Testing Hydrodynamics Schemes in Galaxy Disc Simulations

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Submitted

ABSTRACT

We examine how three fundamentally different numerical hydrodynamics codes follow the evolution of an isothermal galactic disc with an external spiral potential. We compare an adaptive mesh refinement code (\textsc{ramses}), a smoothed particle hydrodynamics code (\textsc{sphNG}), and a volume-discretised meshless code (\textsc{gizmo}). Using standard refinement criteria, we find that \textsc{ramses} produces a disc that is less vertically concentrated and does not reach such high densities as the \textsc{sphNG} or \textsc{gizmo} runs. The gas surface density in the spiral arms increases at a lower rate for the \textsc{ramses} simulations compared to the other codes. There is also a greater degree of substructure in the \textsc{sphNG} and \textsc{gizmo} runs and secondary spiral arms are more pronounced. By resolving the Jeans’ length with a greater number of grid cells we achieve more similar results to the Lagrangian codes used in this study. Other alterations to the refinement scheme (adding extra levels of refinement and refining based on local density gradients) are less successful in reducing the disparity between \textsc{ramses} and \textsc{sphNG}/\textsc{gizmo}. Although more similar, \textsc{sphNG} displays different density distributions and vertical mass profiles to all modes of \textsc{gizmo} (including the smoothed particle hydrodynamics version). This suggests differences also arise which are not intrinsic to the particular method but rather due to its implementation. The discrepancies between codes (in particular, the densities reached in the spiral arms) could potentially result in differences in the locations and timescales for gravitational collapse, and therefore impact star formation activity in more complex galaxy disc simulations.

Key words: methods: numerical – hydrodynamics – galaxies: evolution – galaxies: structure

1 INTRODUCTION

It is well known that galactic dynamics play an important role in the formation of star-forming regions (e.g. Dobbs et al. 2014), with the gravitational potential of the spiral structure competing with hydrodynamical forces, large-scale differential rotation, and energetic feedback. The interplay of these processes is particularly difficult to study observationally and the hydrodynamical complexity means that a full understanding is analytically intractable. For these reasons, numerical simulations are a dominant tool for furthering our understanding of gas dynamics in a galactic context. However the results can be quite different depending on the particular methodology for solving these equations. Hydrodynamics codes (which by now are extremely complex) may give conflicting results due to the respective strengths and weaknesses of the different implementations.

Code comparisons seek to quantify how different methodologies reproduce fluid flow and where weaknesses lie. These comparisons tend to concentrate on idealised test conditions (Agertz et al. 2007; Tasker et al. 2008; Price 2008; Hopkins 2015) which are easy to compare objectively, or on reproducing turbulent behaviour (Klessen, Heitsch & Mac Low 2000; Kitsionas et al. 2009; Price & Federrath 2010; Kritsuk et al. 2011), while others consider cosmological galaxy formation (Frenk et al. 1999; Pearce et al. 1999; O’Shea et al. 2005; Torrey et al. 2012; Kereš et al. 2012; Scannapieco et al. 2012). To date, almost no work (excepting Hopkins (2015)) has compared the behaviour of isolated galaxy discs with different hydrodynamical codes.

In this work we compare the Adaptive Mesh Refinement (AMR) code \textsc{ramses} (Teyssier 2002), the Smoothed Parti-
1.1 Hydrodynamical Methods

1.1.1 Smoothed Particle Hydrodynamics

SPH methods use the movement and concentration of particles to provide automatic refinement in dense areas. This means that low density regions are more poorly resolved and the resolution of phenomena in areas where density gradients are steep are not guaranteed. Despite these issues, overdensities are almost always our regions of interest, and SPH methods are extremely useful.

We will not describe here the details of all SPH codes employed in this field but simply note works using them (e.g. Dobbs, Bonnell & Pringle 2006; Dobbs et al. 2008; Saitoh et al. 2008; Robertson & Kravtsov 2008; Hopkins et al. 2012; Williamson & Thacker 2012; Grand, Kawata & Cropper 2012; Dobbs & Pringle 2013; Mata-Chávez, Gómez & Puerari 2014; Williamson et al. 2014) and that typically resolution is given by particles numbers of $\sim 10^6 - 10^7$ and mass resolutions of order $10^4 - 10^5 M_\odot$. Other than deliberate choices of cooling, star formation, and feedback etc, the key differences between these codes include how the smoothing length is determined, whether the SPH equations are used in pressure-energy or density-entropy form, and how artificial viscosity and conductivity are treated.

1.1.2 Grid-based Hydrodynamics

The majority of other work in simulating isolated galactic discs employ some form of grid method. The simplest of these is a fixed cartesian grid which may fit around a single spiral arm in the fashion of Kim & Ostriker (2002, 2006) or around the entire disc (e.g. Wada & Norman 2007; Wada 2008; Khoperskov et al. 2013) for which spatial resolution depends upon the size of disc that is simulated but can feasibly reach $\sim 7$ pc. Of the works mentioned here only Khoperskov et al. (2013) simulates a disc with size comparable to the MW.

To reach greater resolution, grid simulations often use adaptive mesh refinement (Berger & Oliger 1984; Berger & Colella 1989), whereby grids are subdivided to some appropriate resolution based on local hydrodynamical properties. The key strength of this approach is that it gives the user almost unlimited flexibility and control over which parts of the simulation volume are to be resolved and that large scale effects can be incorporated into simulations where very fine spatial phenomena are to be studied. It should be noted though that this means that different works can employ different refinement schemes and parameters which influence the simulations.

One approach requires that the Jeans’ length be resolved by a minimum number of cells at each level. The minimum number is often set to the limit derived in Truelove et al. (1997), i.e. 4 grid cells per Jeans’ length. This refinement criterion is most important when considering self-gravitating gas and the Truelove et al. (1997) limit is intended to prevent numerical fragmentation, although resolving the Jeans’ length is also necessary to capture physical fragmentation of the gas. The Jeans’ length refinement criterion is employed in Tasker & Tan (2009); Renaud et al. (2013); Fujimoto et al. (2014); Petit et al. (2015); Tasker, Wadsley & Pudritz (2015). Of particular note here is the work of Petit et al. (2015) in which the number of grid cells per Jeans’ length is increased from the typical 4 to a more rigorous value of 32.

Another approach, sometimes termed ‘quasi-Lagrangian’, resolves cells based on the local density such that the mass enclosed in a grid cell is roughly the same on each refinement level (e.g. Bournaud et al. 2010; Fujimoto et al. 2014; Agertz, Romeo & Grisdale 2015). One strength of this technique is that the mass per grid cell may also include stellar or dark matter mass which are usually gravitationally dominant. In this way, a stellar substructure can be well resolved for the gas phase even before the gas density increases. Note that even for isothermal runs in the absence of non-gaseous mass (such as presented in this work), a fixed mass threshold for refinement is not equivalent to a fixed Jeans’ length threshold.

The typical finest resolution for AMR runs ranges from 9 pc to 1.5 pc (considered sufficient to resolve the formation of the largest giant molecular clouds) with Renaud et al. (2013) achieving 0.05 pc resolution, albeit only for the last 50 Myr of the 780 Myr run. However, while oft quoted as “the resolution”, stating the minimum grid size is only slightly more informative than is the minimum smoothing length of an SPH simulation unless the phenomenon in which one is interested is entirely comprised of grid cells on that ultimate refinement level. Other refinement criteria are available, see Khokhlov (1998), but those described here cover the approaches currently employed in the simulation of galactic discs.

In addition to the criteria for refinement, and limits on the permitted levels, refinement usually also takes place in a cubic buffer around refined cells, this smooths the transition between different levels and prevents a noisy mesh structure from forming; this buffer can vary in size and only ever increase resolution, refining cells that are not ordinarily qualified. Finally, some codes require de-refinement criteria for the grid, but as RAMES is not among them, we do not discuss this here.

