Selected Methods in N-body Simulations

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1 Introduction

The dominant force over large distances is the gravitational force. The force exerted on a body with mass m_2 at the point \mathbf{x}_2 by a body with mass m_1 located at \mathbf{x}_1 can be expressed by the relation

$$\mathbf{F} = -G \frac{m_1 m_2}{|\mathbf{x}_{21}|^3} \mathbf{x}_{21}$$

where G is the gravitational constant $6.674 \times 10^{-11} \text{m}^3 \text{kg}^{-1} \text{s}^{-2}$ and $\mathbf{x}_{21} = \mathbf{x}_2 - \mathbf{x}_1$. Therefore, the evolution of a system of N bodies is described by N equations

$$\ddot{\mathbf{x}}_i = -G \sum_{j \neq i} \frac{m_j}{|\mathbf{x}_{ij}|^3} \mathbf{x}_{ij}. \tag{1}$$

for each i = 1, ..., N. Direct application of Equation 1 is the basis of the so-called particle-particle method. The method is characterized by $O(N^2)$ time complexity (more precisely, it requires (N-1)N/2 operations if Newton's 3rd law is used in the computation). Assuming that 100ns are required to perform the floating-point operations under the summation symbol, N = 30,000, and 150 iterations, the simulation would take approximately 2 hours to complete. Therefore, it is evident that more efficient algorithms are needed to make simulations of this scale feasible.

The particle-mesh (PM) technique, introduced around 1985 by Hockney and Eastwood, was an early improvement over the PP method. In the PM approach, the space is divided into a rectangular grid (or mesh) of cells. Each cell is assigned a portion of the mass of nearby particles, creating a density distribution $\rho(\mathbf{x})$. The relation between the density and gravitational potential ϕ , in the form of Poisson's equation

$$\nabla^2 \phi = 4\pi G \rho, \tag{2}$$

is then used to obtain the potential at each cell center. The gravitational field \mathbf{g} can then be calculated as $\mathbf{g} = -\nabla \phi$. Since \mathbf{g} equals the acceleration due to gravity, we get $\ddot{\mathbf{x}}_i = \mathbf{g}(\mathbf{x}_i)$.

The drawback of the PM method is its poor modeling of forces over short distances. Eastwood and Hockney proposed a remedy for this problem: the particle-particle-mesh method (or P^3M in short). In the P^3M method, the force on the i-th particle is split into two components: short-range and long-range force. The long-range force is calculated using the PM method, whereas the short-range force can be found by direct summation of the forces due to nearby particles.

The computational complexity of the PM and P^3M methods depends on the implementation of the potential solver used to calculate ϕ from Equation 2. For instance, if a fast Fourier transform is used,

then the complexity of the PM algorithm is $O(N+N_g^3 \log N_g)$, where N_g is the number of cells in a single dimension of the grid (note it is linear in N). For the P³M method, the worst-case scenario happens when all particles are clustered closely together, which causes the short-range $O(N^2)$ correction part to become dominant.

2 Particle-mesh method

The particle-mesh method can be described as the following sequence of four steps:

- 1. Assign masses to mesh points,
- 2. Solve the field equation (Equation 2) on the mesh,
- 3. Calculate the field strength at mesh-points,
- 4. Find forces applied to individual particles by interpolation.

In this section, each of these steps will be described in more detail.

2.1 Mass assignment

The specifics of assigning mass from particles to mesh points depend on the density profile (or *shape*) associated with the particles. In general, the particles need not be represented as idealized dimensionless points; indeed, it is possible to construct a hierarchy of shapes, where each successive member covers a larger number of mesh points and whose application leads to smaller numerical errors.

An infinite hierarchy of shapes with this property, as described by Hockney and Eastwood in [6], can be generated by successive convolutions with the "top-hat" function Π , defined as

$$\Pi(x) = \begin{cases} 1, & |x| < \frac{1}{2} \\ \frac{1}{2}, & |x| = 1 \\ 0, & \text{otherwise} \end{cases}$$

The three most popular assignment schemes that hail from this family (and the ones implemented in our program) are the nearest grid point (NGP), cloud in cell (CIC), and triangular shaped cloud (TSC) schemes, with shapes S given by

$$S_{\mathrm{NGP}} = \delta(x), \quad S_{\mathrm{CIC}} = \delta(x) * \frac{1}{H} \Pi\left(\frac{x}{H}\right) = \frac{1}{H} \Pi\left(\frac{x}{H}\right), \quad S_{\mathrm{TSC}} = \frac{1}{H} \Pi\left(\frac{x}{H}\right) * \frac{1}{H} \Pi\left(\frac{x}{H}\right) = \frac{1}{H} \Lambda\left(\frac{x}{H}\right),$$

where Λ is the triangle function

$$\Lambda(x) = \begin{cases} 1 - |x|, & |x| < 1\\ 0, & \text{otherwise.} \end{cases}$$

For illustrative purposes, the shape $S_{\rm CIC}$ is depicted in Figure 1.

In the one-dimensional case, the fraction of mass W_p assigned to mesh-point p from particle at position x is given by

$$W(x - x_p) = W_p(x) = \int_{x_p - H/2}^{x_p + H/2} S(x' - x) dx'.$$

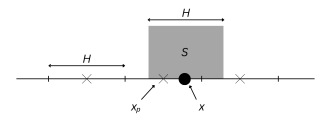


Figure 1: The CIC shape centered at x (particle position). The particle is within cell x_p , however, the cell x_{p+1} gets non-zero density contribution from the particle.

A simple rule for relating the assignment function W defined above with shape S can be found by noticing that

$$W(x) = \int_{-H/2}^{H/2} S(x'-x)dx' = \int_{-\infty}^{\infty} \Pi\left(\frac{x'}{H}\right) S(x'-x)dx' = \Pi\left(\frac{x}{H}\right) * S(x).$$

This implies that

$$W_{\rm NGP}(x) = \Pi\left(\frac{x}{H}\right), \quad W_{\rm CIC}(x) = \Lambda\left(\frac{x}{H}\right), \quad W_{\rm TSC}(x) = \Pi\left(\frac{x}{H}\right) * \frac{1}{H}\Lambda\left(\frac{x}{H}\right) = (\Pi * \Lambda)\left(\frac{x}{H}\right). \quad (3)$$

Splitting the domain of integration in the expression for W_{TSC} into five disjoint intervals shows that

$$(\Pi * \Lambda)(x) = \begin{cases} \frac{1}{8}(3 - 2|x|)^2, & \frac{1}{2} \le |x| < \frac{3}{2} \\ \frac{3}{4} - x^2, & |x| < \frac{1}{2} \\ 0, & \text{otherwise.} \end{cases}$$

Two- and three-dimensional versions of the assignment functions in Equation 3 are products of the assignment functions in each dimension. For example, the three-dimensional assignment function W is

$$W(\mathbf{x}) = W(x)W(y)W(z).$$

Hence, the mass assigned at mesh-point at $\mathbf{x}_{\mathbf{p}}$ is

$$m(\mathbf{x}_{\mathbf{p}}) = \sum_{i} m_i W_{\mathbf{p}}(\mathbf{x}_i),$$

or, in terms of density ρ ,

$$\rho(\mathbf{x}_{\mathbf{p}}) = \frac{1}{V} \sum_{i} m_i W_{\mathbf{p}}(\mathbf{x}_i), \tag{4}$$

where $V = H^3$ is the volume of a cell and i indexes the particles.

Obviously, Equation 4 is not suitable for direct application in the actual algorithm. Instead, we iterate over all particles, identify the parent cell \mathbf{p} of each particle (and its neighborhood) and update ρ . This process is illustrated in Algorithm 1. The set $\mathcal{C}_S(\mathbf{x}_i)$ of cells that have to be considered while

Algorithm 1 Density assignment algorithm

- 1: for each particle i do
- 2: **for each** cell **q** in $C_S(\mathbf{x}_i)$ **do**
- 3: $\rho(\mathbf{x_q}) \leftarrow \rho(\mathbf{x_q}) + m_i W(\mathbf{x}_i \mathbf{x_q}) / V$

assigning density from the i-th particle, depends on the shape S of the particle. Specifically, we have

 $C_{NGP}(\mathbf{x}) = \{ [\mathbf{x}/H] \}, C_{CIC}(\mathbf{x}) = \{ [\mathbf{x}/H] + \mathbf{t} \mid t_i = 0, 1 \}, \text{ and } C_{TSC}(\mathbf{x}) = \{ [\mathbf{x}/H] + \mathbf{t} \mid t_i = -1, 0, 1 \}.$ It follows that $|C_{NGP}(\mathbf{x})| = 1$, $|C_{CIC}(\mathbf{x})| = 8$, and $|C_{TSC}(\mathbf{x})| = 27$ which illustrates the increasing computational cost resulting from using higher-order assignment schemes. We note that Algorithm 1 can be parallelized if atomic increments are used in line 3.

2.2 Solving the field equation

The Poisson equation (Equation 2) can be restated in integral form

$$\phi(\mathbf{x}) = \int \mathcal{G}(\mathbf{x} - \mathbf{x}') \rho(\mathbf{x}') dV',$$

which has the following discrete analogue

$$\phi(\mathbf{x}_{\mathbf{p}}) = V \sum_{\mathbf{p}'} \mathcal{G}(\mathbf{x}_{\mathbf{p}} - \mathbf{x}_{\mathbf{p}'}) \rho(\mathbf{x}_{\mathbf{p}'}), \tag{5}$$

where \mathcal{G} is the Green's function (potential due to unit mass). The right-hand side of Equation 5 is a convolution sum that runs over a finite set of mesh points. If we assume periodic boundary conditions, we can apply the discrete Fourier transform to both sides and use the convolution theorem to conclude that¹

$$\hat{\phi}(\mathbf{k}) = \hat{\mathcal{G}}(\mathbf{k})\hat{\rho}(\mathbf{k}). \tag{7}$$

An approximation to $\hat{\mathcal{G}}$ can be found using a discretized version of the Laplacian in Equation 5. Specifically, for a 7-point stencil,

$$4\pi G \rho(\mathbf{x}_{ijk}) = \frac{\phi(\mathbf{x}_{i-1,j,k}) - 2\phi(\mathbf{x}_{ijk}) + \phi(\mathbf{x}_{i+1,j,k})}{H^2} + \frac{\phi(\mathbf{x}_{i,j-1,k}) - 2\phi(\mathbf{x}_{ijk}) + \phi(\mathbf{x}_{i,j+1,k})}{H^2} \cdot + \frac{\phi(\mathbf{x}_{i,j,k-1}) - 2\phi(\mathbf{x}_{ijk}) + \phi(\mathbf{x}_{i,j,k+1})}{H^2}.$$

Applying the discrete Fourier transform to both sides and using the shift theorem we get

$$4\pi G \hat{\rho}(\mathbf{k}) = \frac{1}{H^2} \sum_{i=1}^{3} \left(e^{-iHk_i} + e^{iHk_i} - 2 \right) \hat{\phi}(\mathbf{k})$$
$$= \frac{1}{H^2} \sum_{i=1}^{3} \left(e^{iHk_i/2} - e^{-iHk_i/2} \right)^2 \hat{\phi}(\mathbf{k})$$
$$= -\frac{4}{H^2} \sum_{i=1}^{3} \sin^2 \left(\frac{Hk_i}{2} \right) \hat{\phi}(\mathbf{k}).$$

$$D(x_p) = \frac{1}{L} \sum_{l=0}^{N-1} \hat{D}(k)e^{ikx_p}, \quad \hat{D}(k) = H \sum_{p=0}^{N-1} D(x_p)e^{-ikx_p},$$

where $x_p = pH$. The conversion between this form and another popular definition,

$$\widetilde{D_H}(k) = \sum_{p=0}^{N-1} D_H(p)e^{-i2\pi kp/N},$$
(6)

is given by

$$\widetilde{D_H}(k) = \frac{1}{H} \hat{D} \left(\frac{2\pi}{NH} k \right),$$

where $D_H(p) = D(pH)$.

