“I went on to test the program in every way I could devise. I strained it to expose its weaknesses. I ran it for high-mass stars and low-mass stars, for stars born exceedingly hot and those born relatively cold. I ran it assuming the superfluid currents beneath the crust to be absent – not because I wanted to know the answer, but because I had developed an intuitive feel for the answer in this particular case. Finally I got a run in which the computer showed the pulsar’s temperature to be less than absolute zero. I had found an error. I chased down the error and fixed it. Now I had improved the program to the point where it would not run at all.”

Frozen Star: Of Pulsars, Black Holes and the Fate of Stars
George Greenstein

3

Smoothed Particle Hydrodynamics

3.1 Introduction

The standard approach to solving the equations of fluid dynamics numerically is to define fluid quantities on a regular spatial grid, computing derivatives using finite difference or finite volume schemes. This is an extremely well studied approach and most ‘state of the art’ methods for fluid dynamics have been developed in this manner. In astrophysical fluid dynamics problems frequently involve changes in spatial, temporal and density scales over many orders of magnitude. Thus, adaptivity is an essential ingredient which is absent from a fixed-grid approach. Progress in this area has been rapid in recent years with the development of procedures for adaptive mesh refinement (AMR). The implementation of such procedures is far from trivial, although the availability of libraries and toolkits for grid-based codes eases this burden somewhat. However, a further constraint is that astrophysical problems are frequently asymmetric which can result in substantial numerical diffusion when solving on (fixed or adaptive) Cartesian grids. Other approaches to this problem are to use unstructured grids (where typically the grid is reconstructed at each new timestep) or Lagrangian grid methods, where the grid shape deforms according to the flow pattern.

An alternative to all of these methods is to remove the spatial grid entirely, resulting in methods which are inherently adaptive. In this approach fluid quantities are carried by a set of moving interpolation points which follow the fluid motion. Since each point carries a fixed mass, the interpolation points are referred to as ‘particles’. Derivatives are evaluated either by interpolation over neighbouring particles (referred to as particle methods), or via a hybrid approach by interpolation to an overlaid grid (referred to as particle-mesh methods, typified by the particle-in-cell (PIC) method used extensively in plasma physics.

Smoothed Particle Hydrodynamics (SPH) is a particle method introduced by Lucy (1977) and Gingold
and Monaghan (1977). It has found widespread use in astrophysics due to its ability to tackle a wide range of problems involving complex, asymmetric phenomena with relative ease. Since these features are highly desirable in many non-astrophysical applications, it is unsurprising that SPH is currently finding many applications in other fields such as geophysics and engineering (and even film-making\textsuperscript{1}).

The advantages of SPH over standard grid based approaches can be summarised as follows: Firstly, SPH is conceptually both simple and beautiful. All of the equations can be derived self-consistently from physical principles with a few basic assumptions. As a result complex physics is relatively simple to incorporate. Its simplicity means that for the user it is a very intuitive numerical method which lends itself easily to problem-specific modifications. Secondly, adaptivity is a built-in feature. The Lagrangian nature of the method means that changes in density and flow morphology are automatically accounted for without the need for mesh refinement or other complicated procedures. As a result of its adaptivity, SPH is also very efficient in that resolution is concentrated on regions of high density, whilst computational effort is not wasted on empty regions of space. Thirdly, free boundaries, common in astrophysical problems, are simple and natural in SPH but often present difficulties for grid-based codes (such as spurious heating from the interaction with a low density surrounding medium). This means that no portions of fluid can be lost from the simulation, unlike in a grid based code where fluid which has left the grid cannot return (this has been dubbed the ‘Columbus effect’ by Melvyn Davies, since fluid can fall off the edge of the world). Fourthly, a significant advantage in an astrophysical context is that SPH couples naturally with widely used N-Body codes and techniques, for which there exists a vast amount of literature. Finally (although perhaps many more advantages could be given) visualisation and analysis is also somewhat easier with Lagrangian techniques, since it is a simple matter to track and visualise portions of the flow.

