An improved SPH scheme for cosmological simulations

A. M. Beck, 1,2 G. Murante, 3 A. Arth, 1,4 R.-S. Remus, 1 A. F. Teklu, 1,5 J. M. F. Donnert, 6,7 S. Planelles, 3,8 M. C. Beck, 9 P. Förster, 1 M. Imgrund, 1 K. Dolag 1,2 and S. Borgani 3,8,10

1 University Observatory Munich, Scheinerstr. 1, D-81679 Munich, Germany
2 Max Planck Institute for Astrophysics, Karl-Schwarzschild-Str. 1, D-85741 Garching, Germany
3 INAF - Osservatorio Astronomico di Trieste, via Tiepolo 11, I-34131 Trieste, Italy
4 Max Planck Institute for Extraterrestrial Physics, Giessenbachstr. 1, D-85748 Garching, Germany
5 Excellence Cluster Universe, Boltzmannstr. 2, D-85748 Garching, Germany
6 INAF - Instituto di Radioastronomia, via P. Gobetti 101, I-40129 Bologna, Italy
7 Leiden Observatory, Leiden University, PO Box 9513, NL-2300 Leiden, the Netherlands
8 Department of Physics, University of Trieste, via Tiepolo 11, I-34131 Trieste, Italy
9 Department of Physics, University of Konstanz, Universitätsstr. 10, D-78457 Konstanz, Germany
10 INFN - Sezione di Trieste, via Valerio 2, I-34127 Trieste, Italy

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ABSTRACT

We present an implementation of smoothed particle hydrodynamics (SPH) with improved accuracy for simulations of galaxies and the large-scale structure. In particular, we implement and test a vast majority of SPH improvement in the developer version of GADGET-3. We use the Wendland kernel functions, a particle wake-up time-step limiting mechanism and a time-dependent scheme for artificial viscosity including high-order gradient computation and shear flow limiter. Additionally, we include a novel prescription for time-dependent artificial conduction, which corrects for gravitationally induced pressure gradients and improves the SPH performance in capturing the development of gas-dynamical instabilities. We extensively test our new implementation in a wide range of hydrodynamical standard tests including weak and strong shocks as well as shear flows, turbulent spectra, gas mixing, hydrostatic equilibria and self-gravitating gas clouds. We jointly employ all modifications; however, when necessary we study the performance of individual code modules. We approximate hydrodynamical states more accurately and with significantly less noise than standard GADGET-SPH. Furthermore, the new implementation promotes the mixing of entropy between different fluid phases, also within cosmological simulations. Finally, we study the performance of the hydrodynamical solver in the context of radiative galaxy formation and non-radiative galaxy cluster formation. We find galactic discs to be colder and more extended and galaxy clusters showing entropy cores instead of steadily declining entropy profiles. In summary, we demonstrate that our improved SPH implementation overcomes most of the undesirable limitations of standard GADGET-SPH, thus becoming the core of an efficient code for large cosmological simulations.

Key words: hydrodynamics – methods: numerical.

1 INTRODUCTION

Smoothed particle hydrodynamics (SPH) is a commonly employed numerical method in astrophysics. It solves the fluid equations (Landau & Lifshitz 1959) in a Lagrangian mass-discretized fashion, which ensures Galilean invariance and conservation of mass, momentum, angular momentum, energy and entropy. It was pioneered by Gingold & Monaghan (1977) and Lucy (1977) and has since then become one of the cornerstones of computational astrophysics. The discretization of mass automatically adapts spatial resolution by removing the constraint of handling geometry explicitly. It also easily couples to N-Body schemes for calculation of gravitational forces (Hernquist & Katz 1989). An excessive amount of papers and literature about SPH has been produced over the past decades. We point out the latest reviews by Rosswog (2009), Springel (2010b), Monaghan (2012) and Price (2012a) for the basic concepts and e.g. Ritchie & Thomas (2001) for an extension to multiphase fluids and...
Rosswog (2015) for a special-relativistic adaption. As every numerical method, SPH comes with its own set of benefits and pitfalls, which we address in this paper.

The inability of traditional SPH methods to treat contact discontinuities and to mix different fluid phases is a long-standing problem (Agertz et al. 2007; Wadsley, Veeravalli & Couchman 2008). It leads to a completely numerical spurious surface tension at the discontinuities preventing particle movements. Consequently, it results in a failure of these formulations of SPH to resolve fluid instabilities such as the Kelvin–Helmholtz or Rayleigh–Taylor instabilities (see e.g. Junk et al. 2010; Valcke et al. 2010; McNally, Lyra & Passy 2012; Puri & Ramachandran 2014). In applications to cosmic structure formation it causes entropy profiles to diverge towards the centres of dark matter haloes, at variance with Eulerian codes that predict entropy plateaus to build up (Frenk et al. 1999; Wadsley et al. 2008; Planelles & Quilis 2009; Vazza 2011; Power, Read & Hobbs 2014; Biffi & Valdarnini 2015; Rasia et al. 2015). This difference is due to the lack of mixing in simple SPH, which makes low-entropy gas in merging substructures sink towards the centre of the main structure. Many modifications have been proposed to overcome this problem. For example, Wadsley et al. (2008) propose a mixing solution, which resolves the differences in the entropy profiles of dark matter haloes between Eulerian and SPH codes (Frenk et al. 1999). Further cosmological applications have been performed by Shen, Wadsley & Stinson (2010). First, the equation of motion (EoM) can be re-formulated from a standard ‘density’ approach into a ‘pressure’ based approach (Hopkins 2013; Saitoh & Makino 2013). While the ‘pressure’ formulation correctly treats contact discontinuities, it leads to increased noise at strong shocks. Secondly, considerable effort has been made to unite grid-based solvers for the fluid equations with the Lagrangian nature of SPH. Eulerian Godunov methods (see e.g. Cha, Inutsuka & Nayakshin 2010; Springel 2010a; Murante et al. 2011) and their coupling to Lagrangian methods is a promising alternative. Connecting a Lagrangian moving-mesh with grid-based solvers (Springel 2010a) or mesh-free approaches (Hopkins 2015; Hopkins & Raives 2015) represent more advanced approaches. Thirdly, artificial conduction (AC) of internal energy can be employed to overcome the mixing problem. Most modern SPH codes include AC of some sort to diffuse entropy (Read & Hayfield 2012) or energy (Price 2008) across particles. The use of AC has to be taken carefully and it is only desirable at contact discontinuities in traditional ‘density’ SPH and at shocks in modern ‘pressure’ SPH. The application of AC in other regions can have catastrophic impact on the fluid dynamics and can smear out gravitationally established pressure gradients, thus leading to totally numerically induced transport of internal energy (Valdarnini 2012).

Next, traditional SPH has difficulties treating subsonic turbulence as it experiences a high effective viscosity, which limits the inertial range (see e.g. Bauer & Springel 2012). Thus, traditional SPH cannot achieve high Reynolds’ numbers compared to, for example, Eulerian methods. This high effective viscosity is a function of resolution, but no general solution has yet been proposed to resolve this issue in general. For the correct capturing of shocks numerically motivated artificial viscosity (AV) is commonly employed. It smooths the particle velocity distribution and gives order to the fluid sampling. However, AV is only desired at the shock and the fluid should be inviscid otherwise. Too much AV smears out physical motions and damps subsonic and turbulent motions in isolated tests (Bauer & Springel 2012) as well as in cosmological simulations (Dolag et al. 2005). Therefore, several different implementations of AV reduction are proposed (Morris & Monaghan 1997; Cullen & Dehnen 2010). They are all based around a proper shock detection method and a time-dependent viscosity decay scheme. Application of such advanced schemes give better results in the description of fluid motions (Dolag et al. 2005; Price 2012b).

Finally, it might seem easy to simply reduce the quantitative errors by increasing the number of neighbours, which contribute to the local density and force estimators. However, the standard weighting functions of SPH respond differently to an increase of smoothing neighbours and can possibly become unstable to the pairing instability (Schuessler & Schmitt 1981; Price 2012a). Therefore, recently, alternative kernel functions immune against this instability are proposed for better fluid sampling and convergence (Read, Hayfield & Agertz 2010; Dehnen & Aly 2012). The advantage of flexible geometry of SPH comes with difficulties in creating well-defined initial conditions or sampling analytical profiles, where we use either glass set-ups (White 1996) or Weighted Voronoi-Tesselations (Diehl et al. 2012).

To overcome the named disadvantages, we implement a large set of improvements for SPH into the developer version of the cosmological N-Body/SPH simulation code GADGET-3 (Springel, Yoshida & White 2001; Springel 2005). We include a time-step limiter for strong shocks, a time-dependent viscosity scheme for subsonic turbulence, a high-order gradient estimator and shear flow limiter for shearing motions, an improved kernel function for convergence and a time-dependent artificial conduction scheme to promote fluid mixing. We discuss the accuracy and the performance of our new scheme in hydrodynamical standard test problems, within quiet and violent environments as well as in idealized simulations of galaxy and galaxy cluster formation, in which our new scheme is applied to reasonable astrophysical problems.

The paper is organized as follows. The improved implementation of hydrodynamics is presented in Section 2. In Section 3, we validate our SPH algorithms in a set of hydrodynamical standard tests and we proceed to standard tests with gravity in Section 4. We continue in Section 5 with idealized applications to the evolution of an isolated disc galaxy and a forming galaxy cluster. We summarize our developments and code performance in Section 6.

2 A NEW SPH IMPLEMENTATION

We start with a presentation of the main equations corresponding to a ‘standard’ and our ‘new’ formalism of GADGET-SPH. The formalism of SPH is already well described by a large number of reviews (see e.g. Price 2012a). We refer to the ‘standard’ version of SPH as the implementation within the GADGET-3 code without our modifications. We point out our modifications and discuss the kernel function, the EoM, the particle wake-up scheme and the time-dependent AV and AC.

