

# Vectorization and Parallelization of the Adaptive Mesh Refinement $N$ -body Code

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## Abstract

In this paper, we describe our vectorized and parallelized adaptive mesh refinement (AMR)  $N$ -body code with shared time steps, and report its performance on a Fujitsu VPP5000 vector-parallel supercomputer. Our AMR  $N$ -body code puts hierarchical meshes recursively where higher resolution is required and the time step of all particles are the same. The parts which are the most difficult to vectorize are loops that access the mesh data and particle data. We vectorized such parts by changing the loop structure, so that the innermost loop steps through the cells instead of the particles in each cell, in other words, by changing the loop order from the depth-first order to the breadth-first order. Mass assignment is also vectorizable using this loop order exchange and splitting the loop into  $2^{N_{\text{dim}}}$  loops, if the cloud-in-cell scheme is adopted. Here,  $N_{\text{dim}}$  is the number of dimension. These vectorization schemes which eliminate the unvectorized loops are applicable to parallelization of loops for shared-memory multiprocessors. We also parallelized our code for distributed memory machines. The important part of parallelization is data decomposition. We sorted the hierarchical mesh data by the Morton order, or the recursive N-shaped order, level by level and split and allocated the mesh data to the processors. Particles are allocated to the processor to which the finest refined cells including the particles are also assigned. Our timing analysis using the  $\Lambda$ -dominated cold dark matter simulations shows that our parallel code speeds up almost ideally up to 32 processors, the largest number of processors in our test.

**Key words:** methods:n-body simulations — cosmology: large-scale structure of universe

## 1. Introduction

By introduction and invention of the high resolution cosmological  $N$ -body methods, such as the particle-particle-particle-mesh ( $P^3M$ ) method (Efstathiou & Eastwood 1981; Hockney & Eastwood 1988), cosmological  $N$ -body simulations became fundamental and powerful tools to investigate the structure formation processes in the universe (e.g. Davis et al. 1985). Besides, parallelization schemes for the  $P^3M$  method were developed by some authors (Ferrell & Bertschinger 1994; Ferrell & Bertschinger 1995; MacFarland et al. 1998). Since the  $P^3M$  method is memory efficient, the number of particles in  $P^3M$  simulations reached to  $10^9$  (Evrard et al. 2002). However, the  $P^3M$  method is not almighty. In the  $P^3M$  method, forces from nearby particles are calculated by the direct sum method. Hence, as the system gets clustered, direct sum part dominates the whole CPU time and the computational cost grows rapidly. In order to resolve this defect of the  $P^3M$  method, some methods are proposed, such as the adaptive  $P^3M$  method (Couchman 1991) which uses adaptive meshes in the dense regions, and the tree particle-mesh method (Xu 1995; Bagla 2002; Bode & Ostriker 2003) which uses tree method (Barnes & Hut 1985) instead of the direct summation. The adaptive mesh refinement (AMR) method which places hierarchical meshes where required, is also one of those methods. Some groups place rectangular structured mesh (Villumsen 1989; Anninos et al. 1994; Suisalu & Saar 1995; Gelato et al. 1997; Norman & Bryan 1999) while other groups place non-rectangular structured mesh (Kravtsov et al. 1997; Knebe et al. 2001; Yahagi & Yoshii 2001; Teyssier 2002). A review of cosmological  $N$ -body codes is found in Bertschinger (1998).

For the tree method, there are various vectorization schemes (Barnes 1990; Hernquist 1990; Makino 1990) and parallelization schemes for distributed memory machines (Salmon 1990; Warren & Salmon 1993; Dubinski 1996; Yahagi et al. 1999; Springel et al. 2001). Moreover, there is a parallel tree code (Makino 2004) accelerated by the special purpose computer, GRAPE (Sugimoto et al. 1990; Kawai et al. 2000; Makino et al. 2003). On the other hand, Fryxell et al. (2000) discussed the parallelization scheme of their AMR hydrodynamics code. However, parallelization of the  $N$ -body part is not described extensively. Hence, we describe our vectorization and parallelization scheme in this paper. Some may think that the vectorization is obsolete technique. However, the Earth Simulator which had been the fastest computer in the world until November 2004, is a vector-parallel type computer, and the vectorized code is easily parallelized for shared-memory multiprocessors.

In §2, we review the data structures and algorithms of our AMR  $N$ -body code to define the terminologies used in the following sections. In §3, we describe the vectorization scheme of the AMR  $N$ -body code. In §4, we describe the parallelization scheme of the AMR  $N$ -body code, and the timing analysis of the parallelized code is described in §5. Finally, we summarize this paper in §6.

## 2. AMR $N$ -body code

We vectorized and parallelized the shared time step version of the AMR  $N$ -body code described in Yahagi & Yoshii (2001). We review the AMR  $N$ -body code briefly in the beginning of this section. Then, since adopted data structure determines the algorithm suited to the problem, and vice versa, we review the data structures used in our AMR  $N$ -body code to define terminologies used in the following sections, prior to discussing the vectorization and parallelization schemes.

### 2.1. Structure of the code

The structure of the code is iteration of the following routines: velocity update stage of the leap-frog integration by a half time step, position update by a full time step, reconnection of the linked list of particles, addition and deletion of hierarchical meshes, mass assignment from particles to meshes, the Poisson solver, and velocity update by a half time step. Force on particles at velocity update is calculated by interpolation of force on meshes onto particles. Position update needs only position and velocity of particles, because the leap-frog integration is adopted. After position update is carried out, some particles go across the cell boundaries. Hence, in order to make the linked list consistent, the linked list of particles are reconnected by picking up particles which go across the boundaries and inserting them to the linked list beginning from the finest cells which include them and will not be deleted at the mesh modification stage. Mesh generation and deletion is basically operation using mesh data. However, when cells are generated, particles in the parent cells are inherited to the newly generated cells, and at this stage, particle data are also used. After this mesh modification, force on mesh is calculated, finally. First density on meshes are calculated using the position of particles in the cells. Then, potential on meshes are calculated from density on meshes by the Poisson solver. Force on meshes are calculated by taking difference of potential at neighboring cells.

We keep the particle linked list consistent as follows: First, particles are grouped into the stay particles and the leaked particles. The stay particles do not move across the boundary of the cell during the time step, while the leaked particle move across the boundary. The leaked particles are removed from the linked list, then added to a linked list starting from the neighboring cell. However, the neighboring cell might be a buffer cell although the starting cell of the particle linked lists is restricted to a refined cell. (See §3.3 for the definition of a buffer cell.) In that case, those leaked particles are passed to the parent cell until they reach to a refined cell or the base mesh. On the other hand, the neighboring cell which is to be connected by the leaked particles may have refined child cells. As mentioned above, starting cell of the particle linked list must not have any child refined cells. In that case, those leaked particles are passed to the refined child cell until they reach to the finest refined cell including them. At this stage, linked lists are consistent with the distribution of particles and meshes, but cell classification and the refinement criteria could be inconsistent. As the first step of the next stage, cells satisfying the refinement criteria are flagged, If starting cells of particle linked lists are not flagged or the buffer child cells are flagged, again, particles in such cells are passed to the parent cell or the child cell until they reach to the base mesh or the finest refined cell. Then, at each level from the base level to the deepest level, unused cells are removed and new refined cells are added. At the same time, particles connected from the parent cell of the newly refined cells are handed to the new child refined cells. Finally, hierarchical meshes are sorted in the Morton order. Through these procedures, the linked lists are consistent with the distribution of particles and meshes and meshes classification is also consistent with the refinement criteria.

We adopted the Cloud-in-cell (CIC) scheme (Hockney & Eastwood 1988) for force interpolation and mass assignment. In the CIC scheme, mass of a particle is assigned to eight vertices of the cell including the particle, and force defined at eight vertices of the cell is interpolated on the particle. Mass assigned to cells are restricted to the parent cells by the CIC scheme from the finest mesh level to the base mesh level. For the Poisson solver, we use red-black Gauss-Seidel relaxation. Further details of our serial AMR  $N$ -body code are described in Yahagi & Yoshii (2001).

