field values by a full step dt and then the field values are used to calculate the second derivatives \ddot{f} , which are in turn used to advance the field derivatives by dt . This evolution is done in place, meaning the newly calculated field values and/or derivatives overwrite the old ones.

To implement this scheme on multiple processors CLUSTEREASY uses “slab decomposition,” meaning the grid is divided along a single dimension (the first spatial dimension). For example, in a 2D run with $N = 8$ on two processors, each processor would cast a 4×8 grid for each field. At each processor the variable n stores the local size of the grid in the first dimension, so in this example each processor would store $n = 4$, $N = 8$. Note that n is not always the same for all processors, but it generally will be if the number of processors is a factor of N .

In practice, the grids are actually slightly larger than $n \times N$ because calculating spatial derivatives at a gridpoint requires knowing the neighboring values, so each processor actually has two additional columns for storing the values needed for these gradients. Continuing the example from the previous paragraph, each processor would store a 6×8 grid for each field. Within this grid the values $i = 0$ and $i = 5$ would be used for storing “buffer” values, and the actual evolution would be calculated in the range $1 \leq i \leq 4$, $0 \leq j \leq 7$.

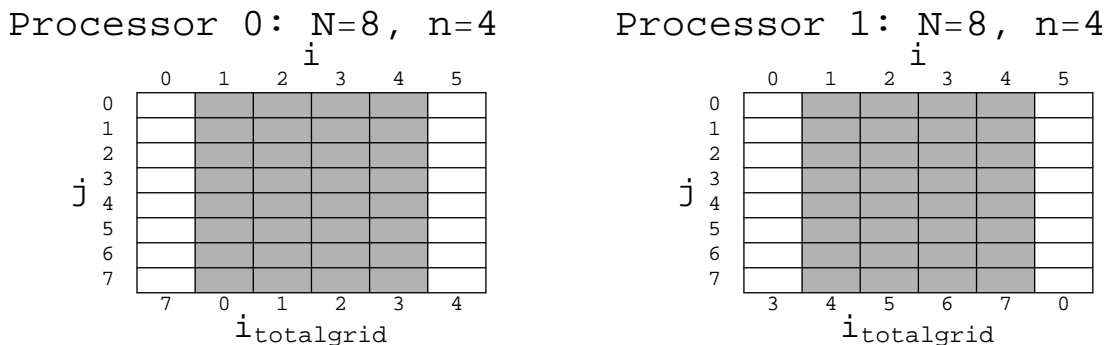


FIG. 1: Data layout in CLUSTEREASY

This scheme is shown in Figure 1. At each time step each processor advances the field values in the shaded region, using the buffers to calculate spatial derivatives. Then the processors exchange edge data. At the bottom of the figure I’ve labeled the i value of each column in the overall grid. During the exchange processor 0 would send the new values

at $i_{totalgrid} = 0$ and $i_{totalgrid} = 3$ to processor 1, which would send the values at $i_{totalgrid} = 4$ and $i_{totalgrid} = 7$ to processor 0.

The actual arrays allocated by the program are even larger than this, however, because of the extra storage required by the Fourier Transform routines. In two and three dimensions CLUSTEREASY uses FFTW. When you Fourier Transform the fields the Nyquist modes are stored in extra positions in the last dimension, so the last dimension is $N+2$ instead of N . The total size per field of the array at each processor is thus typically $n+2$ in 1D, $(n+2) \times (N+2)$ in 2D and $(n+2) \times N \times (N+2)$ in 3D. In 2D FFTW sometimes requires extra storage for intermediate calculations as well, in which case the array may be somewhat larger than this, but usually not much. This does not occur in 3D.

IV. CONCLUSIONS

We have found that the speed of the simulation scales roughly as the number of processors, provided that number is significantly smaller than N , the number of gridpoints along each edge of the lattice. A good rule of thumb is that you probably won't get much benefit from using more processors than $N/4$. Also, you will get slightly better performance per processor if the number of processors is a factor of N so that the processors can divide the lattice up evenly.

CLUSTEREASY offers the opportunity to do simulations of much larger, more complex, and more realistic early universe theories than was possible with serial simulations. We offer it in the hope that it will be useful to the research community.

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[1] G. N. Felder and I. Tkachev, [hep-ph/0011159].