An adaptive multigrid solver for high-resolution cosmological simulations

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ABSTRACT

We have developed an adaptive multigrid code for solving the Poisson equation in gravitational simulations. Finer rectangular subgrids are adaptively created in locations where the density exceeds a local level-dependent threshold. We describe the code, test it in cosmological simulations, and apply it to the study of the birth and evolution of a typical pancake singularity. The initial conditions for the pancake are generated on the basis of the theory of Lagrangian singularities; we follow its evolution for a few collapse times, finding a rich substructure in the final object. We achieve a spatial resolution of 1/1024 of the size of the overall computational cube in the central parts of the pancake, with computing time comparable to that of the FFT-solvers.

Key words: gravitation – methods: numerical – large-scale structure of Universe

1 INTRODUCTION

It is common knowledge that the dynamical and spatial resolution of standard cosmological simulations of smooth (dark) matter evolution is rather low. The basic particle-mesh (PM) code smooths the density on a grid and finds the potential using FFT-type techniques. As shown by Bouchet, Adam \& Pellat (1985), grid effects damp the growth of structure for all scales smaller than at least 6 cell sizes. This restricts considerably the range of scales where we can believe our models.

In order to improve the situation several different methods have been proposed. The best known of them is the PPPM (particle-particle particle-mesh) code, developed by Eastwood \& Hockney (1974) and described in detail in their book (Hockney \& Eastwood 1981). It finds the large-scale forces from the smoothed density, as does the basic PM code, but uses the nearby mass points to calculate the detailed short-range force. This code was adapted for cosmo-
logical simulations by Efstathiou, Davis, Frenk & White (1985), and is presently the standard code for cosmology. Couchman (1991) has modified it for situations where the density range becomes large and most of the time in the PPPM-code would be spent on calculating pair interactions, introducing adaptive mesh refinement in order to reduce the amount of calculations of pairwise forces.

The ultimate high-resolution code is the tree code, proposed by Barnes & Hut (1986) and first used for cosmological simulations by Bouchet & Hernquist (1988). This code treats all interactions between gravitating masses basically on a pairwise basis and is thus free from any grid effects.

Both the PPPM code and the tree code are based on the paradigm of individual clouds of matter. Although a softened force is usually used, it is not clear how well the massive clouds represent the essentially continuous distribution of dark matter. Thus it is necessary to develop a high-resolution code for this class of problems.

The first of such codes, a multiple mesh particle code was proposed by Chan, Chau, Jessop & Jorgensen (1986), who use local higher resolution grid patches in high density regions. Another mesh refinement method has been proposed by Villumsen (1989). These methods differ mainly by their approach to solving field equations on local grids. Chan et al (1986) obtain the local grid potential as a solution to a boundary value problem for the Poisson equation on the local grid, where the boundary values are defined from the solution on a global coarser grid. They find the solution on local grids by an efficient iterative scheme. Villumsen (1989) considers the local potential as a sum of a solution for the distribution of matter on the subgrid, using isolated boundary conditions, and the external global solution for the full matter distribution (without the mass at refined regions). He uses FFT to find the potentials, and special tricks to speed up the solution for the local isolated local patches.

As we see, in the latter approach there is no backreaction from submesh particles to the global grid, the coarser periodical part of the field is computed without subgrid particles. In Chan et al method this backreaction is included.

An important issue intrinsic to all multiple resolution mesh schemes is how well they treat particles that enter subgrids. In principle, when switching to a higher resolution force anisotropies and radial errors should be less than these on a coarser grid, and additional errors will be generated mainly by solution errors near grid interfaces. In case of a boundary value problem, (Chan et al approach), the error depends on interpolation errors on the boundary and on the density estimate at the grid point next to the boundary. For the Villumsen code the errors are generated by an external lower resolution solution from a coarser grid and by the density estimation both at the grid boundary and at neighbouring grid points.

In the Couchman's $AP^3M$ code that uses force splitting to long-range and short-range parts (intrinsic to $P^3M$), the mesh part of the force is approximated by a smooth reference interparticle force $R(r, a_m)$, where $a_m$ is the softening scale appropriate for the current mesh resolution. The potential is calculated using a modified Green's function which minimizes the differences between computed and reference forces. The total mesh part of the force acting on a particle is found by summing the force $R(r, a_m)$ from the base (periodical) mesh and the forces due to particles on a refinement,
calculated with a different softening scale \( a_s \), and the system is locally considered as isolated.

