Parallel Linear General Relativity and CMB Anisotropies

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Abstract

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We have developed a code which links the primeval fluctuations in the early universe with those observable at the present time by integrating the coupled, linearized, Einstein, Boltzmann, and fluid equations governing the evolution of metric perturbations and the density fluctuations; this is the most accurate treatment to date of both the physics and the numerical integration. The results are useful both for calculations of the cosmic microwave background (CMB) anisotropy and the linear power spectrum of matter fluctuations.

The serial code (LINGER) is highly efficient on vector machines. Furthermore, this application is perfectly suited for coarse-grained parallelism. A portable, parallel implementation (PLINGER) using common message-passing libraries (PVM, MPI, MPL, and PVMe) has been completed; it achieves Gflop rates on current parallel supercomputers such as the T3D and SP2. LINGER and PLINGER will soon to be released for general use.

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1 Introduction

In standard "big bang" models of cosmology, the structures seen in the universe today formed due to self-gravity; initially small perturbations in matter and radiation were gravitationally amplified into galaxies and clusters of galaxies. By following how these primordial fluctuations grow in time one can give testable predictions for the large-scale structure of the matter distribution.

Due to the same initial perturbations, there are slight deviations from a pure blackbody spectrum in the Cosmic Microwave Background (CMB) blackbody radiation, a fossil relic of the big bang. The COBE satellite, which confirmed the isotropy and blackbody spectrum (with temperature 2.726 K) of the CMB, also measured these anisotropies. Subsequent balloon-borne and ground-based experiments have measured the amplitude of CMB anisotropies on different angular scales (see Steinhardt 1995 and Lubin 1994 for recent reviews). For a given cosmological model, one can compute the predicted amplitudes of the CMB anisotropies as another test of the theory.

There are a variety of cosmological parameters which are not well constrained: the Hubble constant, neutrino masses, a possible cosmological constant, the initial perturbation spectrum, etc. However, once the initial conditions are set, the input physics is well understood, and since the initial perturbations are small the subsequent evolution is linear system. The goal of cosmological spectral codes is to compute, for a given matter composition and initial spectrum of perturbations, the spectrum for mass perturbations and CMB anisotropies expected at the present time. These predictions can serve as a discriminant of the various models.

In the past varying degrees of approximation have been made in order to carry out the evolution (for example Peebles & Yu 1970, Bond & Efstathiou 1987, Holtzman 1989, Sugiyama and Gouda 1992, among others). The code we discuss here has a highly accurate treatment of both the physics and the numerical integration; we believe it is the most accurate to date. The tradeoff for this accuracy is increased computational cost, making the use of supercomputers necessary.

The serial and parallel versions of the code (called LINGER and PLINGER,
respectively) will soon be made available through the GC3 Software Archive.

2 Equations Governing the Evolution

A number of equations govern the growth of perturbations. The Einstein equations give the effect of gravity, including the rate of universal expansion. Also needed is the Boltzmann equation, which governs the phase space evolution of photons and neutrinos. Baryons, electrons, and cold dark matter follow the equations for a pressureless perfect fluid (though the electrons and baryons are coupled to photons by Thomson scattering). See Bertschinger 1995 for a detailed introduction to the linearized Einstein equations, and Ma & Bertschinger 1994 for a derivation of the equations used in the LI NGER code.

Other physics which is modeled includes accurate treatments of hydrogen and helium recombination, decoupling of photons and baryons, and Thomson scattering (including two photon polarizations and the full angular dependences of the scattering cross section and distribution functions).

The equations are most easily solved in $k$-space. In Fourier space, all the $k$ modes in the linearized Einstein, Boltzmann, and fluid equations evolve independently. In addition to the Fourier transform, there is also an angular expansion of the phase space distributions in terms of Legendre polynomials; this turns the Boltzmann equations into moment hierarchies determined at each time step. At a given time it is also necessary to integrate over the 3-momentum, $q$, of the massive neutrinos. We carry out a full integration down to the final time without use of any free-streaming approximation. The time integration, ending at the present, is carried out using the standard Runge-Kutta integrator DVERK, obtained from netlib@research.att.com.

3 The Serial Code

The serial code, called LINGER, is highly efficient on vector machines; on a single Cray C90 node it runs at 570 Mflop, a significant fraction of the
theoretical 1 Gflop peak performance. While the message passing version discussed below will run on the C90, it is more efficient to use Cray’s Autotasking directives to parallelize the serial code. This has been done, so typical speeds in excess of 8 Gflop should be possible on 16 nodes of the C90, although the timing runs have not yet been carried out.

