Tree-based solvers for adaptive mesh refinement code FLASH - I: gravity and optical depths.

R. Wünsch1, S. Walch2,3, F. Dinnbier1,3,5, A. Whitworth4
1 Astronomical Institute, Czech Academy of Sciences, Boční II 1401, 141 00 Prague, Czech Republic
2 Max-Planck-Institute for Astrophysics, Karl-Schwarzschild-Str. 1, 85741 Garching, Germany
3 1 Physikalisches Institut, Universität zu Köln, Zülpicher Str. 77, 50937 Köln, Germany
4 School of Physics & Astronomy, Cardiff University, The Parade, Cardiff CF24 3AA, Wales, UK
5 Faculty of Mathematics and Physics, Charles University in Prague, V Holešovičkách 2, CZ-180 00 Prague, Czech Republic

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ABSTRACT

We describe an OctTree algorithm for the MPI-parallel, adaptive mesh-refinement code FLASH, which can be used to calculate the gas self-gravity, and also the angle-averaged local optical depth, for treating ambient diffuse radiation. The algorithm communicates to the different processors only those parts of the tree that are needed to perform the tree walk locally. The advantage of this approach is a relatively low memory requirement, important in particular for the optical depth calculation, which needs to process information from many different directions. This feature also enables a general tree-based radiation transport algorithm that will be described in a subsequent paper, and delivers excellent scaling up to at least 1500 cores. Boundary conditions for gravity can be either isolated or periodic, and they can be specified in each direction independently, using a newly developed generalisation of the Ewald method. The gravity calculation can be accelerated with the adaptive block update technique by partially re-using the solution from the previous time-step. Comparison with the FLASH internal multi-grid gravity solver shows that tree based methods provide a competitive alternative, particularly for problems with isolated or mixed boundary conditions. We evaluate several multipole acceptance criteria (MACs) and identify a relatively simple APE MAC which provides high accuracy at low computational cost. The optical depth estimates are found to agree very well with those of the RADMC-3D radiation transport code, with the tree solver being much faster. Our algorithm is available in the standard release of the FLASH code in version 4.0 and later.

Key words: galaxies: ISM – gravitation – hydrodynamics – ISM: evolution – radiative transfer

1 INTRODUCTION

Solving Poisson’s equation for general mass distributions is a common problem in numerical astrophysics. Grid-based hydrodynamic codes frequently use iterative multi-grid or spectral methods for that purpose. On the other hand, particle codes often use tree-based algorithms. The extensive experience with tree gravity solvers in particle codes can be transferred to the domain of grid-based codes. Here we describe an implementation of the tree-based gravity solver for the Adaptive Mesh Refinement (AMR) code FLASH (Pryxell et al. 2000) and show that its efficiency is comparable to the FLASH intrinsic multi-grid solver (Ricker 2008). An advantage of this approach is that the tree code can be used for more general calculations performed in parallel with the gravity; in particular, calculation of the optical depth in every cell of the computational domain with the algorithm developed by Clark et al. (2012) and general radiation transport with the TreeRay algorithm (described in Paper II; Wünsch et al., in prep.). Hierarchically structured, tree-based algorithms represent a well-established technique for solving the gravitational N-body problem at reduced computational cost (Barnes & Hut 1986). Many Lagrangian codes implement trees to compute the self-gravity of both collisionless (stars or dark matter) and collisional (gas) particles, e.g. GADGET-2 (Springel 2005), VINE (Wetzstein et al. 2009, Nelson et al. 2009), EVOl. (Merlin et al. 2010), SÉREN, (Hubber et al. 2011), GANDALF (Hubber et al. 2018). The three most important characteristics of the tree algorithm are the tree structure (also called the grouping strategy), the multipole acceptance criterion (MAC) deciding whether to open a child-node or not, and the order of approximation of the integrated quantity within nodes (e.g. mass distribution).
Tree structure: Each node on the tree represents a part of the computational domain, hereafter a volume, and the child-nodes of a given parent-node collectively represent the same volume as the parent-node. The most common ‘OctTree’ structure is built by a recursive subdivision of the computational domain, where every parent-node is split into eight equal-volume child-nodes, until we reach the last generation. The nodes of the last generation are called leaf-nodes and they cover the whole computational domain.

Tree structures other than the OctTree are also often used. Bentley (1979) constructs a balanced "k-d" binary tree by recursively dividing parent-nodes so that each of the resulting child-nodes contains half (±1) of the particles in the parent-node; this tree structure is used in the codes pKdGRAV (Stadel 2001) and GASOLINE (Wadsley et al. 2004). In contrast, Press (1986) constructs a binary tree, from the bottom up, by successively amalgamating nearest neighbour particles or nodes into parent nodes. This “Press-tree” has been further improved by Jernigan & Porter (1989), and is used, for instance, by Benz et al. (1990) and Nelson et al. (2009). More complex structures have been suggested. For example, Ahn & Lee (2008) describe the "k-means" algorithm, in which a parent-node is adaptively divided into k child-nodes according to the particle distribution in the parent-node.

There seems to be no unequivocally superior tree structure. Waltz et al. (2002) compare OctTrees with binary trees, and find that OctTrees provide slightly better performance with the same accuracy. On the other hand, Anderson (1999) argues, on the basis of an analytical study, that certain types of binary trees should provide better performance than OctTrees. Makino (1990) points out that differences in performance are mainly in the tree construction part, and that the tree-walk takes a comparable amount of time in either type of tree-structure. Therefore, the choice of tree-structure should be informed by more technical issues, like the architecture of the computer to be used, other software to which the tree will be linked, and so on.

Multipole acceptance criterion: Another essential part of a tree code is the criterion, or criteria, used to decide whether a given node can be used to calculate the gravitational field, or whether its child-nodes should be considered instead. This is a key factor determining the accuracy and performance of the code. Since this criterion often reduces to deciding whether the multipole expansion representing the contribution from the node in question provides a sufficiently accurate approximation for the calculation of the gravitational potential, it is commonly referred to as the multipole acceptance criterion (MAC). We retain this terminology even though nodes in the code presented here may possess more general properties than just a multipole expansion.

The original BH86 geometric MAC uses a simple criterion, which is purely based on the ratio of the angular size of a given node and its distance to the cell at which the gravitational potential should be computed. More elaborate methods also take into account the mass distribution within a particular node or even constrain the allowed total acceleration error (Salmon & Warren 1994 SW94; see 2.2.1).

Order of approximation: Springel et al. (2001) suggest that if the gravitational acceleration is computed using multipole moments up to order p, then the maximum error is of the order of the contribution from the $\Phi^{(p+1)b}$ multipole moment. There is no consensus on where to terminate the multipole expansion of the mass distribution in a node. The original BH86 tree code uses moments up to second order ($p = 2$), i.e. quadrupoles, and many authors follow this choice. Wadsley et al. (2004) find the highest efficiency using $p = 4$ in the GASOLINE code. On the other hand, SW94 find that their code using the SumSquare MAC is most efficient with $p = 1$, i.e. just monopole moments. This suggests that the optimal choice of $p$ may depend strongly on other properties of the code and its implementation, and possibly also on the architecture of the computer. Springel (2005) advocates using just monopole moments on the basis of memory and cache usage efficiency. We follow this approach and consider only monopole moments, i.e. $p = 1$ for all implemented MACs.

Further improvements: Tree codes have often been extended with new features or modified to improve their behaviour. Barnes (1990) noted that neighbouring particles interact with essentially the same nodes, and introduced interaction lists that save time during a tree-walk. This idea was further extended by Dehnen (2000) who describes a tree with mutual node-node interactions. This greatly reduces the number of interactions that have to be calculated, leading – in theory – to an $O(N)$ CPU-time dependence on the number of particles, $N$. Dehnen’s implementation also symmetrizes the gravitational interactions to ensure accurate momentum conservation, which is in general not guaranteed with tree-codes. Recently, Potter et al. (2017) develop this so called Fast Multipole Method (FMM) further and implement it into massively parallel cosmological N-body code pKdGRAV3.

Hybrid codes: Tree-codes are also sometimes combined with other algorithms into ‘hybrid’ codes. For example, Xu (1995) describes a TreePM code which uses a tree to calculate short-range interactions, and a particle-mesh method (Hockney & Eastwood 1981) to calculate long-range interactions. The TreePM code has been developed further by Bode et al. (2000), Bagla (2002), Bode & Ostriker (2003), Bagla & Khandai (2009), Khandai & Bagla (2009). There are also general purpose tree codes available, that can work with both N-body and grid-based codes, e.g. the MPI parallel tree gravity solver FLY (Becciani et al. 2007).

In this paper we describe a newly developed, cost-efficient, tree-based solver for self-gravity and diffuse radiation that has been implemented into the AMR code FLASH. This code has been developed since 2008, and since FLASH version 4.0 it is a part of the official release. The GPU accelerated tree gravity solver, based on the early version of the presented code, has been developed by Lukat & Banerjee (2016). The paper is organized as follows: In §2 we describe the implemented algorithm, which splits up into the tree-solver (§2.1), the gravity module (§2.2) and the optical depth module (§2.3). Accuracy and performance for several static and dynamic tests are discussed in §3 and we conclude in §4. In appendix A we provide formulae for acceleration in computational domains with periodic and mixed boundary conditions, and in appendix B we give runtime parameters of the code.

2 THE ALGORITHM

The FLASH code (Fryxell et al. 2000) is a complex framework consisting of many inter-operable modules that can be combined to solve a specific problem. The tree code described here can only be used with a subset of the possible FLASH configurations. The basic requirement is usage of the PARAMESH-based grid unit (see MacNe...
tire computational domain, is called the root-block and represents the refinement level \( l = 1 \). The root-block is divided into eight equal-volume blocks having the same shape and orientation as the root-block, and these blocks represent refinement level \( l = 2 \). This process of block division is then repeated recursively until the blocks created satisfy an adaptive-mesh refinement criterion. The blocks at the bottom of the tree, which are not divided, are called leaf-blocks, and the refinement level of a leaf-block is labelled \( l_b \). In regions where the AMR criterion requires higher spatial resolution, the leaf-blocks are smaller and their refinement level, \( l_b \), is larger (i.e. they are further down the tree).

The number of grid cells in a block (a logically cuboidal collection of cells; see below) must be the same in each direction and equal to \( 2^{3l_b} \) where \( l_b \) is an arbitrary integer number. In practice, it should be \( l_b \geq 3 \), because most hydrodynamic solvers do not allow blocks containing fewer than 8 cells, in order to avoid overlapping of ghost cells. Note that the above requirements do not exclude non-cubic computational domains, because such domains can be created either by setting up blocks with different physical sizes in each direction or by using more than one root block in each direction (Walch et al. 2015).

Within each leaf-block is a local block-tree which extends the amr-tree down to the level of individual grid cells. All block-trees have the same number of levels, \( l_b \geq 3 \). The nodes on a block-tree represent refinement levels \( l_b + 1 \) (8 nodes here), \( l_b + 2 \) (\( 8^2 = 64 \) nodes here), \( l_b + 3 \) (\( 8^3 = 512 \) nodes here), and so on. The nodes at the bottom of the block-tree are leaf-nodes, and represent the grid cells on which the equations of hydrodynamics are solved.

Each node – both the nodes on the amr-tree, and the nodes on the local block-trees – stores collective information about the set of grid cells that it contains, e.g. their total mass, the position of the centre of mass, etc.

Our algorithm consists of a general tree-solver implementing the tree construction, communication and tree-walk, and modules which include the calculations of specific physical equations, e.g. gravitational accelerations or optical depths. The tree-solver communicates with the physical modules by means of interface subroutines which allow physical modules, on the one hand to store various quantities on the nodes, and on the other hand to walk the tree accessing the quantities stored on the nodes. When walking the tree, physical modules may use different MACs that reflect the nature of the quantity they are seeking to evaluate. An advantage of this approach is that it makes code maintenance more straightforward and efficient. Moreover, new functionality can be added easily by writing new physical modules or extending existing ones, without needing to change the relatively complex tree-solver algorithm.

The boundary conditions can be either isolated or periodic, and they can be specified in each direction independently, i.e. mixed boundary conditions with one or two directions periodic and the remaining one(s) isolated are allowed (see \( \S\S 2.1.2 \) and \( 2.1.3 \) respectively, we give descriptions of the gravity module and the module (called \texttt{Orca1,Dsm}) which calculates heating by the interstellar radiation field.

\footnote{If there is more than one root block, the single tree structure becomes a forest. This decreases the efficiency of the gravity solver, and therefore the number of root blocks should be kept as small as possible.}

\section{2.1 Tree-solver}

The tree-solver creates and utilises the tree data structure described above. Maintaining a copy of the whole tree on each processor would incur prohibitively large memory requirements. Therefore, only the amr-tree (i.e. the top part of the tree, between the root-block node and the leaf-block nodes) is communicated to all processors. The block-tree within a leaf-block is held on the processor whose domain contains that leaf-block, and communicated wholly or partially to another processor only if it will be needed by that processor during a subsequent tree-walk. The tree-solver itself stores in each tree-node – with the exception of the leaf-nodes – the total mass of the node and the position of its centre of mass, i.e. four floating point numbers. For leaf-nodes (the nodes corresponding to individual grid cells) only their masses are stored, because the positions of their centres of mass are identical to their geometrical centres and are already known. Additionally, each physical module can store any other required quantity on the tree-nodes.

The tree-solver consists of three steps: tree-build, communication and tree-walk. In the tree-build step, the tree is built from bottom up by collecting information from the individual grid cells, summing it, and propagating it to the parent tree-nodes. The initial stages of this step, those that involve the block-trees within individual leaf-blocks, are performed locally. However, as soon as the leaf-block nodes are reached, information has to be exchanged between processors because parent-nodes are not necessarily located on the same processor. At the end of this step, each processor possesses a copy of the amr-tree plus all the block-trees corresponding to leaf-blocks that are located on that processor.