¹In this work, the Hockney & Eastwood definition of DFT is used, i.e.

and hence

$$\hat{\phi}(\mathbf{k}) = -4\pi G \underbrace{\frac{(H/2)^2}{\sin^2(Hk_1/2) + \sin^2(Hk_2/2) + \sin^2(Hk_3/2)}}_{\hat{\mathcal{G}}(\mathbf{k})} \hat{\rho}(\mathbf{k}), \tag{8}$$

where $\hat{\mathcal{G}}$ can be identified by comparison with Equation 7. It is worth noting that the constant multiplier $(-4\pi G)$ is often left out of $\hat{\mathcal{G}}$ (this is the convention used in [6]). In the implementation, values of $\hat{\mathcal{G}}$ are computed only once and saved for future look-up.

In the one-dimensional case, the above discussion can be easily rephrased in terms of a diagonalization problem [3]. In one dimension, the Poisson equation is

$$\frac{d^2\phi}{dx^2} = \rho. (9)$$

The interval [a,b] on which we wish to find ϕ is assumed to be discretized into N points $x_j = a + jH$, where H = (b-a)/(N-1) and j = 0, ..., N-1. Furthermore, we assume periodic boundary conditions so that $\phi(x_{-1}) = \phi(x_{N-1})$ and $\phi(x_N) = \phi(x_0)$. The approximation of Δ at x_j is

$$(\Delta_H \phi)_j \equiv \frac{\phi_{j-1} - 2\phi_j + \phi_{j+1}}{H^2}.$$

Thus, the discrete version of Equation 9 reads $(\Delta_H \phi)_j = \rho_j$ or

$$\frac{\phi_{j-1} - 2\phi_j + \phi_{j+1}}{H^2} = \rho_j,$$

where $\rho_j = \rho(x_j)$. This gives a system of N equations (one per each sampled value of ρ) with N unknowns (values of ϕ) of the following form

$$\underbrace{\frac{1}{H^2} \begin{bmatrix} 2 & -1 & & & -1 \\ -1 & 2 & \ddots & & \\ & \ddots & \ddots & \ddots & \\ & & \ddots & 2 & -1 \\ -1 & & & -1 & 2 \end{bmatrix}}_{\Delta_H} \underbrace{\begin{bmatrix} \phi_0 \\ \phi_1 \\ \vdots \\ \phi_{N-2} \\ \phi_{N-1} \end{bmatrix}}_{\bar{\phi}} = - \underbrace{\begin{bmatrix} \rho_0 \\ \rho_1 \\ \vdots \\ \rho_{N-2} \\ \rho_{N-1} \end{bmatrix}}_{\bar{\rho}}.$$
(10)

Next we define $\bar{\psi}(k) \in \mathbb{C}^N$ by $\psi_j(k) \equiv e^{i2\pi jk/N} = \omega^{jk}$ for j = 0, ..., N-1. The *j*-th row of $H^2\Delta_H\bar{\psi}$ equals

$$(H^{2}\Delta_{H}\bar{\psi})_{j} = -\psi_{j-1} + 2\psi_{j} - \psi_{j+1} = -\omega^{(j-1)k} + 2\omega^{jk} - \omega^{(j+1)k}$$
$$= -\omega^{jk}(\omega^{-k} - 2 + \omega^{k}) = -\omega^{jk}(\omega^{k/2} - \omega^{-k/2})^{2}$$
$$= 4\omega^{jk}\sin^{2}\left(\frac{\pi k}{N}\right).$$

Hence we have $(H^2\Delta_H \bar{\psi})_j = 4\sin^2(\pi k/N)\psi_j$ which implies

$$\Delta_H \bar{\psi}(k) = \frac{4}{H^2} \sin^2\left(\frac{\pi k}{N}\right) \bar{\psi}(k).$$

This results tells us that for any value of k, $\bar{\psi}(k)$ is an eigenvector of Δ_H with eigenvalue $(4/H^2)\sin^2(\pi k/N)$. For $k=0,\ldots,N-1$ we get N linearly independent eigenvectors $\{\bar{\psi}(k)\}$ which necessarily form the basis of \mathbb{C}^N . This implies that Δ_H is diagonalizable. The matrix F of its eigenvectors is given by $F_{jk} = \omega^{jk}$, and the inverse is easily verified to be $F^{-1} = (1/N)F^{\dagger}$. The eigendecomposition of Δ_H is therefore

$$\Delta_H = F\Lambda F^{-1}$$

where $\Lambda = \text{diag}((4/H^2)\sin^2(\pi k/N))$. Substitution into Equation 10 yields $F\Lambda F^{-1}\bar{\phi} = -\bar{\rho}$ or

$$F^{-1}\bar{\phi} = -\Lambda^{-1}F^{-1}\bar{\rho}.$$

Evaluating the left-hand side leads to the conclusion that $F^{-1}\bar{\phi}$ is nothing else but the DFT of ϕ . Indeed,

$$(F^{-1}\phi)_k = \frac{1}{N}(F^{\dagger}\phi)_k = \frac{1}{N}\sum_{j=0}^{N-1}F_{kj}^{\dagger}\phi_j = \frac{1}{N}\sum_{j=0}^{N-1}\omega^{-kj}\phi_j = \frac{1}{N}\sum_{j=0}^{N-1}e^{-2\pi ijk/N}\phi_j$$

which is exactly the discrete Fourier transform of ϕ (using the standard definition of the DFT). Thus we see that the DFT of the Green's function derived in Equation 8 (or rather the one-dimensional variant thereof) is the k-th eigenfunction of the discretized Laplacian (with slight differences due to a different definition of the DFT being used there). By following the line of reasoning presented above, we observe a deep connection between the DFT and the eigendecomposition of the discretized Laplace operator.

A natural question that arises in the context of this discussion is whether a similar relation holds in the continuous case. The answer turns out to be positive; assuming free-space boundary conditions, the complex exponential $e^{2\pi i \mathbf{k} \cdot \mathbf{x}}$ (kernel of the inverse Fourier transform) is an eigenfunction of the Laplace operator (with eigenvalue $-(2\pi k)^2$) and the Fourier transform allows for a "diagonalization" of the Laplacian [3]. More precisely, we have

$$\Delta = F\Lambda F^{-1}$$
.

where F is the inverse FT, and $\Lambda = -(2\pi k)^2$. Applying this result to the Poisson equation yields $F^{-1}\phi = \Lambda^{-1}Q^{-1}\rho$ or

$$\hat{\phi} = -\frac{1}{4\pi^2 k^2} \hat{\rho},$$

where the circumflex denotes the standard Fourier transform. The function $-(2\pi k)^2$ is sometimes used instead of the Green's function \mathcal{G} defined in Equation 8. This approach is the basis of the "poor man's Poisson solver" (as dubbed by [6]).

2.3 Field strength calculation

The strength \mathbf{g} of the gravitational field at mesh-point $\mathbf{x}_{\mathbf{p}}$ can be approximated using a central difference. Our implementation currently supports two types of finite differences, described below.

The two-point finite difference operator \mathbf{D} , whose x component is given by

$$D_x(\phi)(\mathbf{x_p}) = \frac{\phi(\mathbf{x}_{i+1,j,k}) - \phi(\mathbf{x}_{i-1,j,k})}{2H}$$

(and analogously for the y and z components), is second order accurate.

The fourth-order accurate finite difference is given by

$$D_x(\phi)(\mathbf{x}_{\mathbf{p}}) = \alpha \frac{\phi(\mathbf{x}_{i+1,j,k}) - \phi(\mathbf{x}_{i-1,j,k})}{2H} + (1 - \alpha) \frac{\phi(\mathbf{x}_{i+2,j,k}) - \phi(\mathbf{x}_{i-2,j,k})}{4H},$$

where $\alpha = 4/3$.

The difference between the accuracy of both methods is illustrated in Figure 2. The figure also provides insight into how the error depends on the value of parameter α .

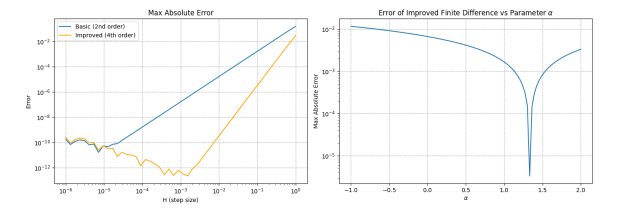


Figure 2: Left pane: Approximation error for second-order and fourth-order schemes. For small values of H, round-off errors dominate. Right pane: Approximation error vs. α in the improved finite difference scheme (H=0.1). Note that the scheme is fourth-order accurate only for $\alpha=4/3$ (cusp in the graph).