The main difference between the Chan et al, Villumsen’s and Couchman’s codes is that in the former two the aim of the mesh refinement is to enhance the resolution of the force field through higher resolution potential field whilst in the last code the refined grid is introduced mainly to reduce the computational overhead in short-range force calculation in case of heavy clustering of particles. A smaller gridsize allows to use a smaller neighbouring radius \( r_c \), reducing the number of particles which should be used for the short range force correction. Another advantage of AP3M is the adaptive dynamic creation of submeshes.

This paper presents another adaptive algorithm for a continuous density distribution, based on the well-known multigrid method (Brandt 1977). We shall give a short overview of the method below. The main positive features of the multigrid method are that for the case of the Poisson equation its computational complexity scales as \( O(N) \), it allows one to control the errors inherent in the solution of the Poisson equation, and it lends itself naturally to adaptive refinement. It is also very flexible in applying different boundary conditions and allowing special treatment if necessary.

Compared with the methods described above, our implementation of the adaptive multigrid solver for Poisson equation is physically closest to that of Chan et al, as it finds the local grid potential as the solution for the boundary value problem defined by a coarser grid solution. The main difference is that in the adaptive multigrid method the creation of subgrids is a natural part of the solution process. The local refinements are introduced during multigrid iteration in locations where predefined error estimates demand it. There is a tight interference between the coarse grid solution and a finer grid solution as the coarse grid is used for correction of the solution on the finer grid and vice versa. At the internal boundaries (subgrid interfaces) the values of the potential change during the search for an overall solution, due to changes in the finer grid solution. This is natural, because they should agree despite of different scales of the discretisation and interpolation errors. The point is that the local grid problem should be solved together with the global problem. If we treat it separately, we contradict with global boundary conditions. Natural boundary conditions for the local problem are the Dirichlet’ conditions, as this guarantees that the potential will be continuous at subgrid boundaries.

Comparison of our approach with the Couchman code is not so straightforward due to different approaches in force calculation. There the global (periodic) part of the potential is calculated coarsely and the local (nonperiodic) part is found using isolated boundary conditions, so there is no backreaction to large scales. However, as the force resolution is enhanced further using local direct summation, the backreaction effects could be less important.

Our code has already been applied to the modelling of gravitational microlensing (Suisalu 1991). We describe here its application to cosmological situations, and follow as an example the birth and the subsequent evolution of a typical pancake singularity.
Multigrid methods were introduced for solving boundary value problems by Brandt (1977). Quite comprehensive reviews of multigrid methods can be found in Press et al. (1992) and Brandt (1984).

As the name implies, in this method the problem is solved iteratively, using several sets of grids with different fineness. In the standard case the mesh size ratio of the grids at neighbouring levels $H/h = 2$ and the points of the coarser grid coincide with every second point of the finer grid. The solutions on the coarser grids are used to estimate the smooth components of the final solution and the increasingly finer grids are used to determine details of the solution. We refer readers to the recent introduction in Press et al. (1992, sec. 19.6) and will not give the details of the multigrid method here, noting here only that we use the Full Multigrid algorithm (FMG), the Gauss-Seidel red-black relaxation to smooth the high-frequency error components, and the Full Approximation Storage (FAS) algorithm in order to be able to adaptively introduce finer grid levels.

In standard multigrid problems the source term (r.h.s. of the equations) is usually well determined. In gravitational simulations, however, the density distribution is sampled by discrete mass points, and the problem of determining the matter density on different grid levels is rather complicated. We shall describe our approach below.

We start with a fixed number of levels of uniform grids $G_h$ which cover all the computing domain. On the finest level we compute the density $D_h$ from particles using the cloud-in-cell (CIC) algorithm in the chosen computational volume and make a linked list of particles inside the volume. At the end of the density computation we find also the boundary values of the potential field $f_h$ using this density. In this way we have defined the r.h.s. term of our differential equation for a grid with mesh size $h$:

$$\mathcal{L}_h F_h = D_h,$$

subject to the boundary condition

$$F_h = f_h,$$

on the boundary of $G_h$, where $\mathcal{L}_h$ is in our case the usual 7-point finite difference operator for the Laplace equation.