1.1.3 Hybrid Hydrodynamical Methods

For a long time the two dominant hydrodynamics schemes were SPH and grid (often AMR) methods, recently however new techniques have been developed. One new method, called Godunov-SPH or GSPH, involves a reformulation of SPH to introduce a Riemann solver that determines the force acting between each particle pair (Inutsuka 2002; Cha &
Whitworth 2003). This does in principle grant the ability to resolve shocks in the absence of artificial viscosity and therefore avoid any side effects that occur as a consequence.

‘Moving-mesh’ techniques are a different approach to hydrodynamical codes which hybirdise Lagrangian and Eulerian schemes. This technique is used in AREPO (Springel 2010). It evolves a finite-volume unstructured Voronoi mesh that moves with the fluid flow. This approach retains the ability of Eulerian codes to resolve shocks by employing a Riemann solver across each boundary between cells. Moving-mesh schemes are Galilean invariant and less noisy and less diffusive than SPH codes. AREPO is applied to the problem of an isolated galactic disc in Pakmor & Springel (2013) and Smith et al. (2014).

Another method that hybridises SPH and grid codes is the so-called ‘meshless’ scheme that is detailed in Larson & Vila (2008a,b); Gaburov & Nitadori (2011); Hopkins (2015). In these codes, the particle ensemble is topologically similar to moving-mesh frameworks but for the lack of a sharply defined boundary between the resolution element domains. These codes are closer in their execution to SPH codes than moving-mesh techniques are but similarly employ a Riemann solver across the interfaces between resolution elements, enabling shock capturing. In this work we consider one of these meshless codes, GIZMO (Hopkins 2015), in our comparisons.

### 1.2 Code Comparisons

The differences between SPH and grid-based codes have been much discussed and we now briefly review the most relevant code comparison literature.

The first main type of comparison uses idealised tests that are simple setups for which analytical solutions exist such as the Sod shock tube or Sedov blast wave or instability tests (e.g. Kelvin-Helmholtz). Agertz et al. (2007) compare AMR and SPH using a similar number of resolution elements in the areas of interest. They found that all grid codes tested produced similar results for blob tests, Kelvin-Helmholtz and Rayleigh-Taylor instability tests and likewise, the SPH codes produced similar results as each other. Agertz et al. (2007) find that contact discontinuities and blob disintegration are less well reproduced by SPH codes, however Price (2008) counters that the inclusion of an artificial thermal conductivity term in SPH formulations reproduces these discontinuities. Likewise, the introduction of thermal diffusion improves instability resolution in SPH codes (Wadsley, Veeravalli & Couchman 2008). The inclusion of artificial conductivity/thermal diffusion is not relevant to this particular study because we are considering isothermal flows.

Idealised tests are presented in Tasker et al. (2008) with the conclusion that SPH codes excel at resolving the behaviour of dense objects while AMR codes are the preferred choice for voids (because they retain resolution in low density regions) and shocks because of the ability to force resolution in areas of steep density contrast. This work also found that SPH codes can struggle to resolve fluid instabilities, while praising grid codes for being able to model multiphase fluids due to the sharp contrasts that can exist across grid boundaries. Despite these differences, the conclusion is that concordance between grid and particle codes is reached when there is one particle per grid cell in the region of interest.

The second main type of code comparison examines the turbulent properties of gas under the influence of some driving mechanism. Price & Federrath (2010) compare the ability of PHANTOM and FLASH to model supersonic driven turbulence and achieve similar results with comparable numbers of resolution elements ($512^3$). Despite similarities, PHANTOM is better at resolving dense structures, reaching densities at $128^3$ resolution that FLASH only achieves at $512^3$. The resolution of high density regions is best achieved with SPH, but the grid resolves low density structures better.

In Kitsionas et al. (2009), decaying turbulence is modelled with a variety of static grid and SPH codes. Grid codes are found to be less dissipative, but for the same number of resolution elements encouragingly similar results emerge. They conclude that resolution rather than the method primarily drives differences in turbulence simulations. Supersonic turbulence decay is also examined in Kritsuk et al. (2011) as a test of magnetodynamics codes. While the nature of the simulation set up is not completely relevant to this work we emphasise the result that even quite similar codes can generate different results due to minor code construction choices.

The third common framework for code comparison is galaxy formation in a cosmological context (Frenk et al. 1999; Pearce et al. 1999; Torrey et al. 2012; Kereš et al. 2012; Scannapieco et al. 2012). In works of this kind, insights into the accuracy of the hydrodynamical methods can be masked by dominant gravitational effects or by the method of analysis which usually focuses on derived physical properties not directly relevant to the scope of this study. Forthcoming comparisons in a cosmological context and with an isolated disc is expected under the AGORA project (Kim et al. 2014).

Finally, we mention Hopkins (2015) because in addition to a very detailed look at a number of idealised tests, the author also provides a comparative analysis of a cold Keplerian disc test and an isolated galactic disc, both of which are relevant to the work presented here. Hopkins (2015) shows the results of the viscous instability that affects SPH realisations due to shear viscosity in the cold Keplerian disc. Grid methods do not suffer from numerical viscosity in this context, but advection errors can cause rings to form in what should be a uniform disc. The work also presents two new methods in the form of a meshless lagrangian code either with finite-mass (MFM) or finite-volume (MFV) resolution elements, which perform well in this test and others. The analysis of isolated disc galaxy simulations is similar to what we present in this work. Those runs have lower particle mass resolution compared to ours but include star formation, stellar feedback, gas cooling and a live dark matter halo, central black hole, disc and bulge initialized in equilibrium. The finding here is that SPH runs are similar to one another and to the MFM realisations, however MFV transfers mass outwards due to an angular momentum advection error. It is particularly noteworthy that the critical flaw seen in SPH when applied to the Keplerian disc problem (viscous instability) does not manifest in this context because the pressure forces are much higher and the large stellar/gas mass ratio allows the stellar component to dominate and stabilise the gas disc.

The problem with determining the strengths and weak-
nesses of different methodologies using idealised tests is that the exact implementation used in those tests is often changed when the codes are applied to real problems. One example that stands out is that in idealised tests of AMR codes, different refinement criteria are applied to that which are used in production runs. In some cases there is a justification for not applying the same refinement criteria, i.e. if the phenomenon being studied requires resolution based on the local density or if the ideal criteria would resolve an impractical amount of the volume which makes running the simulation intractable. This is the case in simulations of isolated galactic disc using AMR where resolution criteria are usually ‘quasi-Lagrangian’ or designed to resolve the Jeans’ length rather than resolving local gradients, as used for example when performing shock tube or blast wave tests.

2 METHOD

In this work we compare three simulation codes: RAMSES, SPHNG, and GIZMO in a common framework. We simulate the evolution of isothermal gas from an initially uniform surface density disc under the influence of a disc galaxy gravitational potential with a fixed spiral perturbation. We now describe each of the codes employed in this work.