SPH also has a number of disadvantages when compared to finite difference codes. The first of these is that, unlike grid-based codes, SPH involves the additional computational cost of constructing the neighbour lists. This is offset somewhat in that N-Body techniques used to calculate the gravitational force (namely via tree-codes) can also be used in constructing the neighbour lists. Secondly, SPH suffers from a lack of algorithm development, since a vast amount of research effort is focussed on finite difference or finite volume techniques. This often means that such techniques, although often applicable in an SPH context, can be slow to filter into mainstream use. Thirdly, although not a disadvantage as such but a point which is often overlooked, is that the setup of initial conditions is often more complicated and requires much greater care. Since particles can be laid down in an infinite variety of ways, choosing an appropriate setup for a given problem requires some experience and usually some experimentation. Inappropriate particle setups can lead to poorer simulation results than might otherwise be expected (we give some examples of this in §3.7.5). Finally, in the case of magnetohydrodynamics and other problems involving anisotropic stresses (as we will discuss in chapter 4), numerical stability can become an issue which must be dealt with appropriately.

In this chapter we provide an overview of the SPH method, including several improvements to the basic method which have been made since the review article of Monaghan (1992) was published (such as improvements in shock-capturing techniques and the treatment of terms related to the use of a variable smoothing

\textsuperscript{1}for example many of the graphics involving fluids in the film ‘Tomb Raider’ were computed using SPH
length). In particular we focus on those aspects of the algorithm that are relevant in an MHD context. The chapter is organised as follows: In section §3.2 we present the basic formalisms inherent to SPH; in §3.3 we derive the SPH equations for compressible hydrodynamics using a variational principle. Formulations of dissipative terms used to capture shocks are presented and discussed in §3.5. In §3.3.4 we discuss the incorporation of terms relating to the spatial variation of the smoothing length and in §3.4 alternative formulations of SPH are examined within the variational framework. Timestepping is discussed in §3.6. Finally, we present numerical tests in §3.7 in support of the previous sections and as preliminaries for the MHD tests described in Chapters 4 and 5.

3.2 Basic formalisms

3.2.1 Interpolant

The basis of the SPH approach is given as follows (Monaghan, 1992). We begin with the trivial identity

$$A(r) = \int A(r') \delta(|r - r'|) \, dr',$$

(3.1)

where $A$ is any variable defined on the spatial co-ordinates $r$ and $\delta$ refers to the Dirac delta function. This integral is then approximated by replacing the delta function with a smoothing kernel $W$ with characteristic width $h$, such that

$$\lim_{h \to 0} W(r - r', h) = \delta(r - r'),$$

(3.2)

giving

$$A(r) = \int A(r') W(|r - r'|, h) \, dr' + O(h^2).$$

(3.3)

The kernel function is normalised according to

$$\int W(r - r', h) \, dr' = 1.$$

(3.4)

Finally the integral (3.3) is discretised onto a finite set of interpolation points (the particles) by replacing the integral by a summation and the mass element $\rho \, dV$ with the particle mass $m$, ie.

$$A(r) = \int \frac{A(r')}{\rho(r')} W(|r - r'|, h) \rho(r') \, dr' + O(h^2),$$

$$\approx \sum_{b=1}^{N} m_b \frac{A_b}{\rho_b} W(|r - r_b|, h),$$

(3.5)

Footnote: It is interesting to note that this equation, with $A = \rho$ is used to define the density of the fluid in terms of the Lagrangian co-ordinates in the Hamiltonian description of the ideal fluid (eq. (94) in Morrison, 1998). Similarly the SPH equivalent of this expression, (3.42), forms the basis for the Hamiltonian description of SPH (see §3.3.2).
where the subscript $b$ refers the quantity evaluated at the position of particle $b$. This ‘summation interpolant’ is the basis of all SPH formalisms. The errors introduced in this step are discussed in §3.2.2. Gradient terms may be calculated by taking the analytic derivative of (3.5), giving

$$
\nabla A(r) = \frac{\partial}{\partial r} \int \frac{A(r')}{\rho(r')} W(|r-r'|, h) \rho(r') dr' + O(h^2), \quad (3.6)
$$

$$
\approx \sum_b m_b \frac{A_b}{\rho_b} \nabla_a W_{ab}, \quad (3.7)
$$

where we have assumed that the gradient is evaluated at another particle $a$ (i.e. $r = r_a$), defining $\nabla_a \equiv \partial / \partial r_a$ and $W_{ab} \equiv W(|r_a - r_b|, h).