Having solved this equation approximately on the level $G_h$, using coarser levels to accelerate the solution process, we proceed by adaptively introducing finer grid levels. We have used the value of the local density as the criterion for deciding if there was a need for finer grids, but there are other possibilities. One of the most popular criteria used is the local truncation error (the difference between the discrete $\mathcal{L}_h F_h$ and the analytical $\mathcal{L}F$) which is a natural byproduct of any multigrid solution. To apply the local density criteria we mark during the density computation the gridpoints where the density is higher than a chosen threshold. By analyzing this flag field we define the flag clusters which could be bounded by rectangular grid boxes. As we use at present only non-overlapping grids some postprocessing is necessary.

After allocating new grids, we compute the density on these grids using the particle lists from the coarser level that contains the current grid. We also create a new point list for this particular grid, consisting of those points only that are inside the grid, and we subtract this list from the pointlist of the coarser grid in order to keep points divided between grids. This, firstly, eliminates unnecessary scans over the full
Figure 1. The forces obtained from the multigrid simulation compared to the exact $r^{-2}$ force, versus distance (in grid cell units). The force scale is given in decimal logarithm, but the distance is in base two logarithm. The exact force (labeled TH) is shown by a solid line, the basic mean multigrid force (MG) by a dashed line, and the mean force obtained from a two level finer adaptive solution (AMG) by a dotted line. The two latter forces were found by choosing randomly a position of a massive point and calculating then forces at a number of positions around it. The lines labeled RF and TF show the rms relative radial and tangential force fluctuations with respect to the mean radial force (AMG) obtained with adaptive grids.

particle array, and, secondly, these lists are needed later on if we start to move particles around on our collection of grids.

Besides the density we interpolate the boundary values for the potential from those on the coarse grid in order to specify the local Dirichlet problem. We will repeat the multigrid solution process on new grids using coarser grids and generating new finer grids until the desired resolution is achieved. For every iteration we update the density on coarser levels by the fine level density in order to keep our differential equation consistent on different grid levels. It is possible to use the CIC scheme for density computation, but as the cells at different grid levels cover different volumes, the density estimates differ for the same point at different levels. This introduces additional noise that makes the convergence more difficult. Thus one has either to introduce spatial averaging for coarser grids or to invent a density measurement which gives the same value at the same point in space independent of the mesh size of a grid. In the present code we have used the full weighting scheme that finds densities on coarser grids by averaging it over neighbouring points on the finer grid.

The multigrid solution process stops after making a few additional iterations on the finest level and checking that the changes in the solution are less than the truncation error norm on that level times a small factor (usually 0.01). This assures that the discrete solution is solved down to truncation errors. Usually it takes 3-5 multigrid iterations to get the solution to a desired accuracy. Compared to “exact” solvers, as for example FFT, multigrid is an $O(N)$ procedure, only the multiplicative factor in this estimate could be larger. Nevertheless, multigrid is quite comparable in speed to FFT (Foerster & Witsch 1982) for medium $N$ and could be more efficient in large $N$. There exist several versions of parallel multigrid codes (e.g. R. S. Tuminaro 1989 and Gärtel, Krechel, Niestegge & Plum 1991).

We compare the exact ($r^{-2}$) force-radius law with that obtained by the multigrid code in Fig. 1. The figure shows the averaged pairwise force between a massive particle and massless test particles that were homogeneously distributed in logarithmic radial bins and randomly in angular coordinates. The massive particles were randomly distributed on a finest subgrid. The force was computed using the usual second order differencing scheme. The curve labeled TH gives the exact force for point particles, the curve labeled MG is the usual smooth-field force that has to go to zero for zero radius in order to avoid self-forces for a (CIC) particle. The curve labeled AMG is the result of an adaptive refine-
ment by two levels – as we divide the mesh size by 2 for the next finer grid, this curve is shifted to four times smaller coordinate values, as it should be. We have shown also the rms relative radial and tangential force fluctuations that are around 0.1 per cent of the adaptive mesh force. The maximum deviations from the mean mesh force that we found in our experiments were around 10 per cent, but their percentage was extremely small (only a few cases from an average of about 15000 points per radial bin). These deviations arose at the boundaries of subgrids and are typical for the single massive point case; similar deviations should not occur in the case of a continuous density distribution.