The communication step ensures that each processor imports from all other processors all the information that it will need for the tree-walks, which are subsequently called by the physical modules. To this end, the code considers all pairs of processors, and determines what tree information the one processor (say CPU0; see Figure 1) needs to export to the other processor (say CPU1). To do this, the code walks the block-trees of all the leaf-blocks on CPU0, and applies a suite of MACs (required by the tree-solver itself and the used physical modules) in relation to all the leaf-blocks on CPU1. This suite of MACs determines for each leaf-block on CPU0, the level of its block-tree that delivers sufficient detail to CPU1 to satisfy the resolution requirements of all the physical modules that will be called before the tree is rebuilt. Thus, a leaf-block on CPU0 that has very little physical influence on any of the leaf-blocks on CPU1 (for example by virtue of being very distant or of low mass) may only need to send CPU1 the information stored on its lowest (i.e. coarsest resolution) level, \( l_{\text{low}} \). Conversely, a leaf-block on CPU0 that has a strong influence on at least one of the leaf-blocks on CPU1 (for example by virtue of being very close or very massive) may need to send the information stored on its highest (finest resolution) level, \( l_{\text{vhigh}} \). In order to simplify communication, the required nodes of each block-tree on CPU0 are then stored in a one-dimensional array, ordered by level, starting at \( \ell = l_{\text{low}} \) and proceeding to higher levels (see Figure 2). Finally, the arrays from all the block-trees on CPU0 are collated into a single message and sent to CPU1. This minimizes the number of messages sent, thereby ensuring efficient communication, even on networks with high latency.

Note that this communication strategy in which tree-nodes are communicated differs from a commonly used one in which particles (equivalents of grid cells) are communicated instead (e.g. Gadget, Springel 2005). In this way, the communication is completed before the tree-walk is executed and the tree-walk runs locally, i.e. separately on each processor. The communication strategy adopted
in this work provides a significant benefit for the OpticalDepth and the TreeRay modules as they work with a large amount of additional information per grid cell (or particle), which does not have to be stored and communicated (see §3.5).

The final step is a tree-walk, in which the whole tree is traversed in a depth-first manner for each grid cell or in general for an arbitrary target point (e.g. the position of a sink particle). During the process, the suite of MACs is evaluated recursively for each node and if it is acceptable for the calculation, subroutines of physical modules that do the calculation are called, otherwise its child-nodes are opened.

The tree-solver itself only implements a simple geometric MAC (Barnes & Hut 1986), which accepts a node if its angular size, as seen from the target point, \( r \), is smaller than a user-set limit, \( \theta_{\text{lim}} \). Specifically, if \( h \) is the linear size of the node and \( r_\ell \) is the position of the centre of mass of the node, the node is accepted (and so its child-nodes need not be considered) if

\[
\frac{h}{|r - r_\ell|} < \theta_{\text{lim}}.
\]

It has been shown by Salmon & Warren (1994) hereafter SW94 that the BH86 MAC can lead to unexpectedly large errors when the target point is relatively far from the centre of mass of the node but very close to its edge. Several alternative geometric MACs were suggested to mitigate this problem (Salmon & Warren 1994; Dubinski 1996). Following Springel (2005), we extend the geometric MAC by setting the parameter \( \eta_h \) such that a node is only accepted if the target point lies outside a cuboid \( \eta_h \) times larger than the node (with the default value \( \eta_h = 1.2 \)). Additional MACs specific to the physical modules are implemented by those modules (see §2.2).

The tree-walk is the most time consuming part of the tree-solver. Typically it takes more than 90% of the computational time spent by the whole tree-solver. We stress that the tree-walk does not include any communication; the tree is traversed in parallel independently on each processor for all the grid cells in the spatial domain of that processor. The tree-solver exhibits very good scaling, with speed-up increasing almost linearly up to at least 1500 CPU cores (see §3.5).

### 2.2 Gravity module

This module calculates the gravitational potential and/or the gravitational acceleration. We use the same approach as Springel (2005) and store only monopole moments in the tree, because this substantially reduces memory requirements and communication costs. Since masses and centres of mass are already stored on the tree-nodes by the tree-solver, the gravity module does not contribute any extra quantities to the tree.

In §2.2.1 we describe three data-dependent MACs which can be used instead of the geometric MACs of the tree-solver: Maximum Partial Error (MPE), Approximate Partial Error (APE) and mixed boundary conditions (§2.2.4). Finally in §2.2.6 we describe a technique called the Adaptive Block Update to save computational time by re-using the solution from previous time-step when possible.

![Figure 1. Determining the block-tree levels that need to be exported from the leaf-blocks in the spatial domain of processor CPU0 to processor CPU1. In this case the spatial domains of the two processors are adjacent, and are separated by the thick dotted line. For each leaf-block on CPU0 (for example, the one enclosed by a thick dashed line) its block-tree is traversed and the MAC is evaluated in relation to all the leaf-blocks on processor CPU1; for this purpose the code uses the distance from the centre of mass of a node of the block-tree on CPU0, to the closest point of a leaf-block on CPU1, as illustrated by the coloured arrows. The level of detail communicated to CPU1 is then set by the finest level reached during this procedure. In the case illustrated, the leaf-block on CPU0 that is furthest from the leaf-blocks on CPU1 (the one enclosed by a thick dashed line) exports only the first two levels of its block-tree, i.e. from level \( l_{\text{b}} \) to \( l_{\text{b}} + 1 \). In contrast, the leaf-blocks on CPU0 that are closest to the leaf-blocks on CPU1 export their full block-trees, i.e. from level \( l_{\text{b}} \) to level \( l_{\text{b}} + 3 \).](image-url)

#### 2.2.1 Data-dependent MACs

A general weakness of the purely geometric MACs is that they do not take into account the amount and internal distribution of mass in a node. This can make the code inefficient if the density is highly non-uniform. For example, if the code calculates the gravitational potential of the multi-phase interstellar medium, the contribution from nodes in the hot rarefied gas is very small, but it is calculated with the same opening angle as the much more important contribution from nodes in dense molecular cores.

MPE MAC (Maximum Partial Error): To compensate for the above problem, SW94 propose a MAC based on evaluating the maximum possible error in the contribution to the gravitational acceleration at the target point, \( r \), that could derive from calculating it using the multipole expansion of the node up to order \( p \) (instead of adding directly the contributions from all the constituent grid cells)

\[
\Delta \rho_{\text{max}}(p) = \frac{G}{d^3} \frac{1}{(1 - B_{\text{max}}/d)^2} \left( (p+2) \frac{B_{(p+1)}}{d^{p+1}} -(p+1) \frac{B_{(p+2)}}{d^{p+2}} \right),
\]

\[
B_{(p)} = \sum_i m_i |r_i - r|^p.
\]

Here, \( r \) is the mass centre of the node; \( d = |r - r_\ell| \) is the distance from \( r \) to the target point; \( B_{\text{max}} \) is the distance from \( r \) to the furthest point in the node; \( B_{(p)} \) is the \( p^\text{th} \)-order multipole moment, obtained
by summing contributions from all the grid cells \( i \) in the node: \( m_i \) and \( \mathbf{r}_i \) are the masses and positions of these grid cells. The node is then accepted only if \( \Delta a_{\text{lim}}^{(p)} \) is smaller than some specified maximum allowable acceleration error. This threshold can either be set by the user as a constant value, \( a_{\text{lim}} \), in the physical units used by the simulation

\[
\Delta a_{\text{lim}}^{(p)} < a_{\text{lim}}, \tag{4}
\]

or it can be set as a relative value, \( \epsilon_{\text{lim}} \), with respect to the acceleration from the previous time-step \( a_{\text{old}} \)

\[
\Delta a_{\text{lim}}^{(p)} < \epsilon_{\text{lim}} a_{\text{old}}. \tag{5}
\]

APE MAC (Approximate Partial Error): An alternative way to estimate the partial error of a node contribution was suggested by [Springel et al. 2001]. It takes into account the node total mass, but it ignores the internal node mass distribution. It is therefore faster, but less accurate. Using multipole moments up to order \( p \), the error of the gravitational acceleration is then of order \( \epsilon_{\text{lim}} \) the multipole moment

\[
\Delta a_{\text{lim}}^{(p)} \approx \frac{GM}{d^2} \left( \frac{h}{d} \right)^{p+1}, \tag{6}
\]

where \( M \) is the mass in the node and \( p = 1 \) in our case since we only store monopole moments. Similar to the MPE MAC, the APE error limit can be either set absolutely as \( a_{\text{lim}} \) (Equation 4), or relatively through \( \epsilon_{\text{lim}} \) (Equation 5).

**SUMSquare MAC:** SW94 argue that it is unsafe to constrain the error using the contribution of a single node only, since it is not known a priori how these contributions combine. They suggest an alternative procedure, which limits the error in the total acceleration at the target point; one variant of this procedure is the SUMSquare MAC which sums up squares of \( a_{\text{lim}}^{(p)} \) given by Equation (2) over all nodes considered for the calculation of the potential/acceleration at a given target point. In this way, the SUMSquare MAC controls the total error in acceleration resulting from the contribution of all tree-nodes. This MAC requires a special tree-walk which does not proceed in the depth-first manner. Instead it uses a priority queue, which on-the-fly reorders a list of nodes waiting for evaluation according to the estimated error resulting from their contribution. This feature is still experimental in our implementation, nevertheless we evaluate its accuracy and performance and compare it to other MACs in §3.4.

### 2.2.2 Isolated boundary conditions

In the case of isolated boundary conditions (BCs), the gravitational potential in a target point given by position vector \( \mathbf{r} \) is

\[
\Phi(\mathbf{r}) = -\sum_{a=1}^{N} \frac{GM_a}{|\mathbf{r} - \mathbf{r}_a|}, \tag{7}
\]

where index \( a \) runs over all nodes accepted by the MAC during the tree-walk, \( M_a \) and \( \mathbf{r}_a \) are the node mass and position. The gravitational acceleration is then obtained either by differentiating the potential numerically, or it is calculated, as

\[
a(\mathbf{r}) = -\sum_{a=1}^{N} \frac{GM_a (\mathbf{r} - \mathbf{r}_a)}{|\mathbf{r} - \mathbf{r}_a|^3}. \tag{8}
\]

The first approach needs less memory and is slightly faster. The second approach results in less noise, because numerical differentiation is not needed.

### 2.2.3 Periodic boundary conditions

In the case of periodic boundary conditions in all three directions, the gravitational potential is determined by the Ewald method [Ewald 1921, Kleessen 1997], which is designed to mitigate the very slow convergence in case one evaluates contributions to the potential, essentially \( 1/d \) where \( d = |\mathbf{r} - \mathbf{r}_a| \), over an infinite number of periodic copies, by brute force. This is achieved by splitting it into two parts

\[
1/d = \frac{\text{erf}(\alpha d)}{d} + \frac{\text{erf}(\alpha d)}{d} \tag{9}
\]

and summing the term \( \text{erf}(\alpha d)/d \) in Fourier space; \( \alpha \) is an arbitrary constant controlling the number of nearby and distant terms which have to be taken into consideration. In this section, we present formulae only for the potential. The expressions for acceleration are straightforward to derive, and we list them in appendix A.

The computational domain is assumed to be a rectangular cuboid, with sides \( L_x \), \( L_y = b L_x \) and \( L_z = c L_x \), where \( b \) and \( c \) are arbitrary real numbers. The gravitational potential \( \Phi \) at the target

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**Figure 2:** Organization of a block-tree within a block in memory. It is a 1D array sorted by levels, starting from \( l = l_b \).
The number of horizontal oscillations is proportional to the value of index $\Phi$. Note that the corresponding formula in Klessen (1997, ; their Equation 13) are integer numbers; $r$, and the density in the computational domain at $L_0 < z < 2L_0$ is set to zero (i.e. $n = 2$); (c) The material in the periodic copies is displaced further $(n = 4)$. The box to the left of the computational domain shows the shortest wavelength in the direction $e_x$ fulfilling condition (13). The number of horizontal oscillations is proportional to the value of index $l_1$ for given $n$. The condition (13), which is now fulfilled all mass in the computational domain to interval $(0 \leq z \leq L_0)$, (i.e. strict all mass in the computational domain to interval $(0 \leq z \leq L_0)$, and with nor-

dals pointing in direction $e_x$. As $n$ increases, the layers move away from one another, but Equation (11) still holds. In the limit $n \rightarrow \infty$, the periodic copies of the computational domain are touching one another in x- and y-directions, however, neighbouring layers in the z-direction are at infinite distance and hence they do not contribute to the gravitational field in the original computational domain.

As $n$ increases, the short-range contributions are zero for all $i \neq 0$, because the argument of the complementary error function in Equation (11) tends to infinity. The long-range term $\phi_{ll}(r - r_a)$ in the limit $n \rightarrow \infty$ becomes

$$\lim_{n \rightarrow \infty} \frac{n!}{(\pi L_c)^{n}} \frac{1}{\alpha} \frac{\pi}{L_c} \frac{\beta}{L_c} \frac{\gamma}{L_c} = \frac{2\pi l_1}{cL_0} \cos \left( \frac{2\pi l_1(x - x_0)}{L_x} + \frac{2\pi l_1(y - y_0)}{L_y} \right) \left( \cos \left( \frac{2\pi l_1(z - z_0)}{L_z} + \frac{2\pi l_1(z - z_0)}{cL_0} \right) \right).$$

The condition (13), which is now fulfilled all mass in the computational domain to interval $(0 \leq z \leq L_0)$, (i.e. $|z - z_0| \leq cL_0$) is independent of $n$, because we restrict all mass in the computational domain to interval $(0 \leq z \leq L_0)$, (i.e. $|z - z_0| \leq cL_0$) at any target point at $r$ and node at $r_a$. Bearing this in mind, the term after the limit sign in Equation (14) corresponds to a Riemann sum over interval $(-\sqrt{10}, \sqrt{10})$ with equally spaced partitions of size $1/nc$. Using the identity $\cos(A + B) = \cos(A) \cos(B) - \sin(A) \sin(B)$, where $B = 2\pi l_1(z - z_0)/(cL_0)$, the...
integral can be transformed into a 1D integral, but we have not been able to express it in a closed form. In this case (1P2I), we arrive at

\[ \Phi(r) = -G \sum_{a=1}^{N} m_a \left( \sum_{i,j} \frac{\text{erfc}(a(r-r_a - i_e L_a - j_e b L_a))}{|r-r_a - i_e L_a - j_e b L_a|} + \frac{2}{L_a} \sum_{i,j} \frac{\exp(-\zeta l_i^2)}{l_i^2} \frac{2 \pi l_i (x-x_a)}{L_a} \right) \times \]

\[ K(l_i, y-y_a, z-z_a), \]

(21)

where function \( K(l_i, y-y_a, z-z_a) \) is given by

\[ K(l_i, y-y_a, z-z_a) = \int_{0}^{\infty} J_0(\eta q) \exp(-\zeta q^2) q dq, \]

and \( \eta = 2 \pi \sqrt{(y-y_a)^2 + (z-z_a)^2}/L_a \). Function \( J_0 \) is the Bessel function of the first kind and zeroth order.