We can alternatively define the finite difference operators in terms of the delta function to get rid of the dependence on the differenced function. (Technically, the resulting quantities are functions rather than operators.) Consider for example the two-point finite difference in Equation 11. The definition can be generalized beyond mesh points by letting

$$D_j(\phi)(\mathbf{x}) = -\frac{\phi(\mathbf{x} + H\mathbf{e}_j) - \phi(\mathbf{x} - H\mathbf{e}_j)}{2H} = -\int \left[\frac{\delta(\mathbf{x} + H\mathbf{e}_j - \mathbf{x}') - \delta(\mathbf{x} - H\mathbf{e}_j - \mathbf{x}')}{2H} \right] \phi(\mathbf{x}') d\mathbf{x}',$$

where \mathbf{e}_{j} is the j-th standard basis vector. This motivates us to define

$$D_j(\mathbf{x}) = \frac{\delta(\mathbf{x} + H\mathbf{e}_j - \mathbf{x}') - \delta(\mathbf{x} - H\mathbf{e}_j - \mathbf{x}')}{2H}$$
(11)

and

$$D_{j}(\mathbf{x}) = \frac{\delta(\mathbf{x} + H\mathbf{e}_{j} - \mathbf{x}') - \delta(\mathbf{x} - H\mathbf{e}_{j} - \mathbf{x}')}{2H} + (1 - \alpha)\frac{\delta(\mathbf{x} + 2H\mathbf{e}_{j} - \mathbf{x}') - \delta(\mathbf{x} - 2H\mathbf{e}_{j} - \mathbf{x}')}{4H}$$
(12)

as the two-point as four-point finite difference operators respectively.

If ϕ denotes the gravitational potential, then the field ${f g}$ is approximated at mesh point ${f x_p}$ as

$$\mathbf{g}(\mathbf{x}_{\mathbf{p}}) = -\mathbf{D}(\phi)(\mathbf{x}_{\mathbf{p}}).$$

2.4 Interpolation

The value of the field strength $\mathbf{g}(\mathbf{x})$ at the position particle's position \mathbf{x} is calculated by interpolating the values of \mathbf{g} from the neighboring mesh-points. Formally,

$$\mathbf{g}(\mathbf{x}) = \sum_{\mathbf{p}} W(\mathbf{x} - \mathbf{x}_{\mathbf{p}}) \mathbf{g}(\mathbf{x}_{\mathbf{p}}).$$

Algorithm 2 Field strength interpolation

- 1: for each particle i do
- 2: **for each** cell \mathbf{q} in $\mathcal{C}_S(\mathbf{x}_i)$ **do**
- 3: $\mathbf{g}(\mathbf{x}_i) \leftarrow \sum_{\mathbf{q}} W(\mathbf{x}_i \mathbf{x}_{\mathbf{q}}) \mathbf{g}(\mathbf{x}_{\mathbf{q}})$

In practice, there is no need to sum over all mesh points. Instead, we use an algorithm analogous to Algorithm 1 to only include the cells with non-zero contribution to the sum. The method is illustrated in Algorithm 2. It is important to note that in order to retain correct physical behavior, the interpolation and mass assignment schemes must use the same shape to represent the particles. The procedure in Algorithm 2 is trivially parallelized by converting the sequential loop into a parallel one.

The procedures of density assignment and interpolation presented in Algorithm 1 and Algorithm 2 are high level description. More concrete formulations suitable for direct use in an implementation are given in [6] and [7].

2.5 Code units

Implementation of the PM (and P^3M) methods can be simplified by switching to a system of dimensionless units, often called *code units*. The natural units of time and length in a PM simulation are H and DT, respectively. Hence, length in a PM code is conveniently expressed in terms of multiples of H, and similarly time intervals are given as a multiple of DT, i.e. the conversion relations are

$$x' = \frac{x}{H}$$
 and $t' = \frac{t}{DT}$.

From there, it follows that

$$v' = \frac{DT}{H}v \quad \text{and} \quad a' = \frac{DT^2}{H}a.$$

The expected relation $\mathbf{g}' = -\nabla'\phi'$ leads to the definition $\phi' = (DT^2/H^2)\phi$. By stipulating that we have $\nabla'^2\phi = \rho'$, we get $\rho' = DT^2 \cdot 4\pi G\rho$, $m' = (DT^2 \cdot 4\pi G/H^3)m$, and $G' = 1/(4\pi)$.

2.6 Properties of the calculated field

The field produced by the PM method is neither homogenous nor isotropic. Anisotropy can be observed by measuring the field generated by a particle in two different directions. In Figure 3, the field strength due to a single source at x = H is shown in two variants: when measured along the x-axis (blue line) and the x = y line (orange line). The difference between these two graphs illustrates anisotropy of the calculated field. The figure also shows the field strength measured along the x-axis when the source was shifted by H/2 in the -x direction (green line). Its deviation from the case when the source was placed at x = H (the blue graph) exemplifies inhomogeneity of the field computed using the PM method. As expected, the PM-calculated approximation gets better with increasing distance from the source; the inverse-square law is reproduced accurately for $r \gtrsim 4H$.

The single-source case analysis can be extended by considering the effect of choice of finite difference and mass assignment schemes on the relative error between the actual and approximated field strength.

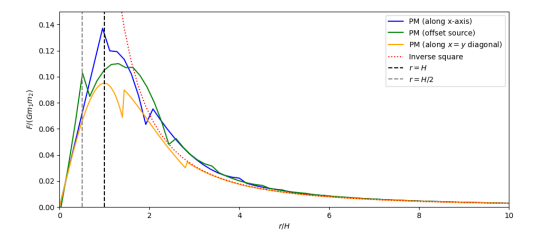
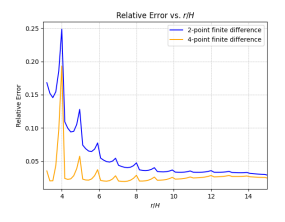
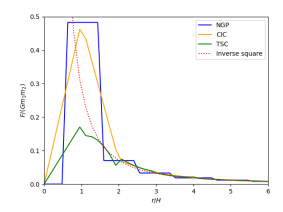


Figure 3: Anisotropy and inhomogeneity of the field as calculated by the PM method (TSC assignment, second order finite difference).

Figure 4 shows a comparison of (a) the relative error as a function of distance from the source for PM with second-order and fourth-order finite differences, and (b) the field strength computed using different mass assignment schemes discussed in subsection 2.1. The conclusion that can be drawn from Figure 4





- (a) Relative error of field strength approximation in different PM variants.
- (b) Field strength calculated with the PM method using different assignment schemes.

Figure 4: Comparison of PM method performance. (a) Relative error due to finite difference accuracy. (b) Effect of mass assignment schemes on computed field strength (4-point finite difference).

is that, as expected, the TSC mass assignment scheme and the four-point finite difference yield best results.

3 Particle-particle particle-mesh method

The P³M algorithm is a hybrid method: Forces between distant particles are calculated using the PM method, whereas, for particles lying closely together, the PP method is used. The total force applied to

particle i is

$$\mathbf{F}_{i}^{\mathrm{SR}} + \mathbf{F}_{i} = \sum_{j \neq i} (\mathbf{f}_{ij}^{\mathrm{tot}} - \mathbf{R}_{ij}) + \mathbf{F}_{i}, \tag{13}$$

where $\mathbf{F}_i \approx \sum_{j \neq i} \mathbf{R}_{ij}$ is the force computed using the PM method and $\mathbf{R}_{ij} = \mathbf{R}(\mathbf{x}_i - \mathbf{x}_j)$ is a prescribed reference force. The reference force is defined as the force between two particle-clouds, i.e. each particle is represented by a sphere with diameter a and a given density profile. The two examples of reference forces described in [6] are

$$R(r) = Gm_1m_2 \times \begin{cases} \frac{1}{35a^2}(224\xi - 224\xi^3 + 70\xi^4 + 48\xi^5 - 21\xi^6), & 0 \le \xi \le 1\\ \frac{1}{35a^2}(12/\xi^2 - 224 + 896\xi - 840\xi^2 + 224\xi^3 + 70\xi^4 - 48\xi^5 + 7\xi^6), & 1 < \xi \le 2\\ \frac{1}{r^2}, & \xi > 2 \end{cases}$$

where $\xi = 2r/a$ for a sphere with uniformly decreasing density (shape S_2) and

$$R(r) = Gm_1m_2 \times \begin{cases} \frac{1}{a^2} (8r/a - 9r^2/a^2 + 2r^4/a^4), & r < a \\ \frac{1}{r^2}, \end{cases}$$
 (14)

for a solid sphere (shape S_1). The reference force vector lies along the line joining the two bodies.

3.1 Optimal Green's function

As it is apparent from Equation 13, the method's validity depends on how well the reference force is approximated by the mesh force. The average deviation between the two forces can be minimized by a suitable choice of Green's function given (in k space) by

$$\hat{\mathcal{G}}(\mathbf{k}) = \frac{\hat{\mathbf{D}}(\mathbf{k}) \cdot \sum_{\mathbf{n}} \hat{U}^{2}(\mathbf{k}_{\mathbf{n}}) \hat{\mathbf{R}}(\mathbf{k}_{\mathbf{n}})}{|\hat{\mathbf{D}}(\mathbf{k})|^{2} \left[\sum_{\mathbf{n}} \hat{U}^{2}(\mathbf{k}_{\mathbf{n}})\right]^{2}}.$$
(15)

The derivation is mathematically involved and is detailed in [6]. We now proceed to examine the terms appearing in Equation 15.

3.1.1 Finite difference operator

The Fourier transform $\hat{\mathbf{D}}$ of the two-point finite difference operator defined in Equation 11 has the components

$$\begin{split} \hat{D}_{j}(\mathbf{k}) &= \int D_{j}(\mathbf{x})e^{-i\mathbf{k}\cdot\mathbf{x}}d\mathbf{x} \\ &= \frac{1}{2H}\left(\int \delta(\mathbf{x} + H\mathbf{e}_{j})e^{-i\mathbf{k}\cdot\mathbf{x}}d\mathbf{x} - \int \delta(\mathbf{x} - H\mathbf{e}_{j})e^{-i\mathbf{k}\cdot\mathbf{x}}d\mathbf{x}\right) \\ &= \frac{1}{2H}(e^{ik_{j}H} - e^{-ik_{j}H}) = \frac{i\sin(k_{j}H)}{H}. \end{split}$$

Similarly, for the four-point finite difference (Equation 12) we have

$$\hat{D}_j = \alpha \frac{i \sin k_j H}{H} + (1 - \alpha) \frac{i \sin 2k_j H}{2H},$$

where j = 1, 2, 3.