Special care has to be taken with calculation of forces near boundaries of adaptive refinements. As was mentioned above, the boundary values of the potential for local fine grids are obtained by interpolation from the coarser grid. This guarantees the continuity of the solution across the boundary, but not necessarily the continuity of its derivative (force). To smooth possible jumps in force estimates on boundaries of subgrids, we found these forces also by interpolation from the coarser grid. (We used cubic interpolation if there were enough data points, otherwise linear interpolation was used.) As grids are simply a mathematical artefact, particles should not feel the crossing over between grids of different resolution. Our procedure enables particles to change grids smoothly.

Apart from the multigrid Poisson equation solver, other parts of our code are similar to those in a usual PM code. We move particles with a standard leapfrog integrator (we change the timestep, though) on every grid, where the particles are taken from the point list that belongs to that grid. The time step is controlled by the Courant condition: the maximum change of a coordinate should be less than a fraction of the mesh size (we have used one-half). We can have different masses for particles, and for our cosmological sphere we can guarantee mass conservation by reflecting the leaving particles back from the opposite side. The code is not too large, the main modules sum up to about 6200 lines of C.

3 TESTING THE CODE

The code we developed first is meant to be used for studying the evolution of generic types of density singularities. There are only a few of these, all listed in Arnold (1980), and they could be thought of as typical progenitors of large-scale structure, describing the regions of the first collapse. On scales where there has yet been no significant interaction of neighbouring elements of structure (superclusters), the models of singularities could describe the actual dynamics and geometry of structure.

In order to understand the evolution of specific singularities we have to study first the case of isolated singularities, using vacuum boundary conditions. In order to minimize the influence of the geometry of the computational volume on the results we have to work in a sphere. Of course, working in a cube would be much simpler, but we have not been able to get rid of the ghosts of the cube in the final configurations.

Using isolated boundary conditions on a sphere could seem to be contrary to the usual cosmological practice of periodic boundary conditions. In the case of the tree-code, where vacuum boundaries arise naturally, people have taken enough trouble to modify the code to better mimic periodic
boundaries (Bouchet & Hernquist 1988, Hernquist, Bouchet & Suto 1991). Thus we have to check if our isolated region models the evolution of structure in cosmology fairly enough.

As there are really no perfect numerical methods to use for comparison, one should use the exact solutions for the evolution of structure. There are only three of these—one for the linear regime and two nonlinear solutions, one for a spherical top-hat collapse and the other for a one-dimensional plane wave. The latter solution clearly cannot be used in our case, so only two remain.

We use the usual cosmological equations for the evolution of structure for the $\Omega = 1$ universe in comoving coordinates $x$, connected with the physical coordinates $r$ by $r = a(t)x$, where $a(t)$ is the scale factor that describes the expansion of the universe. We choose this function as the new time coordinate, and write our basic equations as

$$\Delta \Phi = \delta,$$

where $\Phi$ is a suitably normalized gravitational potential and $\delta$ is the usual density contrast, and

$$\frac{du}{da} + \frac{3}{2a} u = -\frac{3}{2a^2} \text{grad} \Phi,$$

where $u = dx/da$. Our equations are similar to those used by Matarrese, Lucchin, Moscardini & Saez (1992).

In order to test for linear evolution, we have to generate appropriate initial data—a realization of a gaussian random field with a given power spectrum. One starts usually with the power spectrum of the density contrast:

$$P(k) = |\tilde{\delta}(k)|^2 |k|^{-n},$$

where $k$ are the wavenumbers and $\tilde{\delta}$ are the Fourier amplitudes of the density contrast. We proceed following the method described by Nusser & Dekel (1990).

In the linear approximation the movement of particles can be described by the formula derived by Zeldovich (1970):

$$x = q + a u(q),$$

where $q$ are the initial (Lagrangian) coordinates and the velocity $u(q)$ depends on $q$ only. This leads to the (linear) relation between density and velocity

$$\delta(q) = a \text{div}_q u.$$

The latter relation can be satisfied if the Fourier amplitudes of the velocity are

$$\tilde{u}(k) = \frac{k}{k^2} \tilde{\delta}(k).$$

Having chosen the complex Fourier amplitudes for the density contrast $A(k) + iB(k)$ as random Gaussian numbers with the distribution $N(0, P(k))$ on an appropriate grid $k_i$ in wavenumber space ($-N/2 \leq k_i \leq N/2$, where $N$ is the resolution of the grid), we can form the Fourier amplitudes of the velocity by (4) and find the velocity field in real space by an inverse Fourier transform. We use the Fourier transform algorithm for real 3-D data from Press et al. (1992).