Formulae for accelerations corresponding to potentials Equation (11), Equation (20) and Equation (21) are listed in appendix A.

2.2.5 Look-up table for the Ewald array

Since the explicit evaluation of \( \phi_2(r-r_a) \) and \( \phi_3(r-r_a) \) at each time-step would be prohibitively time consuming, these functions are pre-calculated before the first hydrodynamical time step, and their values are stored in a look-up table. We experiment with two approaches to approximate the above functions from the look-up table at the time when the gravitational potential is evaluated.

In the first approach, the function \( \Phi(r-r_a) = \Phi_2(r-r_a) + \Phi_3(r-r_a) \) is pre-calculated on a set of nested grids, and particular values are then found by trilinear interpolation on these grids. Coverage of the grids increases towards the singularity at the origin \((r-r_a) \to 0\). The gravitational potential at target point \( r \) is then calculated as

\[ \Phi(r) = -\sum_{a=1}^{N} GM_a \phi(r-r_a). \]

(23)

In the second approach, we avoid the singularity of \( \phi(r-r_a) \) by subtracting the term \( 1/|r-r_a| \) from \( \phi(r-r_a) \). This enables us to use only one interpolating grid with uniform coverage for the whole computational domain. Moreover, for mixed BCs, \( \phi(r-r_a) \) can be approximated at some parts of the computational domain by analytic functions. The function \( \phi(r-r_a) \) converges to \( 2\pi|z-z_a|/(6L_a^2) \) with increasing \((z-z_a)/L_a\) for configuration 2P1I, and it converges to \( 2\pi \sqrt{(y-y_a)^2 + (z-z_a)^2}/L_a \) for configuration 2P1I and 1P2I respectively. Accordingly, we use the analytic expression in these regions and pre-calculate \( \phi(r-r_a) \) only at the region where \((z-z_a) < 2L_a \) or \( \sqrt{(y-y_a)^2 + (z-z_a)^2} > 2L_a\), so the grid covers only a fraction of the computational domain if the computational domain is elongated. In combination with using only one interpolating grid this results in smaller demands on memory while it retains the same accuracy as in the first approach.

In the second approach, we pre-calculate not only \( \phi(r-r_a) \) but also its gradient. The actual value of \( \phi(r-r_a) \) at a given location is then estimated by a Taylor expansion to the first order. This is faster than the trilinear interpolation used in the first approach, and leads to a speed up in the Gravity module by a factor of \( 1.4 \) to \( 1.9 \) depending on the shape of the computational domain, the adopted BCs, and whether the potential or acceleration is used. Thus the
second approach appears to be superior to the first one. In each approach, if gravitational accelerations rather than the potential are required, we adopt an analogous procedure for each of its Cartesian components.

Note that in a very elongated computational domain, the evaluation of $\phi(r - r_0)$ can be accelerated by adjusting the parameter $\alpha = 2/L_e$. Since $\phi(r - r_0)$ is pre-calculated, the choice of $\alpha$ is of little importance in our implementation and we do not discuss it further in this paper.

2.2.6 Adaptive block update

Often, it is not necessary to calculate the gravitational potential/acceleration at each grid cell in each time-step. Since the FLASH code uses a global time-step controlled by the Courant-Friedrichs-Lewy (CFL) condition, there may be large regions of the computational domain where the mass distribution almost does not change during one time-step. In such regions, the gravitational potential/acceleration from the previous time step may be accurate enough to be used also in the current time-step. Therefore, to save the computational time, we implement a technique called the Adaptive Block Update (ABU). If activated, the tree-walk is modified as follows. For each block, the tree-walk is at first executed only for the eight corner grid cells of the current block. Then, the gravitational potential or acceleration (or any other quantity calculated by the tree-solver, e.g. the optical depth) in those eight grid cells is compared to the values from the previous time-step. If all the differences are smaller than the required accuracy (given e.g. by Equation 3 or 5), the previous time-step values are adopted for all grid cells of the block.

For some applications, the eight test cells in the block corners may not be sufficient. For instance, if the gas changes its configuration in a spherically symmetric way within a block, the gravitational acceleration at the block corners does not change, even though the acceleration may change substantially in the block interior. Such situation is more probable if larger blocks than default 8$^3$ cells are used. Therefore, it is easily possible to add more test cells by editing array gr_bhTestCells in file gr_bhData.f90, where test cells are listed using cell indices within a block, i.e. in a form (1,1,1), (1,1,8)...(8,8,8).

ABU can save a substantial amount of the computational time, however, on large numbers of processors it works well only if a proper load balancing among processors is ensured, i.e. each processor should be assigned with a task of approximately the same computational cost. FLASH is parallelized using a domain decomposition scheme and individual blocks are distributed among processors using the space filling Morton curve (see Pryxell et al. 2000 for details). Each processor receives a number of blocks estimated so that their total expected computational time measured by a workload estimate is approximately the same as the one for the other processors. By default, FLASH assumes that processing each leaf-block takes approximately the same amount of time to compute, and it assigns workload weight 2 to each leaf-block (because it includes active grid cells) and workload weights 1 to all other blocks (they are used only for interpolations between different AMR levels).

The assumption of the same workload per leaf-block cannot be used with ABU, because if the full tree-walk is executed for a given block less often, the average computational time spent on it is substantially lower in comparison with more frequently updated blocks. It is generally hard to predict whether a given block will be fully updated in the next time-step or not without additional information about the calculated problem. Therefore, we implement a simple block workload estimate that leads in most cases to better performance than using the uniform workload, even though it may not be optimal. It is based on the assumption that the probability that the block will be updated is proportional to the amount of work done on the block during several previous time-steps. This assumption is motivated by considering that a typical simulation includes on one hand regions where the density and the acceleration change rapidly (e.g. close to fast moving dense massive objects), and on the other hand, regions where the acceleration changes slowly (e.g. large volumes filled with hot rarefied gas). Consequently, the past workload of a given block provides an approximate estimate its current workload. However, this information is valid only until the density field evolves enough to change the above property of the region. The time at which this happens can be approximately estimated as the gas crossing time of a single block. Due to the CFL condition, the corresponding number of time-steps is approximately a number of grid cells in a block along one direction. Specifically, the block workload estimate works as follows. For each leaf-block, a total number of node contributions during the tree-walk to all its grid cells, $N_{\text{int}}$, is determined. Then, the workload weight, $W_b^{(\omega)}$, of that block is calculated as

$$W_b^{(\omega)} = W_b^{(\omega-1)} \exp \left( -\frac{1}{\tau_{\text{int}}} + 1 - \exp \left( -\frac{1}{\tau_{\text{int}}} \right) \right) \left( 2 + \omega_{\text{ark}} \frac{N_{\text{int}}}{N_{\text{max}}} \right)$$

where $W_b^{(\omega-1)}$ is the workload weight from the previous time-step, $\tau_{\text{int}}$ is a characteristic number of time-steps on which the workload changes, $\omega_{\text{ark}}$ is a dimensionless number limiting the maximum workload weight, and $N_{\text{max}}$ is the maximum $N_{\text{int}}$ taken over all leaf-blocks in the simulation. In this way, the block workload weight depends on its tree-solver computational cost during the last several (~ $\tau_{\text{int}}$) time-steps and is between 2 (zero cost) and 2 + $\omega_{\text{ark}}$ (maximum cost). By default, we set two global parameters $\tau_{\text{int}} = 10$ and $\omega_{\text{ark}} = 8$. The workload weight of non-leaf blocks remains equal to 1.

2.3 OpticalDepth module

The OpticalDepth module is used to evaluate the simplified solution to the radiative transfer equation

$$I_\nu = I_{\nu,0} e^{-\tau_\nu},$$

where $I_\nu$ is the specific intensity at frequency $\nu$, $I_{\nu,0}$ is the specific intensity at the source location, and $\tau_\nu$ is the optical depth along a given path through the computational domain at frequency $\nu$. In this form, the problem of evaluating what radiation intensity reaches a given point in the computational domain, i.e. a given target point, is reduced to computing the optical depth in between a radiation source and the target point. The optical depth is proportional to the absorption cross-section and the column density along the path.

Hence, the OpticalDepth module calculates the total and/or specific column densities (e.g. of molecular hydrogen) for each cell in the computational domain, and can therefore be used to compute the local attenuation of an arbitrary external radiation field. The implementation presented here follows the idea of the Treecol method (Clark et al. 2012), which has been implemented in the Gadget code (Springel et al. 2001). It has been established as a fast but accurate enough approximative radiative transfer scheme to treat the (self)-shielding of molecules –on-the-fly – in simulations of molecular cloud formation (e.g. Clark & Glover 2014).

Recently, the method has also been applied in larger-scale simulations of Milky-Way like galaxies (Smith et al. 2014) with the Arepo
For each grid cell, the module constructs a Hsion to the accumulation of the node masses during the tree-walk. In principle, the OpticalDepth module adds another dimension to the accumulation of the node masses during the tree-walk. For each grid cell, the module constructs a Healpix sphere (Gorski et al. 2005) with a given number of pixels, \( N_{\text{pix}} \), each representing a sphere surface element with index \( i_{\text{pix}} \) corresponding to polar and azimuth angles \( \theta \) and \( \phi \), respectively. This temporary map is filled while walking the tree, as only the tree-nodes in the line of sight of a given pixel contribute to it, and are added accordingly. At the end of the tree-walk, one has acquired a column density map of a given quantity, e.g. total mass.

Since the tree-walk in FLASH is executed on a block-by-block basis, the additional memory requirement for the local pixel maps is \( 2^{n_1} \times N_{\text{pix}} \times L_{\text{pix}} \), where \( L_{\text{pix}} \) is the number of quantities that are mapped and stored. For this paper, we map \( L_{\text{pix}} = 3 \) variables: (1) the total mass giving the total hydrogen column density, \( N_H\text{pix} \); (2) the H2 column of molecular hydrogen, which is used to compute its self-shielding and which contributes to the shielding of CO; and (3) the CO column of carbon-monoxide, which is necessary to compute the self-shielding of CO. We store three separate maps because we actually follow the relative mass fractions of multiple species in the simulation using the FLASH MultiSpecies module. After the tree-walk for a given block has finished, the local maps are erased and the arrays can be re-used for the next block. This approach is only possible because the tree-walk is computed locally on each processor (see §2.1).

When using the OpticalDepth module, there are two major modifications with respect to the usual tree-walk (as described above). First, the intersection of a given tree-node with the line of sight of each pixel has to be evaluated during the tree-walk. Second, at the end of the tree-walk for a given block, the acquired column density maps have to be evaluated for each cell.

Node-Ray Intersection: The mapping of tree-nodes onto the individual pixels represents the core of all additional numerical operations that have to be carried out when running OpticalDepth in addition to the gravity calculation. It has to be computationally efficient in order to minimise additional costs. At this point, we do not follow the implementation of Clark et al. (2012), who make a number of assumptions about the shape of the nodes and their projection onto the pixels, which are necessary to reduce the computational cost. Instead, we pre-compute the number of intersecting Healpix rays and their respective, relative weight for a large set of nodes at different angular positions (\( \theta, \phi \)) and different angular sizes \( \phi \). These values are stored in a look-up table, which is accessed during the tree-walk. In this way, the mapping of the nodes is highly efficient. Since \( \theta, \phi, \) and \( \psi \) are known, we can easily compute the contribution of a node to all intersecting pixels by simply multiplying the mass (or any other quantity that should be mapped) of the node with the corresponding weight for each pixel and adding this contribution to the pixel map. For better accuracy, we over-sample the Healpix tessellation and construct the table for four times more rays than actually used in the simulation.

Radiative Heating and Molecule Formation: The information that is obtained by the OpticalDepth module is necessary to compute the local heating rates and the formation and dissociation rates of H2 and CO. At the end of the tree-walk for a given block, the mean physical quantities needed by the Chemistry module calculating the interaction of the radiation with the gas are determined.

\[
A_v = \frac{1}{2.5} \ln \left( \frac{1}{N_{\text{pix}}} \sum_{i_{\text{pix}}} \exp \left( -2.5 \frac{N_{H\text{pix}}}{1.87 \times 10^{21} \text{ cm}^{-2}} \right) \right)
\]

where the constant \( 1.87 \times 10^{21} \text{ cm}^{-2} \) comes from the standard relation between the hydrogen column density, \( N_H\text{pix} \), and the visual extinction in a given direction (Draine & Bertoldi 1996). The weighted mean is calculated in this fashion, because the photodissociation rates of molecules such as CO and the photoelectric heating rate of the gas all depend on exponential functions of the visual extinction (see Clark et al. 2012 for details). Additionally, the shielding coefficients, \( f_{\text{shield,H}_2} \) and \( f_{\text{shield,CO}} \) (Glover & Mac Low 2007; Glover et al. 2010), as well as the dust attenuation, \( I_{\text{att}} \) (Glover & Clark 2012; Clark et al. 2012), are computed by averaging over the Healpix maps in a similar way. These quantities are stored as globally accessible variables and can be used by other modules. In particular, we access them in the Chemistry module, which locally (in every cell) evaluates a small chemical network (Glover et al. 2010) on the basis of its current density and internal energy and re-computes the relative mass fractions of the different chemical species. The evaluation of the chemical network is operator split and employs the Dvořák solver (Brown et al. 1989) to solve a system of coupled ODEs that describes the chemically reactive flow for the given species, i.e. their creation and destruction within a given time step. Here, we explicitly follow the evolution of five species, i.e. the different forms of hydrogen (ionised, H+, atomic, H, and molecular, H2) as well as ionised carbon (C+) and carbon-monoxide (CO). Details about the chemical network, e.g. the considered reactions and the employed rate coefficients in the current implementation can be found in Glover et al. (2010) and Walch et al. (2015).