### 3.2.2 Errors

The errors introduced by the approximation (3.3) can be estimated by expanding $A(r')$ in a Taylor series about $r$ (Benz, 1990; Monaghan, 1992), giving

$$
A(r) = \int \left[ A(r) + (r' - r)^\alpha \frac{\partial A}{\partial r^\alpha} + \frac{1}{2} (r' - r)^\beta (r' - r)^\gamma \frac{\partial^2 A}{\partial r^\beta \partial r^\gamma} + O((r - r')^3) \right] W(|r-r'|, h) dr',
$$

$$
= A(r) + \frac{\partial A}{\partial r^\alpha} \int (r' - r)^\alpha W(r) dr' + \frac{1}{2} \frac{\partial^2 A}{\partial r^\beta \partial r^\gamma} \int (r' - r)^\beta (r' - r)^\gamma W(r) dr' + O((r' - r)^3),
$$

where $r \equiv |r' - r|; \alpha, \beta$ and $\gamma$ are indices denoting co-ordinate directions (with repeated indices implying a summation) and we have used the normalisation condition (3.4). The odd error terms are zero if $W$ is an even function of $(r - r')$ (i.e. depending only on its magnitude), which, since $|r - r'|$ is always less than the smoothing radius ($2h$ in most cases), results in an approximation to $O(h^2)$. In principle it is also possible to construct kernels such that the second moment is also zero, resulting in errors of $O(h^4)$ (discussed further in §3.2.7). The disadvantage of such kernels is that the kernel function becomes negative in some part of the domain, resulting in a potentially negative density evaluation. The errors in the summation interpolant differ slightly since the approximation of integrals over summations over particles no longer guarantees that these terms integrate exactly. Starting from the summation interpolant evaluated on particle $a$, we expand $A_b$ in a Taylor series around $r_a$, giving

$$
\sum_b m_b \frac{A_b}{\rho_b} W_{ab} = A_a \sum_b m_b W_{ab} + \nabla A_a \cdot \sum_b m_b (r_b - r_a) W_{ab} + O((r_b - r_a)^2). \quad (3.9)
$$

From this we see that the summation interpolation is exact for constant functions only when the interpolant is normalised by dividing the interpolation of unity. In practical calculations the summation interpolant is only used in the density evaluation (§3.3.1), resulting in a slight error in the density value. More important are the errors resulting from the SPH evaluation of derivatives, since these are used throughout in the discretisation of the fluid equations (§3.3).

The errors resulting from the gradient evaluation (3.6) may be estimated in a similar manner by again
expanding $A(r')$ in a Taylor series about $r$, giving

$$
\nabla A(r) = \int \left[ A(r) + (r' - r) \frac{\partial A}{\partial r^\alpha} + \frac{1}{2} (r' - r)^\beta (r' - r)^\gamma \frac{\partial^2 A}{\partial r^\beta \partial r^\gamma} + \mathcal{O}[(r' - r)^3] \right] \nabla W(|r - r'|, h) dr',
$$

where we have used the fact that $\int \nabla W d\mathbf{r}' = 0$ for even kernels, whilst the second term integrates to unity for even kernels satisfying the normalisation condition (3.4). The resulting errors in the integral interpolant for the gradient are therefore also of $\mathcal{O}(h^2)$. The errors in the summation interpolant for the gradient (3.7) are given by expanding $A_b$ in a Taylor series around $r_a$, giving

$$
\nabla A_a = \sum_b m_b \frac{A_b - A_a}{\rho_b} \nabla_a W_{ab},
$$

where the summations represent SPH approximations to the integrals in the second line of (3.10).