3.1.2 Assignment function

The quantity \hat{U} is defined as \hat{W}/V . For the mass assignment scheme hierarchy described in subsection 2.1 we have

$$\hat{U}(\mathbf{k}) = \left(\prod_{i=1}^{3} \frac{\sin(k_i H/2)}{k_i H/2}\right)^p,$$

where p = 1, 2, 3, ... with p = 1 corresponding to NGP assignment, etc. In particular, for the TSC assignment scheme, the alias sum^2

$$\sum_{\mathbf{n}} \hat{U}^2(\mathbf{k_n}) \equiv \sum_{\mathbf{n}} \hat{U}^2 \left(\mathbf{k} + \mathbf{n} \frac{2\pi}{H} \right)$$

can be rewritten as

$$\sum_{\mathbf{n}} \hat{U}^{2} \left(\mathbf{k} + \mathbf{n} \frac{2\pi}{H} \right) = \sum_{\mathbf{n}} \prod_{i=1}^{3} \left[\frac{\sin(k_{i}H/2 + n_{i}\pi)}{k_{i}H/2 + n_{i}\pi} \right]^{6}$$

$$= \prod_{i=1}^{3} \sum_{n_{i}} \left[\frac{\sin(k_{i}H/2 + n_{i}\pi)}{k_{i}H/2 + n_{i}\pi} \right]^{6}$$

$$= \prod_{i=1}^{3} \left[\sin^{6} \left(\frac{k_{i}H}{2} \right) \sum_{n_{i}} \frac{1}{(k_{i}H/2 + n_{i}\pi)^{6}} \right]$$

Using the the partial fractions expansion of the cotangent function [2],

$$\frac{(-1)^s}{s!} \frac{d^s}{dx^s} \cot x = \sum_{n = -\infty}^{\infty} \frac{1}{(x - n\pi)^{s+1}},$$

we can simplify the sum over n_i to

$$\frac{-1}{5!} \frac{d^5}{dx^5} \cot\left(\frac{k_i H}{2}\right) = 1 - \sin^2\frac{k_i H}{2} + \frac{2}{15} \sin^4\frac{k_i H}{2}.$$

Hence,

$$\sum_{\mathbf{n}} \hat{U}_{TSC}^{2}(\mathbf{k}_{\mathbf{n}}) = \prod_{i=1}^{3} \left(1 - \sin^{2} \frac{k_{i}H}{2} + \frac{2}{15} \sin^{4} \frac{k_{i}H}{2} \right).$$

Using the same approach, we can obtain similar results for the CIC and NGP schemes, namely

$$\sum_{\mathbf{n}} \hat{U}_{\text{CIC}}^2 = \frac{1}{3} \prod_{i=1}^{3} \left(1 + 2\cos^2 \frac{k_i H}{2} \right) \quad \text{and} \quad \sum_{\mathbf{n}} \hat{U}_{\text{NGP}}^2 = 1.$$

3.1.3 Reference force

The quantity $\hat{\mathbf{R}}$, the transformed reference force, is related to the shape S of the particle-cloud by

$$\hat{\mathbf{R}}(\mathbf{k}) = \frac{i\mathbf{k}\hat{S}^2(k)}{k^2},\tag{16}$$

where $k = |\mathbf{k}|$. This can be shown by recalling that $\mathbf{R}(\mathbf{x}_i - \mathbf{x}_j)$ is the force applied to cloud i due to cloud j. Hence $\mathbf{R}(\mathbf{x})$ is the force on cloud centered at \mathbf{x} due to a cloud at the origin. The force acting on mass element $d\mathbf{x}'S(\mathbf{x}' - \mathbf{x})$ of the cloud centered at \mathbf{x} is therefore

$$-Gd\mathbf{x}'S(\mathbf{x}'-\mathbf{x})\int d\mathbf{x}''S(\mathbf{x}'')\frac{\mathbf{x}'-\mathbf{x}''}{|\mathbf{x}'-\mathbf{x}''|^3}$$

$$\sum_{\mathbf{n}} \tilde{U}^2(\mathbf{k_n}) \equiv \sum_{\mathbf{n}} \tilde{U}^2(\mathbf{k} + \mathbf{n}N) = \frac{1}{H} \sum_{\mathbf{n}} \hat{U}^2 \left(\mathbf{k} \frac{2\pi}{NH} + \mathbf{n} \frac{2\pi}{H} \right)$$

instead.

 $^{^2\}mathrm{To}$ get the alias sums compatible with the DFT definition given in Equation 6, one has to compute

and the total force is

$$\mathbf{R}(\mathbf{x}) = -G \int d\mathbf{x}' S(\mathbf{x}' - \mathbf{x}) \int d\mathbf{x}'' S(\mathbf{x}'') \frac{\mathbf{x}' - \mathbf{x}''}{|\mathbf{x}' - \mathbf{x}''|^3}.$$
 (17)

The expression on the right-hand side of Equation 17 is a double convolution, $\mathbf{R} = -S * (S * \mathbf{g})$, where $\mathbf{g}(\mathbf{x}) = \mathbf{x}/|\mathbf{x}|^3$. The x-coordinate of \mathbf{g} is $x/|\mathbf{x}|^3$ which coincides with $\partial h/\partial x$ for $h(\mathbf{x}) = -1/|\mathbf{x}|$. The Fourier transform of h is given in [4] (p. 363):

$$\hat{h}(\mathbf{k}) = \frac{4\pi}{|\mathbf{k}|^2}.$$

The formula for the transform of a derivative yields

$$\hat{g}_x(\mathbf{k}) = \frac{4\pi i k_x}{|\mathbf{k}|^2}.$$

Applying the convolution theorem twice and factoring the constant $(-4\pi G)$ out of the Green's function in Equation 15 leaves us with the desired Equation 16.

Since the shapes S are spherically symmetric, the calculation of \hat{S} (appearing in Equation 16) can be simplified. The Fourier transform of S is

$$\hat{S}(\mathbf{k}) = \int S(\mathbf{x})e^{-i\mathbf{k}\cdot\mathbf{x}}d\mathbf{x}.$$

Observe that $\mathbf{k} \cdot \mathbf{x} = kr \cos \theta$, where θ is the angle between \mathbf{k} and \mathbf{x} , and $r = |\mathbf{x}|$. Since S is invariant under rotations, θ can be chosen to be the angle between \mathbf{x} and the z-axis. Thus, the integral, rewritten in spherical coordinates becomes

$$\hat{S}(k) = \int_0^{2\pi} d\phi \int_0^{\pi} d\theta \int_0^{\infty} dr S(r) e^{-ikr\cos\theta} r^2 \sin\theta = 2\pi \int_0^{\infty} r^2 dr \int_0^{\pi} d\theta e^{-ikr\cos\theta} \sin\theta.$$

The θ -integral upon the substitution $-kr\cos\theta \to u$ becomes

$$\frac{1}{kr} \int_{-kr}^{kr} e^{iu} du = \frac{1}{kr} \int_{-kr}^{kr} (\cos u + i \sin u) du = \frac{2\sin kr}{kr}$$

and hence

$$\hat{S}(k) = 4\pi \int_0^\infty r^2 S(r) \frac{\sin kr}{kr} dr.$$

This integral, evaluated for the S_1 and S_2 shapes respectively, gives

$$\hat{S}_1(k) = \frac{3}{(ka/2)^3} \left(\sin \frac{ka}{2} - \frac{ka}{2} \cos \frac{ka}{2} \right)$$

and

$$\hat{S}_2(k) = \frac{12}{(ka/2)^4} \left(2 - 2\cos\frac{ka}{2} - \frac{ka}{2}\sin\frac{ka}{2} \right).$$

Finally, we note that the infinite sum in the numerator of Equation 15 does not have a closed form but this does not pose a problem since the summand decays rapidly with **n** moving further away from zero.

The result of applying the optimal Green's function in Equation 7 is illustrated in Figure 5a. As can be seen, the PM force closely follows the reference force, especially for distances r > a, where the reference force nearly matches the inverse-square law. Interestingly, for r slightly smaller than a, the reference force, and its mesh-based PM approximation, still provides a good fit to the inverse-square behavior.

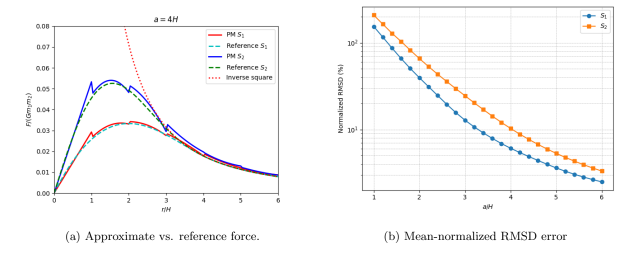


Figure 5: Comparison of PM approximation using S_1 and S_2 shape functions. The mesh approximation to the reference force was computed using the PM method with a TSC assignment scheme, two-point finite difference, and Green's function optimal for each shape.

This observation allows the cutoff radius r_e , which defines the boundary for direct summation, to be chosen smaller than a (e.g., $r_e = 0.7a$), improving performance without sacrificing accuracy. Figure 5a indicates that using the S_2 shape function in the PM calculation results in a force that more closely approximates the inverse-square law at small r, whereas the S_1 -based method begins to show noticeable deviations. However, Figure 5b shows that the S_2 -based approximation exhibits higher mean-normalized RMSD error compared to S_1 , especially at intermediate distances. Therefore, the choice between S_1 and S_2 shapes involves a trade-off between accuracy near the source and overall error, making the optimal choice context-dependent.

3.2 Identifying close pairs of particles

In the P^3M method, in addition to the mesh used in the PM algorithm (the "potential mesh"), a second mesh (the *chaining mesh*) is used. The chaining mesh is sparser than the potential mesh. Its sole purpose is to partition the space into cells so that particles "close" to the ones in a given cell can be found efficiently. In this context, two particles are considered to lie close to one another if their separation is less than the cutoff radius.

The number of chaining mesh cells in a single dimension is given by $M_i = \lfloor L_i/r_e \rfloor$, where L_i is the side length of the computational box (i = 1, 2, 3). This implies that the side length of a chaining mesh cell is $HC_i = L_i/M_i \ge r_e$. Thus, for every particle i in a given cell \mathbf{p} , it is sufficient to search through the immediate neighborhood of \mathbf{p} to find all the particles within the cutoff radius from i.