For runs determining the CMB anisotropy, we desire a high degree of accuracy, with errors < 0.1% for angular degree \( l < 3000 \). This requires the inclusion of up to 10,000 moments \( l \), and the integration of up to 5000 points in \( k \). Thus this problem is computationally intensive; despite getting 570 MFlop on the Cray C90, a full run still requires roughly 75 C90 CPU-hours.

4 The Parallel Code

One important feature of the treatment in the previous section is that each mode characterized by a given \( k \) evolves independently. This problem is thus perfect for coarse-grained parallelization, since each node can work on solving the equations for a particular value of \( k \) without the need to communicate with other nodes.

Another important feature is that for any value of \( k \), the computation necessary to evolve a given mode to the present is much larger than that required for the initial and final message passing. For example, with the smallest values of \( k \) required, the CPU time is at least two minutes on an IBM Power2 chip, while the results are gathered as a single message of roughly 150 bytes. (The largest \( k \)-values, corresponding to smaller scales requiring a larger number of moments \( l \), can take up to half an hour of CPU time; the message length increases roughly in proportion to the CPU time, to a maximum of 80 kbyte). Thus the overhead from message passing is insignificant.

The main loop of the serial code is in \( k \); the obvious method of parallelization is to use a master/worker approach. The message passing required is quite straightforward. At the beginning of a run, the master process needs to broadcast a few quantities to all the workers, such as the time at which to end the evolution and the maximum number of angular moments \( l \) to compute; it then waits for a message from any worker process. When a worker receives
this information it then requests a value of $k$ from the master, which replies
with the appropriate value. When the worker completes the computation, it
sends an array containing the values of interest back to the master, which
prints out these values and sends the next $k$ value to the worker (or, if no
further work is to be done, a message to stop). Thus only a few basic mes-
ringe routines are required: broadcasting to all other nodes, sending,
receiving, and checking for an incoming message (either from a particular
process or from any process), as well as the ability to tag messages.

See Appendix A to see the algorithm in more detail.

In the parallel code, calls to wrapper routines are made; these routines in
turn invoke the actual message passing libraries. The wrapper routines are
provided in a separate file, tailored to the particular library of choice. To
date, we have used PVM (see Geist et al. 1994), MPI (see Gropp et al. 1994),
MPL, and PVMe (available from IBM). Given the computationally intensive
nature of this code, the choice of which library to use has no effect on the
efficiency of the code and is simply a matter of which is most convenient to
the user.

The parallel code, called PLINGER, has been run on the DEC Alpha
Cluster and the C90/T3D at the Pittsburgh Supercomputing Center, and
the IBM SP2 at the Cornell Theory Center. On the SP2, MPL requires
that messages be received in the order in which they arrive, but this does
not create difficulties. On some machines, PVM allows the master process
to cohabit a particular node along with a worker process; this is desirable
because the master process requires little CPU time compared to the workers.
Thus PVM has a slight edge on these machines.

5 Timing of the Parallel Code

5.1 Flop Rates

SP2 On a single IBM Power 2 chip, about 15 times as much CPU is needed
as for a single Cray C90 node, so the serial code runs without special opti-
mization at 40 Mflop, or a seventh of the 266 Mflop peak performance of the
Power 2. Thus on the IBM SP2 PLINGER achieves sustained speeds of 2.4 Gflop using 64 nodes, and 9.6 Gflop on 256 nodes. We have recently found that the use of the MASS library, inlining of subroutines, and higher-order loop transformations can significantly improve the PLINGER performance on Power 2 processors, up to 58 Mflop on a single node; further improvements may be possible. Thus 15 Gflop or more should be achievable on the SP2.

T3D Rather than just using the T3D, the message passing code is most suitable for PSC’s C90/T3D heterogeneous computing environment. The master process resides on the C90, handling the input/output and controlling the T3D processes; this master process uses a negligible amount of CPU time. PLINGER runs at 15 Mflop on a single T3D node, or a tenth of the theoretical peak rate (the flop rate was found by comparison with the C90). For 256 nodes on the T3D the total rate is 3.7 Gflop.