2.1 Original code platform

We implement our SPH modifications into the developer version of the cosmological N-Body/SPH code GADGET-3 (Springel et al. 2001; Springel 2005). We evolve entropy as the thermodynamical variable (Springel & Hernquist 2002) and use the prescriptions for radiative cooling, supernova feedback and star formation following Springel & Hernquist (2003). In the following sections, we compare two different SPH schemes (see also Table 1), which are distinguished as follows. The ‘standard’ implementation corresponds to the developer version of GADGET-3 without our modifications (Springel 2005). The ‘new’ implementation includes all the SPH improvements presented in this section. In principle, we always use the
entire new scheme and we also employ the same set of all numerical parameters throughout our entire simulation test suite. Thus, unless otherwise stated, we do not tune individual standard tests or astrophysical applications. However, if necessary we sometimes switch off some of the modifications to analyse their individual and isolated impact on several of the test problems.

### 2.2 Kernel functions and density estimate

Foremost, there is the question in a Lagrangian method how to derive fluid field quantities from a given set of point masses. In particular, the estimation of the gas density is crucial as many further equations rely on it. We employ the standard estimator of SPH and calculate the density \( \rho(x_i) = \rho_j \) of an individual particle \( i \) at the position \( x_i \), by summing the contributions of \( N \) neighbouring particles \( j \) within a smoothing radius \( h(x_i) \) at a distance \( x_{ij} \) in a mass-weighted \( (m_i) \) and distance-weighted \( (W_{ij}(x_{ij}, h_i)) \) fashion

\[
\rho_i = \sum_j m_j W_{ij}(x_{ij}, h_i). \tag{1}
\]

Simultaneously, the smoothing length \( h_i \) is a function of density

\[
h(x_i) = \eta \left( \frac{m_i}{\rho_i} \right)^{1/3}, \tag{2}
\]

where \( \eta \) defines the ratio of smoothing length to the mean distance between particles. Equations (1) and (2) roughly ensure constant mass resolution throughout the simulation and have to be solved in parallel. This mimics the evolution of spheres of the same mass \( 4/3\pi h_i^3 \rho_i = N_i m_i \) but with varying number of neighbours. The number of neighbours varies across space and time with an increase or decrease of smoothing length and local quality of fluid sampling by the point masses. The weighting function is commonly chosen to decrease monotonically with distance, yield smooth derivatives, is symmetric with respect to \( x_{ij} = x_{ji} \) and has a flat central portion. A historical choice (Monaghan & Lattanzio 1985) of kernel function

\[
W_{ij}(x_{ij}, h_i) = w(q)/h_i^3, \tag{3}
\]

is the cubic B-spline function with \( q = x_{ij}/h_i \) and

\[
w(q) = \frac{8}{\pi} \begin{cases} 1 - 6q^2 + 6q^3 & 0 \leq q \leq \frac{1}{2} \\ 2(1 - q)^3 & \frac{1}{2} \leq q \leq 1 \\ 0 & 1 \leq q \end{cases}, \tag{4}
\]

which we commonly employ with a choice of 64 neighbours in three dimensions. However, this traditional kernel function is subject to the pairing (or clumping) instability when the number of neighbours is too large (see Price 2012a). An alternative choice to achieve better numerical convergence is necessary and an entire new family of kernels is needed. In a kernel stability analysis Dehnen & Aly (2012) show that the Wendland kernel functions are a much better choice. We choose the Wendland \( C^4 \) (WC4) kernel with 200 neighbours in three dimensions as our smoothing function without the pairing instability problem. The functional form of the \( C^4 \) is given by

\[
w(q) = \frac{495}{32\pi}(1 - q)^6 \left( 1 + 6q + \frac{35}{3} q^2 \right). \tag{5}
\]

For values of \( q > 1 \), it is set to \( w(q) = 0 \). The Wendland functions require similar computational effort as the cubic spline kernel but nevertheless, the total computational time increases due to larger number of neighbours. Therefore, we do not employ the higher order \( C^6 \) functions because of the required 295 neighbours. In summary, the total computational cost of the density and hydrodynamical force calculation increases by a factor of about 2 in comparison to a cubic spline with 64 neighbours. However, a better estimate of the kernel will result in a more accurate density estimate and improved gradient estimators. These estimators are the cornerstones of the SPH formalism and determine the accuracy and convergence rate in all our test simulations.

### 2.3 Equation of motion

The EoM for a system of point masses are derived (see e.g. Price 2012a) from a discretized version of the fluid Lagrangian

\[
L = \sum_i m_i \left[ \frac{1}{2} v_i^2 - u \right], \tag{6}
\]

where \( v \) denotes velocities and \( u \) internal energy of individual particles. The Lagrangian nature of SPH, when complemented with a symplectic time integration scheme, automatically conserves mass, momentum, angular momentum, energy and entropy. We use the standard kick-drift-kick Leapfrog time integration of \textsc{gadget} (Springel 2005). The EoM then follows from the principle of least action, where the spatial derivative of internal energy comes (if constant entropy is assumed) from the first law of thermodynamics \( dU = -PdV \). We choose a volume element depending on density \( (V = m_i/p) \) and an adiabatic equation of state for the pressure \( P = A\rho^\gamma \), which is defined individually for every particle. We integrate

\[
\left\{ \begin{array}{ll}
1 - 6q^2 + 6q^3 & 0 \leq q \leq \frac{1}{2} \\
2(1 - q)^3 & \frac{1}{2} \leq q \leq 1 \\
0 & 1 \leq q
\end{array} \right.
\]

\[
w(q) = \frac{8}{\pi} \begin{cases} 1 - 6q^2 + 6q^3 & 0 \leq q \leq \frac{1}{2} \\
2(1 - q)^3 & \frac{1}{2} \leq q \leq 1 \\
0 & 1 \leq q
\end{cases}
\]
entropy $A$ as the thermodynamical variable of choice and thus employ what is commonly called ‘density-entropy’ SPH.

The EoM in the ‘density-entropy’ (for a derivation see Springel & Hernquist 2002) for the hydrodynamical force part of an individual particle reads

$$\frac{dv_i}{dt}_\text{hyd} = -\sum_j m_j \left[ f_{ij}^\text{co} \frac{P_i}{\rho_i^2} \nabla W_j(h_i) + f_{ij}^\text{co} \frac{P_j}{\rho_j^2} \nabla W_i(h_j) \right],$$

(7)

where the factor $f^\text{co}$ is a correction factor, which accounts for the mutual co-dependence of smoothing length $h(\rho)$ and density $\rho(h)$ and their corresponding derivatives. Its functional form is given by

$$f_{ij}^\text{co} = \left[ 1 + \frac{h_i}{3\rho_i} \frac{\partial \rho_j}{\partial h_j} \right]^{-1}.$$  

(8)

Equation (7) leads to a non-vanishing force at contact discontinuities even when pressure is constant. This is the artificial ‘surface tension’ of SPH, which suppresses particle movement across contact discontinuities. In the following sections, we present our equations in notation of internal energy $u$, which is related to the entropic function $A = (\gamma - 1)u/\rho^\gamma - 1$.

### 2.4 Artificial viscosity

#### 2.4.1 Smoothing of jumps

By construction, SPH solves the ideal Euler equation and no dissipative terms are included but those are necessary to describe discontinuities correctly. In highly dynamical regions (e.g. in shocks) fast particles commonly penetrate into regions of resting particles causing unwanted particle disorder and oscillations in the sampling of the fluid. However, SPH already contains an intrinsic remeshing force but to re-establish particle order and capture shocks properly an additional dissipative term in velocity is needed. This AV aims to remove post-shock oscillations and noise and helps to smooth the velocity field (see Monaghan & Gingold 1983). We include AV in an energy conserving way with a contribution to the EoM of the form

$$\frac{dv_i}{dt}_\text{visc} = \frac{1}{2} \sum_j m_j \left( v_i - v_j \right)^2 \alpha_{ij}^\text{loc} f_{ij}^\text{shear} v_{ij}^\text{sig,v} \mathbf{T}_{ij},$$

(9)

and with a contribution to the energy equation of the form

$$\frac{du_i}{dt}_\text{visc} = -\frac{1}{2} \sum_j m_j \left( v_i - v_j \right)^2 \alpha_{ij}^\text{loc} f_{ij}^\text{shear} v_{ij}^\text{sig,v} \mathbf{T}_{ij},$$

(10)

where the symmetrized variables represent $\rho_{ij} = (\rho_i + \rho_j)/2$ for the density, $\alpha_{ij} = (\alpha_{ij}^\text{loc} + \alpha_{ij}^\text{glob})/2$ as a numerical coefficient to include AV (see below) and $f_{ij}^\text{shear} = (f_{ij}^\text{shear} + f_{ji}^\text{shear})/2$ as the Balsara (1995) shear flow limiter (see Section 2.3.2 below), which aims to ensure the application of AV only in strong shocks (high-velocity divergence) and not in rotating or shearing flows (high-velocity curl). Furthermore, in the above equation $\mathbf{T}_{ij} = (F_i(h_i) + F_j(h_j))/2$ is the symmetrized scalar part of the kernel gradient terms $\nabla^\text{s} W_j(h_i) = F_{ij} r_{ij}/r_{ij}$, which are used to linearly interpolate the second-order Laplacian derivative in the velocity field diffusion equation. The pairwise signal velocity $v_{ij}^\text{sig,v}$ (first introduced by Monaghan 1997, and already used in GADGET-2) determines the strength of AV and directly includes a quantitative measure of particle disorder

$$v_{ij}^\text{sig,v} = c_i^v + c_j^v - \beta \mu_{ij},$$

(11)

where $c^v$ is the sound speed of the particles and $\mu_{ij} = v_{ij} \cdot x_{ij}/x_{ij}$ with a commonly chosen pre-factor of $\beta = 3$. AV is only applied between approaching pairs of particles (i.e. $\mu_{ij} < 0$) and otherwise switched off. The local signal velocity $v_{ij}^\text{sig,v}$ (also used by the time-step criterion, see Section 2.6) represents the maximum value of $v_{ij}^\text{sig,v}$ between all particle pairs $ij$ within the entire smoothing sphere of particle $i$.