2.1 RAMSES

RAMSES (Teyssier 2002) is an AMR code in which gas dynamics are computed with a second-order unsplit Godunov scheme. This scheme is inherently shock capturing with no need to invoke artificial viscosity. Time steps are advanced using a midpoint method with time centered fluxes at cell boundaries used to update the hydrodynamical variables. The time centered fluxes are determined using a second-order Godunov method (otherwise known as the Piecewise Linear Method). The duration of the time steps themselves is limited by a modified Courant-Friedrichs-Lewy (CFL) condition such that the time step is

$$\Delta t_{\text{CFL}} = \frac{\Delta x_{\ell}}{\sum_{i=1}^{N_{\text{dim}}}(|u_i|+c_s)} \sqrt{1 + 2C_{\text{CFL}}G_{\text{SR}} - 1} - 1$$

where $\Delta x_{\ell}$ is the linear extent of a grid cell at level $\ell$ with $N_{\text{dim}}=3$ dimensional velocity $u$ and sound speed $c_s$. The right-hand part of equation 1 replaces the more traditional multiplication by the CFL factor, $C_{\text{CFL}}$. This is changed here to vary with the gravitational strength ratio,

$$\epsilon_{\text{GSR}} = \frac{\Delta x_{\ell}}{\sum_{i=1}^{N_{\text{dim}}}|g_i|} \left(\sum_{i=1}^{N_{\text{dim}}}(|u_i|+c_e)\right)^2$$

where $g$ is the gravitational acceleration experienced by each grid cell. The right-hand part of equation 1 is equal to $C_{\text{CFL}}$ for $\epsilon_{\text{GSR}} \to 0$ and is smaller (thus shortening the timestep) when the gravitational acceleration is large relative to the local gas velocity. The RAMSES runs presented here use $C_{\text{CFL}}=0.8$. RAMSES supports different time steps for each level of the grid but we have enforced complete synchronicity so that the time step of coarse levels is identical to, and thus limited by, lower levels.

RAMSES is equipped with a number of solver options, the choices of which are rarely mentioned in literature. In this work we follow Renaud et al. (2013) and use the acoustic Riemann solver with the MinMod slope limiter, however we also include a single run that uses the exact Riemann solver and the MonCen slope limiter. We set the mesh-smoothing parameter $n_{\text{expand}}=1$.

In this work we use different combinations of refinement criteria which are now described. The first criterion ensures that the Jeans’ length ($\lambda_J$) is resolved by $N_j$ cells. Each cell on level $\ell$ is marked for refinement if

$$\frac{\lambda_J}{\Delta x_{\ell}} < N_j$$  \hspace{1cm} (3)

An alternative criterion forces refinement where the local gradient in a variable $q$ exceeds a fraction of the value of that variable. Refinement occurs if the following condition is satisfied,

$$C_q < 2 \cdot \text{MAX} \left[ \frac{q_{i-1} - q_i}{q_{i-1} + q_i + f_q} \right], \frac{q_i - q_{i+1}}{q_i + q_{i+1} + f_q} \right]$$  \hspace{1cm} (4)

where $i-1$ and $i+1$ are the cells that bound cell $i$ in each dimension. The two user defined parameters here are the threshold $C_q$ and a pseudo-floor value $f_q$. In this work we use only the gradient in density for grid refinement. In principle the gradient in any hydrodynamical variable may be used, but as our simulations are isothermal, density and pressure criterion are degenerate and we have found that for our simulation set up, using velocity gradients as a criterion for refinement adds very little if density gradients are already being used.

One refinement scheme that we do not examine is the quasi-Lagrangian scheme whereby the mass enclosed in a given cell is compared to a threshold to determine if the cell is massive enough to warrant refinement. We do not examine this method because it is commonly used by combining the mass from the gas phase and stellar particles, the latter of which are not included in our simulations.

2.2 SPHNG

The SPH code used here (referred to as SPHNG) is a modified version of the code presented in Benz (1990). The density of a particle $i$ is estimated through a weighted sum of the mass ($m$) of itself and its neighbours;

$$\rho_i = \sum_j m_j W(|x_i - x_j|, h_{ij})$$  \hspace{1cm} (5)

where the weighting function where $W$ is the cubic spline kernel which is a function of the particle positions $x$ and the mean smoothing length for each particle pair, $h_{ij}$. Particles are assigned variable smoothing lengths according to the local particle density (Price & Monaghan 2004). The smoothing length and density are solved iteratively via the Newton-Raphson method according to

1. RAMSES is a publicly available code and can be found at http://www.ics.uzh.ch/~teyssier/ramses/RAMSES.html
\[ h = \eta (m/\rho)^{1/N_{\text{dim}}} \]  
\[ \Delta t_{\text{CFL}} = \frac{C_{\text{CFL}} \Delta t}{c_s + h |\nabla \mathbf{v}| + 1.2(\alpha_v c_s + \text{MIN}[\beta_v h |\nabla \mathbf{v}|])} \]  
The code employs a second-order Runge-Kutta-Fehlberg integrator (Fehlberg 1969) to evolve the hydrodynamics equations. Particles are assigned individual timesteps, where the timestep of an individual particle is the minimum value from several limiters: a conventional CFL condition (\(C_{\text{CFL}}=0.3\)):

\[ \Delta t_a = C_{\text{CFL}} \sqrt{\frac{h}{|\mathbf{a}|}} \]  

and a third requirement that changes in a particle’s velocity, acceleration and smoothing length do not exceed a given tolerance, full details are given in Bate, Bonnell & Price (1995).

2.3 GIZMO (Hopkins 2015) is a hydrodynamics code, based on the SPH code GADGET-3, designed to accommodate the benefits of both grid and particle based hydrodynamics schemes. The method is based on the works of Lanson & Vila (2008a,b) and Gaburov & Nitadori (2011), and has some common features with moving-mesh codes such as AREPO (Springel 2010). GIZMO uses a Lagrangian-like formulation, where the volume is discretised using a weighting function. The weighting function is similar to the kernel in SPH though in contrast, the kernel gradients play no role in the equations of motion as they do in SPH. The discretisation of the fluid is defined by an ensemble of particles that trace the motion of the cells. Shocks are captured with a Riemann solver, eliminating the need for artificial viscosity (similar to the approach of Godunov SPH codes). There are two new methods available in GIZMO (as well as two different versions of SPH with a number of viscosity switches); meshless finite-volume (MFV) and meshless finite-mass (MFM) methods. The methods differ in whether the particles/cells are allowed to experience a mass flux between their neighbours (MFV) or whether their masses are fixed (MFM). The methods appear to differ only slightly in the test problems shown in Hopkins (2015), especially relative to the differences seen when compared to pure grid or SPH methods.

The time integration scheme used in GIZMO is a second order leapfrog integrator very similar to AREPO and GADGET2 (Springel 2010) and is described in detail in the appendix G of Hopkins (2015). Local time-steps are used in all modes so that for each particle the timestep is

\[ \Delta t_{\text{CFL}} = 2C_{\text{CFL}} \frac{h}{|v_{\text{sig}}|} \]  

where \(h\) is the kernel smoothing length and \(|v_{\text{sig}}|\) is the signal velocity (Whitehurst 1995; Monaghan 1997; Hopkins 2015). For the GIZMO runs we use a CFL factor of \(C_{\text{CFL}}=0.1\), however note that the CFL factor is used differently by each of the codes and in particular the way that GIZMO and SPHNG employ this value is not directly comparable with the CFL factor used in RAMSES due to the different way in which resolution is defined for each framework. Time-steps in GIZMO are also limited to prevent spurious events of particle interpenetration when neighbouring particles have very different in time-steps (Saitoh & Makino 2009; Durier & Dalla Vecchia 2012; Hopkins et al. 2014). The time-step may be further restricted depending on the acceleration of particles, with \(\Delta t_a = (2\alpha_\epsilon g_{\text{grav}}/|\mathbf{a}|)^{1/2}\) after Power et al. (2003), where \(g_{\text{grav}}\) is the force softening length (4 pc) and \(\alpha_\epsilon=0.02\).

The MF and MFV GIZMO modes use a standard Harten-Lax-van Leer-Contact (HLLC) Riemann solver (Toro 1999; Miyoshi & Kusano 2005) as the default method. In the rare cases where the HLLC solver returns a non-physical result, the code automatically falls back on the slower but more accurate exact solver described in Toro (1997). Flux-limiting is used for the purposes of maintaining numerical stability but we direct the interested reader to appendix B of Hopkins (2015) for a complete and detailed description of how this is implemented in GIZMO.