5.2 Scaling with number of Processors

Figure 1 shows wallclock and CPU time as a function of the number of processors for a test run on the SP2. The filled circles show the total CPU time (as measured by calls to etime) divided by 100. The open squares show the wallclock time. The parallel efficiency, (total CPU time)/(wallclock time x number of nodes)), is 95\% dedicated mode. The ‘X’ show the wallclock time for a 256-node T3D run. The line shows the curve expected if the wallclock time scaled exactly as the inverse of the number of processors.

There is practically no overhead to adding more processors, so the CPU time does not change as the number of processors is increased. Once the final value of \( k \) has been given to a worker process, the other nodes will no longer have any work to do (once they have completed the \( k \)-value they are currently working on). Thus there will be a period of time at the end of each run when not all the processors are working, so doubling the processors does not quite halve the wallclock time. Since larger wavenumbers require greater computation, one simple method by which we minimized this idle time was to compute the largest \( k \) first. For productions runs, which are much longer than these test runs, this idle time will be less significant.
6 Sample Results

6.1 Power spectrum of the anisotropies

The anisotropies in the CMB can be characterized in terms of multipole moments. The two-point temperature autocorrelation function, $C$ compares the temperatures at points in the sky separated by some angle. Roughly speaking, the y-axis of this plot shows the power in the spectrum on the angular scale of the multipole $l$.

The points in Figure 2 are experimental measurements of the CMB anisotropy. The two leftmost points are the COBE first- and second-year data, probing an angular scale of ten degrees. The other points are from balloon flights or ground-based experiments; these data are available as part of the COSAPP software package made available by Rahul Dave and Paul Steinhardt at the University of Pennsylvania. The curve shows the output of a PLINGER run using standard Cold Dark Matter initial conditions and normalized to the COBE data. The PLINGER run took 20 hours on 64 nodes of the SP2. Increased accuracy in the measurement and theoretical prediction of the power spectrum will help to discriminate between cosmological models.

Figure 3 shows a simulated sky map, analogous to the COBE sky map, made using the output of PLINGER. There is much greater detail here because this map has not been smoothed like the COBE map; the angular resolution is one-half degree, compared to ten degrees for COBE. The maximum temperature differences are +/- 200 micro-K (with the average temperature equal to 2.726 K).

An mpeg movie in the HTML version of this paper shows the evolution of the potential psi of the conformal Newtonian gauge; psi plays the role of the gravitational potential in the Newtonian limit. The square is a comoving 100 Mpc across (1 pc = 3.3 light years). Standard Cold Dark Matter initial conditions are used. The movie ends shortly after recombination, at conformal time 250 Mpc (expansion factor $1/a = 1028$). The potential oscillates at early times due to the acoustic oscillations of the photon-baryon fluid. These same oscillations produce the small angular scale features in the CMB anisotropy map shown above.
7 Appendix A: The Message-Passing Algorithm

7.1 Overview

Here we briefly outline the message passing algorithm used.

Main Routine

Initialize message passing routines.
If master, call master subroutine; else call worker subroutine
Exit message passing routines

The Master Subroutine

Do initialization.
Broadcast initial data to workers.
While there are wavenumbers not yet complete:
  receive message from worker
  print out the data
  send a wavenumber, or a message to stop

The Worker Subroutine

Receive initial data from master.
Ask for wavenumber from master.
Receive from master: next wavenumber or message to stop
While a message to stop has not been received:
  integrate the equations
  send the results to the master
  receive from master: next wavenumber or message to stop

Message Passing Wrapper Routines

Certain basic message passing elements are required by PLINGER.
We have implemented the following routines in PVM, MPL, MPI, and PVMe.

initpass -initialize message passing
endpass -exit from message passing

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mybcastreal - send a message to all other processes
mysendreal - send a message to a given process
mycheckany - check for message of any type from any process
mycheckone - check for message of a given type from a given process
mychecktid - check for message of any type from a given process
myrecvreal - receive a message

7.2 In More Detail

Tags Each message carries a tag which reveals its function.

| tag | type |
|-----|------|
| 1   | - first message from master to workers |
| 2   | - from worker; asking for a wavenumber |
| 3   | - from master; giving worker a wavenumber to work on |
| 4   | - from worker; giving first set of data and lmax |
| 5   | - from worker; giving data (length = 2*lmax+8) |
| 6   | - from master; telling worker to stop |

Message Passing Wrapper Routines Here we show what the wrapper routines look like when using the MPI library.