The calculation of the viscosity coefficient $\alpha_{ij}^\text{loc}$ is based on an approach developed by Cullen & Dehnen (2010) but modified for more efficient computation as follows. The presence of a shock is indicated via computation of velocity divergence contributions across the entire smoothing length by

$$R_i = \frac{1}{\rho_i} \sum_j \text{sign}(\nabla \cdot v)_j m_j W_i,$$

(12)

where a shock corresponds to $R_i \approx -1$. In principle, an accurate calculation of $R_i$ for every particle requires the previous computation of $(\nabla \cdot v)$, for every particle. Therefore, an extra SPH summation loop added between the calculation of density (where velocity divergence can also be calculated) and hydro forces would be necessary. For computational reasons, we use the velocity divergence calculated in the previous time-step. Furthermore, a convergent flow is also indicated by a high-velocity divergence but that condition does not distinguish between pre-shock and post-shock regions. Therefore, we employ the time derivative of velocity divergence to determine a directional shock indicator

$$A_i = \xi_i \max(0, -|\nabla \cdot v|),$$

(13)

which is able to distinguish between pre-shock and post-shock regions. We calculate $(\nabla \cdot v)$, via interpolation between the current and the previous time-step (as suggested by Cullen & Dehnen 2010) in the time interval $\Delta \tau_i$.

Subsequently, we use the shock indicator $R_i$ to determine the ratio $\xi_i$ of strength of shock and shear of strength in quadratic form via

$$\xi_i = \frac{2(1 - R_i)^2 |\nabla \cdot v|_i^2}{2(1 - R_i)^2 |\nabla \cdot v|_i^2 + |\nabla \times v|_i^2},$$

(14)

which is proposed by Cullen & Dehnen (2010) as an additional limiting factor for AV in equation (13) and was experimentally determined. Now, for every particle we can define and set the target value $\alpha_{ij}^\text{loc,v}$ of AV with the help of the directional shock indicator to

$$\alpha_{ij}^\text{loc,v} = \alpha_{\text{max}} \frac{h_i^2 A_i}{h_i^2 A_i + (v_{ij}^\text{sig,v})^2},$$

(15)

In the case, where the viscosity coefficient $\alpha_{ij}^\text{loc}$ is smaller than $\alpha_{ij}^\text{loc,v}$, we set the coefficient to $\alpha_{ij}^\text{loc,v}$. Otherwise, we let it decay with time according to

$$\dot{\alpha}_{ij}^\text{loc} = (\alpha_{ij}^\text{loc} - \alpha_{ij}^\text{loc,v}) \frac{v_{ij}^\text{sig,v}}{h_i},$$

(16)

which we integrate in time together with the hydrodynamical quantities. The constant $\ell$ specifies the decay length and in our test problems we find a numerical value of $\ell = 4.0$ to give reasonable results.

#### 2.4.2 Gradient estimators

We use the Balsara (1995) form of the shear viscosity limiter

$$f_{ij}^\text{shear} = \frac{|\nabla \cdot v|_i + |\nabla \times v|_i + \sigma_j}{|\nabla \cdot v|_i + |\nabla \times v|_i + \sigma_j},$$

(17)
with \( \sigma_i = 0.0001c_i^2/\rho_i \) for numerical stability reasons. At this point, the question arises how to calculate the divergence and vorticity. The common curl estimator of SPH reads

\[
(\nabla \times \mathbf{v})_i = -\frac{1}{\rho_i} \sum_j m_j (\mathbf{v}_j - \mathbf{v}_i) \times \nabla_i W_{ij},
\]

(18)

which takes the lowest order error term into account. Since higher order error terms are neglected, this formulation performs very poorly in the regime of strong shear flows. Therefore, we resort to a higher order calculation scheme of the velocity gradient matrix (see similar approaches by Cullen & Dehnen 2010; Price 2012a; Hu et al. 2014). We follow the approach presented in Price (2012a) for the computation of the gradient matrix. We expand \( \mathbf{v}_j \) for every vector component \( k \) in a Taylor-series around \( i \) with

\[
\mathbf{v}_j = \mathbf{v}_i + (\partial_i \mathbf{v}_j)((\mathbf{x}_j - \mathbf{x}_i)^h + O(h^2)).
\]

Inserting equation (19) into equation (18) yields an easy solution for the linear term \( \partial_i \mathbf{v}_j \) and the velocity gradient matrix by solving the following system of equations:

\[
\chi^{\alpha\beta} = \sum_j m_j (\mathbf{x}_j - \mathbf{x}_i)^\alpha \nabla^\beta W_{ij},
\]

(20)

which requires a matrix inversion for \( \chi^{\alpha\beta} \). Conveniently, the estimator is also independent of density and thus, can be calculated in the same computational loop along with densities. Subsequently, the updated estimates of velocity divergence and curl are calculated directly from the full velocity gradient matrix via

\[
(\nabla \cdot \mathbf{v})_i = \frac{\partial \mathbf{v}_i^\alpha}{\partial x^\alpha},
\]

(22)

\[
(\nabla \times \mathbf{v})_i^\alpha = \epsilon_{\alpha\beta\delta} \frac{\partial \mathbf{v}_i^\beta}{\partial x^\delta}.
\]

(23)

In our test problems, we find the low-order estimator of velocity divergence to give already satisfying results (see also appendix A2 in Schaye et al. 2015). In contrast, the low-order estimator of velocity curl performs very poorly and we obtain significantly improved results with the high-order curl estimator of equation (23). The high-order estimators are not restricted to the AV scheme but they also enter various other modules of the code, where their precise calculation is required. For example, this additionally greatly improves the approximation of fluid vorticity written into the simulation snapshots.

2.5 Artificial conductivity with gravity correction

2.5.1 Smoothing of jumps

We move on to address the mixing problem in SPH by introducing a kernel-scale exchange term for internal energy transport. We include AC for purely numerical reasons to treat discontinuities in the internal energy (similar to the capturing of velocity jumps by AV), which arise from our ‘density-entropy’ formulation of SPH. We note that a ‘pressure-entropy’ formulation of the EoM is also able to address the mixing problem but it also requires the presence of AC in order to smooth noise in internal energy behind shocks (Hopkins 2013; Hu et al. 2014). Thus, in either flavour of SPH the inclusion of AC is recommended and many different formulations of the AC diffusion equation have been investigated so far. Although their precise details vary across the literature, they all ensure conservation of internal energy within the kernel. Price (2008), Price (2012a) and Valdarnini (2012) propose the diffusion of internal energy, while Read & Hayfield (2012) propose the diffusion of entropy. Wadsley et al. (2008) propose a first mixing formulation to resolve the differences in entropy profiles within cosmological comparison simulations (Frenk et al. 1999) between grid and SPH codes. The diffusion coefficient is approximately proportional to \( \alpha \nu_{\text{sig}}^2 \mathbf{x}_{ij} \) and the numerical coefficient \( \alpha' \) is commonly treated as constant through space and time. We adapt the formulation of a spatially varying coefficient of Tricco & Price (2013) and additionally calculate a limiter depending on the local hydrodynamical and gravitational states. We compute the gradient of internal energy as

\[
(\nabla u_i) = \frac{1}{\rho_i} \sum_j m_j (u_j - u_i) \nabla_i W_{ij}
\]

(24)

and approximate the AC coefficient

\[
\alpha'_i = \frac{h_j}{3} |\nabla u_i|, \frac{\nu_{\text{sig}}^2}{\rho_i}
\]

(25)

as a measure of noise of internal energy sampling on kernel scale. The time evolution (i.e. spatially varying SPH discretization of the second-order diffusion equation) of the internal energy for each particle and its neighbours is then given by

\[
\frac{du_i}{dt}_{\text{cond}} = \sum_j \frac{m_j}{\rho_{ij}} (u_j - u_i) \alpha'_{ij} \nu_{\text{sig}}^2 \mathbf{T}_{ij},
\]

(26)

where we employ the choice of Price (2008) for signal velocity depending on the pressure gradient of the form

\[
\nu_{ij}^{\text{sig,c}} = \sqrt{|P_i - P_j|/\rho_{ij}}
\]

(27)

and \( \alpha'_{ij} = (\alpha_i' + \alpha_j')/2 \) is the symmetrized conduction coefficient, which are individually limited to the interval [0, 1]. In the literature several other forms of AC (see e.g. Wadsley et al. 2008; Valdarnini 2012) or approaches to the mixing problem (see e.g. Hopkins 2013) have been proposed.