Parameters: The main parameters controlling both the accuracy and the speed of the calculation are the number of pixels per map \( N_{\text{pix}} \), and the opening angle, \( \theta_{\text{lim}} \), with which the tree is walked (see Equation (1)). Both should be varied at the same time. A high number of \( N_{\text{pix}} \) used with a relatively large opening angle will not improve the directional information since the nodes that are mapped into each solid angle will not be opened and thus, a spatial resolution that is sufficient for a fine-grained map cannot not be achieved. Therefore we vary both \( N_{\text{pix}} \) and \( \theta_{\text{lim}} \) at the same time.

The number of Healpix pixels is directly related to the solid angle of each element on the unit sphere

\[
\Omega_{\text{pix}} = \frac{4\pi}{N_{\text{pix}}} \text{sr}.
\]

Tests in §3.3.1 show, in agreement with Clark et al. (2012), that the code efficiency is optimal if \( \theta_{\text{lim}} \) is approximately the same as the angular size Healpix elements, i.e.

\[
\theta_{\text{lim}} = \sqrt{\Omega_{\text{pix}}}. \tag{28}
\]

Therefore, for \( N_{\text{pix}} = 12, 48, 192 \) pixels we recommend to use \( \theta_{\text{lim}} \approx 1.0, 0.5, 0.25 \).

3 ACCURACY AND PERFORMANCE

Since more computational time is needed to reach higher accuracy when solving numerical problems, accuracy and performance are connected and therefore, these two properties should always be evaluated at the same time. However, they are often highly dependent on the specific type of the problem and finding a test that allows one to objectively measure both accuracy and performance
is hard. Another complication is that the tree-solver saves time by using the information from the previous time-step (if ABU is switched on), and thus any realistic estimate of the performance must be measured by running a simulation in which the mass moves in a similar way as in real applications and by integrating the computational time over a number of time-steps. Unfortunately, such simulations are unavoidably too complex to have an analytic solution against which the accuracy could be easily evaluated.

Therefore we perform two types of tests: static tests that measure accuracy using simple problems and dynamic tests that evaluate accuracy and performance together. The static tests need substantially less CPU time and thus allow for a higher number of parameter sets to be tested. Furthermore, analytic or semi-analytic solutions are known and the results can be compared to them. On the other hand, the dynamic tests represent more complex simulations which are more similar to problems that one would actually want to solve with the presented code. They also show how well the tree-solver is coupled with the hydrodynamic evolution (where we use the standard PPM Riemann solver of the Flash code) and how the error accumulates during the evolution. In this section, we describe four static and two dynamic tests of the Gravity module and one test of the OpticalDepth module.

When possible, i.e. for fully periodic of fully isolated boundary conditions, we compare the results obtained with the new tree-solver to the results obtained with the default multi-grid Poisson solver of FLASH (Ricker 2008). The multi-grid solver is an iterative solver and the accuracy is controlled by checking the convergence of the L2 norm of the Poisson equation residual $R(\mathbf{r}) \equiv 4\pi G \rho(\mathbf{r}) - \nabla \Phi(\mathbf{r})$. The iteration process is stopped when $\|R_n\|/\|R_1\| \leq \epsilon_{\text{res},\text{lim}}$, where $\|R_n\|$ is the residual norm in the n-th iteration and $\epsilon_{\text{res},\text{lim}}$ is the limit set by user. If isolated boundary conditions are used, the gravitational potential at the boundary is calculated by a multipole Poisson solver expanding the density and potential field into a series up to a multipole of order $m_{\text{mp}}$. By default $m_{\text{mp}} = 0$ in FLASH version 4.4. However, using this value we found unexpectedly high errors close the boundaries (see test §3.1.1 and Figures 4 and 5), and therefore we use $m_{\text{mp}} = 15$ (the highest value allowed for technical reasons) in most tests because it yields the smallest error.

In general, the calculated gravitational acceleration deviates from the exact analytical solution due to two effects. The first one is the inherent inaccuracy of the gravity solver (either the tree gravity solver or the multi-grid solver), the second one is caused by an imperfect discretisation of the density field on the grid. Since we are mainly interested in evaluating the first effect, we measure the error by comparing the calculated accelerations to the reference solution obtained by direct “$N^2$” summation of all interactions of each grid cell with all the other grid cells in the computational domain. We additionally give the difference between the analytical and the “$N^2$-integrated” acceleration when possible.

We define the relative error $e_a$ of the gravitational acceleration $\mathbf{a}$ at the point $\mathbf{r}$ as

$$e_a(\mathbf{r}) = \frac{|\mathbf{a}(\mathbf{r}) - \mathbf{a}_{\text{ref}}(\mathbf{r})|}{a_{\text{ref,max}}},$$

where $\mathbf{a}_{\text{ref}}$ is the acceleration of the reference solution and $a_{\text{ref,max}}$ is its maximum taken over the whole computational domain.

In most of the gravity module tests, we control the error by setting the absolute limit $a_{\text{lim}}$ on the acceleration, which is calculated from the initial maximum acceleration in the computational domain, $a_{\text{max}}$, as $a_{\text{lim}} = a_{\text{max}} \times a_{\text{max}}$; typically, $a_{\text{lim}} = 10^{-2}$ or $10^{-3}$.

Table 1. Results of the marginally stable Bonnor-Ebert sphere test.

| mod. | solver | quan. | MAC | $e_{\text{lim}}$ | $\theta_{\text{lim}}$ | $m_{\text{mp}}$ | $e_{\text{max}}$ | $t_{\text{grav}}$ |
|------|--------|-------|-----|-----------------|---------------------|-------------|-----------------|-----------------|
| (a)  | tree   | accel. | APE | $10^{-3}$       | -                   | -           | 0.0009          | 83              |
| (b)  | tree   | accel. | APE | $10^{-2}$       | -                   | -           | 0.0057          | 35              |
| (c)  | tree   | accel. | BH  | -               | 0.5                 | -           | 0.0008          | 110             |
| (d)  | tree   | pot.  | APE | $10^{-3}$       | -                   | -           | 0.0085          | 80              |
| (e)  | tree   | pot.  | APE | $10^{-2}$       | -                   | -           | 0.031           | 38              |
| (f)  | tree   | pot.  | BH  | -               | 0.5                 | -           | 0.0095          | 106             |
| (g)  | mg     | pot.  | -   | -               | -                   | 0           | 0.058           | 21              |
| (h)  | mg     | pot.  | -   | -               | -                   | 15          | 0.077           | 20              |

We give the model name in column 1. The following columns are:

- solver: indicates whether the tree-solver or the multi-grid solver (mg) is used
- quan.: quantity calculated by the gravity solver (acceleration or potential which is then differentiated)
- MAC: Multipole Acceptance Criterion (Barnes-Hut or Approximate Partial Error)
- $e_{\text{lim}}$: requested accuracy of the solver as given by Equation (4)
- $\theta_{\text{lim}}$: maximum opening angle when the Barnes-Hut MAC is used
- $e_{\text{max}}$: maximum relative error in the computational domain given by Equation (5)
- $t_{\text{grav}}$: time (in seconds) to calculate a single time-step on 8 cores

The difference between using the absolute or the relative error control is discussed in §3.3.

Most of the tests were carried out on cluster Salomon of the Czech National Supercomputing Centre IT4I. A few static tests that do not need larger computational power have been run on a workstation equipped with a 4-core Intel Core i7-2600 processor.

3.1 Static tests of gravity module

In order to test all combinations of the boundary conditions implemented in the Gravity module, we present four static tests. A marginally stable Bonnor-Ebert sphere is used to test the code with isolated boundary conditions (see §3.1.1) and a density field perturbed by a sine wave not aligned with any coordinate axis is used to test setups with fully periodic boundary conditions (§3.1.2). For mixed boundary conditions, periodic in two directions and isolated in a third one, or periodic in a single direction and isolated in the remaining two, we use an isothermal layer in hydrostatic equilibrium (§3.1.3) and an isothermal cylinder in hydrostatic equilibrium, respectively (§3.1.4). Finally, in §3.1.5 we test how the code accuracy depends on the alignment or non-alignment of the gas structures with the grid axes using a set of parallel cylinders lying in the xy-plane inclined at various angles with respect to the x-axis.

3.1.1 Bonnor-Ebert sphere

We calculate the radial gravitational acceleration of a marginally stable Bonnor-Ebert sphere (Ebert 1955; Bonnor 1956) with
Figure 4. Error in the gravitational acceleration for the Bonnor-Ebert sphere as a function of radius. At a given radius, \( r \), the error \( e_{a,r} \) is calculated as a maximum over all angular directions \( \phi \) and \( \theta \). The vertical black line shows the BE sphere edge. The solid black line shows the difference between the acceleration obtained analytically and the reference solution calculated using the \( N^2 \) summation. Left panel: shows tests where the acceleration was calculated directly using Equation (8), the green, blue and red lines show errors of runs (a), (b) and (c), respectively, with parameters given in Table 1. Right panel: displays tests where the tree-solver calculates the gravitational potential using Equation (7) and the acceleration is obtained by numerical differentiation. The green, blue and red line denote models (d), (e) and (f). The magenta lines show tests calculated with the multi-grid solver using Equation (6). Figure 5. Error in the gravitational acceleration, \( e_{a,r} \), displayed in the \( z = 0 \) plane for the Bonnor-Ebert sphere test. The four panels show four selected runs with parameters given in Table 1. Top left corresponds to model (b) using the tree-solver calculating the gravitational acceleration; top right shows model (e) where the tree-solver calculated the potential; bottom left is model (g) calculated using the multi-grid solver with \( m_{\text{up}} = 0 \); and bottom right is model (h) calculated using the multi-grid solver with \( m_{\text{up}} = 15 \). The grid geometry (boundaries of \( 8^3 \) blocks) is shown in the top right panel.

Figure 5 shows the relative error in the gravitational acceleration, \( e_{a,r} \), as a function of radial coordinate, \( r \), and Table 1 lists all models, their maximum relative error, \( e_{a,\text{max}} \), and the time to calculate one time step, \( t_{\text{cpu}} \). We compare the solutions calculated with the tree gravity solver using the geometric (BH) MAC with \( \theta_{\text{lim}} = 0.5 \) (red curves) to the ones calculated using the APE MAC with \( \epsilon_{\text{lim}} = 10^{-2} \) (green lines) and \( \epsilon_{\text{lim}} = 10^{-3} \) (blue lines), respectively. The APE MAC and \( \epsilon_{\text{lim}} = 10^{-3} \) as well as the geometric MAC with \( \theta_{\text{lim}} = 0.5 \) always give a maximum relative error which is smaller than 0.1%. In case of the APE MAC and \( \epsilon_{\text{lim}} = 10^{-2} \), the maximum relative error reaches \( < 1\% \). Note that the error due to the discretisation of the density field is also of the order of 1% (black line; the jumps are due to changes in the refinement level in the AMR grid).

With the tree gravity solver, the user may choose to directly compute the gravitational accelerations (left panel of Figure 5) or to calculate them by numerical differentiation of the gravitational potential (right panel of Figure 5). Usually, the latter is the standard practice in grid-based 3D simulations, also because only one field variable, the potential, has to be stored instead of three, the accelerations in three spatial directions. However, for the tree-solver we generally find that the error in the gravitational accelerations is significantly smaller (about a factor of 10 in the test presented here) if they are computed directly. This is independent of the used MAC.

For comparison, we also show the results obtained with the multi-grid solver (magenta lines) using \( \epsilon_{\text{lim,mg}} = 10^{-4} \) and \( m_{\text{up}} = 0 \) (solid lines) or \( m_{\text{up}} = 15 \) (dotted lines), respectively. Although the mass distribution is spherically symmetric, the order of the multipole expansion of the boundary condition affects the accuracy of the multi-grid solver relatively far away from boundaries, even inside the BES. The error of the multi-grid solver is very low in the central region, it reaches \( < 1\% \) in regions where the refinement level changes (due to numerical differentiation of the potential), and increases to relatively high values at the border of the computational domain (\( \sim 1\% \) for \( m_{\text{up}} = 15 \) and \( \sim 5\% \) for \( m_{\text{up}} = 0 \)).
to inaccuracy of the boundary conditions calculated by the multi-pole solver. We note that a direct calculation of the gravitational acceleration is not possible with the multi-grid solver.

The distribution of the relative error $e_{r}$ in the $z = 0$ plane through the centre of the BES is depicted in Figure 5. The results show that the acceleration obtained with the tree gravity solver using the APE MAC with $\epsilon_{\mathrm{lim}} = 10^{-2}$ has a substantially smaller error if it is calculated directly (top left panel; see Table 1 model (b)) instead of by numerical differentiation of the potential (top right panel; model (c)). The bottom panels show the results for the multi-grid solver with $m_{\mathrm{sup}} = 0$ (model (d)) and $m_{\mathrm{sup}} = 15$ (model (e)), respectively. The default setting of $m_{\mathrm{sup}} = 0$ gives errors of $\sim 5\%$ near the domain boundaries due to the low accuracy of the multi-pole solver. This error propagates into a large fraction of the computational domain.

### 3.1.2 Sine-wave perturbation (Jeans test)

In a computational domain with purely boundary conditions we calculate the gravitational acceleration of a smooth density field with a harmonic perturbation,

$$\rho(r) = \rho_{0} + \rho_{1} \cos(k \cdot r),$$

(30)

where $\rho_{0} = 1.66 \times 10^{-24}$ is the mean density and $\rho_{1} = 0.99\rho_{0}$ is the amplitude of the perturbation. The computational domain is a cube of size 500 pc with 128 grid cells in each direction. The wave-vector $k = 6\pi(3,2,1)/L$ was chosen such that it is not aligned with any of the coordinate axes. The gravitational acceleration can be obtained analytically with the help of the Jeans swindle (Jeans 1902, Kiessling 1999)

$$g(r) = -4\pi G \rho_{0} \frac{k}{k^{2}} \sin(k \cdot r).$$

(31)

Figure 6 shows the maximum relative error $e_{r,d}$ as a function of the position $x_{l}$ on a line parallel to the perturbation wave-vector $k$. The maximum error is computed from all points projected to a given position on the line. It can be seen that the error of the multi-grid solver (magenta curve) is very small, almost the same as the difference between the analytical solution and the reference solution (black line). This is because without the need to calculate the boundary conditions separately, and on a uniform grid, the FFT accelerated multi-grid method is extremely efficient. Again, the results for the tree-solver simulations show that direct calculation of the acceleration (solid curves) leads to a much lower error than the calculation of the potential and subsequent differentiation (dashed lines). In particular, the calculation of the potential with the geometric MAC that does not take into account the different mass density in the tree-nodes leads to a relative error greater than $10\%$. However, a direct calculation of the acceleration gives very accurate results for both, the geometric MAC and the APE MAC with $\epsilon_{\mathrm{lim}} = 10^{-3}$. In Table 2 we list all models with their respective $\epsilon_{r,\max}$ and $t_{\mathrm{gyr}}$.