As described in the next subsection, particles are connected to the finest refined cell. Density at cells which do not have a child cell octet but include particles are calculated by the CIC scheme. On the other hand, density at cells which have a child cell octet is calculated by the full weighting fine-to-coarse operator from the child cells and their neighbor cells:

$$\begin{aligned} \rho_{2l,2m,2n}^{L-1} &= F(\rho^L)_{2l,2m,2n} \\ &= \frac{1}{8} \rho_{2l,2m,2n}^L + \\ &\quad \frac{1}{16} (\rho_{2l-1,2m,2n}^L + \rho_{2l+1,2m,2n}^L + \rho_{2l,2m-1,2n}^L + \rho_{2l,2m+1,2n}^L + \rho_{2l,2m,2n-1}^L + \rho_{2l,2m,2n+1}^L) + \end{aligned}$$

**Table 1.** Position of 1-th child cell

1	Position of cell
0	( 2*i , 2*j , 2*k )
1	( 2*i , 2*j , 2*k+1 )
2	( 2*i , 2*j+1, 2*k )
3	( 2*i , 2*j+1, 2*k+1 )
4	( 2*i+1, 2*j , 2*k )
5	( 2*i+1, 2*j , 2*k+1 )
6	( 2*i+1, 2*j+1, 2*k )
7	( 2*i+1, 2*j+1, 2*k+1 )

$$\begin{aligned}
& \frac{1}{32}(\rho_{2l-1,2m-1,2n}^L + \rho_{2l+1,2m-1,2n}^L + \rho_{2l-1,2m+1,2n}^L + \rho_{2l+1,2m+1,2n}^L + \\
& \quad \rho_{2l-1,2m,2n-1}^L + \rho_{2l+1,2m,2n-1}^L + \rho_{2l-1,2m,2n+1}^L + \rho_{2l+1,2m,2n+1}^L + \\
& \quad \rho_{2l,2m-1,2n-1}^L + \rho_{2l,2m+1,2n-1}^L + \rho_{2l,2m-1,2n+1}^L + \rho_{2l,2m+1,2n+1}^L) + \\
& \frac{1}{64}(\rho_{2l-1,2m-1,2n-1}^L + \rho_{2l+1,2m-1,2n-1}^L + \rho_{2l-1,2m+1,2n-1}^L + \rho_{2l+1,2m+1,2n-1}^L + \\
& \quad \rho_{2l-1,2m-1,2n+1}^L + \rho_{2l+1,2m-1,2n+1}^L + \rho_{2l-1,2m+1,2n+1}^L + \rho_{2l+1,2m+1,2n+1}^L).
\end{aligned}$$

It is possible to prove the following equation:

$$\begin{aligned}
\rho_{\text{CIC}}^L &= F(\rho_{\text{CIC}}^{L+1}) \\
&= F(F(\rho_{\text{CIC}}^{L+2})) \\
&\vdots \\
&= F^N(\rho_{\text{CIC}}^{L+N}) \\
&\vdots
\end{aligned}$$

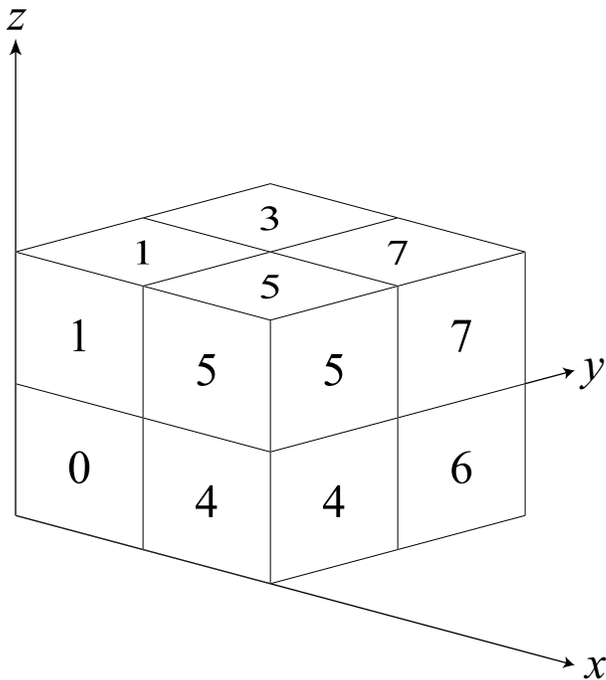
where  $\rho_{\text{CIC}}^L$  is the density field derived by the cloud-in-cell scheme directly.

## 2.2. Data structure used in the code

Next, we describe the data structure used in our code. First, we divide data into two categories: particle and mesh. Data structure of particles is fairly simple. Particles are just stored in arrays. Data structure of meshes is much more complicated. First, meshes are divided into two groups further: base mesh and hierarchical mesh. Data structure of the base mesh is simple. The base mesh is stored in arrays. Accessing neighboring cells at the base mesh is trivial, if they have a frame, or ghost points. For example, if the size of base mesh is  $N^3$ , arrays of  $(N+2)^3$  are declared for the base mesh, and inside  $N^3$  cells are used for calculation. Data structure of the hierarchical meshes is more complicated. Eight cells refined from the same parent cell are bundled into a single data structure (Khokhlov 1998), we call it cell octet. Cells and cell octets are also stored in arrays. Thus, a cell whose index is  $\text{ic}$  belongs to a cell octet whose index is  $\text{ic} \gg N_{\text{DIM}}$ , where  $N_{\text{DIM}}$  is the number of dimension and  $\gg$  represents a right shift operator. For example, when  $N_{\text{DIM}}=3$ , a cell  $\text{ic}$  belongs to a cell octet  $\text{ic}/8$ . Since cell octets have indices to the six neighboring cell octets, it is easy to access the neighboring cell octets from a cell octet. However, it is not so simple to access six neighboring cells from a cell. First, we numbered siblings in a cell octet in the Morton order (see §4.1). For example, consider a cell octet placed at  $(x, y, z) = (2*i, 2*j, 2*k)$ , where  $i, j$ , and  $k$  are arbitrary integers. At this time, 1-th child cell of the cell octet is placed at  $((2*i \mid ((1 \& 4) \gg 2), 2*j \mid ((1 \& 2) \gg 1), 2*k \mid (1 \& 1))$ , as given in Table 1 (see also Fig. 1). Here,  $\&$  and  $\mid$  represent a bitwise AND operator and a bitwise OR operator, respectively. Then, six neighboring cells of the 0th child are given in Table 2. It is trivial to extend the above relation to the neighbors of other seven child cells (see also Fig. 2).

**Table 2.** Neighbor cells of 0th child cell

Neighbor's position	Position of cell octet	index of cell
$(2*i-1, 2*j, 2*k)$	$(2*i-2, 2*j, 2*k)$	4
$(2*i+1, 2*j, 2*k)$	$(2*i, 2*j, 2*k)$	4
$(2*i, 2*j-1, 2*k)$	$(2*i, 2*j-2, 2*k)$	2
$(2*i, 2*j+1, 2*k)$	$(2*i, 2*j, 2*k)$	2
$(2*i, 2*j, 2*k-1)$	$(2*i, 2*j, 2*k-2)$	1
$(2*i, 2*j, 2*k+1)$	$(2*i, 2*j, 2*k)$	1

**Fig. 1.** Position of eight cells in a cell octet. Eight sibling cells are numbered by the Morton order and their position value is given in Table 1.

There is yet another data structure: the linked list of particles. Particles in the same cell are connected by a linked list. If a refined cell whose parent cell satisfies the refinement criteria does not have any child refined cells, such refined cell has the index of the head particle of the linked list consisting of particles in the cell.

### 3. Vectorization

With pipeline processing, vector processors calculate loops in codes much faster than usual scalar processors on personal computers and work stations. Moreover, there are many vector processors in Japan, including the super-computer system of the National Astronomical Observatory of Japan which is exclusively served to the astronomical community, and the Earth Simulator. Thus, we decided to vectorize our AMR  $N$ -body code. In order to bring out the performance of those vector processors, we must eliminate as many unvectorizable loops in the code as possible. Most often the loops that are unvectorizable are recursive loops, in which store operation to a certain address precedes load operation from the same address during the loop. We describe the vectorization schemes of our AMR code in this section. Hereafter, C-notation is used for code description unless otherwise mentioned, and some codes are rewritten for this paper to make clear the argument.

Incidentally, some may think that vectorization is an obsolete technique. However the scheme described in this section to eliminate the unvectorized loops is in common with the parallelization scheme for shared-memory multiprocessors.

#### 3.1. Loop types

Before discussing the vectorization scheme, we classify loops in our code into three categories from the data structure point of view. The first category consists of those loops which treat particle data only. Loops in the integration of

	1	3	1	3	1	
	0	2	0	2	0	
		$(2*i, 2*j, 2*(k+1))$				
	1	3	1	3	1	
	0	2	0	2	0	
$(2*i, 2*(j-1), 2*k)$		$(2*i, 2*j, 2*k)$		$(2*i, 2*(j+1), 2*k)$		
	1	3	1	3	1	
	0	2	0	2	0	
		$(2*i, 2*j, 2*(k-1))$				

**Fig. 2.** Position of neighboring cells in  $y$ -direction and  $z$ -direction of cells in a cell octet placed at  $(2*i, 2*j, 2*k)$ . Position of cells and cell octets is defined at their lower left corner. For example, four of six neighboring cells of the 0-th child cell in the cell octet placed at the center in this figure are shown (see also Table 2).

particle trajectory are classified into this group. We call loops categorized into this group particle loops. These loops are vectorized easily, and usually compilers vectorize them automatically. The second category includes those loops which treat mesh data only. We call those loops mesh loops. Loops in the multigrid Poisson solver are grouped here. Loops treating only the base mesh are vectorized automatically. Loops treating the hierarchical meshes need some tricks to be vectorized. However schemes for those loops are not as elaborate as schemes for loops categorized into the third group, that treat particles and mesh simultaneously in the same loop. Mass assignment and force interpolation belong to this category. We call this group particle-mesh loops. In the rest of this section, we discuss the vectorization schemes for each type of loops.