In our program, the chaining mesh is implemented as a head-of-chain (HOC) array, depicted in Figure 6. The basic version of the HOC array is very cheap to build. Given a particle at point $\mathbf{x} = (x, y, z)$, we can determine in which chaining mesh cell \mathbf{c} it lies by simply performing integer division on each of its coordinates, i.e. $\mathbf{c} = (x/\text{HC}_1, y/\text{HC}_2, z/\text{HC}_3)$. Then, to get a HOC table index i corresponding to \mathbf{c} , standard index flattening is used. Subsequently, a new linked-list node is allocated and inserted at

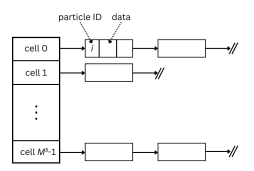


Figure 6: Head-of-chain data structure used for mapping particles to their parent cells in the chaining mesh. Here $M_1 = M_2 = M_3 = M$.

the beginning of the list HOC[i]. Clearly, the whole process is linear in the number of particles and the array can be constructed anew at each time step without degrading performance. In our implementation, additional savings are made by preallocating a memory pool large enough to store N nodes of the linked lists and reusing it for the HOC array initialization.

A potential optimization discussed in [6] involves sorting each linked list by a selected particle coordinate, such as the y-coordinate. This ordering enables early termination of the direct summation loop when the condition $y_i - y_j > r_e$ is met, effectively reducing unnecessary pairwise computations as particle i "sweeps through" a cell containing particles j. Our experiments indicate that constructing the linked lists in sorted order is substantially more expensive than the unsorted variant. This is to be expected; in the worst-case scenario where all particles are in a single chaining-mesh cell, and are inserted into the list in decreasing order of y coordinates, the complexity is $O(N^2)$. For a system of 50,000 particles, incorporating y-sorting increased the HOC (head-of-chain) construction time from approximately 180 milliseconds to over 13,000 milliseconds, a 70-fold slowdown. Despite this added cost, we observed no significant performance improvement in the short-range correction phase of the computation.

3.3 Short-range correction

The short-range correction, which takes place immediately after the mesh forces are found using the PM method, is at the heart of the P^3M algorithm. Since it scales with a square of the number of particles in each neighborhood, further optimizations are highly desirable.

By Newton's 3rd law, $\mathbf{f}_{ji}^{\mathrm{SR}} = -\mathbf{f}_{ij}^{\mathrm{SR}}$, which allows us to do the calculation of the short-range interparticle force for any pair (i,j) of particles only once, leading to the reduction of the total running time by half. Informally, the particle i updates its total short-range force $\mathbf{F}_i^{\mathrm{SR}}$ as well as the total short-range force $\mathbf{F}_j^{\mathrm{SR}}$ of its neighbor j. To avoid double-counting, the particle i residing in cell \mathbf{q} has to look for its neighbors in a subset \mathcal{N} of the immediate neighborhood of \mathbf{q} . More specifically, define

$$\mathcal{N}(\mathbf{q} = (q_1, q_2, q_3)) = \{(q_1 + t, q_2 - 1, q_3 + s), (q_1 + s, q_2, q_3 - 1), (q_1 - 1, q_2, q_3) \mid s, t = -1, 0, 1\}.$$
(18)

Thus $|\mathcal{N}| = 13$, which is half of the size of the immediate neighborhood. The set \mathcal{N} given in Equation 18 is not easily to illustrate. Since for the purposes of the later discussion, it is beneficial to have a clear picture in mind, we depict its two-dimensional analog in Figure 7.

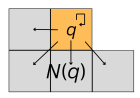


Figure 7: Set \mathcal{N} in two dimensions.

The short-range correction part of the P³M method is shown in Algorithm 3. For each chaining mesh

Algorithm 3 Short-range correction

```
1: for each chaining cell q do
        for each q_n \in \mathcal{N}(\mathbf{q}) \cup \{\mathbf{q}\} do
2:
            for each i \in HOC(q) do
3:
                 for each j \in HOC(\mathbf{q}_n) do
4:
                      if |y_i - y_j| > r_e then
5:
                          break
6:
                      UPDATESHORTRANGE(i, j, \mathbf{q}, \mathbf{q}_n)
7:
```

cell \mathbf{q} , we compute the pair-wise interactions between the particles in \mathbf{q} and the particles in the chaining mesh cells \mathbf{q}_n from the reduced neighborhood $\mathcal{N}(\mathbf{q})$ plus \mathbf{q} . The inclusion of \mathbf{q} allows us to take into account forces between particles within q.

The UPDATESHORTRANGE procedure is defined in Algorithm 4. In line 2, we exclude self-forces.

Algorithm 4 Updating short-range forces

```
1: procedure UPDATESHORTRANGE(i, j, \mathbf{q}, \mathbf{q}_n)
                 if i = j then return
  2:
                 \mathbf{r}_{ij} \leftarrow \mathbf{r}_i - \mathbf{r}_j
  3:
                 if |\mathbf{r}_{ij}|^2 > r_e^2 then return
  4:
                 r_{ij} \leftarrow |\mathbf{r}_{ij}|
  5:
                 \hat{\mathbf{r}}_{ij} \leftarrow \mathbf{r}_{ij}/r_{ij}
                 \mathbf{R}_{ij} \leftarrow -m_i m_j R(r_{ij}) \mathbf{\hat{r}}_{ij}
  7:
                 \mathbf{f}^{\text{tot}} \leftarrow -Gm_i m_j / r_{ij}^2 \mathbf{\hat{r}}_{ij}
                 \mathbf{f}_{ij}^{\mathrm{SR}} \leftarrow \mathbf{f}^{\mathrm{tot}} - \mathbf{R}_{ij}
                 \mathbf{f}_{ji}^{\mathrm{SR}} \leftarrow -\mathbf{f}_{ij}^{\mathrm{SR}}
10:
                 \mathbf{F}_{i}^{\mathrm{SR}} \leftarrow \mathbf{F}_{i}^{\mathrm{SR}} + \mathbf{f}_{ii}^{\mathrm{SR}}
11:
                 if q_n \neq q then
                                                                                                                                                              ▷ Avoid double-counting in the parent cell
12:
                          \mathbf{F}_{j}^{\mathrm{SR}} \leftarrow \mathbf{F}_{j}^{\mathrm{SR}} + \mathbf{f}_{ji}^{\mathrm{SR}}
13:
```

The check in line 4 assures that the correction happens only for particles with separation less than the cutoff radius r_e . The pair-wise short-range force is calculated in lines 5–10, with the calculation in line 10 exploiting the Newton's third law, as described previously. The accumulation of total short-range force for a given particle takes place in line 11. The short-range force is also added to the total short-range force of the other particle in the pair (particle j) but only if $\mathbf{q}_n \neq \mathbf{q}$. This stipulation is crucial, as otherwise we would be double counting the forces between particles within \mathbf{q} .

As suggested in [6], the computational burden of operations in lines 5–8 in Algorithm 4 can be greatly reduced by storing the values of $f^{\rm SR}(r)/r = (f^{\rm tot}(r) - R(r))/r$ in a lookup table T at uniform intervals Δ^2 of $[0, r_e^2]$ and interpolating them later. The schematic depiction of the interpolation is shown in Figure 8. If we define $\xi = r^2/\Delta^2$ and $t = \lfloor \xi \rfloor$, then

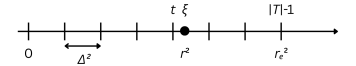


Figure 8: Interpolation of short-range force values.

$$\frac{f^{SR}(r)}{r} \approx T[t] (1 - (\xi - t)) + T[t + 1](\xi - t) = T[t] + (\xi - t)(T[t + 1] - T[t]).$$

The value $\mathbf{f}_{ij}^{\mathrm{SR}}$ can then be obtained by multiplying the interpolated quantity $f^{\mathrm{SR}}(r_{ij})/r_{ij}$ by $Gm_im_j\mathbf{r}_{ij}$, eliminating the use of the square root operations and reducing the total number of floating-point operations to just four.

In our implementation, the procedure outlined in Algorithm 3 is parallelized by splitting the work done in the outmost loop between some number of threads. In doing so, extra care has to be taken to avoid data races. A thread t that is currently processing cell \mathbf{p} and its neighbors (we say that t is assigned to \mathbf{p}) may "clash" with a different thread assigned to a nearby cell \mathbf{q} (because possibly $\mathbf{p} \in \mathcal{N}(\mathbf{q})$). However, by the construction of the set \mathcal{N} , it is possible to split the short-range force into 14 parts, each of which is accessed by only one thread. For example, consider a particle i in cell $\mathbf{p} = (p_1, p_2, p_3)$ (in other words, \mathbf{p} is the parent cell of i). If thread t is currently assigned to this cell, t will update the part of \mathbf{F}_i^{SR} corresponding to updates of i coming from within the same cell as the parent cell of i. Possibly at the same time, thread t' assigned to cell $\mathbf{q} = (p_1 + 1, p_2, p_3)$ will update a different part of $\mathbf{F}_{i}^{\mathrm{SR}}$, i.e. the part corresponding to updates of i coming from the cell "to the right" of the parent cell of i. Since only one thread is responsible for updates to particle i coming "from the right," (or any other "direction") no data races can occur. This approach presents two major drawbacks. First, it significantly increases memory usage, requiring storage for an additional 13N three-dimensional vectors. Second, it offers no guarantee of uniform workload distribution across threads. This imbalance, combined with the substantial variation in operations performed by individual threads, leads to severe thread divergence, rendering the parallelization scheme unsuitable for GPU execution.

4 Barnes-Hut algorithm

The idea behind Barnes-Hut algorithm differs substantially from previously described mesh-based methods. The algorithm deals with gravitational forces directly instead of deriving them from the mesh-defined potential, as was the case with the PM and P^3M methods. Significant reduction of time complexity, from quadratic to $O(N \log N)$, is achieved by approximating the potential due to "far enough" groups of particles by the initial terms of its multipole expansion [10]. The grouping of particles is hierarchical in nature and is thus best understood as a tree. The entire set of particles comprises the top-level group, represented by the root of the tree; the eight children of the root node are representative of groups of particles residing in each of the octants of the computational domain, etc. The process of subdividing the space into eight smaller volumes at each node continues recursively until there is only one or zero particles left in a given volume. Nodes which satisfy this condition are the leafs of tree and are sometimes called the external nodes. The remaining nodes, each of which has eight children, are called internal nodes.