For tests we used a low-resolution model, 28 cells for the diameter of a sphere (32 for the surrounding cube that we use to fix the boundary conditions), in order to clearly see the influence of a discrete grid in both methods. If we wish to get a good representation of small perturbations the density has to be generated from a regular grid. We chose 8 particles per grid cell which made the total particle number rather large, $64^3$ for the PM-cube and about 100000 for the sphere.

For the initial density spectrum we chose white noise, in order to see better the damping of high-frequency modes. We generated the initial velocity field as described above,
and generated the coordinate displacements from a regular \( q \)-grid, choosing them to be proportional to velocities and normalized to a fixed (small) rms displacement amplitude. As we work in a sphere, but the FFT works in rectangular regions, we first find the displacements in a \( 32^3 \) cube and then set the displacements outside our sphere to zero.

We have to take a little more trouble about the velocities. As the high-frequency modes of the velocity that we generated cannot be caused (and changed) by the potential found from the same grid, our initial state is too hot. To remedy this we used the quiet start recipe proposed by Efstathiou et al. (1985). We recalculated the velocities, finding the density, solving for the potential and using the linear approximation relation between acceleration and velocity, which for our case \((\Omega = 1)\) reads:

\[
\mathbf{u} = \frac{1}{a^2} \nabla \Phi.
\]

(5)

In the linear case we can check for the evolution of the velocity and density fields. If the initial velocities are given by the above formula, it is easy to see that the velocities have to remain constant, \(d\mathbf{u}/dt = 0\), (see also (3)). The continuity equation

\[
\frac{d\delta}{da} + (1 + \delta) \text{div} \mathbf{u} = 0
\]

tells us that the density contrast has to grow linearly with \(a\).

We started from the rms amplitude 0.025 for the density contrast and followed the evolution of structure from \(a = 1\) until \(a = 10\). The results are not too good – see Figs. 2 and 3. In Fig. 2 we show the ratio of our rms density contrast to the expected value from linear theory (the curve labeled MG). As is seen, the evolution lags behind the true rate and the difference reaches about 2.5 times at the scale factor \(a = 10\). A similar picture can be seen in Fig. 3 – while the rms velocity is expected to remain constant, it actually drops in time (although the differences are smaller than these for the density).

This discrepancy is typical for smooth-field simulations, and is mainly caused by damping of high-frequency modes. Bouchet et al. (1985) have studied it extensively in the case of PM-codes. Most of the reasons for the damping, the CIC
density assignment scheme and a finite grid size, are present in our code too. In order to have standard errors to compare with, we solved the same problems by a standard PM-code. There are, certainly, better codes around, but simple PM is a pure smooth-field algorithm, similar to multigrid, in contrast to improved PPPM-type codes. We used a $32^3$ grid with periodic boundary conditions, the same number of particles (8) per cell, and started from the same initial state. The corresponding curves in Figs. 2 and 3 (labeled PM) look similar, the density contrast behaves a little better in the PM-code, the velocity drops a little faster, but differences between the two codes are small.

We can understand what is happening a little better if we look at the evolution of the density on different scales. In Fig. 4 we give the ratio of the density spectra at the epoch $a = 10$ to that at the start of the calculations, normalized by the theoretical growth factor. If we look at the PM-code curve we see that the largest scales closely follow the linear growth law, but the difference comes in at smaller scales. The MG-curve shows us that this damping is also present in multigrid, although to a lesser extent. As we calculated our spectra in a $32^3$ cube, the largest wavelengths present in a cube do not fit into our sphere and we have a slight drop in the MG-case these scales, but otherwise the MG-code with isolated boundary conditions describes the evolution of small perturbations at least as well as does the periodic PM-code.

The second test we made is the highly nonlinear spherical tophat collapse. This is also a textbook problem that can be solved exactly (see, e.g., Padmanabhan 1993, sec. 8.2). We shall check for the moment of collapse $a_{\text{coll}}$ that is predicted to be

$$a_{\text{coll}} = \left( \frac{3\pi}{2} \right)^{2/3} a_i \delta_i,$$

where $\delta_i$ is the initial density contrast for the sphere (at the scale $a_i$) and the only approximation used to get this result is the requirement $\delta_i \ll 1$.