### 3.2. Particle loops

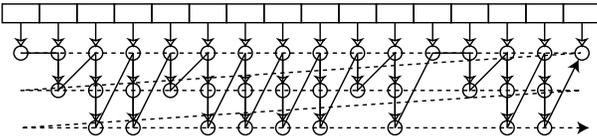
There is only one function which has a particle loop. That is the position step of the leap-frog time integration of particles. Usually, this part is vectorized automatically by a compiler.

### 3.3. Mesh loops

Vectorization of loops for the base mesh is trivial. The mesh loops for the hierarchical meshes is more complicated than that for the base mesh. This complexity is partly due to the data structure mentioned in §2. Another source of complexity is the difference of the boundary condition. The boundary of the base mesh is periodic, while the boundary of hierarchical meshes is arbitrarily shaped and non-periodic. There are two types of hierarchical meshes. The first type is refined cells whose parent cells satisfy the refinement criteria, and the second type is buffer cells which are placed to wrap the refined cells to provide boundary conditions for refined cells. Since refined cells are always surrounded by other refined or buffer cells, all potential values assigned to the refined cells are updated at each iteration of the red-black Gauss-Seidel iteration, for example. On the other hand, potential is updated only in a fraction of buffer cells. Why is the potential updated in some buffer cells and not others? Because potential is defined at cell-corners. Each refined cell octet has potential of only 8 points, while each cell octet has 27 cell-corners. If eight neighboring cell octets of a refined cell octet are also refined cell octets, potential at remained 19 points are updated by those cell octets. However, if there are neighboring buffer cell octets, it is not so simple. For clarity, consider a one dimensional case and potential is defined at left boundary of cells. In this case, when the right neighboring cell of a refined cell is also refined cell, there is no problem. However, if the right neighboring cell is a buffer cell, potential of that buffer cell must be updated during the iteration of the Poisson solver. On the other hand, for a buffer cell whose right neighboring cell is a refined cell, potential at that buffer cell need not be updated. Thus, potential at buffer cells whose left neighboring cell is a refined cell must be updated, while potential at other buffer cells need not. Since potential at some buffer cells are updated while potential at other buffer cells are not, we need flags to indicate which buffer cells must be updated. These flags are stored as an integer at each cell as given in Table 3. Using these flags, the code of the Poisson solver for the hierarchical meshes is written as in Appendix 1.1.

**Table 3.** Meanings of the nine least significant bits of the index which is used by the Poisson solver for the hierarchical meshes (See Appendix 1.1).

macro variable	value	meaning
UPDATE_ALL	000000001	Refined cell. (All must be updated.)
UPDATE0	000000010	0th child must be updated.
UPDATE1	000000100	1st child must be updated.
UPDATE2	000001000	2nd child must be updated.
UPDATE3	000010000	3rd child must be updated.
UPDATE4	000100000	4th child must be updated.
UPDATE5	001000000	5th child must be updated.
UPDATE6	010000000	6th child must be updated.
UPDATE7	100000000	7th child must be updated.
UPDATE_RED	011010010	Updated in the red iteration.
UPDATE_BLACK	100101100	Updated in the black iteration.



**Fig. 3.** Schematic illustration of the depth-first order sweep and the breadth-first order sweep. Particles (*circles*) in the same cell (*rectangle*) are connected by a linked list (*downward arrows*). *Vertical solid zigzag* arrow represents the access pattern of the depth-first order sweep which is unvectorizable. *Horizontal dashed zigzag* arrow represents the access pattern of the breadth-first order sweep which is vectorizable.

### 3.4. Particle-mesh loops

It is particle-mesh loops that is most difficult to vectorize among three types of loops described in the beginning of this section. In the scalar code, particles in a same cell are accessed by rolling up the linked list (Fig. 3). The code containing such loops are given in Appendix 1.2. These loops are not vectorizable, because neither the length of the loops nor which particles are to be accessed are not known beforehand. This access pattern of particles is called depth-first order and blocks vectorization frequently. For example, tree code had also confronted with this difficulty. Hernquist (1990) and Makino (1990) overcame this problem by exchanging the depth-first order loops by the breadth-first order ones. We followed this strategy. The code of the exchanged loops are shown in Appendix 1.3. Most of particle-mesh loops are vectorized by this order exchange. The vectorization schemes of three main functions which contain particle-mesh loops are discussed below.

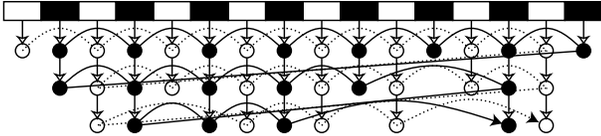
#### 3.4.1. Force interpolation

If each particle has an index of the cell including it, force interpolation is vectorizable even without the order exchange, because only particle velocity is updated and it is not overlapped in the memory during a sweep. However, if each particle does not have an index of the parent to save the memory usage, the loop order of force interpolation must be exchanged to be vectorized.

#### 3.4.2. Mass assignment

In plasma physics, the particle-in-cell (PIC) method is widely used and the PIC method includes charge assignment which is almost identical to mass assignment in the PM method. There is a vectorization scheme for charge assignment named the Abe-Nishihara method (Nishihara et al. 2000). In the Abe-Nishihara method, particles assign their charge assuming that particles reside in different cells, and at the same time, for each cell, the number of particles which assign their charge to the cell is counted. If the number of particles in cells is zero or one, the loop is not recursive. However, if the number of particles in a cell exceeds one, recursive memory access has occurred. In that case, it is necessary to correct the result and the charge is assigned by a scalar loop. If the loop length of charge assignment is well short compared with the number of particles, this correction occurs scarcely.

However, we developed another vectorization scheme for mass assignment which does not have neither the scalar parts, nor the correction part, using the data structure of particles and meshes. First, if the nearest grid point scheme is adopted for mass assignment, it is vectorized by the order exchange, because mass of each particle is assigned to only one cell, and the cells do not overlap during a sweep. However, the higher order assignment schemes, such as the cloud-in-cell scheme or the triangular shaped cloud scheme, cannot be vectorized because those schemes update more than one cells data per particle during a sweep of the loop. Such data accesses are prohibited for vectorization.



**Fig. 4.** Schematic illustration of the loop splitting for vectorization of the mass assignment. Adopted the cloud-in-cell mass assignment scheme, the loop is recursive, if black and white particles assign their mass simultaneously during breadth-first ordered loops. This is because the density at a node could be updated more than once during a sweep. This recursion is avoidable if black particles assign their mass (*solid wavy arrow*) after white particles have done (*dashed wavy arrow*).

**Table 4.** CPU time of functions including a particle-mesh loop

	Part	CPU time [s]
vectorized mass assignment (a)		0.48
particle sieve (b) .....		0.32
(a) + (b) .....		0.80
scalar mass assignment .....		29.04

However, this difficulty is removed by splitting the loop into  $2^{N_{\text{dim}}}$  loops, where  $N_{\text{dim}}$  is the number of dimension if the cloud-in-cell scheme is adopted. Figure 4 shows a one dimensional case. White particles and black particles assign their mass to cells in the breadth-first order, separately. Such loops are vectorizable, because density at a node is updated once per sweep at most.

We classify particles into white and black as follows. First we classify cells into black and white. This is possible because cells having the same parent cell are bundled together and those cells' data are aligned closely. Thus we can classify the first child cell is white and the second child cell is black. Then, particles which are linked from white cells or black cells are classified as white particles or black particles, respectively.