In the basic variant of the algorithm, the potential due to a group of particles is approximated using only the monopole term with respect to the center of mass of the group, i.e.

$$\phi_{\rm mon}(r) = -\frac{GM}{r},$$

where M is the group's total mass. Since the potential is expanded about the center of mass, the dipole moment $\mathbf{p} = \sum_i m_i \mathbf{r}_i$ vanishes. Hence, the next possible improvement comes from including the quadrupole term

$$\phi_{\text{quad}}(r) = -\frac{G}{2r^5} \mathbf{r} \cdot (\mathbf{Qr}),$$

where \mathbf{Q} is the quadrupole moment tensor defined as

$$Q_{ij} = \sum_{k} (3r_{ki}r_{kj} - 3r_k^2 \delta_{ij}) m_k.$$

In theory, we could keep on adding more terms to improve the quality of the approximation. In our implementation however, we restrict ourselves to the quadrupole term.

4.1 Building the tree

The data structure that fits the description given in the introduction is called an *octree*. An internal node of the octree stores the COM vector, the total mass of the group it represents, and the quadrupole tensor, whereas an external node stores a reference to the actual particles (or is empty if no particle was found in its associated volume). The recursive procedure of building the tree is shown in Algorithm 5. The quadrupole moment tensor for each node is calculated once the whole tree is already built. The recursive relation used in this calculation is given in [5] and reads

$$\mathbf{Q} = \sum_{\text{child } c} \mathbf{Q}_c + \sum_{\text{child } c} m_c (3\mathbf{R}_c \otimes \mathbf{R}_c - R_c^2 \mathbf{I}),$$

where $\mathbf{R}_c = \mathbf{x}_c^{\text{COM}} - \mathbf{x}^{\text{COM}}$ is the displacement vector from the COM of child c to the COM of the parent, \mathbf{I} is the identity matrix, and \otimes denotes the outer product.

4.2 Acceleration calculation

In the Barnes-Hut algorithm, the net acceleration of a particle p is calculated by summing the contributions from single particles or groups of particles while traversing the tree. The decision whether the

Algorithm 5 Insert a particle into the Barnes-Hut tree

```
1: function Insert(n, p)
       if n is an internal node then
2:
3:
           Update n.COM and total mass n.M of n with p
           Insert(child of n that should contain p, p)
 4:
       else if n is empty then
 5:
           Assign p to n
 6:
 7:
       else
                                                                                 \triangleright Occupied external node
           Subdivide n into child nodes
 8:
           Move existing particle p' in n into child that should contain p'
9:
           Update center of mass and total mass of n with p and p'
10:
           Insert(child of n that should contain p, p)
11:
```

acceleration can be approximated using the information stored in an internal node n depends on the relative distance from p to n.COM (the center of mass of group represented by n). The distance is relative to the width H of the node, i.e. the side length of the cubical volume encompassed by the node. More concretely, the approximation takes place if $n.H/|n.COM - p.\mathbf{x}| < \theta$, where θ is the so-called opening angle. In the extreme case when θ is set to zero, no approximations take place, and the algorithm reduces to the PP method. The procedure described above is illustrated in Algorithm 6. In the implementation,

Algorithm 6 Compute gravitational force on a particle using Barnes-Hut approximation

```
1: function FINDACCELERATION(n, p, \theta)
         if n is an external node then
 2:
3:
              if n contains a particle q \neq p then
                   p.\mathbf{a} \leftarrow p.\mathbf{a} + \text{GravitySoft}(q.\mathbf{x}, q.m, p.\mathbf{x})/p.\text{mass}
 4:
              return
 5:
         if n.H/|n.COM - p.\mathbf{x}| < \theta then
 6:
              p.\mathbf{a} \leftarrow p.\mathbf{a} + \text{Gravity}(n.\text{COM}, n.M, p.\mathbf{x})
 7:
              return
 8:
         for each child n_c of n do
9:
              FINDACCELERATION(n_c, p, \theta)
10:
```

the GravitySoft function calculates the gravitational force softened by ϵ , i.e. it returns the value

$$\mathbf{F}_{ij}^{\text{soft}} = -G \frac{m_i m_j}{(r_{ij}^2 + \epsilon^2)^{3/2}} \mathbf{r}_{ij}.$$

The pairwise potential energy associated with this force is given by

$$\Phi_{ij}^{\text{soft}} = -\frac{Gm_i m_j}{\sqrt{r_{ij}^2 + \epsilon^2}}.$$
(19)

The GRAVITY function returns the approximation (up to the quadrupole term) of the acceleration due to a group of particles represented by a given node, i.e.

$$\mathbf{a} = -GM\frac{\mathbf{r}}{r^3} + \frac{G}{r^5}\mathbf{Qr} - \frac{5G}{2}(\mathbf{r} \cdot (\mathbf{Qr}))\frac{\mathbf{r}}{r^7}$$

(see [5]).

One possible way to quantify the quality of approximation for a given value of θ is to consider the relative error of calculated force. We set the same initial conditions of the system for both the PP direct summation method and the Barnes-Hut algorithm, compute the deviation of Barnes-Hut forces from PP forces acting on each particle, and take the average over all particles. In other words, the error calculated is

$$\frac{1}{N} \sum_{i} \frac{|\mathbf{F}_{i}^{\mathrm{BH}} - \mathbf{F}_{i}^{\mathrm{PP}}|}{|\mathbf{F}_{i}^{\mathrm{PP}}|}.$$

The dependence of the error on the opening angle θ and the corresponding execution time are shown in Figure 9. The figure includes plots for two cases: when only the monopole term is used in the Barnes-Hut approximation, and when both the monopole and quadrupole terms are included. As can be seen, the quadrupole-based algorithm exhibits significantly improved error scaling with increasing θ . Naturally, this raises the question of the additional computational cost incurred by the inclusion of the quadrupole term. Our tests showed that this impact is minimal. The results (for N = 10,000 particles and $0 \le \theta \le 2$) support this observation. Both tests described above were conducted on a uniform disk

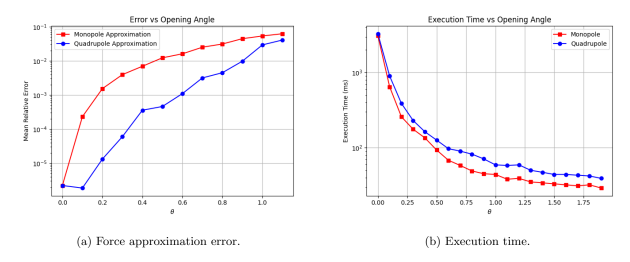


Figure 9: Comparison of error and execution time in the Barnes-Hut algorithm using monopole and quadrupole approximations.

particle distribution.

We note that direct calculation of total potential energy is infeasible as $O(N^2)$ operations would be required. Instead, we use an approximation based on the values stored in the tree. The approximate value of the potential energy is accumulated for each particle using a procedure analogous to force calculation. Indeed, the only difference between the two is the replacement of gravitational force calculation in Algorithm 6 with potential energy calculation according to Equation 19.

It is also noteworthy that the procedure outlined in Algorithm 6 is embarrassingly parallel. In our CPU implementation, the workload is split between an arbitrary number of threads on particle-by-particle basis.

5 Time integration

In the previous sections we described various methods of calculating forces applied to particles in the simulation. Once these forces are found, the evolution of the system in time can be tracked by integrating Newton's 2nd law of motion,

$$\ddot{\mathbf{x}}_i = \frac{\mathbf{F}_i}{m_i}.\tag{20}$$

5.1 Euler's method

Possibly, the simplest numerical method that could be used is Euler's method described by the update rules

$$\mathbf{v}_{i}^{(k+1)} = \mathbf{v}_{i}^{(k)} + \mathrm{DT} \frac{\mathbf{F}_{i}^{(k+1)}}{m_{i}},$$

$$\mathbf{x}_{i}^{(k+1)} = \mathbf{x}_{i}^{(k)} + \mathrm{DT} \mathbf{v}_{i}^{(k)}.$$
(21)

The method defined in Equation 21 is not suitable for physical simulations, however. Its shortcomings are best illustrated by an example of an undamped pendulum of length l in gravitational field of magnitude g. Although it is a very simple system, it illustrates the numerical challenges faced in gravitational simulations over long timescales, particularly the issue of energy preservation.

The motion of the pendulum is governed by the differential equation

$$\ddot{\theta} = -\frac{g}{l}\sin\theta,$$

and its kinetic and potential energy are given by $KE = (1/2)ml^2\dot{\theta}$ and $PE = -mgl\cos\theta$ respectively. As

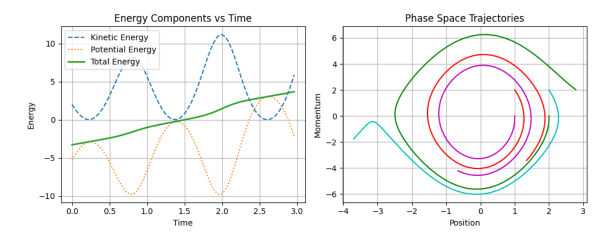


Figure 10: Behavior of Euler's method: lack of conservation of energy and phase space trajectories spiraling out.

shown in Figure 10, Euler's method fails to conserve total energy (PE + KE) and produces trajectories

in phase space that are not closed, contrary to expectations for periodic systems. Additionally, the evolution of an area element in phase space violates Liouville's theorem, as described in [8], making the method unsuitable for long-term physical simulations (see Figure 11).

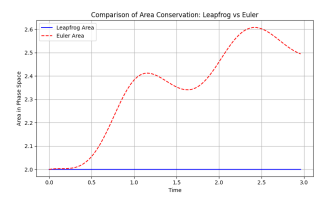


Figure 11: Area in phase space over time. Violation of Liouville's theorem by Euler's method.

5.2 Leapfrog algorithm

The leapfrog algorithm, given by the update rule [12]

$$\mathbf{v}_{i}^{(1/2)} = \mathbf{v}_{i}^{(0)} + \frac{1}{2} DT \frac{\mathbf{F}_{i}^{(0)}}{m_{i}^{(0)}},$$

$$\mathbf{x}_{i}^{(k+1)} = \mathbf{x}_{i}^{(k)} + DT \mathbf{v}_{i}^{(k+1/2)},$$

$$\mathbf{v}_{i}^{(k+3/2)} = \mathbf{v}_{i}^{(k+1/2)} + DT \frac{\mathbf{F}_{i}^{(k+1)}}{m_{i}}.$$
(22)

is the preferred way of integrating Equation 20. When applied to the same pendulum system, it conserves energy much more faithfully and preserves the area in phase space, consistent with Liouville's theorem (see Figure 12 and Figure 11). Given its simplicity and excellent long-term energy behavior, we adopt the leapfrog algorithm to integrate Newton's equations in our program.