initpass - initialize message passing
C Returns process ID in mytid and the ID of the master in mastid. SUBROUTINE initpass(mytid, mastid)
call MPI_INIT( ierr )
call MPI_COMM_RANK( MPI_COMM_WORLD, mytid, ierr )
call MPI_COMM_SIZE( MPI_COMM_WORLD, nproc, ierr )
mastid = 0

endpass - exit from message passing
C Exits MPI SUBROUTINE endpass()
call MPI_FINALIZE(ierr)

mybcastreal -send a message to all other processes
C The master process sends a message with tag=msgtype to all other processes.
C Sends length double precision numbers starting at position buffer.

SUBROUTINE mybcastreal(buffer, length, msgtype)
DO i=1, nproc-1
   call MPI_SEND(buffer, length, MPI_DOUBLE_PRECISION, 
   i, msgtype, MPI_COMM_WORLD, ierr)
END DO

mysendreal -send a message to a given process
C Sends a message with tag=msgtype to the process with ID=target.
C Sends length double precision numbers starting at position buffer.

SUBROUTINE mysendreal(buffer, length, msgtype, target)
call MPI_SEND(buffer, length, MPI_DOUBLE_PRECISION, 
   target, msgtype, MPI_COMM_WORLD, ierr)

mycheckany -check for message of any type from any process
C Waits for a message of any type from any process.

SUBROUTINE mycheckany(msgtype, target)
call MPI_PROBE(MPI_ANY_SOURCE, MPI_ANY_TAG, 
   MPI_COMM_WORLD, status, ierr)
msgtype = status(MPI_TAG)
target = status(MPI_SOURCE)

mycheckone -check for message of a given type from a given process
C Waits for a message of type msgtype from process target.

SUBROUTINE mycheckone(msgtype, target)
call MPI_PROBE(target, msgtype, MPI_COMM_WORLD, status, ierr)

mychecktid -check for message of any type from a given process
C Waits for a message of any type from process target.
C Returns the message tag in msgtype.

SUBROUTINE mychecktid(msgtype, target)
call MPI_PROBE( target, MPI_ANY_TAG, MPI_COMM_WORLD, status, ierr)
msgtype = status(MPI_TAG)

myrecvreal - receive a message
SUBROUTINE myrecvreal( buffer, length, msgtype, target)
  Receives a message of type msgtype from process target.
  length double precision numbers are copied, starting at address of buffer.
  call MPI_RECV(buffer, length, MPI_DOUBLE_PRECISION, target,
    & msgtype, MPI_COMM_WORLD, status, ierr)

The Main Routine

PROGRAM plinger
C mytid = the process ID
C mastid = the ID of the master process
INTEGER mytid,mastid
C initialize message passing routines
  CALL initpass(mytid, mastid)
IF( mytid.EQ.mastid ) THEN
  CALL parentsub(mytid, mastid)
ELSE
  CALL kidsub(mytid, mastid)
ENDIF
C exit message passing routines
  CALL endpass
STOP
END

The Master Subroutine

SUBROUTINE parentsub(mytid, mastid)

{The master initializes various quantities: the number of k values, the maximum value of k, etc. The values needed by the workers are placed into the array y.}
C broadcast data to all node programs
 msgtype = 1
 imsglen = 5
 CALL mybcastreal( y, imsglen, msgtype)

C ik is the next wavenumber to send
 ik = 1
C keep track of how many ik have been received from workers
 ikdone = 0

C Start checking for messages from the workers
 100 CONTINUE
    CALL mycheckany( msgtype, itid)

    IF( msgtype.EQ.2 ) THEN
      C msgtype=2: the worker is ready for its first ik.
      C dispose of the message since it contains no data.
      CALL myrecvreal( deltat, 1, msgtype, itid)
    ENDIF

    IF( msgtype.EQ.4 ) THEN
      C msgtype=4: receive first part of data from worker
      C the length of the next message depends on lmax
      CALL myrecvreal( y, 21, msgtype, itid)
      ikold = INT(y(1))
      lmax = INT(y(21))
      C this data is written to an ascii file
      WRITE(unit_1,*)(y(i),i=1,20)