### 3.1.3 Isothermal layer in hydrostatic equilibrium

In order to test the accuracy of the tree gravity module with mixed boundary conditions (periodic in two directions and isolated in the third one), we calculate the gravitational acceleration of an isothermal layer in hydrostatic equilibrium. The vertical density distribution of the layer is (Spitzer 1942)

$$\rho(z) = \rho_{0} \text{sech}^{2} \left( \frac{2\pi G \rho_{0}}{c_{s}^{2}} z \right),$$

(32)

where $\rho_{0} = 1.6 \times 10^{-24} \text{ g cm}^{-3}$ is the mid-plane density and $c_{s} = 117.1 \text{ km s}^{-1}$ is the isothermal sound speed. The corresponding vertical component of the gravitational acceleration is

$$g_{z}(z) = 2\sqrt{2\pi G \rho_{0} c_{s}^{2}} \tanh \left( \frac{2\pi G \rho_{0}}{c_{s}^{2}} z \right).$$

(33)


### Table 3. Results of the second static test: isothermal layer in hydrostatic equilibrium. The meaning of columns is the same as in Table 4.

| model | solver | quan. | MAC | $\varepsilon_{\text{lim}}$ | $\theta_{\text{lim}}$ | $\epsilon_{\text{a,max}}$ | $t_{\text{ref}}$ |
|-------|--------|-------|-----|-----------------|----------------|------------------|-------------|
| (a)   | tree   | acc.  | APE | $10^{-3}$       | -               | 0.00017          | 170          |
| (b)   | tree   | acc.  | APE | $10^{-2}$       | -               | 0.0035           | 106          |
| (c)   | tree   | acc.  | BH  | 0.5             | 9.0 x $10^{-5}$ | 180              |              |
| (d)   | tree   | pot.  | APE | $10^{-3}$       | -               | 0.00029          | 99           |
| (e)   | tree   | pot.  | APE | $10^{-2}$       | -               | 0.0028           | 45           |
| (f)   | tree   | pot.  | BH  | 0.5             | 0.00043         | 107              |              |

### Table 4. Results of the fourth static test: isothermal cylinder in hydrostatic equilibrium. The meaning of columns is the same as in Table 4.

| model | solver | quan. | MAC | $\varepsilon_{\text{lim}}$ | $\theta_{\text{lim}}$ | $\epsilon_{\text{a,max}}$ | $t_{\text{ref}}$ |
|-------|--------|-------|-----|-----------------|----------------|------------------|-------------|
| (a)   | tree   | acc.  | APE | $10^{-3}$       | -               | 0.00082          | 270          |
| (b)   | tree   | acc.  | APE | $10^{-2}$       | -               | 0.0053           | 110          |
| (c)   | tree   | acc.  | BH  | 0.5             | 0.00111         | 280              |              |
| (d)   | tree   | pot.  | APE | $10^{-3}$       | -               | 0.0095           | 180          |
| (e)   | tree   | pot.  | APE | $10^{-2}$       | -               | 0.015            | 73           |
| (f)   | tree   | pot.  | BH  | 0.5             | 0.010           | 175              |              |

(Ostriker 1964)

\[
\rho(R) = \rho_0 \left(1 + \frac{\pi G \rho_0 R^2}{2 c_s^3}\right)^{-2}
\]

where $\rho_0 = 3.69 \times 10^{-23}$ g cm\(^{-3}\) is the central density and $c_s = 0.2$ km s\(^{-1}\) is the isothermal sound speed. The density distribution is cut off at radius $R_{\text{cy}} = 1.62$ pc and embedded in an ambient gas with $c_{\text{amb}} = 10$ km s\(^{-1}\) and the same pressure as the pressure at the cylinder boundary. The corresponding gravitational acceleration is

\[
g(R) = 2\pi G \rho_0 R \left(1 + \frac{\pi G \rho_0 R^2}{2 c_s^3}\right)^{-1}
\]

The computational domain has dimensions $3.6$ pc $\times$ $1.8$ pc $\times$ $1.8$ pc and contains $256 \times 128 \times 128$ grid cells.

Figure 8 shows the relative error of the gravitational acceleration for the isothermal cylinder. Meaning of line types is the same as in Figure 4. The black vertical line denotes the edge of the cylinder.

The computational domain is a cube of side length $L = 1000$ pc and a uniform resolution of 128 grid cells in each direction.

Figure 7 shows the maximum relative error $e_{\text{rel}}$ in the acceleration as a function of the $z$-coordinate, where the maximum is taken over all cells with the same $z$-coordinate. It can be seen that the error is almost independent of $z$ and there is only a small difference between the cases where the gravitational acceleration is calculated directly (solid lines) or where it is obtained by differentiation of the potential (dashed lines). The reason is that the density field in this test has relatively shallow gradients (e.g. compared to the Jeans test discussed in the previous section) and numerical differentiation leads to particularly severe errors for steep gradients. We find the largest error for runs with APE MAC and $\varepsilon_{\text{lim}} = 10^{-2}$. All other runs have small errors, which are comparable to the difference between the analytical and the reference solution, resulting from the discretisation of the density field. The results are summarised in Table 3.

#### 3.1.4 Isothermal cylinder in hydrostatic equilibrium

In the next static test, we evaluate the accuracy of the tree gravity module for mixed boundary conditions, which are isolated in two directions and periodic in the third one. We calculate the gravitational acceleration of an isothermal cylinder in hydrostatic equilibrium. The long axis of the cylinder is parallel to $x$-coordinate and the radius is given as $R = \sqrt{\frac{x^2 + y^2}{\pi^2}}$. The density distribution is...
3.2 Dynamic tests of gravity module

We run two dynamic tests of the gravity module. The first one (described in §3.2.1) is a collapse of a cold adiabatic sphere suggested by Evrard (1988) and it tests how well the energy is conserved during the gravitational collapse. The second one, describes the evolution of a turbulent sphere (§3.2.2). Both test the accuracy of the gravity module and its coupling to the hydrodynamic solver.

3.2.1 Evrard test

The Evrard test (Evrard 1988) describes the gravitational collapse and a subsequent re-bounce of an adiabatic, initially cold sphere. It is often used to verify energy conservation in SPH codes (e.g. Springel et al. 2001; Wetzstein et al. 2009). Its application on grid-based codes is unfortunately less common. The initial conditions consist of a gaseous sphere of mass $M$, radius $R$ and density profile

$$\rho(r) = \frac{M}{2\pi R^3 r}.$$  \hfill (36)
The initial, spatially constant temperature is set so that the internal energy per unit mass is

$$u = 0.05 \frac{GM}{R},$$

where $G$ is the gravitational constant. The standard values of the above parameters, used also in this work, are $M = R = G = 1$.

In Figure 10 we show the time evolution of the total mass as well as the gravitational, kinetic, internal, and total energy. On the top panel, we compare the results obtained with the tree gravity solver and the multi-grid solver, both computed on a uniform grid of size $128^3$ corresponding to a constant refinement level equal to 5. The tree-solver run uses the Barnes-Hut MAC with $\theta_{\text{lim}} = 0.5$, the multi-grid run was calculated with the default accuracy $\epsilon_{\text{rel, lim}} = 10^{-6}$ and $m_{\text{ref}} = 0$. The two runs are practically indistinguishable, however, the total energy (that should stay constant) rises by approximately 0.1 during the period of maximum compression. Since the distribution of the error in the gravitational acceleration calculated by the two solvers is very different, the same results indicate that the error in the energy conservation is not caused by the calculation of the gravitational acceleration and that the acceleration errors are below the sensitivity of this test.

The bottom panel of Figure 10 compares runs calculated with the tree-solver at different resolutions. It includes three runs with uniform grids of sizes $64^3$, $128^3$ and $256^3$ (corresponding to constant refinement levels of 4, 5, and 6) and three runs calculated on adaptive grids, which are refined such that the Jeans length is resolved by at least 2, 4, and 8 grid cells, respectively.

We find that low resolution leads to a higher numerical dissipation and artificial heating of the gas. Furthermore, lower resolution does not allow high compression of the sphere centre leading to less pronounced peaks of the internal and gravitational energies. Consequently, the results of this test show that high resolution is needed only in the centre of the sphere where the highest density is reached.

The turbulent sphere represents a proto-typical star formation test. We set up a turbulent, isothermal sphere with a total mass of $10^3 \, M_\odot$, radius 3 pc, and temperature 10 K. The initial density profile is Gaussian with a central density of $\rho_0 = 1.1 \times 10^{-21} \, \text{g cm}^{-3}$ and the density at the edge is $\rho_0 / 3$. It is embedded in a cubic box with side length $L = 10 \, \text{pc}$, which is filled with a rarefied ambient medium of density $\rho_{\text{amb}} = 10^{-23} \, \text{g cm}^{-3}$ and temperature 100 K. We add an initial turbulent velocity field to the sphere with a Kolmogorov spectrum on all modes with wave numbers between $k_{\text{min}} = 2 \times (2\pi / L)$ and $k_{\text{max}} = 32 \times (2\pi / L)$. The magnitude of velocity perturbations is scaled so that the total kinetic energy is 0.7 times the absolute value of the total potential energy.

The sphere is evolved under the influence of self-gravity and hydrodynamics, and since it is gravitationally bound it collapses towards the center and forms stars. We use isolated gravity boundary conditions, while the hydrodynamic boundary conditions are set to "outflow". The spatial resolution on the base grid is $128^3$ (refine-
To model the star formation process, we introduce sink particles according to a Jeans criterion if the gas density is above a constant distance from massive concentrations of gas, resulting from the gravitational acceleration, $e_{a_{\text{max}}}$ (y-axis). The error is determined at $t = 2$ Myr and the maximum is taken over the whole computational domain. The thin dashed lines are iso-lines of constant $I_{\text{grav}} \times e_{a_{\text{max}}}$ assessing the code efficiency. Parameters of the displayed runs are given in Table 5.

The top two panels of Figure 12 show calculations with the tree-solver and the multi-grid solver. The results are very similar to the one in run (d), with a maximum relative error of approximately 4.5 % ($\sim$ 1.5 larger) and the mean time per gravity time-step is 7 seconds (slightly smaller). Panel (12g) shows the run with the same tree-solver parameters, but instead of calculating the acceleration directly, the tree-solver calculates the potential and differentiates it numerically. The relative error exhibits a similar pattern to run (a), however, instead of sudden changes it includes high peaks of the error resulting from a numerical differentiation. Even though the mean error is comparable to runs (b) and (c), the maximum error is much higher, reaching 80 %. The time of the gravity calculation is slightly higher than in run (e), even though calculating the potential is cheaper than the acceleration for a single target cell. It is because a higher number of blocks must be updated in each time-step due to the higher error.

Panel (12h) shows the run with the same tree-solver parameters, but with a reduced accuracy of $e_{\text{lim}} = 10^{-1}$ made to test the limits of the tree-solver usability. The relative error is high, in particular in the outer regions where the blocks are updated less often, reaching a maximum of 20 %, however, it is still a factor of 2 smaller than the error at the boundaries of the computational domain found in the run with the multi-grid solver (see below). On the other hand, the calculation is very fast with a mean time per gravity time-step of 3.5 seconds, which is $\sim 70$ % of the time needed by the hydro solver.

The right panel (12b) shows the same calculation, but the ABU was switched on in this case. The relative error exhibits a rectangular pattern, because some blocks, in particular in the outer regions, were not updated in a given time-step and the error in them is larger. The maximum error is approximately 3 %, i.e. 1.5 times more than in the run with ABU off, and at the same time, the ABU makes the calculation approximately two times faster.

In Figure 14 we show the evolving column density in the xy-plane at times 0.2, 2.0, 4.0, and 6.0 Myr. Although the simulation is interesting in itself, we only focus on the error in the resulting gravitational acceleration. Therefore, we compute the same initial conditions six times with different gravity settings and measure the resulting error of the gravitational acceleration, where the supposedly accurate result compared to which we calculate the error is obtained using $N^2$ integration. The results of our analysis are shown in Figure 13 which depicts the error in the $xy$-plane at $t = 2$ Myr. The maximum and average errors $e_{a_{\text{max}}}$ and $e_{a_{\text{avg}}}$, respectively, and mean times per gravity and hydrodynamic time-step computations $t_{\text{grav}}$ and $t_{\text{hydro}}$, respectively, are given in Table 5. The runs are also shown in the $t_{\text{grav}}$-$e_{a_{\text{max}}}$ plane in Figure 13.

The maximum error is approximately 3 %, i.e. 1.5 times more than in the run with ABU off, and at the same time, the ABU makes the calculation approximately two times faster.

Panel (12f) shows a run with the tree-solver using the APE MAC with $e_{\text{lim}} = 10^{-2}$. The results are very similar to the one in run (d), with a maximum relative error of approximately 4.5 % ($\sim$ 1.5 larger) and the mean time per gravity time-step is 7 seconds (slightly smaller). Panel (12h) shows the run with the same tree-solver parameters, but instead of calculating the acceleration directly, the tree-solver calculates the potential and differentiates it numerically. The relative error exhibits a similar pattern to run (a), however, instead of sudden changes it includes high peaks of the error resulting from a numerical differentiation. Even though the mean error is comparable to runs (b) and (c), the maximum error is much higher, reaching 80 %. The time of the gravity calculation is slightly higher than in run (e), even though calculating the potential is cheaper than the acceleration for a single target cell. It is because a higher number of blocks must be updated in each time-step due to the higher error.

Panel (12g) includes the run with a reduced accuracy of $e_{\text{lim}} = 10^{-1}$ made to test the limits of the tree-solver usability. The relative error is high, in particular in the outer regions where the blocks are updated less often, reaching a maximum of 20 %, however, it is still a factor of 2 smaller than the error at the boundaries of the computational domain found in the run with the multi-grid solver (see below). On the other hand, the calculation is very fast with a mean time per gravity time-step of 3.5 seconds, which is $\sim 70$ % of the time needed by the hydro solver.