As the dependence of the collapse time on the initial density contrast is rather strong, one has to take care when generating the initial distribution. We generated, first, points on a regular mesh inside a sphere (with an initial grid of 28 cells per diameter), 8 points per cell, and left a hollow sphere inside, with a radius of 0.75 of that of the large sphere. We then filled this sphere also with points on a regular mesh, but with a slightly smaller mesh size. In order to compare the results we generated a similar sphere inside a $32^3$ cube, with a radius of 0.75 of the half-cube size, and followed its evolution by our PM-code. Although the central densities were the same in both cases, the initial mean densities and density contrasts were different (the cube had a larger volume) and the predicted collapse times differ also.

The boundary conditions were fixed (zero for the zero
total mass in a sphere) for the multigrid and periodic for
the PM-code. We estimated the moment of collapse by find-
course, the observed supercluster chains and the knots they emerge from also look similar to some extent.

There is a mathematical basis for this similarity – if we agree that visible structure forms in locations of the highest density (for cold dark matter this means an infinite density), then we should look for possible classifications of density singularities. This has been, fortunately, already done, and the corresponding theory is called the theory of singularities of Lagrangian mappings. Matter flows in a gravitating medium follow Lagrangian mappings, so this theory is relevant to the formation of structure. This was realized at least ten years ago, and these mappings have been used in cosmology by Arnold, Shandarin & Zeldovich (1982). A very important point is that the number of different mappings in the generic case (the number of types of structure elements) is surprisingly small, only six in 3-D space and four in 2-D (Arnold 1980). If we look at the evolution of these singularities in time, we are dealing with a metamorphosis of the singularities, and there are from two to five types of evolution for every basic singularity type, which is a small number.

The singularity mappings describe the motion of matter only until flows intersect and it is not clear how useful they are afterwards. And as these mappings are local, we do not know how long they will remain so before being distorted by interaction with neighbouring singularities.

This all is a subject of fascinating study, and we can use the basic types of mappings to find the initial conditions for the emerging structure. As they are generic, these are the structures that must be most common both in the sky and in the simulations. The code that we can use to study the formation of structure has clearly to be as high-resolution as possible. This was the motive for starting the development of the present code.

As usual in a new field, there is a mass of new problems and intricacies here. We shall demonstrate how our code works and shall describe these problems using the most familiar Lagrangian singularity – the Zeldovich pancake. It belongs to so-called type $A_3$, and the birth of a pancake is described by the metamorphosis $A_3(-, +)$.

This mapping can be described by the coordinate transformation (from Lagrangian coordinates to the normal Eulerian space)

\[
\begin{align*}
x_1 &= 4q_1^3 + 2(q_2^2 + q_3^2 - t)q_1, \\
x_2 &= q_2, \\
x_3 &= q_3,
\end{align*}
\]

(6)

where $x_1$ are the Eulerian and $q_1$ the Lagrangian coordinates. This mapping is meant to be used near the zero point of coordinates, that is the place where the pancake is born – the density is

\[
\rho(x(q)) = |\frac{\partial x}{\partial q}|^{-1} = |12q_1^2 + 2(q_2^2 + q_3^2 - t)|^{-1},
\]

and, when time grows from $-\infty$ towards the future, this density first becomes infinite at $t = 0$ in $x = 0$.

The velocities for this mapping can also be found (Su-

\[
\begin{align*}
v_1 &= -2q_1, \\
v_2 &= 0, \\
v_3 &= 0
\end{align*}
\]

(7)

(all $A$-type singularities are essentially one-dimensional).

These formulas are in principle all that one needs to set up a pancake birth simulation. The remainder are technical details, but they are rather important. The main problem is
that the mapping is nonlinear, and if we want to model it, we must restrict the mapping to a finite region – a sphere is the best choice, as it minimally distorts the final results. The mappings (6) and (7) are free to change using any diffeomorphism we want, as this does not essentially change the mapping in the centre of the coordinates. However, if we want to study as large a region around the centre as possible, the modification must be minimal. Our choice is

\[ x = f(q)(1 - q^2/R^2) + q q^2/R^2, \]

(8)

where \( f(q) \) is the mapping (6) and \( R \) is the radius of the sphere. The velocities are changed similarly:

\[ v = w(q)(1 - q^2/R^2), \]