### 3.4.3. Particle sieve

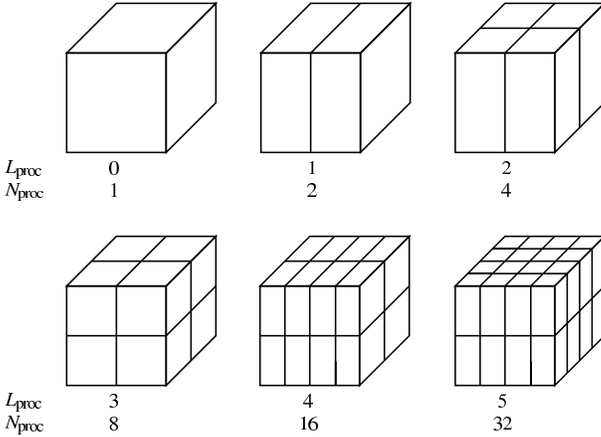
We introduced vectorization schemes for particle-mesh loops so far assuming particles in the same cell are linked listed. However, we must construct the linked list consistently on our own. Particle sieve is the function to construct the linked list. First, particle sieve is divided into two parts: stay particle sieve and leaked particle sieve. Stay particle sieve checks whether particles remain in the same cell from the previous step, or leaked into the neighboring cells. Those particles which remain are added to the linked list of the cell. This process is vectorizable if the consistent linked list is prepared at the previous step. Although, the very first construction of the linked list is unvectorizable, it is called only once in a run. Leaked particle sieve, which added the leaked particle to the neighboring cells, is unvectorizable, either. However, time steps of simulation must be controlled so that the ratio of the number of leaked particles kept small, and the CPU time for leaked particle sieve is negligible.

### 3.4.4. Effect of vectorization

How fast is the vectorized code compared with the the scalar code, especially the particle-mesh loop parts? For example, as an extreme case, consider that nothing is done in a particle-mesh loop. In this case, because of the overhead due to vectorization, the vectorized code is slower than the scalar code. On the other hand, when a lot of work is done in a particle-mesh loop, the vectorized code is advantageous. In order to quantify the advantage of the vectorized code, we measured the CPU time consumed by the vectorized mass assignment, particle sieve, and the scalar mass assignment in the case that there is only a base mesh whose size is  $256^3$  and  $256^3$  particles are scattered randomly and homogeneously. The main parts of the code are given in Appendix 2. The measured CPU times using the VPP5000 are given in Table 4. Even including the particle sieve part, the vectorized mass assignment is 36.3 times faster than the scalar mass assignment. Hence, it is beneficial to vectorize the particle-mesh loops using the loop order exchange.

## 4. Parallelization

The fastest computers in the world adopted distributed memory architecture and have thousands of processors. Thus, in order to carry out large scale simulations, simulation codes must be parallelized for distributed memory machine. In this section, we describe parallelization scheme of our AMR  $N$ -body code. The scheme consists of data decomposition and communication among processors. Our parallelized code described in this section uses Message Passing Interface for a communication library to transfer data among processors.



**Fig. 5.** Data decomposition of the base mesh. Here,  $N_{\text{proc}}$  is the number of processors, and  $L_{\text{proc}} = \log_2 N_{\text{proc}}$ . The shape of the decomposed region depends on the remainder of  $L_{\text{proc}}$  divided by  $N_{\text{dim}}$ . In the three dimensional case as the above, when the remainders are zero, one and two, the decomposed regions are cubes, square slabs and square pillars, respectively.

#### 4.1. Data decomposition

Data decomposition is the most important part of parallelization. We must divide data among processors so that computational load is distributed equally, memory is used uniformly, and communication among processors is kept minimum. Adequate data decomposition realizes these three conditions. We describe data decomposition of the three types of data structure in this subsection.

##### 4.1.1. Base mesh data

Data decomposition of the base mesh is trivial, because the number and position of cells are fixed and do not change during a simulation. Let  $L_{\text{proc}}$  be the logarithm of the number of processors,  $N_{\text{proc}}$ , to the base 2, that is

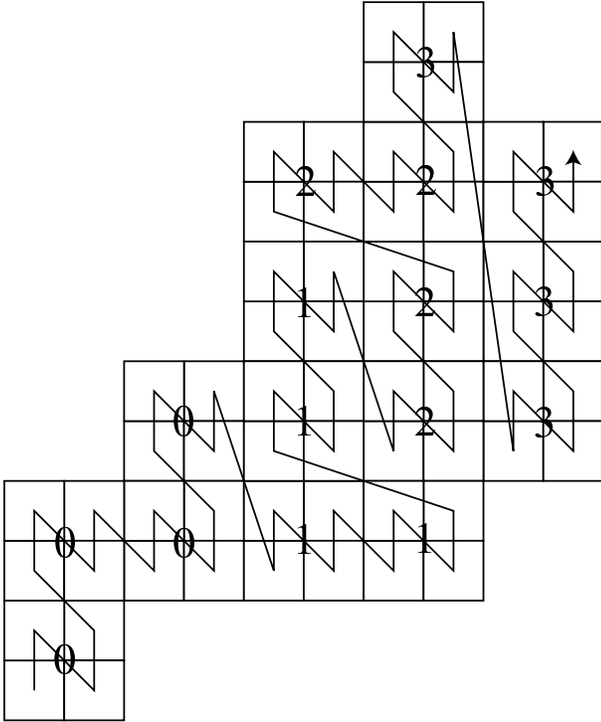
$$L_{\text{proc}} = \log_2 N_{\text{proc}}.$$

Our code assumes that  $L_{\text{proc}}$  is integer. Thus, number of processors is restricted to powers of two. The shape of the decomposed region depends on the remainder of  $L_{\text{proc}}$  divided by  $N_{\text{dim}}$ . In the three dimensional case as shown in Figure 5, when the remainders are zero, one and two, the decomposed regions are cubes, square slabs and square pillars, respectively. Hence, decomposed regions could be elongated. The order of priority for the elongation is z-direction, y-direction and x-direction, because we implemented our code in C whose right most indices are contiguous on the memory. If  $L_{\text{proc}}$  is multiple of  $N_{\text{dim}}$ , the cubic base mesh whose level is  $L_{\text{B}}$  is decomposed into  $N_{\text{proc}}$  cubic base meshes which have  $2^{N_{\text{dim}} \times (L_{\text{B}} - L_{\text{proc}})}$  cells.

##### 4.1.2. Hierarchical mesh data

Among three types of data, the hierarchical meshes uses memory most in our code. and the most time consuming part of our serial code is the AMR Poisson solver. For the  $N$ -body code, other functions including particle-mesh loops also use a significant fraction of the CPU time. However, the ratio of the CPU time for the mesh loops to the total increases if the hydrodynamics blocks are incorporated. Thus, data decomposition of the hierarchical meshes is the most important part of the parallelization. Since the red-black Gauss-Seidel iteration, which is the kernel of the AMR Poisson solver, is iterated for the hierarchical meshes of the same level, we should divide hierarchical mesh data equally in each level. However, we have already sorted the hierarchical meshes in each level by the Morton order (Yahagi & Yoshii 2001). Hence, we simply divide the arrays of hierarchical meshes by  $N_{\text{proc}}$  at each level (Fig. 6). Warren & Salmon (1993) parallelize the Barnes-Hut tree code using the Morton order. Their code, named hashed oct tree method, sorts particles in the Morton order and distributes them to processors. However, we apply the Morton order not to particles but to the hierarchical meshes. Fryxell et al. (2000) also uses the Morton order for data decomposition. Difference between our code and their code is that we sort and divide hierarchical mesh data level by level, while Fryxell et al. (2000) sort and divide all hierarchical meshes at once.

Our decomposition scheme causes that parent and child cells can be on different processors. Hence, communication occurs when accessing child data from parent cell, or vice versa. Details of such communication will be discussed in §4.2.



**Fig. 6.** Hierarchical mesh data are sorted in the Morton order (*connected N-shaped arrow*) at each level. Hierarchical meshes are distributed by dividing the sorted data by the number of processor, in the above case, four. Numbers in the figure indicate the ID of the host processor to which the cell octets allocated.

#### 4.1.3. Particle data

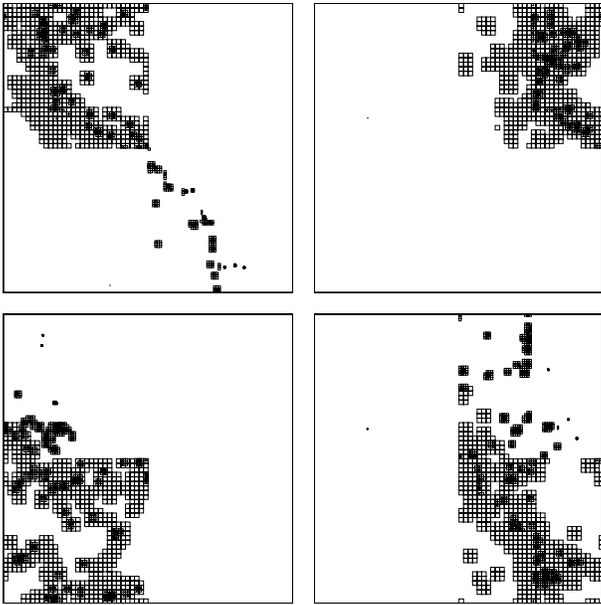
Particles are attached to the finest refined cell including them. Particles belonging to the same cell are connected by a linked list and the cell including those particles keeps an index of the first particle of the linked list. Hence it is natural that the particles are assigned to the processor which also covers cells including them. This data decomposition does not guarantee the balanced decomposition. However, if the numbers of cells which contain one particle, two particles, three particles, etc., are the same among processors, the numbers of particles in processors are balanced. We will return to the issue of particle data decomposition in §5.