One of the SPH modes of GIZMO is the traditional density-weighted approach (such as that of GADGET2 (Springel 2005), upon which GIZMO is partly based) which uses a standard artificial viscosity scheme with no additional measures to allow fluid mixing instabilities (e.g Mittal & Thomas 2001; Price 2008). While operating as an SPH code, a density estimation is required, the density is determined in the same way as for SPHNG (described in §2.2) but with some differences: i) the typical number of neighbours differs with GIZMO particles having \(\sim 32\) rather than \(\sim 58\) and ii) the smoothing length used to scale the kernel is simply \(h_i\) rather than the mean \(h\) of each particle pair.

Another method available in GIZMO is ‘PSPH’ (Saitoh & Makino 2013; Hopkins 2013), in which the equations of motion are rearranged to combat fluid mixing instabilities, and also includes artificial conductivity (Price 2008). In the traditional approach the pressure is calculated using the density estimate, for PSPH the pressure is instead determined from the neighbouring particles using kernel smoothing in the same way as with density.

2.4 Initial Conditions and External Gravitational Potential

We initialise our simulations as a uniform disc with a surface density of \(8 M_\odot\) pc\(^{-2}\) and an outer radius of 10 kpc. Gas is distributed vertically with a \(\text{sech}^2(z/H)\) profile, where the vertical scale-height, \(H=0.18\) kpc. For our RAMSES runs we
set a density floor of \(6.8 \times 10^{-32} \text{ g cm}^{-3}\). The gas is isothermal and has a temperature of 1000 K. We neglect self-gravity in order to investigate the gas response solely to the external gravitational potential. The gas is initially set up with circular orbits according to the underlying gravitational potential. The potential is intended to proxy a rotating stellar mass distribution and is given the logarithmic form from Binney and Tremaine (1987),

\[
\psi(r, z)_{\text{disc}} = \frac{1}{2} \nu_0^2 \log[r^2 + R_c^2 + (z/q_\phi)^2]
\]

which yields a flat rotation curve with \(v_0=220 \text{ km s}^{-1}\). The radial and vertical shape of the potential is set by \(R_c=1 \text{ kpc}\) and \(q_\phi=0.7\). To this potential we add a spiral perturbation of the form given by Cox & Gómez (2002)

\[
\psi(r, \phi, z) = -4\pi G \rho_0 \exp\left(-\frac{r-r_m}{R_s}\right) \sum_{n=1}^{3} \frac{C_n}{K_n D_n} \cos(n\gamma_\alpha) \left[\text{sech}\left(\frac{K_n z}{\beta_n}\right)\right]^{\beta_n}
\]

where

\[\gamma_\alpha = \gamma \int_0^t \left[\Omega_\phi - \frac{\dot{r}}{r \tan(\alpha)}\right] dt\]

\[K_n = \frac{n \gamma}{r \sin(\alpha)}\]

\[\beta_n = K_n H (1 + 0.4 K_n H)\]

\[D_n = \frac{1 + K_n H + 0.3 (K_n H)^2}{1 + 0.3 K_n H}\]

\[C_1 = 8\pi/3, \quad C_2 = 1/2, \quad C_3 = 8\pi/15\]

The parameters \(r_0=8 \text{ kpc}\), \(R_c=7 \text{ kpc}\) and \(H=0.18 \text{ kpc}\) set the scaling of the spiral perturbation in three dimensions, \(N=2\) is the number of spiral arms. The pitch angle is \(\alpha=15^\circ\) and the pattern speed is \(\Omega_\phi=2 \times 10^{-8} \text{ rad yr}^{-1}\). The strength of the spiral perturbation is \(\rho_0 = \rho_{\text{H}} n_{\text{H}}\) with \(n_{\text{H}}=1 \text{ atom cm}^{-3}\). The effective stellar mass of this potential is \(\sim 10^{11} \text{ M}_\odot\).

With these parameters the corotation radius is just beyond the edge of the disc at around 11 kpc. Thus the gas rotation speed within the entire disc exceeds that of the spiral perturbation and gas shocks at the trailing edge of the perturbation.

We have made runs using each code but without applying the spiral perturbation to determine if any significant features arise within a uniform disc. In this case we note some weak concentric rings; these are not related to spurious angular momentum transfer but instead are ripples resulting from the imperfect initial pressure equilibrium. We do not think these ripples have any impact on our results for two reasons: i) the density contrast of the rings is much weaker than that caused by the spiral potential at all radii and ii) we have repeated our baseline runs starting with the disk in vertical pressure equilibrium to reduce the impact of the ripples and see no difference in the results.

\[\text{Figure 1. Resolution vs. density for the different codes employed in this study. Magenta square symbols are RAMSES leaf cells and the resolution is the length of each cell. For SPHNG (red cross symbols) and GIZMO (blue triangular symbols) the resolution is taken as twice the smoothing length (see Hopkins (2015) for an in-depth discussion on the use of the kernel length in GIZMO). For clarity we show only 500 randomly selected resolution elements from each of the simulations.}\]

\[\text{2.5 Full Simulation List}\]

A full list of the simulations used in this work is shown in Table 1 with all the parameters that are varied. Four of the runs represent our ‘baseline’ models which reflect the choices made in most of the work on isolated galaxy simulations with external potentials: these are, RAMSES-1, SPH-1, GIZMO-MFM, and GIZMO-MFV. We selected these runs to represent the typical resolutions found in the literature, \(4 \times 10^9\) particles and 4 grid cells per Jeans’ length for particle and AMR runs respectively.\(^3\) We begin with these runs and later move on to discuss the effect of resolution on a given code.

Given the difficulty in directly comparing resolutions for grid codes with those in particle codes, we illustrate the comparative size of the simulation elements for our baseline simulations in Fig. 1, showing the length of grid cells (for a RAMSES run) and twice the smoothing length (for the GIZMO and SPHNG) versus density.\(^4\) Our goal here is to compare the simulation codes as they are used in the literature, but making sure that anything which can overtly affect the physics of

\(^3\) RAMSES-1 does not refine the grid to the maximum possible resolution, only reaching level 12 due to the high tolerance of its refinement criteria. All other RAMSES runs make use of the relevant maximum level.

\(^4\) Note that the smoothing length in GIZMO is calculated the same as for an SPH code, but is not used in the same fashion for smoothing the particle distribution.
Table 1. Overview of the simulation parameters. The top, middle and bottom sections describe the RAMSES, SPHNG, and GIZMO runs respectively. Column (1): simulation reference name; column (2): maximum refinement level and corresponding physical size of smallest grid cell in pc; column (3): the number of grid cells or particles at the end of the simulation (250 Myr); column (4): number of cells resolving the Jeans’ length; columns (5) and (6): refinement parameters based on density gradients (see Eq. 4); columns (7) and (8): grid cell in pc; column (9): Notes for each run, e.g. code type, solvers and viscosity schemes.