(9)

where \( w(q) \) is the original (Eulerian) velocity in the Lagrangian coordinates (7). This is not a perfect solution, as it gives us a constant density and zero velocities on a Lagrangian sphere \( q^2 = R^2 \), and thus distorts the geometry of our Eulerian computational volume. We have fought this by choosing a small radius, \( R = 0.05 \), as the region inside it maps practically into an Eulerian sphere (our initial time parameter \( t = 0.4 \)). For simulations we choose the scale factor \( a \) as the time variable. The times \( a \) and \( t \) can be connected to each other by any monotonic transformation. We are also at liberty to choose the velocity amplitude – this amounts to changing the time unit. We did this using the quiet start recipe – we built the point distribution by the mapping (8), found the density (the maximum initial density contrast was 0.308), solved for the potential and used the linear evolution formula (5) to find the initial dynamical velocities. We used both the dynamical velocities and the velocities from the mapping scaled down to make the maximum velocities coincide in both cases.

The initial grid is the same as we used for the spherical collapse, a sphere with diameter of 28 cells (in Lagrangian coordinates). In order to better see the details of the structure we choose 27 points per grid cell, distributed regularly, a total of 324609 points. If the number of points per cell gets larger than 70 we create a finer rectangular subgrid with double linear resolution. We have limited the number of refinement levels to five – this limit is imposed mostly by the noise that accumulates during the run. This means that the effective spatial resolution in the central subgrid is \( 1/1024 \). Also, in this case the boundary conditions will change in time – we found them by direct summation over the initial \( (32^3) \) grid. We use the same equations of motion as before, working in the \( \Omega = 1 \) cosmology.

We illustrate the results by a series of figures. As the results for the initial velocities from the mappings and from the dynamics did not differ much, we shall use the case of the mapping (truly unidirectional) velocities. The figures all refer to the time \( a = 13.6 \) (we started with \( a = 1 \)). The first series of figures (Fig. 6a-d) show the distribution of mass points in a thin (one cell of the basic grid) slice along the \( x_1-x_2 \) coordinate plane. We have also shown the borders of the subgrids, and the change of scale can be followed from the coordinate values.

As this is already a well advanced state of collapse, we see a fairly rich substructure in the figures. Fig. 6a covers the whole slice, and the pancake is clearly seen in the centre. The edges of the pancake are formed by the turnback points of the first particle stream that has passed the pancake plane. Only four subgrids can be seen in this figure, the
The central $x_1$-$x_2$ slices of the simulation of the birth of a pancake ($A_3(-, +)$ metamorphosis) at the epoch $a = 13.6$. The simulation started at $a = 1$ with a maximum density contrast $\delta_{\text{max}} = 0.308$ and with the velocity amplitude found from linear dynamics. Panels (a–d) show the particle distribution in a slice on different scales; the boxes show adaptive subgrids.

The lenslike overall geometry of the collapse is not caused by its living in a sphere, this is a generic property of this type of singularity and is determined by the mapping itself. It is possible to distort the shape a little, in principle, but we discovered that such a distortion will not live long — the mapping we used is generic.
Fig. 7. The central $x_2$-$x_3$ slices of the same density distribution as in Fig. 6. Panels (a-d) show the particle distribution in a slice on different scales; the boxes show adaptive subgrids.

Fig. 6b (coinciding with the third subgrid) shows the central part of the collapsed region – we see two perpendicular planes and the formation of ellipsoidal shells – these are the turnback regions of smaller-scale flows, and the collapse tends to become more spherical. This differs from the picture seen in high-resolution 2-D simulations by Beacom et al. (1991), and is probably caused by the fact that 3-D gravitation is in general more effective than 2-D gravitation.
Figure 8. A three-dimensional representation of the same density distribution as in Figs. 6 and 7. A region of space between chosen density levels is cleared to show the inner density levels (see Table 1). Panels (a–d) show the density distribution in increasingly smaller scales - the small cube in the centre shows the size of the next panel.

These shells cannot be caused by the spherical boundary, growing temperature, this effect can be seen in the centre only (Fig. 6d).