2.5.2 Gravity limiter

We note that the amount of AC applied depends on the gradients of internal energy and of pressure. In the case that the thermal pressure gradient is determined by gravitational forces (i.e. hydrostatic equilibrium), this method would incorrectly lead to unwanted conduction. In the following, we determine the contribution of hydrostatic equilibrium to the total thermal pressure gradient and present a method to limit the amount of conduction. First, for every individual active particle, we project the gravitational force \( \mathbf{F}_g \) on to the hydrodynamical force \( \mathbf{F}_h \) and calculate the partial force \( \mathbf{F}_{ig} \) of \( \mathbf{F}_i \), which is balanced by \( \mathbf{F}_{ig} \) to

\[
\mathbf{F}_{ig} = \frac{(\mathbf{F}_g \cdot \mathbf{F}_i)}{||\mathbf{F}_i||^2} \mathbf{F}_i.
\]

(28)

The sign of \( \mathbf{F}_{ig} \) depends on the spatial orientation of the force vectors. Secondly, we subtract/add the partial force \( \mathbf{F}_{ig} \) from/to the hydrodynamical force \( \mathbf{F}_{ih} \) and obtain \( \mathbf{F}_i \), which we call the gravitationally adjusted hydrodynamical force

\[
\mathbf{F}'_i = \mathbf{F}_{ih} + \mathbf{F}_{ig}.
\]

(29)
and which we use to determine a limitation factor $\delta_i^c$ for AC

$$\delta_i^c = \left( \frac{(F_i^c \cdot F_j^c)}{|F_i|^2} \right)^{q}. \quad (30)$$

The limiter ensures that AC is only applied to the part of $F_i^c$ which is not balanced by $F_j^c$. The exponent $q$ represents a scaling for the aggressivity of the gravity correction. We limit our correction factor to the interval $[0, 1]$ and directly multiply it on to the individual AC coefficients $\alpha_i^c$. The limiter performs only as well as the hydrodynamical scheme is able to resolve hydrostatic equilibrium (in the ideal case the angle between force vectors is $180^\circ$). However, in SPH simulations small-scale noise is present at all times within the kernel and thus also in the force vector angles. The exponent $q$ (applied after the boundary verification) can then be understood to account for the noise in the particle distribution and mimics an opening angle of force vectors. After extensive studies and performing a variety of test problems, we settle with $q = 5$. The limiter returns zero in the case no hydrodynamical forces are present and one in the case, where no gravitational forces are present. We are aware that in the presence of strong pressure gradients and rotational forces our approach only marginally limits the amount of AC applied. However, we did not encounter major problems in our simulations performed with the ‘new’ scheme so far. Therefore, we assume this issue to be not too important at the present state.

### 2.6 Particle wake-up and time-step limiter

**GADGET** employs individual time-steps for all of the particles to increase computational efficiency. Thus, the particle population is split into a set of active particles, whose hydrodynamical properties are integrated in the current time-step and a set of inactive particles, which reside on larger time-steps. These individual time-steps are computed from the local thermodynamical properties of each particle. However, the splitting between active and inactive computational regions creates problems, where both sets of particles are overlapping. In the case of a rapid gain in velocity or entropy an active particle can penetrate into a region of inactive particles. The inactive particles do not notice the sudden presence of the highly dynamical particle and therefore large gradients in the time-steps and unphysical results can occur. As a treatment we adopt a time-step limiting particle wake-up scheme as proposed and implemented in the **GADGET**-3 code by Pakmor (2010) and Pakmor et al. (2012) with the help of K. Dolag. It is similar to the time-step limiting scheme described by Durier & Dalla Vecchia (2012) and can be considered an extension of the Saïtoh & Makino (2009) mechanism. Furthermore, our limiter compares signal velocities instead of time-steps and accounts for incorrect extrapolations. For every active particle, in every time-step, the individual time-steps themselves are re-computed according to

$$\Delta t_i = \frac{C h_i}{v_{ij}^{\text{sig}}}, \quad (31)$$

where $C$ is the Courant factor and $v_{ij}^{\text{sig}}$ the maximum signal velocity (see Section 2.4). For the calculation of the time-step the maximum of the signal velocity computed between the active particle $i$ and all its neighbour particles $j$ within the entire kernel is used. **GADGET** employs a check during the hydrodynamical force computation for large differences in signal velocities (see equation 11) within the kernel by evaluating

$$v_{ij}^{\text{sig}} > f_w v_{ij}^{\text{sig}}, \quad (32)$$

with a tolerance factor $f_w$ corresponding to a wake-up triggering criterion, which captures sudden changes in the pairwise signal velocity. From our hydrodynamical standard tests, we find $f_w = 3$ to give reasonable results. Additionally, the fluid quantities of the recently woken-up particles could have already been predicted half a time-step into the future. Therefore, the incorrect extrapolation is removed and the contribution from the real time-step added. These corrections are performed for all particles for which the time-steps are adapted.

### 3 Hydrodynamical tests without gravity

We evaluate the performance and accuracy of the two different SPH implementations with a first set of standard problems. These first test problems are purely hydrodynamical and do not include gravity or more advanced physics, yet. Throughout all the test problems, we use an adiabatic index of $\gamma = 5/3$, the same set of numerical parameters (see Section 2) and we do not specifically tune individual test problems.

#### 3.1 Sod shock tube

We consider the Sod shock tube problem (Sod 1978) to study the SPH behaviour in a simple weak shock test. We set up 630 000 particles of equal masses using a relaxed glass file in a three-dimensional periodic box with dimensions $\Delta x = 140$, $\Delta y = 1$ and $\Delta z = 1$. On the left-half side of the computational domain ($x < 70$), we initialize 560 000 particles with a density of $\rho_L = 1.0$ and a pressure of $P_L = 1.0$. On the right-half side of the computational domain ($x > 70$), we initialize 70 000 particles with a density of $\rho_R = 0.125$ and a pressure of $P_R = 0.1$.

Fig. 1 shows the results of the test problem at time $t = 5.0$. In general, both SPH schemes agree fairly well with a reference solution.
3.2 Sedov blast

We consider the Sedov blast problem (Sedov 1959) to study the SPH behaviour in a simple ultrasonic strong shock test. We set up $130^3$ particles of equal masses using a relaxed glass file in a three-dimensional periodic box with dimensions $\Delta x = \Delta y = \Delta z = 6$ kpc. In the entire computational domain, we initialize the particles with a density of $1.24 \times 10^6 \, M_\odot \, kpc^{-3}$ and one Kelvin as temperature. At the centre of the box, we point-like distribute the energy $E = 6.78 \times 10^{53}$ erg to mimic a supernova explosion among the nearest 102 particles.

Fig. 4 shows thin slices through the centre of the simulation box and Fig. 5 the corresponding particle distribution at time $t = 0.03$. Furthermore, we perform the ‘standard’ scheme test also with the time-step limiter ($f_w = 8000$) because of the very strong shock (Mach $\gg 100$) of the blast and otherwise any comparison will fail. Without the limiter, shocked particles penetrate into quiescent regions causing a highly distorted fluid sampling, which results in an incorrect solution leading to an incorrect propagation of the shock front (see discussions in Saitoh & Makino 2009; Durier & Dalla Vecchia 2012) and corresponding smoothing. The entire ‘new scheme’ reproduces the analytical solution (black line) very well, with the ‘standard’ run (green dots) totally failing, and the ‘new’ run (red dots) capturing the position, density and temperature of the shock fairly well. In addition, we show a partially improved ‘standard’ run (blue dots), where we enabled the time-step limiter but nothing else. This run also yields reasonable good results in this test, but as we see later comes short in other tests. We see that the ‘new’ scheme yields a smooth distribution of particles within the
developed and the ring breaks up. Also, the Balsara limiter does not succeed in limiting AV because of the insufficient calculation of vorticity. In the ‘M&M’ scheme (bottom left-hand panel), we use the implementation of a low-viscosity scheme initially proposed by Morris & Monaghan (1997) and implemented into GADGET by Dolag et al. (2005). Their scheme uses a time-dependent evolution of numerical AV coefficient $\alpha^v$ to suppress AV in the absence of shocks and manages to keep the ring stable for about seven dynamical times. However, the M&M scheme requires a minimum value of $AV$ and also uses a low-order estimator for vorticity, which leads to the ring break-up. At last, we show the results of the ‘new’ scheme (bottom-right panel), which we used without a minimum value for AV. This scheme uses a high-order estimator of velocity gradient matrix, which results in a very accurate calculation of divergence and vorticity. Therefore, also the computation of the Balsara shear flow limiter is very accurate and suppresses AV completely within the entire ring structure. We do not note an artificially induced transport of angular momentum and orbital changes of test particles. Consequently, the ring remains stable for many dynamical times and the initial Gaussian surface density distribution is preserved until we stopped the simulation.

### 3.4 Cold blob test

We consider the blob test (Agertz et al. 2007; Read et al. 2010) set up with publicly available initial conditions$^1$ to study the SPH behaviour in a test problem with interacting gas phases and surfaces. We initialize 9641 651 particles of equal masses using a relaxed glass file in a three-dimensional periodic box with dimensions $\Delta x = 10$, $\Delta y = 10$ and $\Delta z = 30$ in units of the cold cloud radius. A cold cloud is centred at $x, y, z = 5$ and travels at a Mach number of $M = 2.7$. The background medium is set up 10 times less dense and 10 times hotter than the cloud. Spherical harmonics are used to seed large-scale perturbations on to the surface of the cloud. Because of the low Mach number shocks, we expect the time-step limiter to be only of minor importance.

Fig. 7 shows thin slices through the density structure at various times. In the ‘standard’ scheme, the cold gas cloud is prevented from dissociating by the follow major effect (see also e.g. Agertz et al. 2007). The presence of artificial surface tension confines the blob of cold gas. This is clearly visible by the numerically induced stretching of the cloud. Cold material, which should have been mixed into the ambient hot medium is confined within an elongated structure. In the ‘new’ scheme, the presence of AC promotes the mixing with external ambient medium. We also note that the ‘new’ scheme resolves different shock structures propagating through the box both more accurately and smoothly.