6 Galaxy model

The model of a galaxy used as a test bed for the implementation is a simple one. The galaxy is assumed to comprise only two parts: a thin disk and a spherically symmetric halo. The disk comprises a large number of particles, each representing some number of stars. The halo is simulated as a fixed external gravitational field. The schematic illustration of the model is shown in Figure 13.

6.1 Disk

The disk particles are sampled from a radial distribution

$$p(r) = \frac{3}{\pi R_D^2} \left(1 - \frac{r}{R_D} \right), \ z = 0,$$

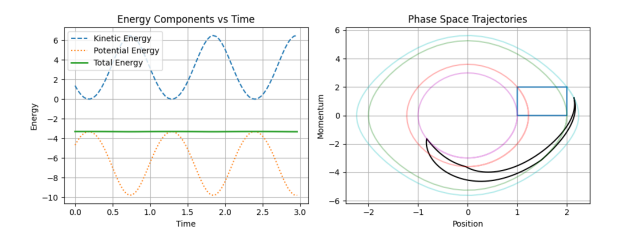


Figure 12: Behavior of the leapfrog algorithm: conservation of energy and phase space trajectories forming closed loops. Evolution of an area element in phase space is shown on the right-hand side: blue rectangle – initial conditions for many copies of the system; black distorted quadrilateral – their state by the end of the simulation.

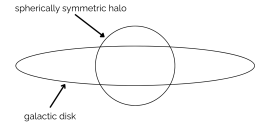


Figure 13: Spiral galaxy model (thin disk and spherical halo).

where R_D is the radius of the disk and $r \leq R_D$. The cumulative distribution function is therefore

$$F(r,\phi) = \int_0^{\phi} \int_0^r p(r')r'dr'd\phi' = \frac{\phi}{2\pi R_D^3} (3R_D r^2 - 2r^3)$$

and the marginal CDFs are

$$F_R(r) = F(r, 2\pi) = \frac{1}{R_D^3} (3R_D r^2 - 2r^3)$$
 and $F_{\Phi}(\phi) = F(R_D, \phi) = \frac{\phi}{2\pi}$.

Now we use inverse transform sampling to generate initial positions (r, ϕ) for the particles, i.e. $\phi = 2\pi u$ and r is given implicitly by $h(r) \equiv 2r^3 - 3R_Dr^2 + uR_D^3 = 0$ with $u \sim U(0,1)$. A straightforward calculation shows that dh/dr < 0 for $0 < r < R_D$ and $h(0)h(R_D) < 0$ implying that h has exactly one zero between 0 and R_D (which can be found for example using Newton's method).

Strength of the gravitational field \mathbf{g}_D due to the disk at point \mathbf{x}_0 lying in the disk is

$$\mathbf{g}_D = G \int_0^{2\pi} \int_0^{R_D} \sigma(r) \frac{\mathbf{x} - \mathbf{x}_0}{|\mathbf{x} - \mathbf{x}_0|^3} r dr d\phi,$$

where $\sigma(r) = \sigma_0(1 - r/R_D)$ describes the density profile of the disk for $r \leq R_D$. If M_D is the total mass of the disk, then $\sigma_0 = 3M_D/(\pi R_D^2)$. By symmetry, the point \mathbf{x}_0 may be chosen to lie on the x-axis, i.e. $\mathbf{x}_0 = (-x_0, 0)$, so that $\mathbf{x} - \mathbf{x}_0 = (x_0 + r\cos\phi, r\sin\phi)$. Letting $\bar{r} = r/R_D$ and $\bar{x}_0 = x_0/R_D$, the integral becomes

$$\mathbf{g}_{D} = G\sigma_{0} \int_{0}^{2\pi} \int_{0}^{1} (1 - \bar{r}) \frac{(\bar{x}_{0} + \bar{r}\cos\phi, \bar{r}\sin\phi)}{|(\bar{x}_{0} + \bar{r}\cos\phi, \bar{r}\sin\phi)|^{3}} \bar{r}d\bar{r}d\phi.$$

By symmetry $g_{D,y} = 0$ and thus the radial component of the field \mathbf{g}_D at distance $R\bar{x}_0$ from the center is

$$g_{D,r} = -|\mathbf{g}_D| = -G\sigma_0 \int_0^{2\pi} \int_0^1 (1-\bar{r}) \frac{\bar{x}_0 + \bar{r}\cos\phi}{(\bar{x}_0^2 + \bar{r}^2 + 2\bar{x}_0\bar{r}\cos\phi)^{3/2}} \bar{r}d\bar{r}d\phi. \tag{23}$$

If the disk had constant density, \mathbf{g}_D could be expressed in terms of elliptic integrals [11]. However, to the best of the author's knowledge, the integral in Equation 23 cannot be further simplified. For this reason, a crude approximation with a quadratic function is used: $g_{D,r} \approx a(r-h)^2 + k$, where the values k = 2.5 and h = 0.66 (the maximum of $g_{D,r}$ and the argument thereof) were estimated based on the graph of $g_{D,r}$ (see Figure 14). The value of $a = -k/h^2$ can be found by setting $g_{D,r}(0) = 0$ in the approximate formula.

6.2 Halo

The density profile of the halo is analogous to the one used for the disk, save for the fact it is 3-dimensional, i.e.

$$\rho(r) = \begin{cases} \rho_0 \left(1 - \frac{r}{R_H} \right), & r \le R_H \\ 0, & \text{otherwise,} \end{cases}$$

where R_H is the radius of the halo. If we let M_H be the mass of the halo, then $\rho_0 = 3M_H/(\pi R_H^3)$. Application of Gauss's law shows that we have

$$g_{H,r} = -GM_H \times \begin{cases} \frac{r}{R_H^3} \left(4 - \frac{3r}{R_H} \right), & r \leq R_H \\ \frac{1}{r^2}, & \text{otherwise.} \end{cases}$$

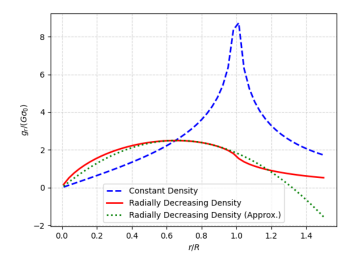


Figure 14: Magnitude of the radial component of the field strength due to a disk. The peak at r = R for the constant density disk is in fact infinite.

6.3 Initial conditions

The total field $\mathbf{g} = \mathbf{g}_D + \mathbf{g}_H$ is used to find initial velocities for the particles with initial positions (x, y, 0). The formula for the centripetal force yields

$$\frac{v^2}{r} = -g_r$$

and thus

$$\mathbf{v} = \left(-v\frac{y}{r}, v\frac{x}{r}, 0\right)$$

with $v = \sqrt{-rg_r}$ for counter-clockwise rotation.

6.4 Disk with a hole

While testing the implemented methods, we observed that for modeling systems comprising multiple galaxies it may be beneficial to replace the fixed halo with a single massive particle. In such case, the simulation turns out to stable if there are no particles in close vicinity to the galaxy center. This means that the galactic disk ought to have a hole of radius r_0 . Using an approach analogous to the one used in subsection 6.1, we obtain the following relations that can be used for sampling points (ϕ, r) in polar coordinates

$$\phi = 2\pi u$$
 and $2(r^3 - r_0^3) - 3R_D(r^2 - r_0^2) + (R_D - r_0)^2(2r_0 + R_D)u = 0$,

where $u \sim U(0,1)$.

Getting rid of the external halo field simplifies the treatment of complex systems significantly, as we do not need to keep track of the movement of the halo. The massive particle in the galaxy center, which effectively replaces it, can be treated just as any other particle in the simulation.

7 Globular cluster model

A globular cluster is a formation comprising a large number stars, closely packed in a spherically symmetric form [9]. In this work, we decided to use the implemented methods to simulate such structure using the Plummer model. The model was chosen due to its simplicity and abundance of dedicated resources.

In the Plummer model, the density of a cluster is given by

$$\rho(r) = \frac{3M}{4\pi a^2} \left(1 + \frac{r^2}{a^2} \right)^{-5/2},\tag{24}$$

where the parameter a controls the spread of the distribution (the size of the cluster core) [1]. Hence, the particles are sampled from a distribution with PDF $p(r) = \rho(r)/M$. Similarly to the galaxy model, we use inverse transform sampling to initialize the positions of the particles. The marginal CDFs are easily calculated as

$$F_R(r) = \frac{r^3}{a^3} \left(1 + \frac{r^2}{a^2} \right)^{-3/2}, \quad F_{\Theta}(\theta) = \frac{1}{2} (1 - \cos \theta), \quad F_{\Phi}(\phi) = \frac{\phi}{2\pi},$$

which means that given a random variable $u \sim U(0,1)$, we can generate random points with spherical coordinates

$$r = a(u^{-2/3} - 1)^{-1/2}, \quad \theta = \cos^{-1}(1 - 2u), \quad \phi = 2\pi u,$$
 (25)

consistent with Equation 24 (assuming equal mass of all particles).

In the Plummer model, the potential is given by

$$\phi(r) = -\frac{GM}{\sqrt{r^2 + a^2}}.$$

This allows us to find the escape velocity v_e at distance r from the center. Using the law of conservation of energy, we get

$$\frac{1}{2}v_e + \phi(r) = 0 \Rightarrow v_e = \sqrt{-2\phi(r)}.$$

For any r, the probability distribution of $q \equiv v/v_e$ is given by [1]

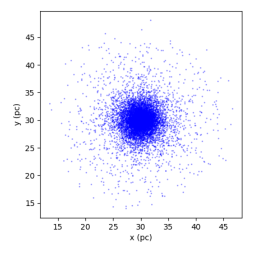
$$q(q) = Nq^2(1 - q^2)^{7/2}, (26)$$

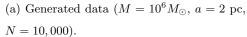
where N is a normalization constant, which can be calculated to be $N=512/(7\pi)$. The CDF of the distribution in Equation 26, $G(q)=\int_0^q g(q')dq'$, can be determined using symbolic integration. The algebraic expression that one obtains in this way is lengthy and we will not cite it here. The random values of q, consistent with the PDF in Equation 26, are again obtained using inverse transform sampling; the equation G(q)=u is solved for q by finding the roots of G(q)-u using Newton's method. Finally, the magnitude v of the velocity vector \mathbf{v} is set to $v=qv_e$. The direction of $\mathbf{v}=(v_x,v_y,v_z)$ is chosen uniformly at random, i.e.

$$v_x = v \sin \theta \cos \phi$$
, $v_y = v \sin \theta \sin \phi$, $v_z = v \cos \theta$,

where θ and ϕ are generated in the same way as in Equation 25.