The flows live in the centre where they do not feel the large-scale symmetry. They cannot either be the result of the

Fig. 6c shows the central matter distribution in more
detail—we see a much smaller pancake in the centre, with a size about 80 times smaller than that of the largest pancake. The last figure in this series, Fig. 6d shows no more detail although the last grid has 13 × 17 cells in this plane. This is probably caused by the (numerical) heating during the collapse. The central density contrast is \(1.6 \cdot 10^5\), this value can be taken to characterize the dynamical resolution of the simulation.

Figs. 7a-d show a slice from the \(x_2-x_3\) plane, using similar scales as the previous series. In Fig. 7a we see the whole plane, where the outer density enhancement is the edge of the main pancake. More inner density ridges can be seen in Fig. 7b. They are rather well resolved in Fig. 7c, and the central part can be seen as a small hot lens at Fig. 7d. The spokes along the coordinate axes that are evident in all these figures are either caused by the anisotropy of the 7-point difference operator, or by the CIC density assignment algorithm that gives enhanced densities along the coordinate planes and axes. These spokes are not too prominent, however.

In the last series of figures (Fig. 8a-d) we have tried to show the 3-D density distribution. Each of these figures is a 3-D representation of three density levels, cut in half by an \(x_1-x_2\) coordinate plane (in these figures the coordinate \(x_1\) is vertical). The smaller inner cube in Figs. 8a-c shows the size of the large cube in the next figure. The density values for the different level surfaces (the outer one, the inner surface of the “shell” and the outer surface of the central detail) are given in Table 1. The surfaces are better seen in Fig. 8a, where the density resolution is \(1/32\) of the cube size. In Figs. 8b-d the resolution is \(1/24\) and is probably a little too low for the IDL (Interactive Data Language—a graphical software package from Research Inc.) shade-volume command to manage. Of course, in the real density distribution there are no holes, they are cut out in the figures only to help to visualize the continuous density distribution.

Fig. 8a starts showing us the basic sphere and the primary pancake inside. The outer shell is, in fact, continuous, and looks striped by the IDL efforts. Fig. 8b shows a second pancake inside the first one (look at the sizing cubes) and a high-density detail at the centre. This detail is better resolved in Fig. 8c, showing a density enhancement that is oriented perpendicularly to the original pancake plane (there is a trace of it in Fig. 6b, the horizontal density enhancement). The plane itself has also rather high density here.

The central core of Fig. 8c is resolved in Fig. 8d—it is a lenslike density concentration along the original pancake plane (the vertical density enhancement in Fig. 6b), and the small pancake of Fig. 6d lives in its centre.

As is seen, the inner regions grow more and more irregular, the smaller the scales and the larger the densities. This could be due to a number of reasons. The first of them is the numerical heating caused mainly by the force fluctuations at the edges of subgrids. Another is the fact that we have probably not taken proper care when arranging the initial mappings, and as a result these are not cool enough.

5 CONCLUSIONS

We have presented here an adaptive multigrid code for gravitational simulations and tested it for cosmological problems. The multigrid approach lends itself naturally to adaptive refinement and does not impose any restrictions on the bound-
Table 1. Density levels in the 3-D configuration

| panel | outer | inner  | central |
|-------|-------|--------|---------|
| a     | 0.02  | 0.7    | 10.5    |
| b     | 14.4  | 33.5   | 173.3   |
| c     | 185.2 | 924.4  | 11090.0 |
| d     | 9244.0| 46220.0| 154000.0|

ary conditions to be used. It does not much use more memory than the popular cosmological codes and it is fast enough to be used on present computers.

The list of possible enhancements is rather long. We have already implemented the case of periodic boundary conditions that are more suitable to simulate the evolution of global structure. We are planning to use better difference operators in order to get more isotropic forces, and we have ideas on how to improve the density assignment algorithm.

The speed of our algorithm is still lower than that of the FFT, but we can use the potential and grids from the previous time step to enhance the convergence – this can be done because we keep the potential separate from the density field. Probably the use of block time will also speed up the code. We determine our boundary conditions at present by direct summation over the grid – we can either use a coarser grid for this or use a FFT solver. I.S. has implemented the code also on a parallel computer (2-D case on a CM-200), but this implementation needs further work.

As for the present application of following the structure of singularities, this has shown us the importance of setting up clean initial conditions and getting rid of noise for truly high-resolution simulations. A typical example is the CIC density assignment scheme – the fact that it may give spurious density enhancements does not worry anybody so far as we are using noisy initial conditions, but in the present case its deficiencies were obvious.

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