Fig. 8 illustrates the dissipation of the gas cloud by tracking the time evolution of cold blob mass. We associate (see also Agertz et al. 2007) particles with the cold cloud with a temperature criterion of $T < 0.9 \cdot T_{\text{ext}}$ (in contrast to the external ambient medium) and a density criterion of $\rho > 0.64 \cdot \rho_{cl}$ (in contrast to the initial density of the cloud). In the ‘standard’ scheme (blue line), only half of the cold gas mass is mixed into the hot ambient medium over five dynamical times. The effects of the improved AV and WC4 kernel (green line) are negligible. The major impact and contribution to cloud dissociation is made by AC (pink line) and the corresponding introduced mixing process. The results are close to a test run performed with the ENZO (Bryan et al. 2014) code in a comparable set-up, which we

$^1$ http://www.astrosim.net/code/doku.php
Figure 7. Blob test. We show thin slices of gas density through the centre of the computational domain at times $t = 0.0, 3.0, 6.0$ and $10.0$. In the ‘standard’ scheme, numerical surface tension prevents mixing between cold and hot phases leading to an artificial stretching of the cloud and an unphysical solution. In the ‘new’ scheme, AC helps to promote cloud dissociation; however, some residual surface tension remains left. Furthermore, the shock structures throughout the box are more defined and better resolved.

Figure 8. Blob test. We show the fraction of cold gas as a function of time for four different SPH schemes. In the ‘standard’ scheme, numerical surface tension prevents mixing between the cold and hot phases. The importance of AC becomes clear as it promotes mixing between gas phases, which allows a dissociation of the cloud. Only some residual surface tension is left. Comparing to a run performed with the enzo grid code, we find our AC scheme to model mixing in a conservative way.

3.5 Kelvin–Helmholtz instability

We consider the Kelvin–Helmholtz instability (Agertz et al. 2007; Read et al. 2010) from publicly available initial conditions 1 to study the SPH behaviour in a simple shearing instability test. We set up 1548 288 particles of equal masses using a cubic lattice in a three-dimensional periodic box with dimensions $\Delta x = 256 = \Delta y = 256$ and $\Delta z = 16$ kpc, which is centred around $(0,0,0)$. In the central half of the box ($|y| < 64$), we initialize 512 000 particles with a density of $\rho_1 = 6.26 \cdot 10^3$ $M_\odot$ kpc$^{-3}$, temperature of $T_1 = 2.5 \cdot 10^6$ K and a velocity in $x$-direction of $v_1 = -40$ km s$^{-1}$. In the outer half of the box ($|y| > 64$), we initialize 1036 288 particles with a density of $\rho_2 = 3.13 \cdot 10^2$ $M_\odot$ kpc$^{-3}$, temperature of $T_2 = 5.0 \cdot 10^6$ K and a velocity in $x$-direction of $v_2 = 40$ km s$^{-1}$. To trigger the instability, we perturb the velocity in $y$-direction with a mode of wavelength 128 kpc and amplitude of 4 km s$^{-1}$ at the boundary layer that is exponentially damped towards the upper and lower edge of the box.

Fig. 9 shows thin projections through the specific entropy structure of the test problem at various times. In the ‘standard’ scheme, the fluid evolves in a laminar fashion and the growth of perturbations is totally suppressed by the artificial surface tension confining the central gas stream and large amounts of AV damping velocity perturbations (see also e.g. Agertz et al. 2007; Price 2008). In
the ‘new’ scheme, the high-order Balsara shear limiter successfully limits AV and allows large-scale perturbations to develop two prominent roll-ups. Additionally, AC nearly removes the artificial surface tension between the two gas phases and promotes mixing within the roll-ups. The entire test set-up does not evolve completely symmetric because of small secondary perturbations caused by the initial set-up on a cubic lattice. Most importantly, the high-order AV and AC prove crucial for this test problem, while the WC4 kernel and time-step limiter are of less importance. At the late stages, the ‘new’ scheme is dominated by diffusion.

3.6 Decaying subsonic turbulence

Recent comparisons of standard SPH implementations with static and moving mesh grid codes have sparked a debate about the capabilities of SPH to model subsonic turbulences (see e.g. Bauer & Springel 2012; Price 2012b; Hopkins 2013, 2015). We study the behaviour of our ‘new’ scheme in idealized simulations of decaying subsonic turbulence. In particular, we are interested in the effective viscosity of the two schemes and the behaviour of the ‘SPH noise’ under conditions appropriate to galaxy formation and cluster simulations, i.e. non-isothermal, decaying motions from solenoidal and compressive modes. As most baryons on cosmological scales are in weakly collisional plasmas, numerical models should aim to minimize viscosity where possible (see e.g. Brunetti & Lazarian 2007).

3.6.1 Grid and particle conversion procedures

We set up $512^3$ particles of equal masses within a periodic box of side length $3 \text{ Mpc \, h}^{-1}$ using carefully relaxed SPH glass files to minimize spurious initial kinetic energy. Subsequently, we define a velocity field on a grid of the same resolution in $k$-space by sampling a spectral distribution using the Box–Mueller method. The velocity field is transformed back to real-space using a fast Fourier transformation (FFT), normalized such that the average velocity is of the desired Mach number. The velocities from the grid are transferred to the particle distribution using the nearest grid point (NGP) sampling kernel.

To assess the impact of random motions near the resolution scale, we need to measure the velocity power spectrum within the SPH kernel. However, the accurate estimation of the velocity power spectrum of a particle distribution close to the Nyqvist frequency is non-trivial, because of aliasing of the velocity power by the binning kernel (see discussions in Jing 2005; Cui et al. 2008; Jasche, Kitauro & Ensslin 2009). This can be compared to a problem in signal processing, where SPH represents an analogue signal, a grid and a digital signal representation of it. Aliasing is strongest at the smallest scales/largest modes, where the velocity power on the particles is modified by the shape of the binning kernel in configuration-space. To understand this effect and compare binning kernels, we take initial conditions with a full Kolmogorov power spectrum ($P_k \propto k^{-11/3}$) without performing a simulation and directly bin the velocity back to a grid using different kernels. After a forward FFT, we radially average the velocity power in $k$-space in 32 logarithmic bins.

Fig. 10 shows the resulting power-spectra where the black line represents the original power spectrum. We show the kernels: NGP (red line), Cloud in Cell (CIC, green line), Triangular Shaped Cloud (TSC, violet line), Daubechies scaling function of 20th order (D20, orange line), and the WC4 SPH kernel with 200 neighbours (SPH, brown line) (Hockney & Eastwood 1988; Daubechies 1992; Dehnen & Aly 2012). We also show the NGP with two times oversampling (blue line), which was used in Bauer & Springel (2012). As vertical lines, we show the Nyqvist frequency $k_{\text{Nyq}} = N k_{\text{min}}$, the WC4 smoothing scale $k_s$, and the WC4 kernel compact support $k_{\text{hsml}} = \pi / h_{\text{sm}}$ (Dehnen & Aly 2012).

During the binning process, the SPH kernel function conserves density to machine precision but not energy, i.e. binning with the SPH kernel is a diffusive process. The other kernels behave
opposite, they conserve mass, scalar velocity and energy to less than 1 per cent but not SPH density and volume. For sufficiently homogeneous particle distributions (Cui et al. 2008). Our comparison also resolves the differences found in Bauer & Springel (2012) and Price (2012b), who use the twice oversampled NGP kernel and the standard SPH kernel, respectively. Prior studies based on the NGP kernel binning overestimated the SPH noise, while SPH kernel based binning suppressed the real noise by aliasing. We conclude that all kernels except the D20 show substantial aliasing and it seems hard to draw definitive conclusions from simulation results under this condition. Thus we make it our fiducial choice for this study. We note that in the presence of strong gradients in density the SPH kernel remains the only viable choice to obtain binned quantities, because it is the only kernel in our comparison that guarantees a non-negative non-zero density in the entire simulation at all grid resolutions. This works reasonably well for a physical interpretation of velocity power-spectra, because motions below the SPH smoothing scale are caused by numerical effects (Price 2012b).

3.6.2 Spectral evolution of turbulence

To compare the ‘standard’ and the ‘new’ schemes we consider decaying turbulence within a periodic box. We seed compressive and solenoidal modes in the range of $k \in [1.6, 3.1]$, to obtain initial conditions appropriate for the galaxy and cluster environment, where turbulence is injected by merger infall on the scale of the halo core radius. We normalize the velocity fluctuations in the box such that the average velocity equals a Mach number of $M = 0.1$ and we do not time-average spectra.

Fig. 11 (left-hand and middle panels) shows the time evolution of velocity power-spectra for the ‘standard’ scheme (left-hand panel) and the ‘new’ scheme (middle panel). Here, we also show the scale of the SPH kernel compact support (black vertical line) and the kernel smoothing scale (dotted vertical line). In-line with previous studies (Bauer & Springel 2012, their fig. 12), the ‘standard’ scheme does not develop a turbulent cascade and damps kinetic energy very quickly. Our ‘new’ scheme develops a cascade at large scales (small $k$) but then shows a depression of kinetic energy close to the kernel scale. This, again, is in-line with prior studies (Hopkins 2013). The damping of the spectrum at the later times appears self-similar, i.e. the shape of the spectrum does not change as energy decreases. Inside the kernel the typical build-up of thermal motions around the smoothing scale $k_{\sigma}$ can be observed, but scales outside the kernel are not affected.

In order to understand if the cause of the velocity depression at $k \approx 10$ is caused by the formulation of $AV$, we perform a test-run without any viscosity (Fig. 11, right-hand panel). Throughout the whole evolution, the spectrum at the smallest $k$ follows the Kolmogorov scaling, as expected. Once the turbulent cascade is established, the spectrum turns over at increasingly smaller scales, which is equivalent to an isotropization of kinetic energy inside the kernel and subsequent filling of larger scales with isotropic motions. This follows from the fact that SPH particles are subject to the pair-wise repulsive force (Price 2012a), and hence behave like a thermal gas below the kernel scale. Eventually, such a system will show a flat power spectrum as expected from the second law of thermodynamics. In our simulation, we also observe the depression outside the smoothing scale, which seems to be an intrinsic feature
of SPH related to an energy transfer from outside the kernel to smaller scales, and not related to our formulation of AV.