### 3.2.3 First derivatives

From (3.11) we immediately see that a straightforward improvement to the gradient estimate (3.7) can be obtained by a simple subtraction of the first error term (i.e. the term in (3.11) that is present even in the case of a constant function), giving (Monaghan, 1992)

$$
\nabla A_a = \sum_b m_b \frac{(A_b - A_a)}{\rho_b} \nabla_a W_{ab},
$$

which is an SPH estimate of

$$
\nabla A(r) = \nabla A - A(\nabla 1).
$$

Since the first error term in (3.11) is removed, the interpolation is exact for constant functions and indeed this is obvious from the form of (3.12). The interpolation can be made exact for linear functions by dividing by the summation multiplying the first derivative term in (3.11), i.e.

$$
\frac{\partial A_a}{\partial r^\alpha} = \chi_{\alpha\beta} \sum_b \frac{m_b (A_b - A_a)}{\rho_b} \nabla^\beta W_{ab}, \quad \chi_{\alpha\beta} = \left[ \sum_b \frac{m_b (r_b - r_a)^\alpha \nabla^\beta W_{ab}}{\rho_b} \right]^{-1}.
$$
where $\nabla^\beta \equiv \partial / \partial r^\beta$. This normalisation is somewhat cumbersome in practice, since $\chi$ is a matrix quantity, requiring considerable extra storage (in three dimensions this means storing $3 \times 3 = 9$ extra quantities for each particle) and also since calculation of this term requires prior knowledge of the density. However, for some applications of SPH (e.g. solid mechanics) it is desirable to do so in order to retain angular momentum conservation in the presence of anisotropic forces (Bonet and Lok, 1999).

A similar interpolant for the gradient follows by using

$$
\nabla A = \frac{1}{\rho} [A \nabla \rho - \nabla (\rho A)]
$$

(3.15)

$$
\approx \frac{1}{\rho_a} \sum_b m_b (A_b - A_a) \nabla_a W_{ab},
$$

(3.16)

which again is exact for a constant $A$. Expanding $A_b$ in a Taylor series, we see that in this case the interpolation of a linear function can be made exact using

$$
\frac{\partial A_a}{\partial r^\alpha} = \chi_{\alpha \beta} \sum_b m_b (r_b - r_a)^\alpha \nabla^\beta W_{ab}, \quad \chi_{\alpha \beta} = \sum_b m_b (r_b - r_a)^\alpha \nabla^\beta W_{ab}^{-1}
$$

(3.17)

which has some advantages over (3.14) in that it can be computed without prior knowledge of the density.

An alternative gradient interpolant is given by

$$
\nabla A(r) = \rho \left[ \frac{A}{\rho^2} \nabla \rho + \nabla \left( \frac{A}{\rho} \right) \right]
$$

$$
\approx \rho_a \sum_b m_b \left( \frac{A_a}{\rho_a^2} + \frac{A_b}{\rho_b^2} \right) \nabla_a W_{ab}
$$

(3.18)

which is commonly used in the SPH evaluation of the pressure gradient since it guarantees conservation of momentum by the pairwise symmetry in the gradient term. It is also the formulation of the pressure gradient which follows naturally in the derivation of the SPH equations from a variational principle ($\S$3.3.2). Expanding $A_b$ in a Taylor series about $r_a$ we have

$$
\sum_b m_b \left( \frac{A_a}{\rho_a^2} + \frac{A_b}{\rho_b^2} \right) \nabla_a W_{ab} = A_a \sum_b m_b \left( \frac{1}{\rho_a^2} + \frac{1}{\rho_b^2} \right) \nabla_a W_{ab} + \frac{\partial A_a}{\partial r^\alpha} \sum_b m_b \frac{1}{\rho_b^2} (r_b - r_a)^\alpha \nabla_a W_{ab}
$$

$$
+ \frac{1}{2} \frac{\partial^2 A_a}{\partial r^\beta \partial r^\gamma} \sum_b m_b \frac{1}{\rho_b^2} (r_b - r_a)^\beta (r_b - r_a)^\gamma \nabla_a W_{ab} + O((r_b - r_a)^3)
$$