#### 4.1.4. Example of data decomposition

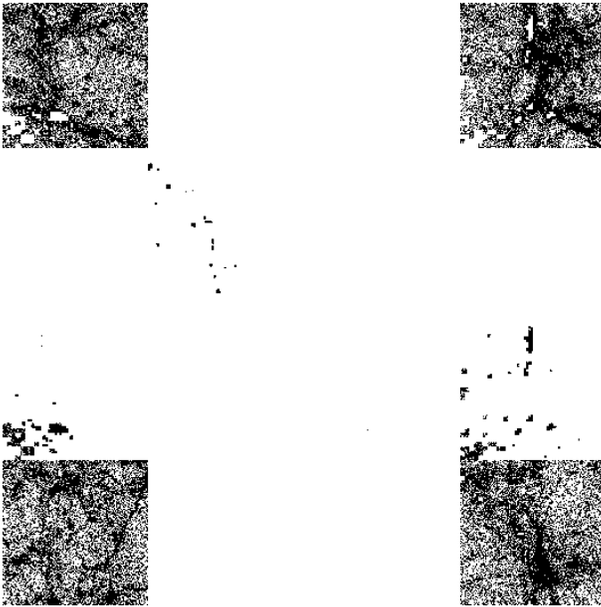
We performed a  $\Lambda$ -dominated cold dark matter (ACDM) simulation with  $64^3$  particles in the  $70 h^{-1}$  Mpc cubic box using four processors. The base level and the dynamic range level, or the deepest level, of the simulation are 6 and 12, respectively. The initial condition is prepared by `grafic2` code provided by Bertschinger (2001), but we modified the code to adopt the fitting function of the transfer function given in Sugiyama (1995) which includes the baryon's effect. Figure 7 shows the distribution of the hierarchical meshes at  $z = 0$  which are assigned to four processors. Each panel corresponds to a processor. For clarity, particles are projected onto a plane, then hierarchical meshes are generated. Note that some cell octets and their parent cell octets are assigned to different processors. Distribution of particles are shown in Figure 8. Distribution of the particles allocated to a processor is similar to that of the hierarchical meshes allocated to the same processor.

#### 4.2. Inter-level communication

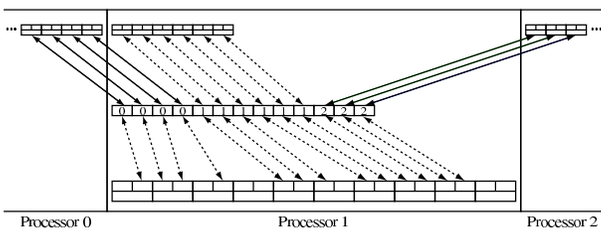
Since we distribute hierarchical meshes by sorting them at each level, some cell octets and their parent cells are attached to the different processors. Therefore, processors must communicate with each other when data of the child cell octets or parent cells are needed, for example, when the coarse-to-fine or the fine-to-coarse operators are called from the AMR Poisson solver. We call such communication among processors inter-level communication. Figure 9 shows how inter-level communication is implemented in our code. When cells (*lower rectangles*) need data of child cell octets (*upper rectangles*), child cell octets' data are transferred to the communication buffer (*rectangles in the middle*) at the processor to which parent cells are attached. Then parent cells read data from the communication buffer. Parent cells' data are transferred to their child cell octets the other way around. Since cells are sorted in accordance with the Morton order, it is guaranteed that child cell octets in the same processor are contiguous on the buffer arrays.



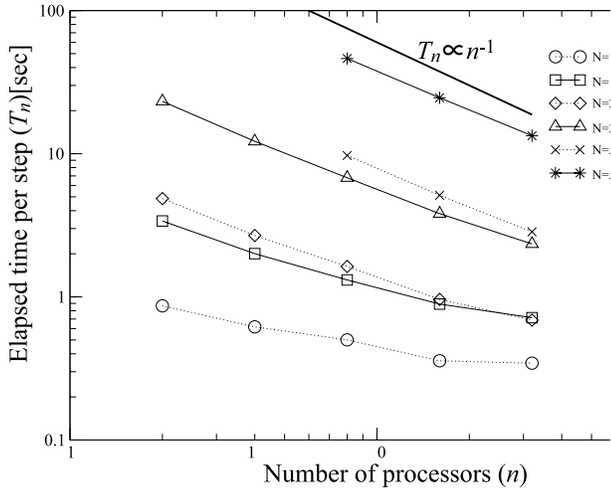
**Fig. 7.** Distribution of the hierarchical meshes among processors. *Squares* indicate cell octets and each panel shows distribution of the cell octets allocated to the corresponding processor. For clarity, cells are placed after projection of particles onto a plane. Note that some cell octets and their parent cell octets are assigned to different processors.



**Fig. 8.** Distribution of particles among processors. Each panel shows distribution of the particles allocated to the corresponding processor. Distribution of particles is similar to that of the hierarchical meshes shown in Fig. 7.



**Fig. 9.** Schematic illustration of the inter-level communication. Cells (*lower rectangles*) collect data from their child cell octets (*upper rectangles*) through the communication buffer (*rectangles in the middle*). Solid arrows and dotted arrows represent inter-processor communication and local load and store, respectively. Cell octets collect data from their parent cells the other way around.



**Fig. 10.** Wall-clock time per step,  $T_n$ , using  $n$  processors. *Circles, diamonds, and crosses* represent the wall-clock time measured during the 38 steps from the beginning of the simulation, using  $128^3$ ,  $256^3$ , and  $512^3$  particles, respectively. *Squares, triangles, and asterisks* indicate the wall-clock time measured during the 40 steps to the end of the simulation, using  $128^3$ ,  $256^3$ , and  $512^3$  particles, respectively. Thick line shows the line,  $T_n \propto n^{-1}$ . Slopes of the lines for the case that the number of particles is  $512^3$ , are close to this  $T_n \propto n^{-1}$  line.

### 4.3. Intra-level communication

Since we simply divide the sorted hierarchical meshes, domain borders could lie across patches of meshes. Thus, neighboring cells could be distributed to different processors after the domain decomposition. However, the Gauss-Seidel iteration needs potential at six neighboring cells to update potential at a cell. Hence, we prepare supplementary cell octets, or ghost cell octets, to store copies of the neighboring cell octets distributed to different processors. After each iteration of the Gauss-Seidel iteration, data from the original cells are transferred to the ghost cells by the intra-level communication.

## 5. Timing analysis

We performed three runs of  $\Lambda$ CDM simulation on a system of vector parallel processors VPP5000 installed at the Astronomical Data Analysis Center, National Astronomical Observatory of Japan (ADAC/NAOJ) to measure the speed of the parallel AMR  $N$ -body code. Each node of VPP5000 has only one CPU and nodes are connected via a crossbar switch. Peak performance of each node of VPP5000 is 9.6 GFlops. Peak data transfer rate is  $1.6 \text{ GB/s} \times 2$  (send and receive). Since in our parallel code the number of processors is restricted to be power of two, the number of processors of the largest queue is 32 in the ADAC/NAOJ system. Hence we measured the speed of our code using from 2 processors to 32 processors. The numbers of particles of three runs are  $128^3$ ,  $256^3$ , and  $512^3$ . We measured the wall-clock time per step during 38 steps from the beginning of the simulation and 40 steps to the end of the simulation ( $z = 0$ ) for each run. Figure 10 shows the measured wall-clock times per step,  $T_n$ , using  $n$  processors. Thick line in the figure shows the line,  $T_n \propto n^{-1}$ . Slopes of the lines for the case that the number of particles is  $512^3$  (*crosses* and *asterisks*), are close to this  $T_n \propto n^{-1}$  line, showing that the speed-up in that case is close to the ideal speed-up.