In our ‘new’ scheme including AV, thermal motions are well controlled inside the kernel scale, commonly referred to as ‘kernel noise’. We argue that these motions are not spurious, because no additional energy is retained in them, as SPH is fully conservative and the spectrum decays roughly in a self-similar manner. If we define the kernel smoothing scale \( k_\sigma \) as the dissipation scale intrinsic to SPH, the ‘new’ scheme does not show a bottle-neck effect as found in grid codes at adjacent larger scales but a depression, roughly in the same range in \( k \). This is in line with the results shown by Hopkins (2015), whose code uses a Riemann solver to formulate noise-free AV on scales of the interparticle separation to obtain grid-code behaviour. We note that the difference in dissipation scale \( d_{\text{max}} \) versus \( \sigma_{\text{kernel}} \) translates into more resolution elements required by SPH compared to Eulerian methods, i.e. slower convergence.

Fig. 12 shows thin slices through the centre of the simulation box after one sound-crossing time. We visualize gas density (left-hand panel) and vorticity (right-hand panel). It can be clearly seen

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**Figure 12.** Decaying subsonic turbulence. We show thin slices through the centre of the simulation box after one sound-crossing time for both schemes. The panels correspond to gas density (left-hand panel) and vorticity (right-hand panel). The velocity field shows well developed turbulence consisting of compressive and shearing motions. The ‘new’ scheme is able to more accurately compute vorticity and suppress AV with the high-order Balsara limiter.
that our ‘new’ scheme resolves compressive and shearing velocity motions better than the ‘standard’ scheme. The high-order derivatives of velocity lead to a more accurate estimation of vorticity and thus, limit the impact of AV and preserve kinetic energy and turbulent motions. Furthermore, the difference in AV and velocity dissipation between the two schemes becomes strikingly evident in the time evolution of kinetic energy (Fig. 13). The ‘standard’ scheme dissipates 90 per cent of the kinetic energy budget within four sound-crossing times, while the ‘new’ scheme preserves energy better by a factor of 5.

We conclude that our code performs comparably to modern implementations of SPH (Price 2012a; Hopkins 2013), even in the case of non-isothermal compressive and solenoidal decaying turbulence found in cosmological simulations. We show that the disagreement between Bauer & Springel (2012) and Price (2012a) is largely caused by technical differences, to solve it we propose a solution based on the D20 binning kernel. We also show that the downturn in the velocity power spectrum is not caused by the AV implementation.

4 HYDRODYNAMICAL TESTS WITH GRAVITY

We continue to evaluate the performance and accuracy of the two different SPH implementations with a second set of standard problems. These second tests include hydrodynamical as well as gravitational forces and also take a cosmological time integration into account.

4.1 Hydrostatic sphere

We consider a sphere in hydrostatic equilibrium to study the SPH behaviour in combination with gravity in an ideally stable system. We set up 88 088 dark matter particles with individual masses of $2 \times 10^9 \, M_\odot$ and 95 156 gas particles with individual masses of $4.75 \times 10^8 \, M_\odot$. The total mass of the sphere is $2.2 \times 10^{14} \, M_\odot$ and we use vacuum boundary conditions and a gravitational softening length of 12 kpc. We set up the initial equilibrium conditions following Komatsu & Seljak (2001), as described in Viola et al. (2008). We evolve the sphere adiabatically and do not include cooling or heating mechanisms in this test.

Fig. 14 shows the results of the test problem at various times. At first, the initial set-up (dotted lines) of the sphere is not yet in hydrostatic equilibrium and requires some time to settle. Once hydrostatic equilibrium is reached around time $t = 2.6$ (dashed lines for the ‘new’ scheme) all hydrodynamical schemes must preserve the structure of the sphere and the radial profiles towards the final simulation time $t = 7.6$. Between all schemes, the thermal pressure profiles are indistinguishable balancing the gravitational pressure. However, the composition of the thermal pressure $P = (\gamma - 1)\rho u$ changes and the radial profiles for density $\rho$ and internal energy $u$ change significantly. The ‘standard’ scheme (blue lines) reaches the highest central density and also features lowest central internal energy and entropy. However, the internal energy drops towards the centre and no stable solution is reached at all. We suggest this behaviour to be a joint impact of pairing instability caused by the cubic spline kernel function and lack of internal energy mixing. When introducing only AC, no gravitational limiter and no further SPH developments (green lines), the results marginally improve. Internal energy still drops towards the centre but this time because too much AC was introduced. In principle, AC leads to entropy cores.

Figure 14. Sphere in hydrostatic equilibrium. We show radial profiles of density (left-hand panel), internal energy (middle panel) and entropy (right-hand panel) at three different times. At first, the initial conditions (dotted lines) must settle into hydrostatic equilibrium (dashed lines), which then remains stable for an extended period of time (solid lines). The stability of the sphere is determined by the differences occurring between the settled (only shown for the ‘new’ scheme) and the evolved state (shown for all schemes). In the ‘standard’ scheme (blue lines), pairing instability caused by the cubic spline kernel and lack of mixing lead to an incorrect central solution in density (too high) and internal energy (too low). The addition of the WC4 kernel (purple lines) prevents the formation of particle clumps at the centre and AC promotes fluid mixing but full AC without gravity limiter (green lines) leads to a numerical transport of internal energy outwards. The gravity limiter treats this behaviour (brown lines). The ‘new’ scheme (red lines) with WC4 and AC and gravity limiter significantly improves the radial profiles in all physical quantities.
but without the gravitational limiter overmixing occurs and internal energy is transported from the centre to the outskirts along the pressure gradient. The addition of the gravitational limiter (brown lines) improves the results significantly. The divergence of profiles towards the centre is removed and a stable core of internal energy and entropy is reached. Furthermore, no numerically induced transport of heat takes place. Additionally, we perform a test run with introducing only the WC4 kernel and no further SPH improvements (purple lines). In this run, the central density is lower than in the ‘standard’ scheme, most probably because the clumping of particles in the centre and the occurrence of the pairing instability is suppressed by the WC4 kernel in contrast to the run with the cubic spline. This also leads to a plateau in internal energy in the centre of the sphere. At last, we show the results of the entire ‘new’ scheme (red lines) and find the results to remain stable with all additional modifications to WC4 kernel, AV and time-step limiter. We find this run to give the most stable radial profiles in time. This test confirms the importance of kernel functions immune to pairing instability and a proper implementation of AC and clearly shows the effects of particle clumping and over and undermixing in gravitationally virialized systems, which are important in cosmological simulations.

4.2 Evrard collapse

We consider the Evrard collapse (Evrard 1988) to study the SPH behaviour in the presence of dynamically important gravitational forces and collapse of gas. We initialize a sphere of gas with mass \( M = 1 \), radius \( R = 1 \) and density profile of \( \rho \sim r \) for \( r < R \) and use vacuum boundary conditions and a gravitational softening length of 0.005. We do not use an external gravitational potential, dark matter particles or radiative cooling and thus the cloud only self-gravitates on the free-fall time-scale. The gas is initially at rest and the thermal energy budget is orders of magnitude smaller than the gravitational binding energy.

Fig. 15 shows the results of the test problem at time \( t = 0.8 \). We compare the SPH results to a reference solution similar to Steinmetz & Mueller (1993). In density (left-hand panel) as well as in velocity (middle panel) all schemes show similar trends. The general structure of the test is well reproduced; however, the shock front is slightly smoothed and broadened. The solution in density deviates slightly from the reference solution at the centre. The most striking differences can be seen in pre-shock entropy (right-hand panel). The ‘new’ scheme (red line) produces higher levels of entropy compared to the ‘standard’ scheme (blue line). We investigate this trend and perform another test simulation (purple line) with the ‘new’ scheme but the cubic spline kernel. This run is closest to the pre-shock reference solution.

4.3 Zel’’dovich pancake

We consider the Zel’’dovich pancake (Zel’dovich 1970) to study the SPH behaviour for the cosmological time integration with Hubble function \( H(t) \) instead of time \( t \). This test describes the evolution of a sinusoidal cosmological perturbation in an expanding Einstein-de-Sitter universe. After an initial linear growth phase, the one-dimensional perturbation collapses and several strong shocks develop. Conveniently, the Zel’dovich pancake has an analytical solution describing the evolution well up to the collapse, which we used to create the initial conditions of the simulation. The comoving position \( x \) of an initially unperturbated coordinate \( q \) at redshift \( z \) is given by

\[
x(q, z) = q - \frac{1 + z_c \sin(kq)}{1 + z} \frac{\sin(kq)}{k},
\]

where \( k = 2\pi/\lambda \) is the wavenumber of the perturbation with a wavelength of \( \lambda \). We numerically invert equation (33) to obtain \( q(x) \). The peculiar velocity corresponding to the initial displacement is given by

\[
v_{pec}(x, z) = -H_0 \frac{1 + z_c}{(1 + z)^{1/2}} \frac{\sin(kq)}{k},
\]

and the comoving density is given by

\[
\rho(x, z) = \frac{\rho_0}{1 - \frac{1}{3z_c} \cos(kq)},
\]

where \( \rho_0 \) is the critical density, \( H_0 \) the present-day Hubble constant and \( z_c \) the redshift of collapse. Furthermore, the temperature evolves adiabatically up to the collapse as

\[
T(x, z) = T_i \left[ \left( \frac{1 + z_i}{1 + z_i} \right)^3 \frac{\rho(x, z)}{\rho_0} \right]^{2/3},
\]

where \( z_i \) is the initial redshift. We follow Bryan et al. (1995), Trac & Pen (2004) and Springel (2010b) in our test set-up and choose \( \lambda = 64 \text{ Mpc} h^{-1} \), \( z_c = 1 \), \( z_i = 100 \) and \( T_i = 100 \text{ K} \). In a fully three-dimensional box, we set up \( 256^3 \) dark matter particles of equal masses as well as \( 256^2 \) gas particles of equal masses.
Fig. 16. Zel’dovich pancake. We show the evolution of density contrast (left-hand column), temperature (middle column) and velocity (right-hand column). We show the state of the pancake at an intermediate redshift $z = 3.6$, while it is still in the linear regime before the collapse and at the final redshift $z = 0$, when it is evolved well into the non-linear regime. Both SPH schemes agree well with the analytical solution during the linear growth phase. At the final redshift, the ‘new’ scheme resolves well the density contrast and yields a broader shock and a slightly smoother velocity.