(3.19)

from which we see that for a constant function the error is governed by the extent to which

$$
\sum_b m_b \left( \frac{1}{\rho_a^2} + \frac{1}{\rho_b^2} \right) \nabla_a W_{ab} \approx 0.
$$

(3.20)

Although a simple subtraction of the first term in (3.19) from (3.18) eliminates this error, the symmetry in the gradient necessary for the conservation of momentum is lost by doing so. Retaining the exact conservation of momentum therefore requires that such error terms are not eliminated. In applications of SPH employing
anisotropic forces (such in the MHD case), these error terms can be sufficient to cause numerical instabilities (§4.4).

Derivatives of vector quantities follow in a similar manner. For example the divergence of a vector quantity \( \mathbf{v} \) can be estimated using

\[
\nabla \cdot \mathbf{v} \approx -\frac{1}{\rho_a} \sum_b m_b (v_a - v_b) \cdot \nabla_a W_{ab},
\]

or

\[
\nabla \cdot \mathbf{v} \approx \rho_a \sum_b m_b \left( \frac{v_a}{\rho_a^2} + \frac{v_b}{\rho_b^2} \right) \cdot \nabla_a W_{ab},
\]

whilst the curl is given by (e.g.)

\[
\nabla \times \mathbf{v} \approx -\frac{1}{\rho_a} \sum_b m_b (v_a - v_b) \times \nabla_a W_{ab}.
\]

### 3.2.4 Second derivatives

Second derivatives are slightly more complicated since for kernels with compact support a straightforward estimation using the second derivative of the kernel proves to be very noisy and sensitive to particle disorder. For this reason it is better to use approximations of the second derivative which utilise only the first derivative of the kernel (Brookshaw, 1985; Monaghan, 1992). For a scalar quantity the second derivative may be estimated using the integral approximation

\[
\nabla^2 A(\mathbf{r}) \approx 2 \int [A(\mathbf{r}) - A(\mathbf{r}')] \frac{(\mathbf{r} - \mathbf{r}') \cdot \nabla W(\mathbf{r})}{|\mathbf{r} - \mathbf{r}'|^2} d\mathbf{r}',
\]

giving the SPH Laplacian

\[
\langle \nabla^2 A \rangle_a \approx 2 \sum_b m_b \frac{(A_a - A_b)}{\rho_b} \frac{\mathbf{r}_{ab} \cdot \nabla_a W_{ab}}{|\mathbf{r}_{ab}|^2},
\]

where \( \mathbf{r}_{ab} \equiv \mathbf{r}_a - \mathbf{r}_b \). This formalism is commonly used for heat conduction in SPH (e.g. Brookshaw 1985; Cleary and Monaghan 1999 and more recently Jubelgas et al. 2004). The integral approximation (3.24) can be derived by expanding \( A(\mathbf{r}') \) to second order in a Taylor series about \( \mathbf{r} \), giving

\[
A(\mathbf{r}) - A(\mathbf{r}') = (\mathbf{r} - \mathbf{r}')^\alpha \frac{\partial A}{\partial \mathbf{r}^\alpha} + \frac{1}{2} (\mathbf{r} - \mathbf{r}')^\alpha (\mathbf{r} - \mathbf{r}')^\beta \frac{\partial^2 A}{\partial \mathbf{r}^\alpha \partial \mathbf{r}^\beta} + \mathcal{O}[(\mathbf{r} - \mathbf{r}')^3].
\]