<table>
<thead>
<tr>
<th>name</th>
<th>ℓmax (pc)</th>
<th>Ngrid</th>
<th>NJ</th>
<th>Cρ</th>
<th>fp</th>
<th>αv</th>
<th>βv</th>
<th>notes</th>
</tr>
</thead>
<tbody>
<tr>
<td>RAMSES-E</td>
<td>14 (3.07)</td>
<td>1.808×10⁶</td>
<td>4</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>AMR, exact solver + MonCen slope limiter</td>
</tr>
<tr>
<td>RAMSES-1</td>
<td>14 (3.07)</td>
<td>9.971×10⁵</td>
<td>4</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>AMR, acoustic solver + MinMod slope limiter</td>
</tr>
<tr>
<td>RAMSES-2</td>
<td>14 (3.07)</td>
<td>5.362×10⁶</td>
<td>8</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>AMR, acoustic solver + MinMod slope limiter</td>
</tr>
<tr>
<td>RAMSES-3</td>
<td>14 (3.07)</td>
<td>2.613×10⁷</td>
<td>16</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>AMR, acoustic solver + MinMod slope limiter</td>
</tr>
<tr>
<td>RAMSES-4</td>
<td>15 (5.14)</td>
<td>2.637×10⁷</td>
<td>16</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>AMR, acoustic solver + MinMod slope limiter</td>
</tr>
<tr>
<td>RAMSES-5</td>
<td>12 (12.3)</td>
<td>3.037×10⁶</td>
<td>4</td>
<td>1.3</td>
<td>5</td>
<td>-</td>
<td>-</td>
<td>AMR, acoustic solver + MinMod slope limiter</td>
</tr>
<tr>
<td>RAMSES-6</td>
<td>13 (6.15)</td>
<td>5.156×10⁶</td>
<td>4</td>
<td>1.3</td>
<td>5</td>
<td>-</td>
<td>-</td>
<td>AMR, acoustic solver + MinMod slope limiter</td>
</tr>
<tr>
<td>RAMSES-7</td>
<td>14 (3.07)</td>
<td>7.676×10⁶</td>
<td>4</td>
<td>1.3</td>
<td>5</td>
<td>-</td>
<td>-</td>
<td>AMR, acoustic solver + MinMod slope limiter</td>
</tr>
<tr>
<td>RAMSES-8</td>
<td>14 (3.07)</td>
<td>2.356×10⁶</td>
<td>4</td>
<td>1.6</td>
<td>5</td>
<td>-</td>
<td>-</td>
<td>AMR, acoustic solver + MinMod slope limiter</td>
</tr>
<tr>
<td>RAMSES-9</td>
<td>14 (3.07)</td>
<td>6.651×10⁶</td>
<td>4</td>
<td>1.0</td>
<td>5</td>
<td>-</td>
<td>-</td>
<td>AMR, acoustic solver + MinMod slope limiter</td>
</tr>
<tr>
<td>SPH-1</td>
<td>-</td>
<td>4×10⁵</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>1</td>
<td>2</td>
<td>SPH + Monaghan &amp; Lattanzio (1985) viscosity</td>
</tr>
<tr>
<td>SPH-2</td>
<td>-</td>
<td>1×10⁵</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>1</td>
<td>2</td>
<td>SPH + Monaghan &amp; Lattanzio (1985) viscosity</td>
</tr>
<tr>
<td>SPH-3</td>
<td>-</td>
<td>8×10⁴</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>1</td>
<td>2</td>
<td>SPH + Monaghan &amp; Lattanzio (1985) viscosity</td>
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<td>-</td>
<td>4×10⁴</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>0.05</td>
<td>0.1</td>
<td>SPH + Monaghan &amp; Lattanzio (1985) viscosity</td>
</tr>
<tr>
<td>GIZMO-MFV</td>
<td>-</td>
<td>4×10⁶</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>mesh-less finite mass</td>
</tr>
<tr>
<td>GIZMO-MFV-2</td>
<td>-</td>
<td>4.593×10⁶</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>mesh-less finite volume</td>
</tr>
<tr>
<td>GIZMO-MFV-3</td>
<td>-</td>
<td>9.188×10⁶</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>mesh-less finite volume</td>
</tr>
<tr>
<td>GIZMO-SPH-NS</td>
<td>-</td>
<td>1.131×10⁶</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>mesh-less finite volume</td>
</tr>
<tr>
<td>GIZMO-SPH-B</td>
<td>-</td>
<td>4×10⁶</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>SPH + no viscosity switch (constant viscosity)</td>
</tr>
<tr>
<td>GIZMO-SPH-C&amp;D</td>
<td>-</td>
<td>4×10⁶</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>SPH + Balsara (1995) viscosity</td>
</tr>
<tr>
<td>GIZMO-PSPH</td>
<td>-</td>
<td>4×10⁶</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>SPH + Cullen &amp; Dehnen (2010) viscosity</td>
</tr>
</tbody>
</table>

3 RESULTS

All the simulations presented here start from an initially flat surface density distribution within the disc region. In the absence of self-gravity the ISM responds to the external potential and pressure forces leading to an increase in the density of arm regions over time, however we note that we do not form a steady state at any point in our simulations. Initially there are only two arms which form around the spiral perturbation. This is shortly followed by the development of second pair of arms between the existing ones. These new features branch from the original arms at ~5 kpc from the center. The appearance of secondary arm features are well documented as arising from the first ultraharmonic (4:1) resonance (Shu, Milione & Roberts 1973; Patsis, Grosbol & Hiotelis 1997; Chakraborti, Laughlin & Shu 2003). We show face-on surface density maps for the four baseline models in Fig. 2 in which the original arms and the bifurcating secondary arms are visible. Both features are most clear for RAMSES-I in the upper left panel.

A frequent criticism of grid codes is that angular momentum is not conserved. For the duration of the runs performed in this work (250 Myr) we calculate the loss of angular momentum in the AMR runs as ~3% and in the SPH and GIZMO runs at ~0.3%. We do not believe that angular momentum loss is a significant cause of the differences we find here, but it may play a more important role in simulations for which self-gravity results in the formation of small-scale eddies.

3.1 Overall structure of the disc with different numerical codes

The different codes tested here respond to the external potential in slightly different ways. Fig. 2 reveals significant differences between the RAMSES-I and SPH-1 runs, with less dense arms and far less fine structure between arms appearing in RAMSES-I. The SPHNG and GIZMO runs clearly show the presence of short inter-arm structures perpendicular to the arms, sometimes referred to as ‘spurs’ or ‘feathers’, in the inner part of the disc. These features have been seen in both grid, SPH and AREPO (Smith et al. 2014) simulations, and have been attributed to a number of potential causes including the Kelvin-Helmholz or wiggle instability (Wada & Koda 2004), feathering instability (Lee 2014) and orbit crossing in the spiral arms (Dobbs, Bonnell & Pringle 2006), although Kim & Ostriker (2006) suggest purely hydrodynamical instabilities disappear or are less evident in 3D. Although there is some slight indication of substructure in the inner part of the RAMSES-I disc, distinct spurs are not visible. The differences between the SPHNG and...
GIZMO runs are largely confined to the strength of the rings found within the central 3 kpc. The two GIZMO runs are not identical: (GIZMO-MFV has less coherent rings than GIZMO-MFM) but are very similar to one another when compared with the other two runs. The growth of rings in a non–self-gravitating gas disc is also found in grid-based simulations by Shetty & Ostriker (2006), in which leading spiral structures develop between the arms near the center of the disc.

Fig. 3 shows the mean density as a function of distance from the mid-plane for the four baseline simulations at the 250 Myr mark. In this comparison we also include a run, labelled as RAMSES-E, which is the same as RAMSES-1 in all respects except the choice of solver and slope limiter. The majority of our RAMSES simulations use quite a diffusive combination of the acoustic Riemann solver with a MinMod slope limiter, the RAMSES-E run uses the exact Riemann solver with the MonCen slope limiter. We use this run to demonstrate the result of using a less diffusive combination. RAMSES-1 does not reach the same mass concentration as the Lagrangian runs, having only approximately half the mid-plane density of the least concentrated Lagrangian run. The RAMSES-E run however has a vertical density profile with a slope that is not too dissimilar from the GIZMO runs.
Figure 3. Mean density as a function of distance from the disc plane. Models RAMSES-E, SPH-I, GIZMO-MFV, and GIZMO-MFM are shown as solid magenta, solid black, dashed red, light blue dot-dashed, and dark blue triple-dot-dashed lines respectively.

The two discs realised with GIZMO are more concentrated than the spgNG run despite having the same particle resolution.

The one-dimensional structure of the ISM can be examined in the form of a density probability distribution function (PDF) as plotted in Fig. 4. It indicates a consistent density peak for all the runs, but differences do appear in the distributions. The maximum density is curtailed at $10^{1.3} \, n_H \, cm^{-3}$ in RAMSES-I while the Lagrangian runs extend continuously to around $10^{1.7} \, n_H \, cm^{-3}$. The PDFs of both RAMSES runs extend down to very low densities in contrast with the Lagrangian simulations, simply because there are resolution elements in the RAMSES runs that are not represented by particles in the other codes. We next describe the evolution of the spiral arms over time and will discuss the difference between the codes further in Section 3.3.