In order to evaluate the efficiency of the ABU, we show in Figure 14 a number of updated blocks in each time-step as a function of time.

Figure 13. Results of the turbulent sphere plotted in the plane of the gravity calculation duration in seconds, $t_{\text{grav}}$ (x-axis) versus the maximum relative error in the gravitational acceleration, $e_{a_{\text{max}}}$ (y-axis). The error is determined at $t = 2$ Myr and the maximum is taken over the whole computational domain. The thin dashed lines are iso-lines of constant $I_{\text{grav}} \times e_{a_{\text{max}}}$ assessing the code efficiency. Parameters of the displayed runs are given in Table 5.

Figure 14. Number of updated blocks, i.e. blocks for which the tree walk was executed for all grid cells in a given time-step, as a function of time (top x-axis) or time-step number (bottom x-axis). The figure shows first 2 Myr of the evolution of the turbulent sphere test. Individual curves represent models (a) – (e) as given in the legend (see also Table 5 for model parameters). Note that blue and magenta lines (models (b) and (c)) are on the top of each other.

The right panel (12b) shows the same calculation, but the ABU was switched on in this case. The relative error exhibits a rectangular pattern, because some blocks, in particular in the outer regions, were not updated in a given time-step and the error in them is larger. The maximum error is approximately 3 %, i.e. 1.5 times more than in the run with ABU off, and at the same time, the ABU makes the calculation approximately two times faster.

Panel (12f) shows a run with the tree-solver using the APE MAC with $e_{\text{lim}} = 10^{-2}$. The results are very similar to the one in run (d), with a maximum relative error of approximately 4.5 % ($\sim$ 1.5 larger) and the mean time per gravity time-step is 7 seconds (slightly smaller). Panel (12h) shows the run with the same tree-solver parameters, but instead of calculating the acceleration directly, the tree-solver calculates the potential and differentiates it numerically. The relative error exhibits a similar pattern to run (a), however, instead of sudden changes it includes high peaks of the error resulting from a numerical differentiation. Even though the mean error is comparable to runs (b) and (c), the maximum error is much higher, reaching 80 %. The time of the gravity calculation is slightly higher than in run (e), even though calculating the potential is cheaper than the acceleration for a single target cell. It is because a higher number of blocks must be updated in each time-step due to the higher error.

Panel (12g) includes the run with a reduced accuracy of $e_{\text{lim}} = 10^{-1}$ made to test the limits of the tree-solver usability. The relative error is high, in particular in the outer regions where the blocks are updated less often, reaching a maximum of 20 %, however, it is still a factor of 2 smaller than the error at the boundaries of the computational domain found in the run with the multi-grid solver (see below). On the other hand, the calculation is very fast with a mean time per gravity time-step of 3.5 seconds, which is $\sim 70$ % of the time needed by the hydro solver.

In order to evaluate the efficiency of the ABU, we show in Figure 14 a number of updated blocks in each time-step as a function of time.
of time / time-step number. The red curve corresponds to model (a) where ABU was switched off, i.e. it shows the number of all blocks in the simulation. It grows from 4096 to almost 6000, as the AMR creates more blocks in regions of high density formed by the gravitational collapse. The number of all blocks is the same for all simulations, as they run the identical model. For model (b) (blue curve), the number of updated blocks stays very small for the first ~ 10 time-steps, because the initial time-step is very low and the density and gravitational acceleration fields almost do not change. As soon as the time-step reaches a value given by the CFL condition, the number of updated blocks quickly rises up to 1000 and then it increases slowly to almost 2000 at 2 Myr. Throughout the evolution, the number of updated blocks is approximately three times lower than in model (a) with ABU off. As a result, model (b) runs more than twice as fast as model (a) and the maximum error in the acceleration is approximately 1.5 times larger (see Table 5 and Figures 12 and 13). For model (c), the fraction of updated blocks is almost the same as for model (b). However, the model runs ~ 30% faster as the APE MAC needs less interaction than the BH MAC of model (b) and consequently, the maximum error is ~ 1.5 larger. Model (e) with larger error limit updates less than 10% of blocks in each time-step and as a result it runs 8x faster than model (a) and its maximum error is almost 10x larger. Model (d) calculating the potential instead of the acceleration behaves in a different way. The number of updated blocks exceeds 3000 shortly after the start of the simulation, their fraction stays above 50% and reaches 100% in the last quarter of the time. It is because the numerical differentiation of the potential at the border between updated and not-updated blocks tends to give high error in the acceleration. Therefore we do not recommend the use of ABU together with calculating the potential.

Note that the efficiency of the ABU test is highly problem dependent. In this regard, the used turbulent sphere setup is a relatively hard one, because the sphere quickly forms dense filaments with large density gradients and they move supersonically as the whole structure collapses (i.e. the time-step is given mainly by the gas velocity, not the sound-speed). On the other hand, there are still regions where the gravitational acceleration changes slowly, e.g. in the computational domain corners, and these regions can be updated less often making the ABU efficient. If the volume with fast moving dense objects is larger, the ABU can be less efficient and vice versa.

### 3.3 Test of the OpticalDepth module

In order to evaluate the accuracy of the OpticalDepth module, we perform two tests. For both of them, we repeat the calculation of the turbulent sphere described in §3.2.2 using an adiabatic equation of state with $\gamma = 5/3$ (instead of the isothermal equation of state used previously). Additionally, we switch on the OpticalDepth and Chemistry modules calculating the gas cooling and heating, and the mass fractions of various species. The sphere is heated from the outside assuming a typical interstellar radiation field (ISRF) of $0.7 \times 10^3$ K, while the interior of the sphere is cold and thus it collapses to form stars as in runs with the isothermal equation of state in §3.2.2. A detailed description of the chemical network, the heating & cooling processes it includes, the dust temperature it calculates, and how the OpticalDepth module is coupled to it can be found in Walch et al. (2015). Here, we are only concerned with the workings of the OpticalDepth module and with the column density (or optical depth) it delivers.

In the first test, we evaluate how accurately the OpticalDepth module determines the column density depending on the chosen angular resolution; and in the second test, we compare the resulting optical depth with the optical depth computed using the Monte Carlo radiative transfer code RADMC-3D.

#### 3.3.1 Column density with increasing $N_{\text{pix}}$

We perform a test similar to the one by Clark et al. (2012) their §3.2, and calculate the “sky map” of hydrogen column density, $N_H$, as seen from the centre of the computational domain, for the turbulent sphere simulation at time $t = 2$ Myr (see the top right panel of Figure 11). The hydrogen column density determined by the OpticalDepth module, $N_{H,\text{pix}}$, is compared to the “actual” reference hydrogen column density, $N_H(\theta, \phi)$, obtained using a direct integration over individual grid cells of the simulation and very high HEALPix resolution $N_{\text{pix}} = 3072$. The angular resolution of the OpticalDepth module is controlled by two parameters: number of HEALPix elements $N_{\text{pix}}$ and tree maximum opening angle $\theta_{\text{max}}$ determining the maximum angular size of tree nodes. We calculate five models with $N_{\text{pix}} = 12, 48$, and 192, and two maximum opening angles $\theta_{\text{max}} = 0.5$ and 0.25 (see Table 6). We define a relative error in the hydrogen column density

$$e_{N_{\text{pix}}} = \frac{|N_{H,\text{pix}} - \langle N_H(\theta, \phi)\rangle_{\text{pix}}|}{\langle N_H(\theta, \phi)\rangle_{\text{pix}}}$$

(38)

where $\langle N_H(\theta, \phi)\rangle_{\text{pix}}$ is the mean value of reference hydrogen column density, $N_H$, in element $i$. In Table 6 we give for each model the maximum error $e_{N_{\text{pix},\text{max}}} = \max(e_{N_{\text{pix}}})$, where the maximum is taken over the whole sphere.

However, directionally dependent $N_{H,\text{pix}}$ does not directly enter calculations of the gas-radiation interaction. Instead, the Chemistry module uses quantities averaged over all directions, e.g. the mean visual extinction, $A_V$, given by Equation (26). Therefore, we further define a relative error in the mean visual extinction

$$e_{A_V} = \frac{|A_V - A_{V,\text{ref}}|}{A_{V,\text{ref}}}$$

(39)

where $A_V$ is the mean visual extinction calculated by the OpticalDepth module and $A_{V,\text{ref}}$ is the reference value obtained by averaging the high-resolution reference hydrogen column density $N_H(\theta, \phi)$ using Equation (26). Values of $e_{A_V}$ for the five calculated runs are also given in Table 6 and in top right corners of right panels in Figure 15.

The results are summarised in Figure 15 showing, in the Hammer projection, the reference hydrogen column density $N_H(\theta, \phi)$ (top panel), values of $N_{H,\text{pix}}$ calculated by the OpticalDepth module (left panels), and relative errors, $e_{N_{\text{pix}}}$ (right panels). Our findings are generally in agreement with those of Clark et al. (2012). Even run (a) with $N_{\text{pix}} = 12$ recovers approximately the overall structure of the cloud and results in $e_{N_{\text{pix},\text{max}}} = 0.16$ and $e_{A_V} = 0.19$. Increasing the HEALPix angular resolution to $N_{\text{pix}} = 48$ (run b) with the approximately same size of tree nodes and HEALPix elements, see the last column in Table 6, leads to a smaller error in $A_V$ while keeping $e_{N_{\text{pix},\text{max}}}$ approximately the same. Since runs (a) and (b) take nearly the same time to calculate, their comparison shows that it is not worth to degrade the HEALPix resolution by decreasing $N_{\text{pix}}$ below the tree-solver resolution (given by $\theta_{\text{max}}$). Similarly, run (c) with better tree solver resolution ($\theta_{\text{max}} = 0.25$) and the
even though it is still higher than in runs (a)-(c).

\[ V = t \theta, \]

\[ N_{\text{pix}} = 48 \]

\[ N_{\text{pix}} = 48 \]

about the approximations adopted when the mass of the real pixels corresponds to the angular resolution \( N_{\text{pix}} \). Since the variations of the dust density field and dust temperature provided by Weingartner & Draine (2001) (table for \( N_{\text{pix}} \)), we have \( \kappa_{\text{abs}}(\lambda_0) = 6.555 \times 10^{4} \text{ cm}^{2} \text{ g}^{-1} \). We obtain

\[ \tau_F = \kappa_{\text{abs}}(\lambda_0) \times N_{\text{dust,F}}. \]

Using the dust density field and dust temperature provided by the simulation, we compute the optical depth at the same wavelength using the RADMC-3D code, \( \tau_R \). With RADMC-3D it is possible to provide an external radiation field, in which case the photon packages are launched from the borders of the computational domain and pass through the grid in random directions. In each cell, they interact with the present dust according to its opacity. Using a typical Milky Way dust opacity provided by Weingartner & Draine (2001). The incoming intensity \( \lambda_0 \) is \( \lambda_0 = 9.547 \times 10^{-2} \text{ erg s}^{-1} \text{ cm}^{-2} \text{ Hz}^{-1} \text{ sr}^{-1} \). We run RADMC-3D in the mode mcmono to compute the intensity field at \( \lambda_0 \) in every cell of the computational domain. Then we convert this

\[ \kappa_{\text{abs}}(\lambda_0) = 6.555 \times 10^{4} \text{ cm}^{2} \text{ g}^{-1} \].

We obtain

\[ \tau_F = \kappa_{\text{abs}}(\lambda_0) \times N_{\text{dust,F}}. \]

Using the dust density field and dust temperature provided by the simulation, we compute the optical depth at the same wavelength using the RADMC-3D code, \( \tau_R \). With RADMC-3D it is possible to provide an external radiation field, in which case the photon packages are launched from the borders of the computational domain and pass through the grid in random directions. In each cell, they interact with the present dust according to its opacity. Using a typical Milky Way dust opacity provided by Weingartner & Draine (2001). The incoming intensity \( \lambda_0 \) is \( \lambda_0 = 9.547 \times 10^{-2} \text{ erg s}^{-1} \text{ cm}^{-2} \text{ Hz}^{-1} \text{ sr}^{-1} \). We run RADMC-3D in the mode mcmono to compute the intensity field at \( \lambda_0 \) in every cell of the computational domain. Then we convert this

\[ \kappa_{\text{abs}}(\lambda_0) = 6.555 \times 10^{4} \text{ cm}^{2} \text{ g}^{-1} \].

We obtain

\[ \tau_F = \kappa_{\text{abs}}(\lambda_0) \times N_{\text{dust,F}}. \]

Using the dust density field and dust temperature provided by the simulation, we compute the optical depth at the same wavelength using the RADMC-3D code, \( \tau_R \). With RADMC-3D it is possible to provide an external radiation field, in which case the photon packages are launched from the borders of the computational domain and pass through the grid in random directions. In each cell, they interact with the present dust according to its opacity. Using a typical Milky Way dust opacity provided by Weingartner & Draine (2001). The incoming intensity \( \lambda_0 \) is \( \lambda_0 = 9.547 \times 10^{-2} \text{ erg s}^{-1} \text{ cm}^{-2} \text{ Hz}^{-1} \text{ sr}^{-1} \). We run RADMC-3D in the mode mcmono to compute the intensity field at \( \lambda_0 \) in every cell of the computational domain. Then we convert this

\[ \kappa_{\text{abs}}(\lambda_0) = 6.555 \times 10^{4} \text{ cm}^{2} \text{ g}^{-1} \].

We obtain

\[ \tau_F = \kappa_{\text{abs}}(\lambda_0) \times N_{\text{dust,F}}. \]

Using the dust density field and dust temperature provided by the simulation, we compute the optical depth at the same wavelength using the RADMC-3D code, \( \tau_R \). With RADMC-3D it is possible to provide an external radiation field, in which case the photon packages are launched from the borders of the computational domain and pass through the grid in random directions. In each cell, they interact with the present dust according to its opacity. Using a typical Milky Way dust opacity provided by Weingartner & Draine (2001). The incoming intensity \( \lambda_0 \) is \( \lambda_0 = 9.547 \times 10^{-2} \text{ erg s}^{-1} \text{ cm}^{-2} \text{ Hz}^{-1} \text{ sr}^{-1} \). We run RADMC-3D in the mode mcmono to compute the intensity field at \( \lambda_0 \) in every cell of the computational domain. Then we convert this

\[ \kappa_{\text{abs}}(\lambda_0) = 6.555 \times 10^{4} \text{ cm}^{2} \text{ g}^{-1} \].