Expanding this expression into (3.24), the integral is given by

\[
\frac{\partial A}{\partial \mathbf{r}^\alpha} \int (\mathbf{r} - \mathbf{r}')^\alpha \frac{(\mathbf{r} - \mathbf{r}') \cdot \nabla W(\mathbf{r})}{|\mathbf{r} - \mathbf{r}'|^2} d\mathbf{r}' + \frac{1}{2} \frac{\partial^2 A}{\partial \mathbf{r}^\alpha \partial \mathbf{r}^\beta} \int (\mathbf{r} - \mathbf{r}')^\alpha (\mathbf{r} - \mathbf{r}')^\beta \frac{(\mathbf{r} - \mathbf{r}') \cdot \nabla W(\mathbf{r})}{|\mathbf{r} - \mathbf{r}'|^2} d\mathbf{r}'.
\]
The first integral is zero for spherically symmetric kernels, whilst the second term integrates to a delta function, giving $\nabla^2 A$. A generalisation of (3.25) is derived for vector quantities by Español and Revenga (2003). In three dimensions the integral approximation is given by

$$\frac{\partial^2 \mathbf{v}}{\partial \mathbf{r}^\alpha \partial \mathbf{r}^\beta} \approx \int \left[ \mathbf{v}(\mathbf{r}) - \mathbf{v}(\mathbf{r}') \right] \left[ 5(\mathbf{r} - \mathbf{r}')^\alpha (\mathbf{r} - \mathbf{r}')^\beta - \delta^{\alpha\beta} \right] \frac{(\mathbf{r} - \mathbf{r}') \cdot \nabla W(\mathbf{r})}{|\mathbf{r} - \mathbf{r}'|^2} d\mathbf{r}',$$

which in SPH form becomes

$$\left( \frac{\partial^2 \mathbf{v}}{\partial \mathbf{r}^\alpha \partial \mathbf{r}^\beta} \right)_a \approx \sum_b m_b \frac{(\mathbf{v}_a - \mathbf{v}_b)}{\rho_b} \left[ 5 \mathbf{r}_{ab}^\alpha \mathbf{r}_{ab}^\beta - \delta^{\alpha\beta} \right] \frac{\mathbf{r}_{ab} \cdot \nabla_a W_{ab}}{r_{ab}^2}. \quad (3.29)$$

### 3.2.5 Smoothing kernels

The smoothing kernel $W$ must by definition satisfy the requirement that it tends to a delta function as the smoothing length $h$ tends to zero (3.2) and the normalisation condition (3.4). In addition the kernel is usually chosen to be an even function of $r$ to cancel the first error term in (3.8) and may therefore be written in the form

$$W(r, h) = \frac{\sigma}{h^\nu} f \left( \frac{r}{h} \right), \quad (3.30)$$

where $r \equiv |\mathbf{r} - \mathbf{r}'|$ and $\nu$ is the number of spatial dimensions. Written in this form the normalisation condition (3.4) becomes

$$\sigma \int f(q)dV = 1, \quad (3.31)$$

where $q = r/h$ and the volume element $dV = dq, 2\pi q dq$ or $4\pi q^2 dq$ in one, two and three dimensions. The simplest kernel with this property is the Gaussian

$$W(r, h) = \frac{\sigma}{h^\nu} e^{-q^2}, \quad (3.32)$$

where $q = r/h$ and $\sigma = [1/\sqrt{\pi}, 1/\pi, 1/(\pi \sqrt{\pi})]$ in [1,2,3] dimensions. This has the advantage that the spatial derivative is infinitely smooth (differentiable) and therefore exhibits good stability properties (Figure 3.2). For practical applications, however, using a Gaussian kernel has the immediate disadvantage that the interpolation spans the entire spatial domain (with computational cost of $O(N^2)$), despite the fact that the relative contribution from neighbouring particles quickly become negligible with increasing distance. For this reason it is far more efficient to use kernels with finite extent (ie. having compact support), reducing the calculation to a sum over closely neighbouring particles which dramatically reduces the cost to $O(nN)$ where $n$ is the number of contributing neighbours (although there is also the additional cost of finding the neighbouring particles). Kernels which are similar to the Gaussian in shape generally give the best performance (see, e.g. Fulk and Quinn, 1996). Of these the most commonly used kernel is that based on cubic splines (Monaghan...
3.2 Basic formalisms

and Lattanzio, 1985), given by

\[ f(q) = \begin{cases} 
1 - \frac{3}{2}q^2 + \frac{3}{7}q^3, & 0 \leq q < 1; \\
\frac{1}{4}(2-q)^3, & 1 \leq q < 2; \\
0, & q \geq 2.
\end{cases} \]  