Fig. 16 shows the results of the test problem at two redshifts. In the top row, we show the evolution of the pancake before the collapse at redshift $z = 3.6$, while it is still in the linear phase. The ‘standard’ (blue lines) and ‘new’ (red lines) schemes give comparable results in density contrast (left-hand column), temperature (middle column) and velocity (right-hand column). The simulated evolution agrees well with the analytical solution (green lines), which describes the linearized evolution of initial sinusoidal perturbation. At this stage of the collapse the test is dominated by gravitational forces and hydrodynamical forces are negligible. Therefore, we do not expect striking differences to arise between both schemes. Both capture the linear growth and adiabatic evolution well.

In the bottom row, we show the pancake at the final redshift $z = 0$. Again, we compare to the analytical solution in the regions outside the central shock. In general, both schemes agree but we note the following differences. The peak density contrast is marginally lower in the ‘new’ scheme because of additional smoothing introduced by AC. However, the evolution of density contrast in the low-density regions is described better by the ‘new’ scheme and is resolved with less noise. We recall the Sod shock tube (Section 3.1) and the Sedov blast wave (Section 3.2) problems, where we also find more accurately resolved density fields and lower peak densities. Concerning temperature, we find the central shock to be slightly broader in the ‘new’ scheme, which is caused by two effects. First, the higher amount of viscosity within the shock leads to an earlier heating of particles and thus broadens the shocks. Secondly, the time-step limiting particle wake-up scheme captures highly active particles before they penetrate into inactive regions, which leads to a better fluid sampling and also shock broadening. This can also be noticed in velocity, where the profile is slightly smoothed in the central region. In summary, our ‘new’ scheme gives reasonable results in this cosmological test problem and the differences between both schemes are very small at redshift $z = 0$. Therefore, our new implementation is ready to be applied to idealized astrophysical problems.

5 ASTROPHYSICAL APPLICATIONS

We complete the evaluation of the performance and accuracy of the two different SPH implementations in idealized simulations of galaxy and galaxy cluster formation.

5.1 Idealized galaxy formation

In order to check if the ‘new’ scheme is numerically stable when coupled with a simple effective description of the interstellar medium, we consider the formation of an isolated disc from a cooling gas cloud embedded within a rotating dark matter halo. This idealized application also includes a prescription for cooling, supernova feedback and star formation of Springel & Hernquist (2003). We focus on the differences between both hydrodynamical schemes and therefore, we do not consider a cosmological environment or more advanced physical processes such as black holes, stellar evolution or metals. Numerical comparison simulations (see e.g. Scannapieco et al. 2012) are a common tool to study the impact of numerical schemes and physical modules.
Figure 17. Idealized galaxy formation. We show the spatial distribution of stars, where the colours visualize the age of stars (top panels) and the spatial distribution of gas, where the colours visualize the star formation rate (bottom panels) at time $t = 7.5$ Gyr. In the ‘standard’ scheme (left-hand panels), the distribution of star formation is clumpy and the object appears bulgy. In the ‘new’ scheme (right-hand panels), the distribution of star formation as well as the stellar component are more extended and pronounced in a disc-like structure. Furthermore, the size of the bulge is smaller and the distribution of young stars is more extended. We show a more quantitative comparison in Figs 18 and 19.

Within a computational domain of roughly $1 \text{ Mpc}^3$, we set up 4041 345 particles resembling a Milky Way-like dark matter halo with a total mass of $1.8 \times 10^{12} \text{ M}_\odot$. We include 4466 429 gas particles with a total mass of $2.2 \times 10^{11} \text{ M}_\odot$, which corresponds to a baryon fraction of approximately 11 per cent. The gravitational softening length is 450 pc. Initially, the distribution of dark matter follows a Navarro–Frenk–White profile (Navarro, Frenk & White 1997) and subsequently, we add the gaseous component similar to the set-up of the hydrostatic test (Section 4.1). The only change is that here, we give the gas and dark matter a rotational velocity which peaks at $180 \text{ km s}^{-1}$. Obviously, the initial hydrostatic equilibrium is broken by the onset of the gas cooling and we follow the evolution of the cloud for 10 Gyr.

Fig. 17 visualizes the spatial distribution of stars, where the colours represent the age of stars (top panels), and the spatial distribution of star-forming gas, where the colours represent the star formation rate (bottom panels) at time $t = 7.5$ Gyr. We use the ray-tracing programme SPLOTCH (Dolag et al. 2008; Jin et al. 2010) to create the images and choose a linear colour bar for stellar age, where the red end of the colour bar corresponds to stars older than 3 Gyr and the blue end to recently formed stars. We visualize the star formation rate since it traces the (cold) gas within the disc and
Figure 18. Idealized galaxy formation. We show the vertical (\(z\)) density profile (left-hand panel), the radial density profile (middle panel) and vertical velocity dispersion (right-hand panel) of the stellar component. For the radial plots, we use cylindrical bins. In the ‘new’ scheme (red lines), the galactic disc is more defined, extended and colder.

In the ‘standard’ scheme, the galaxy shows a prominent bulge containing a large fraction of the stellar population. The entire galaxy appears more spheroidal with a dominant bulge and the stellar disc is not well pronounced. We find similar features in the distribution of star formation. The gas disc is asymmetric and only shows little spiral structure in the face-on view. In the edge-on projection the disc shows a rolling pin morphology. Both discs are dominated by bulges, but in the ‘new’ scheme the bulge is significantly less dominant. The bulge contains a smaller fraction of the stellar population and might eventually be associated with an elliptical bar structure. More as well as younger stars are present within the disc. The gas disc is symmetric and shows a defined spiral structure. At this stage of code testing it is difficult to track down the impact of individual code changes. However, we assume the most significant differences are caused as follows. In the ‘standard’ scheme, large amounts of \(A_V\) might lead to dissipation of kinetic energy, loss of rotational support and numerical angular momentum transport. Additionally, the mixing problem and its associated numerical surface tension tend to confine cold and dense gas blobs.

The ‘new’ scheme, significantly smaller amounts of \(A_V\) are applied and rotational support can be provided. Furthermore, the inclusion of AC promotes gas mixing between hot and cold phases.

We continue with a more quantitative comparison of both schemes, which confirms our previous findings. Fig. 18 shows density in vertical and radial direction (\(\rho_z\) and \(\rho_r\)) of the stellar component as well as the associated vertical velocity dispersion (\(\sigma_z\)) at times 2.5 (dashed lines), 5.0 (dotted lines) and 7.5 Gyr (solid lines). As seen in \(\rho_z\) (left-hand panel), the ‘new’ scheme produces a thinner stellar disc and as seen in \(\rho_r\) (middle panel) the disc also extends to significantly larger radii. This trend is confirmed by \(\sigma_z\) (right-hand panel), which for the ‘standard’ scheme truncates at smaller radii than for the ‘new’ scheme.

Fig. 19 shows vertical and radial profiles of gas density (\(\rho_z\) and \(\rho_r\)) as well as the vertical gas velocity dispersion (\(\sigma_z\)) of the cold gas at times 2.5 (dashed lines), 5.0 (dotted lines) and 7.5 Gyr (solid lines). We employ a temperature criterion of \(T < 10^5\) K to distinguish between cold and hot gas. \(\rho_z\) decreases towards the centre since the gas within the bulge is hot and exceeds our temperature threshold. In the ‘new’ scheme, the distribution of cold gas is slightly more extended in vertical as well as radial direction. However, \(\sigma_z\) indicates a colder gas disc. Most probably, these features are a result of less numerically induced \(A_V\) and angular momentum transport and depression of rotational support. Furthermore, the inclusion of
5.2 Santa Barbara cluster

We carried out the Santa Barbara galaxy cluster (Frenk et al. 1999), which is a common reference simulation for cosmological hydrodynamical simulation codes. Although no analytic solutions exists, the cluster has been simulated with a large variety of different codes. The simulation describes the formation of a massive dark matter halo, with virial mass of $1.2 \times 10^{15} M_{\odot}$ and virial radius of 2.8 Mpc. It is evolved in an Einstein-de Sitter cold dark matter cosmology with parameters of $\Omega_M = 1.0$, $\Omega_{\Lambda} = 0.0$, and $H_0 = 50$ km s$^{-1}$ Mpc$^{-1}$. We choose an initial redshift of $z = 20$ with a perturbed distribution of $256^3$ dark matter particles and $256^3$ gas particles, each of equal masses (see also Frenk et al. 1999, for a detailed description of the initial conditions), and follow the formation until redshift $z = 0$.

Fig. 20 shows thin slices of gas density (left-hand panels), temperature (middle panels) and entropy (right-hand panels) defining the thermodynamical state of the hot intracluster medium (ICM) at redshift $z = 0$. For the definition of entropy, we use $S = T/n_e^{2/3}$, which is commonly used in X-ray studies of the ICM (e.g. Kravtsov & Borgani 2012). From the maps, we note the following interesting features.