The comparison of particle positions generated using the aforementioned method with an example of a real-world globular cluster is shown in Figure 15.







(b) Real-world globular cluster (Messier 13). Credit: Giuseppe Donatiello

Figure 15: The particle positions generated using the described model compared to a real-world globular cluster.

8 Results

The parameters used in the simulation of a spiral galaxy are shown in Table 1. The galaxy is simulated

Parameter	Value
Halo radius	3 kpc
Halo mass	$60 \times 10^9 M_{\odot}$
Disk radius	15 kpc
Disk mass	$15 \times 10^9 M_{\odot}$
Disk thickness	$0.3~\mathrm{kpc}$
Disk density profile	Uniformly decreasing
Mass assignment scheme	TSC
Finite difference	Two-point
Time integration method	Leapfrog

Table 1: Galaxy model parameters used in the simulation.

as an isolated system, however, in deriving Equation 7, periodic boundary conditions were assumed. The simplest way (and the one used) to obtain a free-space solution from the PM method is to extend the computational domain twice in every dimension and fill the space unused in mass distribution with zeros. The total size of the potential mesh used was $128 \times 128 \times 64$ with the region of interest occupying a box of size $60 \,\mathrm{kpc} \times 60 \,\mathrm{kpc} \times 30 \,\mathrm{kpc}$ located in a $64 \times 64 \times 32$ octant of the mesh.

8.1 Particle-mesh method

In the PM method, N = 50,000 particles were used. Cell size H and time-step length were set to 60/64 = 0.9375 kpc, and 1 Myr respectively. The system's evolution over 200 Myr is shown in Figure 16.

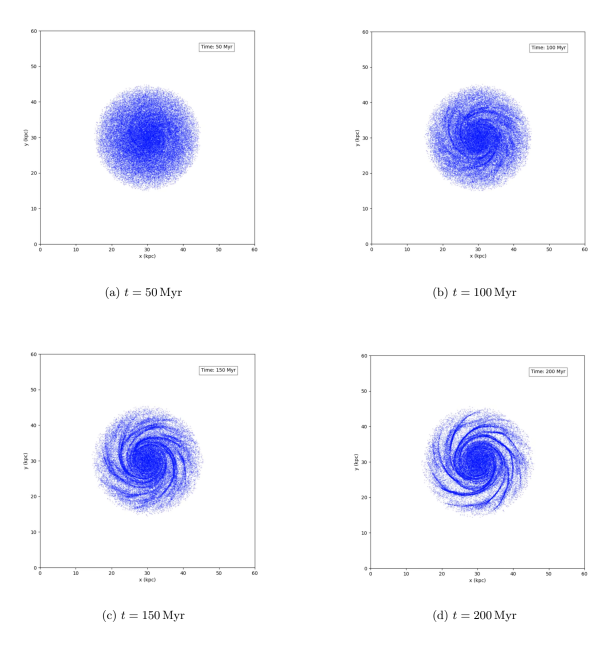


Figure 16: Evolution of a spiral galaxy as predicted by the PM method.

During the simulation, total energy E = KE + PE, angular momentum \mathbf{l} , and the z-component of the momentum vector \mathbf{p} should stay constant. The x- and y-components of momentum change due to the presence of an external gravitational field (representing the halo). We can verify if this variation satisfies the expected relation

$$\dot{\mathbf{p}} = \mathbf{F}^{\text{ext}} \tag{27}$$

by finding the initial total momentum $\mathbf{p}(t=0)$ and incrementing the value of \mathbf{p} in each time-step by

 $\mathbf{F}^{\mathrm{ext}}\mathrm{DT}.$

The exact calculation of the potential energy [8] using the formula

$$PE = -\sum_{i=1}^{N} \sum_{j=i+1}^{N} \frac{Gm_i m_j}{r_{ij}}$$

is computationally infeasible considering the $O(N^2)$ cost. An approximation based on the potential values at mesh points,

$$PE \approx \frac{V}{2} \sum_{\mathbf{p}} \rho(\mathbf{x}_{\mathbf{p}}) \phi(\mathbf{x}_{\mathbf{p}}),$$

is used instead (for derivation refer to [6]).

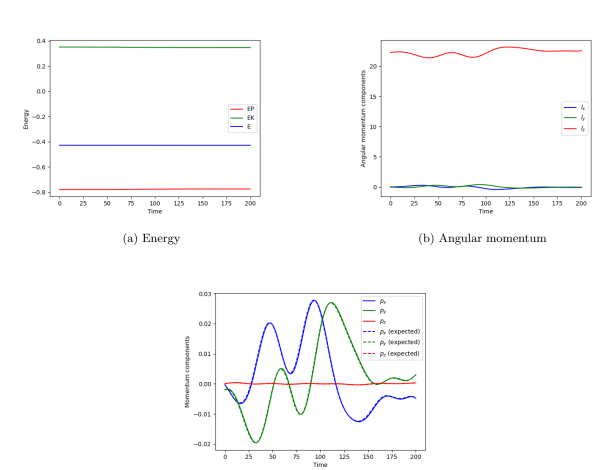


Figure 17: Fundamental physical quantities describing the system over time in the PM simulation. Time is in Myr and the quantities are expressed in units consistent with Table 1

(c) Momentum; broken lines represent the expected

momentum following Equation 27

8.2 Particle-particle particle-mesh method

The P³M based simulation uses the same parameters as the PM method. The reference force was calculated using the S_1 shape formula (Equation 14) with particle diameter a = 3H. The cutoff radius

was set to $r_e = 0.7a$. One extra free parameter is the softening length ϵ which modifies the universal law of gravitation so that division by zero can be avoided, i.e. the modified law is

$$F_{ij}^{\text{soft}}(r) = \frac{Gm_i m_j}{r_{ij}^2 + \epsilon^2}.$$

In the simulation, ϵ was set arbitrarily to 1.5 kpc. The system's evolution is presented in Figure 18. Graphs of energy, angular momentum, and momentum components vs. time are shown in Figure 19.

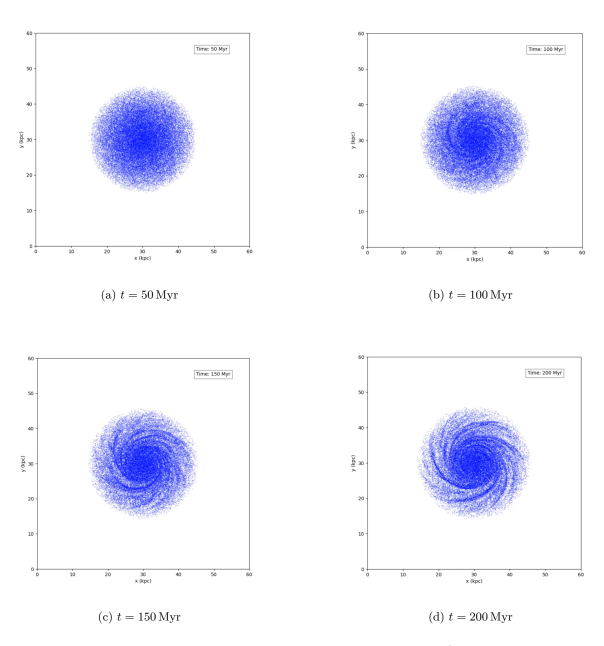


Figure 18: Evolution of a spiral galaxy as predicted by the P³M method.

8.3 Performance analysis

The PM and P³M methods were implemented exactly as described in the previous sections. The PM method was developed for both CPU and GPU architectures, using C++ and CUDA C++, respectively.

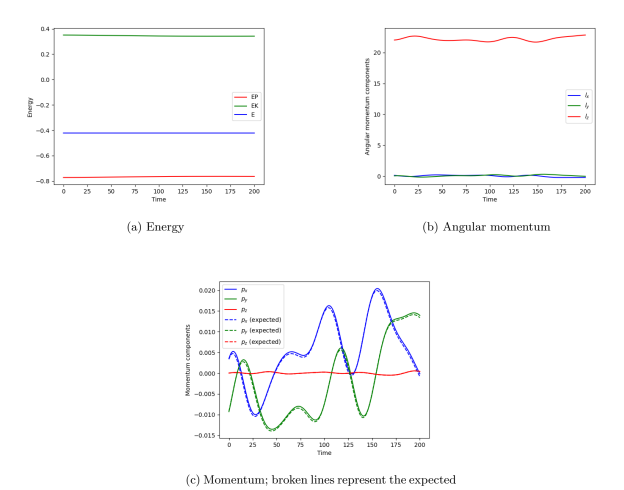


Figure 19: Fundamental physical quantities describing the system over time in the P^3M simulation. Time is in Myr and the quantities are expressed in units consistent with Table 1

momentum following Equation 27

The implementation relies on external libraries for fast Fourier transform computations: FFTW for the CPU version and cuFFT for the GPU version. A performance comparison of the PM method was conducted with N ranging between 50,000 and 1,000,000 over 200 iterations (TSC mass assignment and two-point finite difference). The tests were run on a system equipped with an Intel(R) Core(TM) i7-9750H CPU @ 2.60GHz and an NVIDIA GeForce GTX 1650 GPU. For the purposes of performance evaluation, the parts of the code responsible for diagnostics collection (energy, momentum, etc.) were switched off. Since disk IO (saving simulation state) was the most time-consuming part of both the CPU and GPU implementation, only the final state of the simulation was saved in these tests. This means that data transfers from device to host were also not taken into account. The results of the test are displayed in Figure 20. Both plots in the figure exemplify linear complexity of the PM method. The

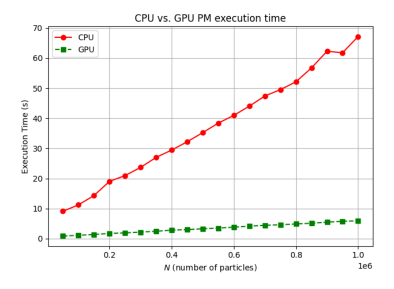


Figure 20: PM method: CPU and GPU implementation running time comparison.

GPU speedup can be estimated at around 1200%.

For the P³M method, performance was measured using N = 50,000 particles on a $128 \times 128 \times 64$ mesh with the TSC assignment scheme. The total runtime was approximately 1 minute and 30 seconds, with the time distribution among key algorithm components as follows:

• HOC table initialization: 12%

• Short-range force calculations: 80%

• PM step: 7.5%

The code is available at https://github.com/AleksyBalazinski/ParticleSimulation under the MIT license.

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