We obtain

\[ \tau_F = \kappa_{\text{abs}}(\lambda_0) \times N_{\text{dust,F}}. \]
Figure 15. Results of the OpticalDepth module test evaluating the accuracy of the hydrogen column density calculation as seen from the centre of the turbulent sphere, as a function of the used angular resolution. Top panel: shows the reference hydrogen column density, $N_H(\theta, \phi)$, displayed in the Hammer projection. Left panels below: show $N_{H,\text{pix}}$ determined by the OpticalDepth module with $N_{\text{pix}} = 12, 48, 192$, and $\theta_{\text{lim}} = 0.5$ and 0.25 as denoted in the top left corner of each panel. Right panels below: show the relative error in the column density, $e_{N_H,\text{pix}}$, calculated using Equation (38). The relative error in the mean visual extinction, $e_{A_V}$, is given in top right corners of the panels.

It is necessary to use a large number of photon packages in order to reduce the noise in the RADMC-3D calculation to an acceptable level. Specifically, we use 200 million photon packages and therefore it takes $\sim 53$ minutes on one 10-core Intel-Xeon E5-2650 CPU to simulate one wavelength on the given uniform grid with $128^3$ resolution, while the calculation with the OpticalDepth module took 24 seconds on 4-core Intel i7-2600, i.e. it was $\sim 330\times$ faster when normalising both calculations by number of cores.

In Figure 16 we show a slice at $z = 0$ of the resulting optical
depths (shown in logarithmic scale), \(T_\tau\) from the FLASH calculation (top left panel) and \(T_\tau\) from the RADMC-3D calculation (top right panel), as well as the difference between the two, normalized to the maximum \(T_{\tau,max}\) of the FLASH optical depth in the \(xy\)-plane (bottom right panel). The resulting gas temperature calculated by FLASH in displayed in the bottom left panel. The overall agreement is very good and on the level of the remaining noise of the RADMC-3D calculation of a few per cent. Although there is a tendency for the OpticalDepth module to slightly overestimate the optical depth in the densest regions, the difference is always < 10% and is ~ 1% for most cells in the computational domain. The result improves slightly if we use \(N_{pix} = 192\) and \(\theta_{lim} = 0.25\), but the additional expense of the calculation is generally not worth the effort.

### 3.4 Comparison of various MACs

We compare all available MACs with their typical parameters for a simple calculation similar to the static Bonnor-Ebert sphere test described in \[3.1.1\] however, carried out on a uniform 128\(^3\) grid. The aim is to provide an approximate measure of the code behaviour. A rigorous analysis of the efficiency of individual MACs, which would need many more tests, since it is highly problem dependent, is beyond the scope of this paper. The time of the gravity calculation, \(t_{grav}\), was measured on a single processor core, and since it is a single time-step calculation, the time is meaningful only for a mutual comparison between individual MACs (as is also the case for all the static tests in \[3.1.1\]). For each calculation, we determine the relative error in the gravitational acceleration and find its maximum in the computational domain, \(\epsilon_{a,max}\). The tested MACs are: the geometric (BH) MAC with three maximum opening angles, \(\theta_{lim}\), the APE MAC with both absolute and relative error limit \(\epsilon_{lim} = \epsilon_{lim,max} = 0.01\) (magenta empty square) and \(\epsilon_{lim} = 0.01\) (dark cyan filled square); an MPE MAC with absolute error limit given by \(\epsilon = 0.01\) (dark green empty triangle); and two SumSquare MACs with absolute error limits \(\epsilon_{lim} = 0.01\) (black filled circle) and \(\epsilon_{lim} = 0.1\) (orange empty circle). The violet filled triangle shows the calculation by the multi-grid solver. The thin dashed lines are iso-lines of constant \(t_{grav} \times \epsilon_{a,max}\) assessing the code efficiency.

![Figure 16](image1.png)

**Figure 16.** Test of the OpticalDepth module showing the calculated optical depths in a slice at \(z = 0\) through the turbulent sphere at time \(t = 2\) Myr. Top left: logarithm of the optical depth, \(\tau_\nu\), at wavelength \(\lambda_\nu = 9.36 \times 10^{-2} \mu m\) computed live during the FLASH simulation by the OpticalDepth module with \(\theta_{lim} = 0.5\) and \(N_{pix} = 48\); top right: logarithm of the optical depth, \(\tau_\nu\), at the same wavelength computed using RADMC-3D with 200 million photon packages; bottom left: gas temperature in the FLASH simulation resulting from the radiative heating taking the optical depth as an input; and bottom right: relative difference between the two optical depths, \((\tau_\nu - \tau_\nu)/\tau_\nu\). The difference is always < 10% and typically on the level of a few per cent, where most of it is caused by the noise in the RADMC-3D data.

![Figure 17](image2.png)

**Figure 17.** Comparison of available MACs for a single time-step calculation of the Bonnor-Ebert sphere on a uniform grid 128\(^3\). Each calculation with a different MAC is plotted in the plane of the gravity calculation duration in seconds (x-axis) versus the maximum relative error in the gravitational acceleration, \(\epsilon_{a,max}\) (y-axis). The maximum is taken over the whole computational domain. The tested MACs are: three geometric (BH) MACs with fixed maximum opening angles \(\theta_{lim} = 1.0\) (red plus), \(\theta_{lim} = 0.5\) (olive x), \(\theta_{lim} = 0.2\) (blue star); two APE MACs with absolute error limit \(\epsilon_{lim} = \epsilon_{lim,max} = 0.01\) (magenta empty square) and \(\epsilon_{lim} = 0.01\) (dark cyan filled square); an MPE MAC with absolute error limit given by \(\epsilon = 0.01\) (dark green empty triangle); and two SumSquare MACs with absolute error limits \(\epsilon_{lim} = 0.01\) (black filled circle) and \(\epsilon_{lim} = 0.1\) (orange empty circle). The violet filled triangle shows the calculation by the multi-grid solver. The thin dashed lines are iso-lines of constant \(t_{grav} \times \epsilon_{a,max}\) assessing the code efficiency.
3.5 Scaling tests

We perform both strong scaling and weak scaling tests. For that we use the setup of the turbulent sphere from §3.2.2 and §3.3.2. The strong scaling tests are done for the Gravity module only, the weak scaling is done for both Gravity and OpticalDepth modules.

For the strong scaling tests, we use three code configurations: tree solver with BH MAC and ABU off (model (a) from Table 5), tree solver with APE MAC and ABU on (model (c)), and multi-grid solver with default parameters (model (f)). We also show the scaling of the FLASH PPM hydrodynamic solver with default parameters. All tests have been run for 10 time-steps on the IT4i/Salomon supercomputer using 48 – 1536 cores. The speed-up on n processor cores, $S_n$, is determined with respect to the run with 48 cores.

$$ S_n = \frac{t_{48}}{t_n} $$  

(42)

where $t_{48}$ is the time spent by the evaluated module on 48 cores and $t_n$ is the time spent by the same module on n cores.

We see that the run with the tree-solver and BH MAC gives the best behaviour (speed-up closest to linear). On the other hand, this model is also the slowest one out of the three on 96 cores (see Table 5 and Figure 18). This can be understood by noting that most of the computational time is spent in the tree-walk, which runs completely in parallel without any communication. Additionally, without ABU there is no problem with load balancing, because the computational time is more or less directly proportional to the number of leaf-blocks, and thus each core receives the same number of leaf-blocks. The run with the APE MAC and ABU exhibits slightly worse scaling, however, the test in §3.2.2 shows that on 96 cores, it is almost 3 times faster than the BH MAC run. This is partially because, due to its more efficient MAC the code spends less time in fully parallel parts and partially because the ABU does not save the time equally on each processor core. The APE MAC scaling is still very good, comparable to the scaling of the hydrodynamic solver, which is highly parallel and needs only to communicate information at the boundaries between domains belonging to different processor cores. The multi-grid solver is very fast on 96 cores, comparable to the tree-solver with APE MAC and ABU, however, its efficiency decreases on higher number of cores.

The weak scaling test have been done for two configurations: (i) the tree solver with the gravity only runs using the APE MAC, $\varepsilon_{\text{lim}} = 0.01$ and ABU switched on, and (ii) runs calculating the gravitational acceleration and column densities of the three components (total, H$_2$ and CO; see §3.3.3) using the BH MAC, $\varepsilon_{\text{lim}} = 0.5$ with ABU switched off. Each configuration is run for four different grid resolution ranging from 64$^3$ to 512$^3$, with the number of cores, $n$, proportional to the number of grid cells ($n = 3, 24, 192, 1536$).

Results of the weak scaling tests are shown in Figure 19, where
a single time-step runtimes of the two configurations are compared with each other and with runtimes of the FLASH internal hydrodynamic solver. The hydrodynamic solver times (blue curve) follow approximately the \( n^{0.09} \) power-law, which is slightly worse than the "ideal" constant scaling. The tree solver using both the Gravity and OpticalDepth modules (green curve) exhibits a similar \( n^{0.15} \) scaling. On the other hand, runs with only the Gravity module (red curve) show the \( n^{0.15} \) scaling only between 3 and 24 cores, and for higher number of cores the scaling gets worse approaching \( n^{0.33} \). This is due to two reasons. Firstly, the gravity only runs are cheaper and the communication making the scaling worse is relatively more important. Note that the communication is negligible for runs on up to 24 cores, since a node on the Solomon computer comprises 24 cores with shared memory. Secondly, the load balancing needed by the ABU becomes worse on a high number of cores. We can also see that the additional calculations of column densities in the OpticalDepth module make the code approximately 10 times slower than the calculation of the gravity on a small number of cores, but it becomes only a factor of \( \sim 3.5 \) for 1536 cores due to better scaling of the more expensive runs with the OpticalDepth module.

4 SUMMARY

We have developed an MPI parallel, general purpose tree solver for the adaptive mesh refinement hydrodynamic code FLASH, that can be used to calculate the gas gravitational acceleration (or potential), optical depths enabling inclusion of the ambient diffuse radiation, and in future also general radiation transport (Paper II). The code uses an efficient communication strategy predicting which parts of the tree need to be sent to different processors allowing the whole tree walk to be executed locally. The advantage of this approach is a relatively low memory requirement, important in particular for the optical depth calculation, which needs to process information from different directions. This also makes the implementation of the general tree-based radiation transport possible. In addition to commonly implemented, fully isolated and fully periodic boundary conditions, the code can handle mixed (i.e. isolated in some directions and periodic in others) boundary conditions using a newly developed generalisation of the Ewald method. The gravity module implements several Multipole Acceptance Criteria (MACs) that increase the code efficiency by selecting which tree-nodes are acceptable for the calculation on the basis of the mass distribution within them. Using the Adaptive Block Update (ABU) technique, the code is able to re-use information from the previous time-step and thus further save computational time.

We have run a series of tests evaluating the code accuracy and performance, and compared them to the in-built multi-grid gravity solver of FLASH. The simpler "static" tests of the gravity module show that the code provides good accuracy for all combinations of boundary conditions. Comparison with the FLASH default multi-grid solver suggests that the tree-solver provides better accuracy for the same computational costs in the case of fully isolated boundary conditions, while with fully periodic boundary conditions the multi-grid solver seems to be more efficient.

Further, we run two more complex dynamical tests. The EUV test (gravitational collapse and re-bounce of the adiabatic, initially cold gaseous sphere) shows that it is critical to resolve well the dense centre, in order to ensure energy conservation. We found that in order to limit the error in the total energy to less than a few percent, it is necessary to resolve the Jeans length with at least 4 grid cells, a result similar to that of [Truelove et al. (1997)] where the same resolution is needed to prevent artificial fragmentation. In general, the EUV test turns out to be harder for grid codes in comparison with Lagrangian (e.g. SPH) hydrodynamic codes which reach almost perfect energy conservation with very small numbers of particles.

In the second dynamical test we simulate a turbulent sphere which collapses, fragments and forms sink particles (representing newly formed stars). We find the the tree-solver performs well and runs with accuracy of order several percent if it calculates the accelerations directly, and if it is used with the BH or APE MAC with typical parameters. Calculating the gravitational potential instead of the acceleration results in high (up to 80 %) errors due to numerical discretisation, and may result in numerical artifacts. The Adaptive Block Update accelerates the calculation by a factor of several (\( \sim 2 \)) for a given test. The multi-grid solver exhibits relatively high error (\( \sim 20 \% \)) close to computational domain boundaries, resulting from an inaccurate multipole solver. If the boundary regions are excluded, the accuracy of the tree-solver and multi-grid solver are comparable, while the tree-solver is approximately 2 times faster for the given test.

We run two tests of the OpticalDepth module. In the first one we measure the direction-dependent optical depths as a function of the angular resolution, and we find (in agreement with [Clark et al. (2012)]) that the code runs most efficiently if the angular resolution given by the number of Healpix rays is similar to the opening angle used by the tree-solver. In the second test we compare the optical depth calculated with the tree-solver with those calculated with the accurate radiation transport code RADMC-3D and we find an excellent agreement even for relatively low angular resolution – \( \sim 40 \) Healpix rays.

Further, using a simplified turbulent sphere test with uniform resolution, we compare the efficiency of all available MACs with their typical parameters. Generally, the BH MAC provides better accuracy for higher computational costs, while APE and MPE MACs result in lower (but often still acceptable) accuracy and are substantially faster. For applications, where an accuracy of order \( 10^{-2} \) is sufficient, the fastest choice seems to be the APE MAC with the absolute limit on the error.

Finally, we run strong scaling tests and show that the code scales up very well up to at least 1536 processor cores. We conclude that the presented tree-solver is a viable method for calculating self-gravity and other processes in astrophysics and that it is competitive with more commonly used iterative multi-grid methods.