### 3.2 Growth of Arm Densities

We now consider the surface density of the gas within two annuli, analysing the azimuthal profile at a number of snapshots throughout the runs. The time evolution of azimuthal variations in surface density for the SPH-I run is shown in Fig. 5 for annuli at $4 \pm 0.1$ kpc and $7 \pm 0.1$ kpc. We show this run as an example which reflects the typical behaviour of all the runs present here. The 4 kpc annulus (upper panel in Fig. 5) illustrates the increasing surface density of the arms until $\sim 150$ Myr, whereupon smaller variations in the peak surface density are present for the remainder of the simulation. The annulus at 7 kpc (lower panel in Fig. 5) shows the same initial increase but with larger variations in the peak arm surface density throughout the run as well as exhibiting the later formation of a secondary arm feature not seen at smaller radii. The effect of the delay between the development of the first and second set of arms is to create oscillations in the maximum surface density. The secondary arms grow downstream in the gas flow and as its density increases, the density of the earlier arm dwindles. This temporarily reduces the maximum surface density until the second arm becomes dominant and the maximum surface density increases again, now representing the secondary arms.

Oscillations in the density of the arms are also expected due to the abrupt activation of the spiral potential. In Woodward (1975) the steepening of spiral waves under the influence of an arm potential that grows over different timescales is examined, finding that there is an initial period of density oscillation which decays over time. Woodward (1975) also demonstrate that the more extreme and persistent oscillations occur in the runs where the spiral potential is activated over shorter timescales. A gradually introduced potential reduces the problem of these initial oscillations but for simplicity, and because it is not common practice to do so, we have not employed such a measure. A gradually introduced potential does occur naturally in simulations with a live stellar component without initial spiral structure.

Fig. 6 shows two annuli at $4 \pm 0.1$ kpc and $7 \pm 0.1$ kpc and the maximum surface density within each annulus as a function of time for each of our baseline models. Fig. 6 thus illustrates the growth of the density of the arms over time. We include only one of the GIZMO runs here (GIZMO-MFV) because its counterpart (GIZMO-MFM) is extremely similar, see Fig. 2. There are substantial differences in the surface density of the arms attained for each of the codes. In Fig. 6 we see that particularly at smaller radii the spgNG runs differ from the GIZMO runs in the timing and amplitude of the oscillations in arm surface density. Lastly we note the

Figure 4. Mass-weighted probability density function for the baseline simulations. Linestyles are the same as in Fig. 3.
considerable difference between \textit{RAMSES-1} and the sphNG and \textsc{gizmo} runs, with much lower densities found in the arm regions and lower amplitude oscillations, indeed these oscillations are virtually absent in the \textit{RAMSES-1} run. When one considers \textit{RAMSES-E} there is an interesting difference in the evolution at different radii. At 4 kpc the oscillations are now to some degree apparent and the arm surface density is only slightly lower than the runs produced with the other codes, however at 7 kpc \textit{RAMSES-E} is not very different to \textit{RAMSES-1} apart from a slight increase in the surface density.

3.3 Dependence of disc evolution on numerical code

Our comparisons of the baseline galaxy disc simulations highlight a number of differences between the codes, particularly the sphNG and \textsc{gizmo} runs compared with \textit{ramses}. The maximum density in the spiral arms, maximum density in the midplane and the degree of interarm structure including the secondary branches are greater with \textsc{gizmo} and sphNG, more so in the \textsc{gizmo} runs. We now consider which characteristics of the codes might lead to these differences. The most obvious possibilities are that the codes reach different effective resolutions, or the inclusion of artificial viscosity in sphNG. We discuss viscosity next, and describe resolution tests of the codes in Sections 3.4 and 3.5.

sphNG runs employ artificial viscosity to allow shock capturing, but artificial viscosity is not required in either \textit{ramses} or \textsc{gizmo} as both use a Riemann solver. For this reason we do not believe that the discrepancy between \textit{RAMSES-1} and the other runs in Fig. 6 is due to different viscosity schemes. Additionally we have run sphNG with reduced viscosity parameters, $\alpha_v=0.05$ and $\beta_v=0.1$. These viscosity parameters yield an arm surface density growth rate (without oscillations) similar to \textit{RAMSES-1} but shock capturing is compromised by reducing the artificial viscosity so harshly.

\textsc{gizmo} can also be operated as an SPH code with various viscosity switches. In order to further eliminate the influence of viscosity as the cause of differences between the inviscous
MFM and MFV methods of GIZMO and the artificially viscous sphNG we have run our model with a number of these modes (detailed in Table 1).

These models are compared with GIZMO-MFV and SPH-1 in Fig. 7. It is clear that the artificial viscosity scheme makes little difference and all GIZMO modes behave in roughly the same way. Therefore, the cause of the discrepancy found between the sphNG and GIZMO runs is not a basic difference in the SPH or the meshless Lagrangian methods of Hopkins (2015). The fact that, despite some minor differences, the evolution of the GIZMO SPH runs are far more similar to GIZMO-MFV than they are to SPH-1 means that it is likely that some aspect of the code is responsible for the offset between the two codes that is separate from the fundamental methodologies. We have not explored the codes in sufficient detail to offer a definitive explanation of this but note that, apart from the core hydrodynamics solver, all components of GIZMO are commons to both the MFV/MFM and SPH modes. The different results found with sphNG and GIZMO may therefore be due to differences in the neighbour finding process, the time-step criteria or time integration.

We note that the SPH kernel for GIZMO and sphNG is a cubic spline but that the smoothing length is defined differently with the smoothing lengths in GIZMO being smaller (see Fig. 1). Exploring this, we have rerun GIZMO-MFM and GIZMO-SPH-Český with the same particle smoothing lengths as our sphNG runs and find that it makes very little difference to the arm surface density and density PDF, however the vertical concentration is slightly reduced and therefore closer to the profile of SPH-1. We also observe a slight weakening of interarm structures when using larger smoothing lengths.

3.4 Resolution of sphNG and GIZMO Runs

In this section we consider how our two Lagrangian codes (sphNG and GIZMO) evolve differently with varying mass resolution. The evolution of the arm surface density for runs that initially have $1 \times 10^6$, $4 \times 10^6$, and $8 \times 10^6$ particles are shown in Fig. 8. The number of particles in sphNG and GIZMO-MFM runs is fixed, but the GIZMO-MFV runs allow particle splitting and increase the number of particles over the course of the simulation. The final number of particles for GIZMO-MFV-3, GIZMO-MFV, and GIZMO-MFV-2 are $1.131 \times 10^6$, $4.593 \times 10^6$, and $9.188 \times 10^6$ respectively.

For both of the Lagrangian codes, the evolution of the peak surface density is invariant with resolution, i.e. the rate of arm growth is unaffected. sphNG and GIZMO runs do not show much difference in the surface density of the galaxy arms as a function of resolution, but we do find differences between the codes themselves. Despite the codes presenting a similar time-averaged growth curve, the oscillations discussed in the previous section are offset in time (see Fig. 8).

Despite the invariance of the maximum surface density with resolution we do see that the vertical density profiles are steeper for runs with higher resolution, as shown in Fig. 9 and consequently the peak volume density is enhanced by resolution. As the particle resolution is increased (and the smoothing length shortens) the vertical density profiles is improved. The GIZMO runs exhibit steeper density profiles than do the sphNG, this is partially due to the different definition and use of the smoothing length/particle domain.