On the other hand, parallel efficiency,  $p_n$ , is widely used as an indicator of parallel codes, and defined as follows:

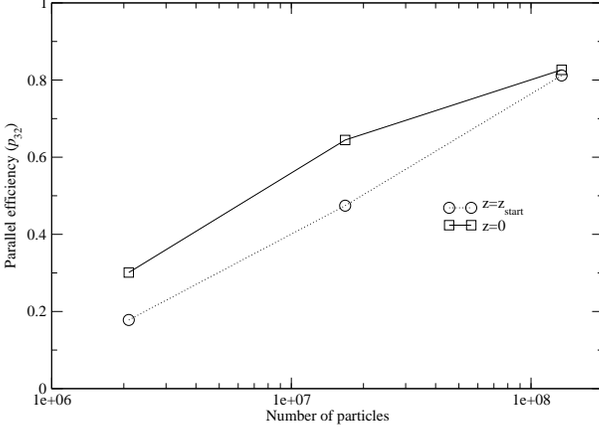
$$p_n = \frac{1}{n} \left( \frac{T_1}{T_n} \right),$$

where  $T_1$  and  $T_n$  are the wall-clock time using one processor and  $n$  processors, respectively. However, as our case, there are cases that  $T_1$  is unmeasurable. In such cases, parallel efficiency is derived as follows assuming the Amdahl's law:

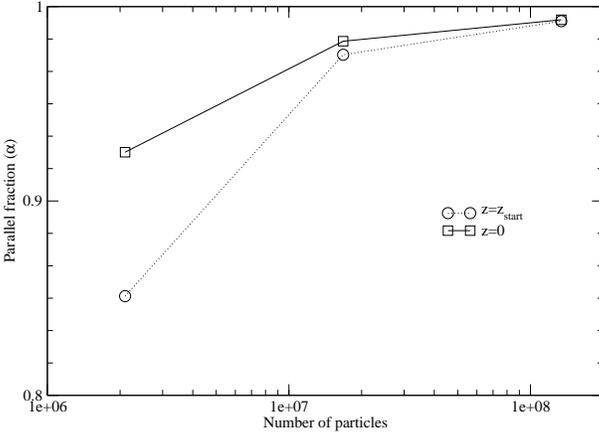
$$\begin{aligned} T_n &= T_1(1 - \alpha + \alpha/n) \\ &= T_1(1 - \beta_n \alpha), \end{aligned}$$

where  $\alpha$  is the parallel fraction, the fraction of time of the serial code which is parallelizable, and  $\beta_n = 1 - 1/n$ . Measuring the wall-clock time using different number of processors,  $\alpha$  is derived as follows:

$$\begin{aligned} T_n &= T_1(1 - \beta_n \alpha) \\ T_m &= T_1(1 - \beta_m \alpha) \end{aligned}$$



**Fig. 11.** Parallel efficiency where the number of processor is 32.  $p_{32}$  estimated by Eq. 2 where  $m = 8$ . Circles and squares indicate the parallel efficiency at the beginning and the end of the simulation, respectively.



**Fig. 12.** Parallel fraction estimated by Eq. 1 where  $n = 32$  and  $m = 8$ . Circles and squares indicate the parallel fraction at the beginning of the simulation and the end of the simulation, respectively.

$$\alpha = \frac{T_m - T_n}{\beta_n T_m - \beta_m T_n}. \quad (1)$$

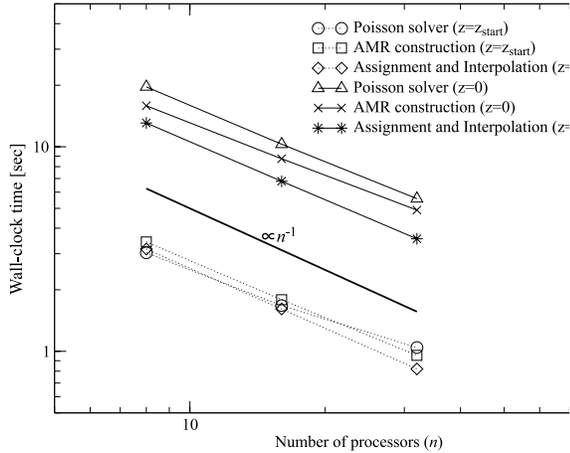
Then, parallel efficiency is:

$$\begin{aligned} p_n &= \frac{1}{n} \left( \frac{1}{1 - \beta_n \alpha} \right) \\ &= \frac{1}{n} \left( \frac{\beta_m T_n - \beta_n T_m}{\beta_m T_n - \beta_n T_n} \right). \end{aligned} \quad (2)$$

We estimated the parallel efficiency at  $n = 32$ ,  $p_{32}$ , in the case that  $m = 8$ . Figure 11 shows  $p_{32}$  of each run at  $z = z_{\text{start}}$  (circles) and  $z = 0$  (squares). Figure 12 shows the parallel fraction estimated by Eq. 1 where  $n=32$  and  $m=8$ . These two figures indicate that our parallel code is more efficient as the size of the simulation increases.

Next, we measured the wall-clock time of three groups of functions: Poisson solver, AMR construction, and mass assignment and force interpolation. Figure 13 shows the measured wall-clock time of each group at  $z = z_{\text{start}}$  and  $z = 0$ . Most groups indicate the expected speed-up. However, speed-up of the Poisson solver at  $z = z_{\text{start}}$  slows down. Since the number of hierarchical meshes at  $z = z_{\text{start}}$  is much smaller than that at  $z = 0$ , wall-clock time of the Poisson solver for the base mesh is dominant. Thus, there might be room for further optimization of the the Poisson solver for the base mesh.

As mentioned in §4.1.3, our data decomposition does not guarantee the balanced decomposition of particle data. Figure 14 shows the numbers of cells which include one particle, two particles, three particles, and more than three particles at the end of the  $\Lambda$ CDM simulation using  $512^3$  particles. The numbers of cells including particles is balanced among processors at the finest level, and the dispersion is large at the coarser levels. Figure 15 shows the number of particles in each level among processors. Again, particles are equally distributed at the finest level, although cells



**Fig. 13.** Wall-Clock time of three groups of functions, Poisson solver (*circles and triangles*), AMR construction including the reconnection of the linked list of particles (*squares and crosses*), and mass assignment and force interpolation (*diamonds and asterisks*). Symbols connected by dotted lines and solid lines are measured at  $z = z_{\text{start}}$  and  $z = 0$ , respectively. The number of particles is  $512^3$ . *Thick line* indicates the line  $\propto n^{-1}$  which shows ideal speed-up. At the end of the simulation, wall-clock times of all three groups of functions shows expected speed-up. However, at the beginning of the simulation, speed-up of the Poisson solver slows down.

at the finest level can contain arbitrarily large number of particles. Figure 16 shows the total number of particles assigned to each processor. The processor which is assigned the largest number of particles has particles about 50% more than the average number. However, mass assignment and force interpolation part of our code, which includes the particle-mesh loops, scales almost ideally as the number of processors is increased as shown in Figure 13. This is because more mesh loops, which assign cells' mass to the parent cells, are included into mass assignment and force interpolation part than particle-mesh loops, which assign particle mass to the cells including it and interpolate force at cells toward particles.

## 6. Summary

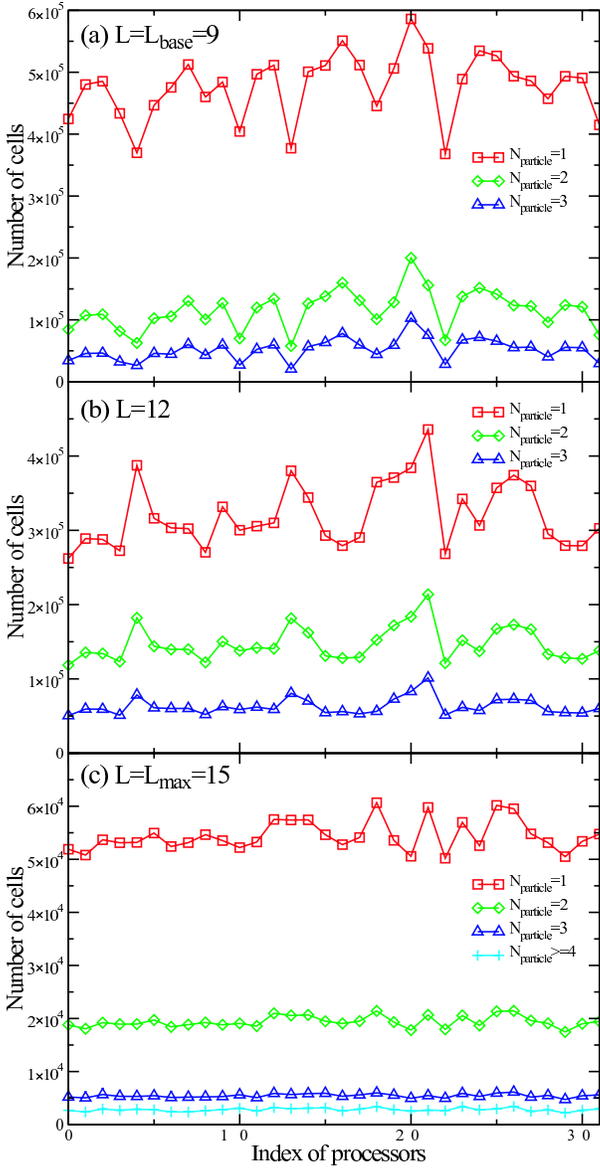
We have vectorized and parallelized the shared time step version of our AMR  $N$ -body code. First, in order to vectorize our code, we grouped loops into three types, mesh loops, particle loops, and particle-mesh loops. It is easy to vectorize mesh loops and particle loops. However, vectorization of particle-mesh loops is not so simple. Particle-mesh loops contain a hash data structure: an array of hierarchical meshes and linked lists of particles beginning from the array. The simplest way to treat a particle-mesh loop containing such a hash data structure is the depth-first order loop, but such loop is unvectorizable. We vectorized particle-mesh loops by changing the loop order to the breadth-first order.