      C msgtype=5: receive second part of data from worker
      msgtype = 5
      CALL mycheckone( msgtype, itid)
      CALL myrecvreal( y, 8+lmax+lmax, msgtype, itid)
      C this data is written to a binary file
      WRITE(unit_2,ikold,y(i),i=7,lmax)
      WRITE(unit_2)(y(8+lmax+i),i=0,lmax)
C done with this wavenumber
   ikdone = ikdone+1
ENDIF

   IF( (msgtype.EQ.2).OR.(msgtype.EQ.5) ) THEN
   IF( ik.LE.last_nk) THEN
   C reply with ik to the worker that sent the last message
   y(1) = DBLE(ik)
   msgtype = 3
   imsglen = 1
   CALL mysendreal( y, imsglen, msgtype, itid)
   C find the next value of ik to be sent
   CALL ik_next(ik)
   ELSE
   C if no more wavenumbers to send, tell the worker to stop (msgtype=6)
   msgtype = 6
   imsglen = 1
   CALL mysendreal( y, imsglen, msgtype, itid)
   ENDIF
ENDIF

C if there are still wavenumbers to receive, go back
IF( ikdone.LE.last_nk) GO TO 100

RETURN
END

The Worker Subroutine

SUBROUTINE kidsub(mytid, mastid)

   C receive initial data from master
   msgtype = 1
   CALL mycheckone( msgtype, mastid)
   CALL myrecvreal( passbuffer, 5, msgtype, mastid)
C ask for wavenumber from master

```plaintext
msgtype = 2
imsglen = 1
CALL mysendreal( passbuffer, imsglen, msgtype, mastid)
```

C receive from master: next ik or message to stop

```plaintext
CALL mychecktid( msgtype, mastid)
CALL myrecvreal( passbuffer, 1, msgtype, mastid)
ik = INT( passbuffer(1) )
```

C if the message is not a wavenumber, then exit

```plaintext
WHILE ( msgtype.NE.3 )
```

C Begin timestep loop.

```plaintext
time = t_start
WHILE( time.LT.end_time)
```

{Here the worker integrates the coupled equations.} </I> <BR>

END WHILE

C send first part of results to master

```plaintext
msgtype = 4
imsglen = 21
CALL mysendreal( passbuffer, imsglen, msgtype, mastid)
```

C send second part of results to master

```plaintext
msgtype = 5
imsglen = 8+lmax+lmax
CALL mysendreal( y, imsglen, msgtype, mastid)
```

C receive from master: next ik or message to stop

```plaintext
CALL mychecktid( msgtype, mastid)
CALL myrecvreal( passbuffer, 1, msgtype, mastid)
ik = INT( passbuffer(1) )
```

END WHILE
8 References

8.1 Articles

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Sugiyama, N., and N. Gouda 1992, Prog. Theor. Phys. 88 803

8.2 Web links

Message passing libraries:
http://www.epm.ornl.gov/pvm/pvm_home.html  
PVM: Parallel Virtual Machine

http://www.mcs.anl.gov/Projects/mpi/index.html  
MPI - Message Passing Interface

Supercomputing Centers (for information on the machines used):

http://www.tc.cornell.edu/UserDoc/Hardware/ 
Cornell Theory Center Hardware Page

http://pscinfo.psc.edu/ 
Pittsburgh Supercomputing Center home page  
(CMB Anisotropy Experiment Data, compiled by Rahul Dave)

Astrophysics Servers:

http://dept.physics.upenn.edu/~www/astro-cosmo  
CMB Window and Bandpower Software Package (COSAPP)

http://xxx.lanl.gov/astro-ph/  
Astro-ph Astrophysics Preprints

The Grand Challenge Cosmology Consortium, GC3: Look for LINGER  
(as part of the COSMICS cosmological initial conditions package) and PLINGER  
to be made available from these sites.
9 Figures

1. Wallclock and CPU time as a function of the number of processors for a test run on the SP2.

2. The points are experimental measurements of the CMB anisotropy, from the COSAPP package. The curve shows the output of a PLINGER run using standard Cold Dark Matter initial conditions and normalized to the COBE $Q_{rms-PS}$.

3. A simulated sky map, analogous to the COBE sky map, made using the output of PLINGER.
This figure "fig1-1.png" is available in "png" format from:

http://arxiv.org/ps/astro-ph/9504040v1
This figure "fig1-2.png" is available in "png" format from:

http://arxiv.org/ps/astro-ph/9504040v1
This figure "fig1-3.png" is available in "png" format from:

http://arxiv.org/ps/astro-ph/9504040v1