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APPENDIX A: EQUATIONS FOR ACCELERATION IN COMPUTATIONAL DOMAINS WITH PERIODIC AND MIXED BOUNDARY CONDITIONS

In this appendix, we provide formulae for acceleration in computational domains with periodic and mixed boundary conditions. These formulae might be interesting particularly for the reader who intends to implement the Ewald method or its modification to a computational domain with mixed BCs. The orientation of symmetric axes is the same as in Section 2.2.

In analog to the equation for potential \(\nabla^2 \phi\), we write acceleration \(a(r)\) at target point \(r\) as

\[
a(r) = -G \sum_{i=1}^{N} m_i \mathbf{a}(r - r_i),
\]

where

\[
A = -\nabla \phi.
\]

### A1 Periodic boundary conditions

#### Defining

\[
e_{l_1,l_2,l_3}(x, y, z) = \frac{\exp(-\zeta(l_1^2 + (l_2/b)^2 + (l_3/c)^2))}{l_1^2 + (l_2/b)^2 + (l_3/c)^2},
\]

\[
u_{l_1,l_2,l_3}(x, y, z) = \left\{ \begin{array}{ll}
\frac{2\pi l_i(x-x_i) + 2\pi l_j(y-y_j) + 2\pi l_k(z-z_k)}{L_x} & \text{if } l = 1, \\
\frac{2\pi l_i(y-y_i) + 2\pi l_j(z-z_j)}{bL_x} & \text{if } l = 2, \\
\frac{2\pi l_i(z-z_i)}{cL_x} & \text{if } l = 3,
\end{array} \right.
\]

one obtains by differencing Equation [11] the components of function \(A\) in the form of

\[
A_x = \sum_{l_1,l_2,l_3} \left\{ \left( \frac{2\alpha}{\sqrt{\pi}} \exp(-\zeta^2 u_{l_1,l_2,l_3}) + \text{erfc}(\alpha \sqrt{\zeta u_{l_1,l_2,l_3}}) \right) \times \right.
\]

\[
(x - x_i - i_1 L_x) + \frac{2}{bL_x^2} \sum_{l_2,l_3} I_1 e_{l_1,l_2,l_3} \sin(v_{l_1,l_2,l_3}),
\]

\[
A_y = \sum_{l_1,l_2,l_3} \left\{ \left( \frac{2\alpha}{\sqrt{\pi}} \exp(-\zeta^2 u_{l_1,l_2,l_3}) + \text{erfc}(\alpha \sqrt{\zeta u_{l_1,l_2,l_3}}) \right) \times \right.
\]

\[
(y - y_i - i_2 b L_x) + \frac{2}{bL_x^2} \sum_{l_1,l_3} I_2 e_{l_1,l_2,l_3} \sin(v_{l_1,l_2,l_3}),
\]

\[
A_z = \sum_{l_1,l_2,l_3} \left\{ \left( \frac{2\alpha}{\sqrt{\pi}} \exp(-\zeta^2 u_{l_1,l_2,l_3}) + \text{erfc}(\alpha \sqrt{\zeta u_{l_1,l_2,l_3}}) \right) \times \right.
\]

\[
(z - z_i - i_3 c L_x) + \frac{2}{bL_x^2} \sum_{l_1,l_2} I_3 e_{l_1,l_2,l_3} \sin(v_{l_1,l_2,l_3}).
\]

### A2 Mixed boundary conditions of type 2P11

To simplify the formulae below, we define

\[
u_{l_1,l_2} = \frac{(x-x_i-i_1 L_x)^2 + (y-y_i - i_2 b L_x)^2 + (z-z_i)^2}{L_x}.
\]

and

\[
\tilde{I}(l_1,l_2,z-z_k) = I(l_1,l_2,z-z_k) \text{erfc}(-\zeta(l_1^2 + (l_2/b)^2))
\]

\[
\times \exp\left(\frac{\gamma^2}{4\zeta}\right)
\]

\[
\times \exp\left(-\gamma \frac{l_1^2 + (l_2/b)^2}{\zeta}\right)
\]

\[
I'(l_1,l_2,z-z_k) = \frac{\partial I(l_1,l_2,z-z_k)}{\partial y}
\]

\[
\times \exp\left(-\gamma \frac{l_1^2 + (l_2/b)^2}{\zeta}\right)
\]

\[
\times \exp\left(-\gamma \frac{l_1^2 + (l_2/b)^2 - \gamma/2}{\zeta}\right)
\]

where \(l(l_1,l_2,z-z_k)\) is defined by Equation [17].

Function \(A\) then takes the form

\[
A_x = \sum_{l_1,l_2,l_3} \left\{ \left( \frac{2\alpha}{\sqrt{\pi}} \exp(-\zeta^2 u_{l_1,l_2,l_3}) + \text{erfc}(\alpha \sqrt{\zeta u_{l_1,l_2,l_3}}) \right) \times \right.
\]

\[
(x - x_i - i_1 L_x) + \frac{2}{bL_x^2} \sum_{l_2,l_3} I_1 e_{l_1,l_2,l_3} \sin(v_{l_1,l_2,l_3}),
\]

\[
A_y = \sum_{l_1,l_2,l_3} \left\{ \left( \frac{2\alpha}{\sqrt{\pi}} \exp(-\zeta^2 u_{l_1,l_2,l_3}) + \text{erfc}(\alpha \sqrt{\zeta u_{l_1,l_2,l_3}}) \right) \times \right.
\]

\[
(y - y_i - i_2 b L_x) + \frac{2}{bL_x^2} \sum_{l_1,l_3} I_2 e_{l_1,l_2,l_3} \sin(v_{l_1,l_2,l_3}),
\]

\[
A_z = \sum_{l_1,l_2,l_3} \left\{ \left( \frac{2\alpha}{\sqrt{\pi}} \exp(-\zeta^2 u_{l_1,l_2,l_3}) + \text{erfc}(\alpha \sqrt{\zeta u_{l_1,l_2,l_3}}) \right) \times \right.
\]

\[
(z - z_i - i_3 c L_x) + \frac{2}{bL_x^2} \sum_{l_1,l_2} I_3 e_{l_1,l_2,l_3} \sin(v_{l_1,l_2,l_3}).
\]

\[
- \frac{2}{bL_x^2} \sum_{l_1,l_2} \cos(v_{l_1,l_2}) I'(l_1,l_2,z-z_k).
\]
A3 Mixed boundary conditions of type 1P2I

Here, we introduce

\[ u_i = (x - x_o - iL_x)^2 + (y - y_o)^2 + (z - z_o)^2, \]  
\[ v_i = \frac{2m_l(x - x_o)}{L_x}, \]

which simplifies the formula for function A to

\[ A_x = \sum_{i, j < 10} \left\{ \frac{2\alpha \exp(-\alpha^2 u_i)}{\sqrt{\pi}} + \frac{\text{erfc}(\alpha \sqrt{\pi} u_i)}{u_i^{1/2}} \right\} (x - x_o - iL_x), \]
\[ A_y = \frac{4\pi}{L_x^2} \sum_{i, j < 10} l_i \exp(-\zeta_i^2) \sin(v_i) K(l_i, y - y_o, z - z_o), \]
\[ A_z = \frac{4\pi}{L_x^2} \sum_{i, j < 10} \frac{1}{\sqrt{\pi} L_x} \exp(-\zeta_i^2) \cos(v_i) M(l_i, y - y_o, z - z_o). \]

where \( K(l_i, y - y_o, z - z_o) \) is given by Equation (22), and function \( M(l_i, y - y_o, z - z_o) = -\delta K(l_i, \eta(y - y_o, z - z_o))/\delta \eta \) is

\[ M(l_i, y - y_o, z - z_o) = \int_0^\infty J_1(\eta \zeta) \exp(-\eta^2 q^2) dq, \]

where \( J_1 \) is the Bessel function of the first kind and first order. Note that variables \( \zeta, \eta \) and \( \eta \) are defined in section 2.2.4.

APPENDIX B: CODE RUNTIME PARAMETERS

Here we list runtime parameters of the tree solver and Gravrv and OpticalDepth modules that can be set in the flash.par configuration file. Apart from parameters discussed in the main body of this work (e.g. MAC selection, accuracy limit), the code should work well with the default parameters. Additional information is provided in the Flash Users Guide and directly in the source code as comments.

B1 Tree solver parameters

- gr_bhTreeSafeBox – relative (w.r.t. to the block size) size of a cube around each block, \( \eta_{ba} \), in which the target point cannot be located (type: real, default: 1.2)
- gr_bhUnifiedTW – obsolete, will be deleted in future versions
- gr_bhTWMaxQueueSize : maximum number of elements in the priority queue (type: integer, default: 10000)
- gr_bhAcceptAccurateOld – indicates whether Adaptive Block Update (see \( 2.2.6 \)) is active; will be renamed to gr_bhABU in future versions (type: logical, default: false)
- gr_bhLoadBalancing – indicates whether Load Balancing (see \( 2.2.6 \)) is active (type: logical, default: false)
- gr_bhMaxBlkWeight – maximum workload weight, \( \omega_{\text{max}} \) (type: real, default: 10)

B2 Gravity module parameters

- grv_bhNewton – Newton’s constant of gravity; if negative, the value is obtained from the Flash internal database of physical constants (type: real, default: -1)
- grv_bhMAC – type of Multiple Acceptance Criterion (MAC) calculated by the Gravity module if grv_bhPhysMACComm is true; currently accepted values are: “ApproxPartialErr”, “MaxPartialErr” and “SumSquare” (experimental), (type: string, default: “ApproxPartialErr”)
- grv_bhMPDegree – degree of multipole expansion used to estimate the error of a single node contribution with APE and MPE MACs; grv_bhMPDegree corresponds to \( p+1 \) used in Equations (2) and (3) (type: integer, default: 2)
- grv_bhUseRelAccErr – indicates whether the grv_bhAccErr parameter (below) should be interpreted as a relative error limit, \( \epsilon_{\text{lin}} \) (true), or an absolute error limit, \( a_{\text{lin}} \) (false); see Equations (4) and (5) (type: logical, default: false)
- grv_bhAccErr – maximum allowed error set either relatively with respect to the acceleration from the previous time-step, \( \epsilon_{\text{lin}} \), or absolutely, \( a_{\text{lin}} \) (type: real, default: 0.1)
- grav_boundary_type – type of boundary conditions for gravity for all directions; the accepted values are: “isolated” and “periodic” and “mixed”, if set to “mixed”, BCs in individual directions are set by the parameters below (type: string, default: “isolated”)
- grav_boundary_type.x – type of gravity BCs in the x-direction; the accepted values are: “isolated” and “periodic” (type: string, default: “isolated”)
- grav_boundary_type.y – same as grav_boundary_type.x but in the y-direction
- grav_boundary_type.z – same as grav_boundary_type.x but in the z-direction
- grav_bhEwaldSeriesN – number of terms used in the expansion given by Equation (11) to calculate the Ewald field (type: integer, default: 10)
- grav_bhEwaldAlwaysGenerate – indicates whether the Ewald field should be regenerated at the simulation start; if false, it is read from file with name given by parameters grav_bhEwaldFName or grav_bhEwaldFNameAccV42 and grav_bhEwaldFNamePosV42 (type: logical, default: true)
- grav_bhEwaldFieldNxV42 – number of points of the Ewald field lookup table in the x-direction when the first approach described in \( 2.2.5 \) is used (default in Flash versions up to 4.2); (type: integer, default: 32)
- grav_bhEwaldFieldNyV42 – same as the preceding parameter but for the y-direction
grv_bhEwaldFieldNzV42 – same as the preceding parameter but for the z-direction

grv_bhEwaldNRefV42 – number of nested grid levels of the Ewald field when the first approach described in §2.2.5 is used; if negative, the number of nested grid levels is calculated automatically from the minimum cell size (type: integer, default: -1)

grv_bhLinearInterpolOnlyV42 – indicates whether the linear interpolation in the Ewald field is used (with the first approach described in §2.2.5); if false, then the more expensive and accurate quadratic interpolation is used for some calculations (type: logical, default: true)

grv_bhEwaldFNameAccV42 – name of file to store the Ewald field accelerations when the first approach described in §2.2.5 is used (type: string, default: "ewald_field_acc")

grv_bhEwaldFNamePotV42 – name of file to store the Ewald field potential when the first approach described in §2.2.5 is used (type: string, default: "ewald_field_pot")

grv_bhEwaldNPer – number of points in each direction of the Ewald field coefficients when the second approach described in §2.2.5 is used (type: integer, default: 32)

grv_bhEwaldFName – name of file to store the Ewald field coefficients in the case the second approach described in §2.2.5 is used (type: string, default: "ewald_coeffs")

grv_useExternalPotential – indicates whether the external time-independent gravitational potential read from file is used (type: logical, default: false)

grv_usePoissonPotential – indicates whether the potential (or accelerations) computed by the (tree) Poisson solver is used (type: logical, default: true)

grv_bhExtrnPotFile – name of file with the external gravitational potential (type: string, default: "external_potential.dat")

grv_bhExtrnPotType – symmetry of the external gravitational potential; currently, two options are available: "spherical" and "planez" (plane-parallel, varying along the z-direction); (type: string, default: "planez")

grv_bhExtrnPotCenterX – center of the external potential x-coordinate given in the FLD internal coordinates (type: real, default: 0)

grv_bhExtrnPotCenterY – same as the preceding parameter but for the y-direction

B3 OpticalDepth module parameters

tr_nSide – level of the HEALPix grid; number of pixels is \(N_{\text{pix}} = 12 \times 4^{(\text{tr}_{\text{nSide}}-1)}\) (type: integer, default: 1)

tr_i1NR – number of points in the radial direction for the calculation of the fraction of node that intersects with a given ray (type: integer, default: 50)

tr_i1NTheta – number of points in the \(\theta\)-direction of the table recording a fraction of the node that intersects with a ray at a given \(\theta\) (type: integer, default: 25)

tr_i1NPhi – number of points in the \(\phi\)-direction of the node-ray intersection table (type: integer, default: 25)

tr_i1NNS – number of points describing the angular node size in the node-ray intersection table (type: integer, default: 50)

tr_i1FinePix – number of additional pixels in each angular directions used to calculate the node-ray intersection table (type: integer, default: 4)

tr_bhMaxDist – maximum distance from a target point up to which the optical depth is calculated (type: real, default: 10^9)

tr_odCDTOIndex – exponent relating the gas density to the absorption coefficient used during the calculation of the optical depth in a given direction (type: real, default: 1)