We also parallelized our code for distributed memory machines. First, we decomposed data among processors. Data decomposition of the base mesh is trivial. The hierarchical meshes are sorted by the Morton order level by level, then split and distributed among processors. Data of particles are allocated to processors to which the data of cells including them cells are assigned. Since the hierarchical meshes are sorted level by level, some cell octets and their parent cells could be assigned to different processors. Thus we also implemented the inter-level communication to collect the data from child cell octet allocated to different processors and vice versa.

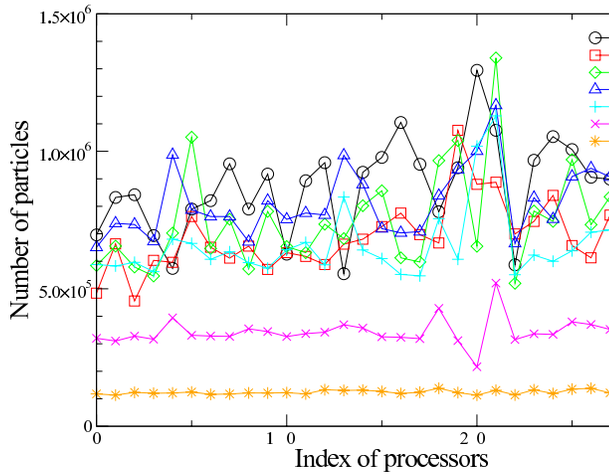
Finally, we measured the parallel efficiency of our parallel AMR  $N$ -body code for three runs of  $\Lambda$ CDM simulation. The parallel efficiency increases as the number of particle of simulation increases. The highest value of the parallel efficiency and the parallel fraction in the case that the number of processors is 32, are 82.6 % and 99.3 %, respectively, and they are measured in the  $512^3$  particles simulation at  $z = 0$ .

With this parallel AMR  $N$ -body code we carried out large scale simulations as in Yahagi et al. (2004), and we are going to investigate the nature of dark halos of galaxies and clusters of galaxies.

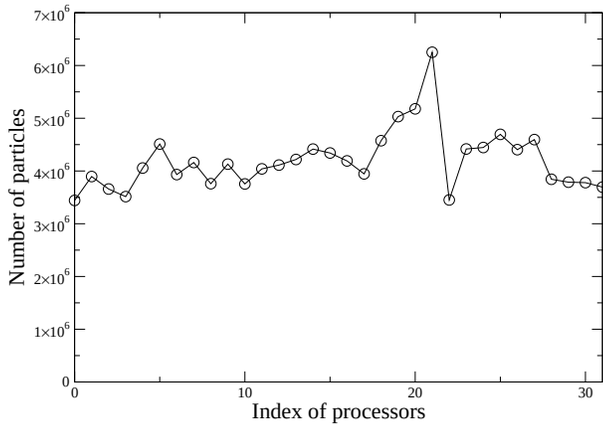
I would like to thank my thesis advisor Y. Yoshii for lasting encouragement and valuable suggestions. I am also grateful to J. Makino for helpful comments which helped to improve this paper a lot, and to an anonymous referee



**Fig. 14.** The numbers of cells which contain one particle (*squares*), two particles (*diamonds*), three particles (*triangles*), and more than three particles (*pluses*) in each processor at the end of the  $\Lambda$ CDM simulation. The number of particles is  $512^3$ . Each panel shows the number of cells in (a) the base level ( $L = 9$ ), (b) 12th level, and (c) the finest level ( $L = 15$ ).



**Fig. 15.** The number of particles assigned to each processor at the end of the  $\Lambda$ CDM simulation. The number of particles in the simulation is  $512^3$ . Circles, squares, diamonds, triangles, pluses, crosses, and asterisks indicate the number of particles at levels from 9 (the base level) to 15 (the finest level), respectively. At the finest level, particles are distributed in balance, while the dispersion



**Fig. 16.** Total number of particles assigned to each processor at the end of the  $\Lambda$ CDM simulation. The number of particles in the simulation is  $512^3$ . The number of particles in each processor should be 4,194,304, if perfectly balanced.

for valuable comments. Simulations described in this paper were carried out using Fujitsu-made vector-parallel processors VPP5000 installed at the Astronomical Data Analysis Center, National Astronomical Observatory of Japan (ADAC/NAOJ), under the ADAC/NAOJ large-scale simulation projects (group-ID: yhy35b, rhy15b). I also acknowledge the support from the Research Fellowships of the Japan Society for the Promotion of Science for Young Scientists.

## Appendix 1. Code

### A.1.1. Poisson solver for the hierarchical mesh

As an example of a mesh loop and in order to show how the flags in Table 3 are used, we show the Poisson solver for the hierarchical meshes here.

```
{
  int ico, jco, ic, i0, i1, i2, i3, i4, i5;
  int number_of_refined, number_of_red, number_of_black;
  int *refined_cellect, *red_cellect, *black_cellect;
  double rho0;
```

```

[...]

number_of_refined=0;
for (ico=head_celloct; ico<tail_celloct; ico++)
  if (celloct[ico].flag & UPDATE_ALL)
    refined_celloct[number_of_refined++] = ico;
number_of_red=0;
for (ico=head_celloct; ico<tail_celloct; ico++)
  if (celloct[ico].flag & UPDATE_RED)
    red_celloct[number_of_red++] = ico;
number_of_black=0;
for (ico=head_celloct; ico<tail_celloct; ico++)
  if (celloct[ico].flag & UPDATE_BLACK)
    black_celloct[number_of_black++] = ico;

/* Red sweep */
/* Refined cells */
for (jco=0; jco<number_of_refined; jco++){
  ico= refined_celloct[jco];
  ic = ico * 8;
  i0 = celloct[ico].neighbor_x_minus * 8;
  i1 = celloct[ico].neighbor_x_plus * 8;
  i2 = celloct[ico].neighbor_y_minus * 8;
  i3 = celloct[ico].neighbor_y_plus * 8;
  i4 = celloct[ico].neighbor_z_minus * 8;
  i5 = celloct[ico].neighbor_z_plus * 8;

  cell[ic ].pot = (cell[i0+4].pot + cell[ic+4].pot +
                  cell[i2+2].pot + cell[ic+2].pot +
                  cell[i4+1].pot + cell[ic+1].pot -
                  cell[ic ].rho * rho0) / 6.0;
  cell[ic+3].pot = (cell[i0+7].pot + cell[ic+7].pot +
                  cell[ic+1].pot + cell[i3+1].pot +
                  cell[ic+2].pot + cell[i5+2].pot -
                  cell[ic+3].rho * rho0) / 6.0;
  cell[ic+5].pot = (cell[ic+1].pot + cell[i1+1].pot +
                  cell[i2+7].pot + cell[ic+7].pot +
                  cell[ic+4].pot + cell[i5+4].pot -
                  cell[ic+5].rho * rho0) / 6.0;
  cell[ic+6].pot = (cell[ic+2].pot + cell[i1+2].pot +
                  cell[ic+4].pot + cell[i3+4].pot +
                  cell[i4+7].pot + cell[ic+7].pot -
                  cell[ic+6].rho * rho0) / 6.0;
}

/* Buffer cells */
for (jco=0; jco<number_of_red; jco++){
  ico = red_celloct[jco];
  ic = ico * 8;
  [...]
  if (celloct[ico].flag & UPDATE0){
    cell[ic ].pot = [...];
  }
  if (celloct[ico].flag & UPDATE3){
    cell[ic+3].pot = [...];
  }
  if (celloct[ico].flag & UPDATE5){
    cell[ic+5].pot = [...];
  }
}

```

```

    if (celloct[ico].flag & UPDATE6){
        cell[ic+6].pot = [...];

    }
}

/* Black sweep */
[...]
```

#### A.1.2. Scalar particle-mesh loop

```

{
    int ip, ico;
    [...]

    for (ico=head_celloct; ico<tail_celloct; ico++){
        for (ip=celloct[ico].head; ip != NULL_PARTICLE; ip=particle[ip].next){
            /* Some operations */
        }
    }
}
```