The gas density (left-hand panels) tends to be smoother in the ‘new’ scheme, which is mostly due to the effect of AC, which introduces entropy mixing among neighbouring gas particles. In contrast, the ‘standard’ scheme produces a clumpy distribution of gas with gas inhomogeneities associated with stripping from merging haloes and cold blobs. These structures are persistent in the hot ICM mainly due to the lack of mixing. In turn, these ‘features’ of the ‘standard’ scheme prevent an efficient action of hydrodynamical instabilities such as Rayleigh–Taylor and Kelvin–Helmholtz instabilities that are spuriously inhibited. Quite remarkably, the clumps are much less evident in the ‘new’ scheme, which also produces a lower value for the central gas density. In the temperatures slices (middle panels), it becomes clear that gas clumps in the ‘standard’ scheme correspond to objects of low temperature. As expected, the effect of introducing AC is the reduction of the degree of ICM thermal complexity.

However, in the ‘new’ scheme the bow-shock, which is induced by the infall of a large substructure on to the main halo is better defined than in the ‘standard’ scheme. The bow-shock is located on the right-hand side of the main halo centre (see Fig. 20). In fact, the WC4 kernel of the ‘new’ scheme captures the entropy discontinuity...
Figure 21. Santa Barbara cluster. We show radial profiles of gas density (left-hand panel), temperature (middle panel) and entropy (right-hand panel). In each panel, we compare the results of the ‘standard’ (blue lines) and the ‘new’ (red lines) scheme to a reference solution (black lines) obtained with a piecewise parabolic grid computation with the MASCLET code (see Quilis 2004). The ‘standard’ scheme does not produce an entropy core but a diverging profile towards the halo centre. In the ‘new’ scheme an entropy core as well as stable temperature and density profiles are reached, which are all in good agreement to the grid code computation.

associated with the shock better. Most importantly, we note from the entropy maps (right-hand panels) that the entropy level in the innermost region of the clusters increases in the ‘new’ scheme.

Fig. 21 shows radial profiles of gas density, temperature and entropy in the same units as shown in Fig. 20 at redshift \( z = 0 \) for both SPH schemes. Furthermore, for a comparison, we also include results obtained with the MASCLET grid code (see Quilis 2004). Both schemes produce quite consistent results at relatively large radii (>200 kpc h\(^{-1}\)) but show striking differences in the innermost regions. The ‘new’ scheme predicts a flatter central density profile, also with no evidence for the inversion of the temperature gradient produced by the ‘standard’ scheme. Density and temperature profiles for the ‘new’ scheme combine to produce a flat entropy core, which extends out to \( \approx\)100 kpc h\(^{-1}\). In the ‘standard’ scheme, the persistence of cold and dense clumps in the cluster atmosphere causes their low-entropy gas to sink to the centre of the cluster, thereby causing the continuous decrease of entropy. In the ‘new’ scheme, AC promotes mixing of gas phases and helps to dissolve low-entropy gas blobs within the hot ICM atmosphere and causes a higher entropy level to be established.

The results for the ‘new’ scheme are remarkably similar to those of the MASCLET code and, more in general, reported by Frenk et al. (1999), Vazza (2011) and Power et al. (2014) for Eulerian codes. We point out that such a close agreement has been obtained without any tuning aimed at producing the entropy core predicted by Eulerian codes in cosmological simulations for the formation of galaxy clusters. The choice of parameters for the ‘new’ SPH scheme was only aimed at preventing the limitations of ‘standard’ SPH in terms of the description of discontinuities and efficiency to capture hydrodynamical instabilities. Note also that these results for the Santa Barbara cluster are in qualitative agreement with those obtained in the hydrostatic sphere (see Section 4.1). The behaviour of both schemes are in the same direction, even if they are less evident than in this Santa Barbara cluster test, due to the lack of the hierarchical process of structure formation within a cosmological environment.

Additionally, we analyse the simulation also at redshifts \( z > 0 \). In general, the profiles of the high-redshift haloes show similar behaviour as their low-redshift counterparts, provided that we choose quiet and virialized objects. Objects, which host dynamically important shocks or undergo merger events show altered radial profiles because the timing of the mergers depends slightly on the simulation scheme. Therefore, a sensible comparison of the entire redshift-evolution and behaviour of both schemes during these violent phases of structure formation is not possible and requires controlled experiments.

6 SUMMARY AND CONCLUSIONS

In this paper, we presented a novel implementation of the SPH scheme in the GADGET code, which provides improved accuracy for simulations of galaxies and large-scale cosmic structures. Since the first development of SPH great advancements have been made to improve the reliability and stability of this hydrodynamical scheme and, in particular, much effort has been spent in a proper treatment of discontinuities. We implemented and improved several of these modifications of SPH into the developer version of GADGET-3, and tested them against a number of standard hydrodynamical problems, as well as first simple astrophysical applications. The main modifications (see also Table 1) of this ‘new’ scheme, when compared to the ‘standard’ (see e.g. Price 2012a) formulation of SPH, can be summarized as follows.

(i) \( A\) \( V\) is introduced for a proper description of shocks. It prevents particle interpenetration into unshocked regions and provides a regularization of the particle field, which supports a proper sampling of the fluid. First, spatially constant low-order formulations of \( A\) \( V\) introduce viscosity not only at shocks, but also within unshocked regions and shearing flows, thereby leading to an overly viscous behaviour and a too fast dissipation of kinetic energy. Most commonly, the so called Balsara switch (Balsara 1995) is used to reduce viscosity in shear flows, while further attempts were made to reduce \( A\) \( V\) where it is unwanted (Morris & Monaghan 1997; Dolag et al. 2005). Recently, modern formulations of \( A\) \( V\) (Cullen & Dehnen 2010; Hu et al. 2014) improved greatly on a correct detection of shocks and use high-order gradient estimators to calculate divergence and curl of velocity from the full-velocity gradient matrix instead of the classical SPH estimators. This allows shear flow limiters, such as the Balsara one, to work more accurately and suppress \( A\) \( V\) outside shocks and in shearing flows. In this way, kinetic energy is better preserved, thus helping simulating turbulent flows or hydrodynamical instabilities with higher accuracy. In our ‘new’ scheme, we compute the velocity gradients from the full velocity gradient matrix instead of low-order classic kernel derivatives.
(ii) Artificial conductivity (AC) is introduced to provide a proper fluid description at contact discontinuities. In fact, in the density–entropy formulation a spurious surface tension arises at discontinuities, which also suppresses the formation of instabilities and prevents mixing of different fluid phases. Lately, AC (see e.g. Price 2008) or pressure–entropy formulations (Hopkins 2013; Saitoh & Makino 2013) were proposed to overcome these issues. AC is applied at contact discontinuities and promotes the transport of heat between particles. However, in the presence of gravitationally induced pressure or temperature gradients, common AC schemes might lead to unwanted transport opposite to the gravitational force. Therefore, numerical limiters are necessary to be included in AC. In our ‘new’ scheme, we include locally adaptive AC to transport heat and treat contact discontinuities in SPH and we limit the amount of AC by correcting for gravitationally induced pressure gradients. While we demonstrated that our AC model is quite efficient at reducing such a surface tension, admittedly a small residual effect is still present, and could potentially impact the long-term stability by overdiffusion.

(iii) As for the choice of the interpolating kernel, the commonly employed cubic spline function has been shown to become easily unstable, which leads to spurious pairs of particles, incorrect gradient estimators and, in general, a poor fluid sampling. Therefore, a change of the kernel function is highly recommended, where commonly the Wendland kernels (Dehnen & Aly 2012) are now used. In our ‘new’ scheme, we employ the Wendland C4 kernel function with 200 neighbours instead of a cubic spline with 64 neighbours. We calculate the density in a classic fashion from the mass distribution of particles and also compute the hydrodynamical forces with the density–entropy formulation.

(iv) At last, within supersonic shocks highly dynamical and computationally active particles can penetrate into regions containing computationally inactive particles causing distortions in the fluid sampling and incorrect hydrodynamical solutions. In our ‘new’ scheme, we use a particle wake-up time-step limiting scheme (see Saitoh & Makino 2009; Pakmor et al. 2012) as a solution, so as to shorten the time-steps whenever necessary and allow particles to become active earlier.

To highlight the improvements associated with this advanced SPH implementation, we investigate both the new and the original scheme in a variety of hydrodynamical standard tests, with and without gravity. Furthermore, we study the behaviour in the cosmological problem of the formation of galaxy cluster and enable simple prescriptions for radiative cooling, supernova feedback and star formation for a test simulation of an isolated rotating disc galaxy. Table 2 presents an overview of our test problems and shows if SPH modules are important in a standard (✓) or improved (✓(imp.)) with respect to GADGET-SPH without our modifications. Furthermore, we list the probed physical features of each test.

The inclusion of AC in SPH changes the thermodynamical evolution of density, internal energy and pressure. Additionally, physical conduction (see e.g. Arth et al. 2014) is also sometimes employed in cosmological SPH simulations to promote (an)isotropic heat transport, which also helps to overcome the limitations of ‘standard’ GADGET-SPH. The joint effect of artificial conduction, introduced for purely numerical reasons, and of physical conduction awaits further investigations. In summary, the ‘new’ GADGET-SPH scheme presented here performs better than the ‘standard’ one in every single of our test simulations. Therefore, it provides a much improved numerical description for weakly collisionless plasmas in cosmological simulations down to galactic scales. We base our future simulations of galaxies and large-scale cosmic structures on this updated formulation of SPH. We will also carry out detailed studies of galactic magnetic fields (see Beck et al. 2013, 2014) and the IC with this advanced method. In view of these applications, it is important to verify how this new SPH implementation performs when compared to other variants of SPH and to Eulerian codes. To this purpose, this SPH implementation participated to the nifty cosmology comparison project (Sembolini et al. 2015), which compares the performances of different hydrodynamical codes in cosmological re-simulations of galaxy clusters. In that comparison project, our code is shown to agree very well to both Eulerian codes and modern SPH implementations on the radial profiles of gas density, temperature and entropy. Given the improvements in the description of hydrodynamics provided by the new SPH implementation presented here, we regard it as the core of an efficient code for modern simulations of cosmic structure formation.

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