The mass-weighted density PDFs for our resolution comparison of Lagrangian runs are shown in Fig. 10. For both sphNG and GIZMO, increased resolution broadens the distribution and increases the fraction of gas at lower densities. GIZMO runs have a consistently narrower distribution in the high density peak but extend to lower densities than their sphNG counterparts. The maximum density achieved by all the runs is quite consistent with the sole exception of SPH-2 (the lowest resolution SPH run) which is truncated around 0.2 dex below the others.

3.5 Resolution of AMR runs

Resolution in AMR simulations is not a linear characteristic. We can change the minimum and maximum refinement levels or the parameters governing refinement. In this section we compare a number of approaches to varying the resolution within RAMSES. In addition to the previously shown run (RAMSES-1) which employs a Jeans’ length refinement criterion with the typical threshold $N_J=4$ we now test the effect of varying $N_J$ and $C_p$, which control grid refinement according to the local Jeans’ length and density gradients respectively. We also vary $\ell_{\text{max}}$, which is the upper limit on the grid level. The specific parameters used in each run are detailed in Table 1. We now discuss these three parameters that control refinement in turn using the mass-weighted density PDFs in Fig. 11 and the maximum surface density
Figure 8. Maximum arm surface density as a function of time for SPHNG (red/orange dashed lines) and GIZMO (blue dot-dashed lines) runs with different resolutions (initially with 1, 4 and 8 million particles). Darker colours represent greater resolutions.

Figure 9. The vertical density profiles of SPHNG (red/orange dashed lines) and GIZMO (blue dot-dashed lines) runs with different resolutions (initially with 1, 4 and 8 million particles). Darker colours representing greater resolutions.

Figure 10. Mass-weighted probability density function for Lagrangian simulations with different resolutions. Lines show SPHNG (red/orange dashed lines) and GIZMO (blue dot-dashed lines) runs that initially have 1, 4 and 8 million particles, darker colours represent greater resolutions.

Within annuli at $4\pm0.1$ kpc and $7\pm0.1$ kpc as a function of time in Fig. 12.

Firstly, we consider the effect of increasing $\ell_{\text{max}}$ which permits the code to refine the grid to higher levels. For this we direct the reader to the runs shown in the top panel of Fig. 11, i.e. we compare RAMSES-3 and RAMSES-4 which have $\ell_{\text{max}} = 14$ and 15 respectively but with all other refinement criteria the same. We also compare three runs that have an alternative set of refinement criteria to the previous two which use $\ell_{\text{max}} = 12, 13$ and 14 (RAMSES-5, RAMSES-6 and RAMSES-7), see Table 1 for details. We see here that increasing $\ell_{\text{max}}$ does not have an enhancing influence on these simulations, indeed RAMSES-3 and RAMSES-4 have identical density PDFs. We further examine the growth of the spiral arm surface density in the left-hand panels of Fig. 12. Again we note that varying $\ell_{\text{max}}$ makes almost no difference to the evolution of the galactic disc, although in this case there is a marginal reduction in arm surface density as $\ell_{\text{max}}$ increases for both the 4 and 7 kpc annuli in the case of RAMSES-5, RAMSES-6 and RAMSES-7.

We now consider whether increasing the number of grid cells that resolve the Jeans' length ($N_J$) has an influence on the simulations. RAMSES-1, RAMSES-2, and RAMSES-3 have $N_J = 4, 8$ and 16 respectively and are shown in the middle panel of Fig. 11. We note a marked increase in the maximum density achieved when $N_J$ takes greater values. In fact the densest end of the density distribution function for RAMSES-3 (for which $N_J=16$) is consistent with that found in the GIZMO and SPHNG runs. The central panels of Fig. 12 illustrate how increasing $N_J$ alters the growth of the spiral arms. We see that greater values of $N_J$ produce higher...
surface densities at 4 kpc. At 7 kpc we see simply that as \( N_J \) increases, the oscillations that are clearly present in sphNG and GIZMO runs (see Fig. 5) become more apparent.

One key strength of grid codes is the ability to resolve sharp density contrasts, achieved partially through the use of refinement criteria based on the local gradient in hydrodynamical variables. The final comparison we make varies the threshold that controls grid refinement based on the density gradients, \( C_\rho \). A lower value of \( C_\rho \) corresponds, in principle, to greater resolution. We compare RAMSES-8, RAMSES-7, and RAMSES-9 (in order of decreasing \( C_\rho \)) in the bottom panel of Fig. 11 and right-hand panel of Fig. 12. In this comparison we also include RAMSES-1 which effectively has an infinite threshold, i.e. no grid refinement is permitted based on density gradients. We recall here that RAMSES-1 does not make use of grid levels 13 and 14 and choosing a finite \( C_\rho \) allows the adaptive grid to make use of these levels.

In Fig. 11 (lower panel) we note a slight increase in the maximum density value as \( C_\rho \) decreases. The lowest \( C_\rho \) run (RAMSES-9) does not follow the trend of increasing peak density but this is likely linked to it having fewer grid cell than RAMSES-7 despite its lower \( C_\rho \). We therefore note that the higher density correlates more with the number of grid cells than with the gradient refinement threshold and likely does not reflect the ability of this refinement scheme to place cells in appropriate locations. Examining the inner annulus in Fig. 12 (upper right-hand panel) gives no clear indication of whether the value of \( C_\rho \) has any impact on the arm surface density. The outer annulus (lower right-hand panel in Fig. 12) however suggests that the lower threshold does enhance the oscillations of the arm surface density.

We find that, in the context of our galactic disc simulations, the grid structure is very sensitive to \( C_\rho \). We therefore find that decreasing \( C_\rho \) can lead to very little increase in resolution (because the grid is already refined to an extent by the Jeans’ length criteria), or it can refine the grid to such a degree that the simulation time increases disproportionately compared with other approaches. We find that despite the lower value of \( C_\rho \) in RAMSES-9 than in RAMSES-7 (which should mean that it refines grid cells more easily) we actually have fewer grid cells by the end of the simulation. For the first 150 Myr, the number of grid cells found in RAMSES-9 is much higher than in RAMSES-7, consistent with its lower refinement threshold, but it then declines gradually to the value found in Table 1.

Surface density maps for our baseline RAMSES run and the two resolution tests that are most similar to the Lagrangian runs (RAMSES-3 and RAMSES-9) are compared in Fig. 13. Both these higher resolution runs have sharper arms but also exhibit more inter-arm structure. RAMSES-3, which resolves the Jeans’ length with 16 grid cells, presents hints of the rings seen in the Lagrangian runs and secondary arms that extend further beyond the branching point (as seen in the Lagrangian runs in Fig. 2). Similar structures are seen in RAMSES-9, although more feathers are seen near the center and the secondary arms are not as extended. The emergence of these structures in the higher resolution runs hints that concordance with the Lagrangian runs is closer.

![Figure 11](http://mnras.oxfordjournals.org/)

**Figure 11.** Mass-weighted probability density functions for RAMSES runs with different refinement schemes. In the top panel we compare runs with differing values of \( \ell_{\text{max}} \). The middle panel compares different \( N_J \) values. The lower panel shows runs with different density gradient refinement thresholds (\( C_\rho \)). Darker colours represent greater resolutions: see Table 1 for details of the parameters used in each run.

4 DISCUSSION AND CONCLUSIONS

We have examined how an isothermal gas disc evolves under the influence of an external spiral potential when realised with different hydrodynamical methods (RAMSES, SPHNG, and GIZMO) and as a function of resolution. With similar resolutions to those found in the literature (and using the ‘acoustic’ solver with a MinMod slope limiter) we find that our AMR code, RAMSES, generates a weaker density contrast between the arm and inter-arm region, less steep vertical profiles and lower arm surface densities than we see when using the Lagrangian codes (SPHNG and GIZMO) as well as less inter-arm structure. When additional refinement measures are used, RAMSES generates results in better agreement with the other codes. If a less diffusive set up is used (i.e. an
Figure 12. Maximum arm surface density as a function of time for RAMSES runs with various refinement schemes. The upper panels illustrate the evolution of the spiral arms at 4±0.1 kpc and the lower panels at 7±0.1 kpc. The left-hand panels compare runs with differing values of $\ell_{\text{max}}$. The middle panels show runs with different $N_J$. The right-hand panels show runs with different density gradient refinement thresholds ($C_\rho$). Darker colours represent greater resolutions: see Table 1 for details of the parameters used in each run.