#### A.1.3. Vectorized particle-mesh loop

```

{
    int mp, np, jp, ico;
    int *list_of_particle;
    [...]

    np = 0;
    for (ico=head_celloct; ico<tail_celloct; ico++){
        if (celloct[ico].head != NULL_PARTICLE)
            list_of_particle[np++] = celloct[ico].head;
    }
    while (np>0){
        for (jp=0; jp<np; jp++){
            /* Some operations */
        }
        for (jp=0; jp<np; jp++){
            list_of_particle[jp] = particle[list_of_particle[jp]].next;
        }
        mp = np;
        np = 0;
        for (jp=0; jp<mp; jp++){
            if (list_of_particle[jp] != NULL_PARTICLE){
                list_of_particle[np++] = list_of_particle[jp];
            }
        }
    }
}
```

## Appendix 2. Code of the scalar mass assignment and the vectorized mass assignment

### A.2.1. Include file

```

#define N_DIM 3
#define BASE_LEV 8
#define BASE_N (1 << BASE_LEV)
#define NBASE (1 << (BASE_LEV * N_DIM))
#define FBASE_N (BASE_N + 2)
```

```

#define NFBASE (FBASE_N * FBASE_N * FBASE_N)
#define BZIND 1
#define BYIND FBASE_N
#define BXIND (FBASE_N * FBASE_N)

#define NPTCL NBASE
#define NULL_PTCL (-1)

#define MALLOC(x, y, z) \
{
    x=(y *) malloc (sizeof(y) * (z));\
    if (!x)\
        fprintf(stderr, "malloc failure (%s) in %s: line %d\n",\
                #x, __FILE__, __LINE__);
}

```

### A.2.2. Scalar mass assignment code

```

double rhobm[NFBASE];
double ppos[NPTCL*N_DIM];

void scalar_assign (double *cpusa)
{
    int im, ip, ix, iy, iz, *prnt;
    double ct0, dx, dy, dz, lmdx, lmdy, lmdz;
    double *tpos;

    ct0 = scnd();
    MALLOC(prnt, int, NPTCL);
    MALLOC(tpos, double, NPTCL * N_DIM);

    for (ip=0; ip<NPTCL; ip++){
        ix = (int) ppos[ip*N_DIM ];
        iy = (int) ppos[ip*N_DIM+1];
        iz = (int) ppos[ip*N_DIM+2];
        prnt[ip] = ix * BXIND + iy * BYIND + iz;
    }

#pragma loop novrec
    for (ip=0; ip<NPTCL*N_DIM; ip++)
        tpos[ip] = ppos[ip] - (double)((int) ppos[ip]);

    for (im=0; im<NFBASE; im++) rhobm[im] = 0.0;
    for (ip=0; ip<NPTCL; ip++){
        im = prnt[ip];
        dx = tpos[ip*N_DIM ]; lmdx = 1.0-dx;
        dy = tpos[ip*N_DIM+1]; lmdy = 1.0-dy;
        dz = tpos[ip*N_DIM+2]; lmdz = 1.0-dz;
        rhobm[im          ] += lmdx * lmdy * lmdz;
        rhobm[im          +BZIND] += lmdx * lmdy * dz;
        rhobm[im          +BYIND ] += lmdx * dy * lmdz;
        rhobm[im          +BYIND+BZIND] += lmdx * dy * dz;
        rhobm[im+BXIND    ] += dx * lmdy * lmdz;
        rhobm[im+BXIND    +BZIND] += dx * lmdy * dz;
        rhobm[im+BXIND+BYIND ] += dx * dy * lmdz;
        rhobm[im+BXIND+BYIND+BZIND] += dx * dy * dz;
    }
    free(prnt);
    free(tpos);
}

```

```

    *cpusa += scnd() - ct0;
}

```

### A.2.3. Vectorized mass assignment code

```

int    hpbm[NFBASE];
int    pnxt[NPTCL];

void vector_assign (double *cpuva)
{
    int ip, im0, im, jm, nm, mm, ix, iy, iz, ich;
    int im0l[] = {0,          BZIND,          BYIND,          BYIND+BZIND,
                 BXIND, BXIND+BZIND, BXIND+BYIND, BXIND+BYIND+BZIND};
    int *lbm, *lpt;
    double ct0, dx, dy, dz, lmdx, lmdy, lmdz;
    double *tpos;
#pragma procedure novrec
    ct0 = scnd();

    MALLOC(lbm, int, NBASE);
    MALLOC(lpt, int, NBASE);
    MALLOC(tpos, double, NPTCL * N_DIM);

    for (ip=0; ip<NPTCL*N_DIM; ip++)
        tpos[ip] = ppos[ip] - (double)((int) ppos[ip]);
    for (ich=0; ich<(1<<N_DIM); ich++){
        im0 = BXIND + BYIND + BZIND + im0l[ich];
        nm=0;
        for (ix=0; ix<(BASE_N/2); ix++){
            for (iy=0; iy<(BASE_N/2); iy++){
                im = im0 + ix * BXIND + iy * BYIND;
                for (iz=0; iz<(BASE_N/2); iz++, im++){
                    if (hpbm[im] != NULL_PTCL) {
                        lbm[nm] = im;
                        lpt[nm] = hpbm[im];
                        nm++;
                    }
                }
            }
        }
    while (nm){
        for (jm=0; jm<nm; jm++){
            im = lbm[jm];
            ip = lpt[jm];
            dx = tpos[ip*N_DIM ]; lmdx = 1.0-dx;
            dy = tpos[ip*N_DIM+1]; lmdy = 1.0-dy;
            dz = tpos[ip*N_DIM+2]; lmdz = 1.0-dz;
            rhobm[im          ] += lmdx * lmdy * lmdz;
            rhobm[im          +BZIND] += lmdx * lmdy *  dz;
            rhobm[im          +BYIND ] += lmdx *  dy * lmdz;
            rhobm[im          +BYIND+BZIND] += lmdx *  dy *  dz;
            rhobm[im+BXIND          ] +=  dx * lmdy * lmdz;
            rhobm[im+BXIND          +BZIND] +=  dx * lmdy *  dz;
            rhobm[im+BXIND+BYIND          ] +=  dx *  dy * lmdz;
            rhobm[im+BXIND+BYIND+BZIND] +=  dx *  dy *  dz;
        }
        for (jm=0; jm<nm; jm++) lpt[jm] = pnxt[lpt[jm]];
        mm=nm; nm=0;
        for (jm=0; jm<mm; jm++){
            if (lpt[jm] != NULL_PTCL){
                lbm[nm] = lbm[jm];
                lpt[nm] = lpt[jm];
            }
        }
    }
}

```

```

        nm++;
    }
}

free(lbm);
free(lpt);
free(tpos);
*cpuva += scnd() - ct0;
}

```

#### A.2.4. Particle sieve code

```

void particle_sieve (double *cpus)
{
    int im, jm, nm, mm, ip, ix, iy, iz, ilk, nlk;
    int *lbm, *lpt, *prnt, *tnxt, *llk;
    double ct0;

#pragma procedure novrec
    ct0 = scnd();
    MALLOC(lbm, int, NBASE);
    MALLOC(lpt, int, NBASE);
    MALLOC(prnt, int, NPTCL);
    MALLOC(tnxt, int, NPTCL);
    MALLOC(llk, int, NPTCL);

    for (ip=0; ip<NPTCL; ip++){
        ix = (int) ppos[ip*N_DIM ];
        iy = (int) ppos[ip*N_DIM+1];
        iz = (int) ppos[ip*N_DIM+2];
        prnt[ip] = ix * BXIND + iy * BYIND + iz;
    }
    nm=0;
    for (im=0; im<NFBASE; im++){
        if (hpbm[im] != NULL_PTCL){
            lbm[nm] = im;
            lpt[nm] = hpbm[im];
            nm++;
        }
    }
    for (im=0; im<NFBASE; im++) hpbm[im] = NULL_PTCL;

    nlk=0;
    while (nm){
        for (jm=0; jm<nm; jm++) tnxt[jm] = pnxt[lpt[jm]];
        for (jm=0; jm<nm; jm++){
            im = lbm[jm];
            ip = lpt[jm];
            if (prnt[ip] == im){
                pnxt[ip] = hpbm[im];
                hpbm[im] = ip;
            }
        }
        for (jm=0; jm<nm; jm++)
            if (prnt[lpt[jm]] != lbm[jm]) llk[nlk++] = lpt[jm];
        mm=nm; nm=0;
        for (jm=0; jm<mm; jm++){
            if (tnxt[jm] != NULL_PTCL){

```

```

        lbm[nm] = lbm [jm];
        lpt[nm] = tnxt[jm];
        nm++;
    }
}

#pragma loop scalar
for (ilk=0; ilk<nlk; ilk++){
    ip = llk[ilk];
    im = prnt[ip];
    pnxt[ip] = hpbm[im];
    hpbm[im] = ip;
}

free(lbm);
free(lpt);
free(prnt);
free(tnxt);
free(llk);
*cpus += scnd() - ct0;
}

```

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