Figure 13. Surface density maps for our baseline RAMSES run (RAMSES-1) and for the two highest resolution runs using $N_J$ and $C_\rho$ criteria (RAMSES-3, and RAMSES-9 respectively) after 250 Myr have passed. Black circles indicate the location of the annuli at 4±0.1 kpc and 7±0.1 kpc used in our analysis. The most significant difference between these runs is the concentration of the arms and the presence of more inter-arm structure in the higher resolution runs.

'exact' solver with a MonCen slope limiter) then a measure of similarity is also achieved in the resolution of physical structures, vertical density profile, and the density PDF. The growth of spiral arm surface density is also enhanced to a similar level as seen in the Lagrangian codes but only at smaller radii: arm surface density is still relatively low at greater radii. In all codes, we also see oscillations in the peak arm surface density, which appear to be associated with the development of secondary arms, but the oscillations tend to be very weak with the baseline RAMSES model. The GIZMO and SPHNG codes also display small differences, namely that GIZMO produces the highest densities and surface densities.
To test why the differences occurred between the codes, we considered the resolution and viscosity. We found resolution had little effect on the Lagrangian runs, except for an increasing steepness of the vertical density profiles with resolution. Whilst artificial viscosity could potentially affect the results with the sph\textsc{ng} code, our fiducial gizmo runs do not include artificial viscosity, so this cannot explain the discrepancies between gizmo and \textsc{ramses}. Furthermore, using gizmo in ‘SPH mode’ (with constant, Balsara (1995), and Cullen & Dehnen (2010) viscosity schemes and with pressure-entropy based SPH) we find very little difference between those runs and the equivalent MFM and MFV gizmo modes (without viscosity). By reducing the artificial viscosity by a factor of 20, the sph\textsc{ng} models produced more similar results to the baseline \textsc{ramses} run, however this dramatic change in viscosity greatly reduces the ability of sph\textsc{ng} to accurately capture shocks. Thus we believe that the differences between \textsc{ramses} and the sph\textsc{ng}/gizmo runs, and between sph\textsc{ng} and gizmo are not due to viscosity. Differences in the choice of smoothing length and the functional form of the kernel partly account for the differences between our sph\textsc{ng} and gizmo runs.

We also investigated varying the resolution in the ramses code using a number of approaches. We first used the most intuitive approach, increasing the maximum refinement level \(N_{\text{max}}\), but surprisingly this made almost no difference. The reason for this was because the code was simply not saturating the maximum refinement level. We stress here though that our simulations are isothermal and do not include self-gravity, gas cooling, or any sources of forcing other than our external potential. These processes could drive the density up and force the grid to refine even without changing the refinement criteria, though it is not clear that this would improve the modelling of spiral arms more generally or, for example, the initial development of Jeans’ instabilities.

We secondly tried using refinement criteria based on density gradients (the kind conventionally employed in idealised code comparison tests), which we might expect would lead to greater refinement and better agreement with the arm densities seen with the other codes. However, we find that the parameters controlling such refinement schemes are not trivial to select \textit{a priori}. We tend to find only slight differences in the density PDF and, observe a small increase in the scale of oscillations in the arm surface density over time. Our model \textsc{ramses-9} shows best agreement with the other codes, but we note that in this run we were refining a significant fraction of the disc, which leads to an large increase in computing time due to refining regions unnecessarily. Again, we note that these results are true for our particular model choices and hydrodynamical gradient criteria may be of benefit under other frameworks, e.g. when supernovae blastwaves are included.

Thirdly we examined increasing refinement by increasing the number of grid cells resolving the Jeans’ length, \(N_{J}\). We find that the greatest, and most reliable, improvement (in the sense of providing more concordance with other methods) in the density PDF, vertical density profile and evolution of the surface density is found by increasing \(N_{J}\). By changing \(N_{J}\) from 4 to 16 we find greater similarities between our AMR and Lagrangian tests. In addition to the spiral arms, the resolution of the inter-arm regions is also important, one reason being that dense structures leaving the refined region should be preserved. To some extent our highest resolution \textsc{ramses} runs do preserve the inter-arm structures found in the Lagrangian runs. Although there are still differences between \textsc{ramses} and the other codes, this finding is in line with other comparison tests that show more similar results tend to be attained with grid and particle-based codes (Tasker et al. 2008; Kitsionas et al. 2009; Price & Federrath 2010) if the resolution is comparable or greater in the grid code. Our results also seem to concur with the idea in Price & Federrath (2010) that grid-based codes tend to be less effective at resolving denser regions of simulations.

Our findings that the density in spiral arms differs according to different numerical codes (we stress that here we mean ‘code’ and not ‘method’), and further that the growth of the maximum density differs, may have implications for studying star formation in spiral arms. The different densities may impact the timescales for gravitational collapse, properties and number of molecular clouds, and the rate and efficiency of star formation. This may be even more relevant for simulations with transient spiral arms, whereby the arms come and go with time, and the time for gas to accumulate into dense structures in the spiral arms may be fundamentally limited. We have demonstrated that by refining adaptive grid simulations further than is usual, similar arm density growth rates are achieved as with Lagrangian codes. We note that a quasi-Lagrangian refinement scheme could also be applied in the case where transient spiral arms arise due to a live stellar disc, but we do not test this idea in this work. We have also not examined how refinement changes once self-gravity is included, but likewise leave this to future studies.

To conclude, our key result is that caution is required when justifying the use of four grid cells per Jeans length as a resolution scheme with AMR (or static grid) simulations. This condition is necessary but may not be sufficient depending on the hydrodynamical solver. For the simulations performed here, the Jeans length needs be resolved by at least 16 grid cells to achieve a similar result to that found with Lagrangian codes. The authors are aware of only one study in the field of isolated galaxy simulations where the refinement criteria is set to a higher value, Petit et al. (2015), wherein 32 cells are used (note the authors also applied a quasi-Lagrangian refinement scheme). We find some dependence on the solver (in particular using a less diffusive solver also produces better resolution), and in general the criteria for the resolution may depend on the exact nature of the code. But given that more diffusive set ups are likely frequently used, we believe that this result is an important one. We find that increasing \(N_{J}\) appears to be the most effective and simplest means of increasing the resolution in regions of interest in a galactic disc, in particular to study processes such as spiral shocks and molecular cloud formation.

5 ACKNOWLEDGMENTS

CGF, CLD and LK acknowledges funding from the European Research Council for the FP7 ERC starting grant project LOCALSTAR. We thank the anonymous referee for a very constructive report. This work used the DiRAC Complexity system, operated by the University of Leicester IT
Services, which forms part of the STFC DiRAC HPC Facility (www.dirac.ac.uk). This equipment is funded by BIS National E-Infrastructure capital grant ST/K000373/1 and STFC DiRAC Operations grant ST/K0003259/1. DiRAC is part of the National E-Infrastructure. Calculations were also performed on Cray XC30 at the Center for Computational Astrophysics, National Astronomical Observatory of Japan.

REFERENCES

Balsara D. S., 1995, Journal of Computational Physics, 121, 357
Berger M. J., Colella P., 1989, Journal of Computational Physics, 82, 64
Dobbs C. L. et al., 2014, Protostars and Planets VI, 3
Fehlberg E., 1969, NASA technical report, R-315
Inutsuka S.-I., 2002, Journal of Computational Physics, 179, 238
Miyoshi T., Kusano K., 2005, Journal of Computational Physics, 208, 315

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