(3.33)

with normalisation \( \sigma = \left[ \frac{2}{3}, \frac{10}{7\pi}, \frac{1}{\pi} \right] \). This kernel satisfies the basic requirements (3.2) and (3.4), is even, has continuous first derivatives and compact support of size \( 2h \). Smoother kernels can be introduced by increasing the size of the compact support region (which correspondingly increases the cost of evaluation by increasing the number of contributing neighbours) and by using higher order interpolating spline functions.

To this end the quartic spline kernel

\[ f(q) = \begin{cases} 
(2.5 - q)^4 - 5(1.5 - q)^4 + 10(0.5 - q)^4, & 0 \leq q < 0.5; \\
(2.5 - q)^4 - 5(1.5 - q)^4, & 0.5 \leq q < 1.5; \\
(2.5 - q)^4, & 1.5 \leq q < 2.5; \\
0, & q \geq 2.5.
\end{cases} \]  

(3.34)

with normalisation \( \sigma = \left[ \frac{1}{24}, \frac{96}{1199\pi}, \frac{1}{20\pi} \right] \) and quintic spline kernel

\[ f(q) = \begin{cases} 
(3 - q)^5 - 6(2 - q)^5 + 15(1 - q)^5, & 0 \leq q < 1; \\
(3 - q)^5 - 6(2 - q)^5, & 1 \leq q < 2; \\
(3 - q)^5, & 2 \leq q < 3; \\
0, & q \geq 3.
\end{cases} \]  

(3.35)

with normalisation \( \sigma = \left[ \frac{1}{120}, \frac{7}{478\pi}, \frac{1}{120\pi} \right] \) can be used (e.g. Morris, 1996). The higher order polynomials have the advantage of smoother derivatives which, in combination with the increased size of compact support, decreases the sensitivity of the kernel to disorder in the particle distribution (§3.2.7).

Note that it is entirely possible to construct kernels based on smoother splines but which retain compact support of size \( 2h \). We derive a class of such kernels and compare their stability properties with the kernels given in this section in §3.2.6. In principle it is also possible to construct higher order kernels where the second error term in (3.8) is also zero. Monaghan (1992) demonstrates that such higher order kernels may be constructed from any lower order kernel such as (3.33) by the simple relation

\[ W_{\text{highorder}} = B(1 - Aq^2)W(q) \]  

(3.36)

where the parameters \( A \) and \( B \) are chosen to cancel the second moment and to satisfy the normalisation condition (3.4). The disadvantage of all such kernels is that the kernel becomes negative in part of the domain which could result in a negative density evaluation. Also it is not clear that such kernels actually lead to significant improvements in accuracy in practical situations (since the kernel is sampled at only a few points).

From time to time various alternatives have been proposed to the kernel interpolation at the heart of
Figure 3.1: Examples of SPH smoothing kernels (solid line) together with their first (dashed) and second (dot-dashed) derivatives. Kernels correspond to those given in the text. The cubic spline (top left) is the usual choice, whilst the quintic (top, middle) represents a closer approximation to the Gaussian kernel (top right), at the cost of increased compact support. The bottom row correspond to various quintic kernels with compact support of $2h$ which we derive in §3.2.6. The stability properties of all these kernels are compared in Figure 3.2.

SPH, such as the use of Delaunay triangulations (Pelupessy et al., 2003) and normalisations of the kernel interpolant (involving matrix inversion) which guarantee exact interpolations to arbitrary polynomial orders (Maron and Howes, 2003; Bonet and Lok, 1999). It remains to be seen whether any such alternative proposals are viable in terms of the gain in accuracy versus the inevitable increase in computational expense and algorithmic complexity.
Finally we note that in most SPH codes, the kernel is evaluated by linear interpolation from a pre-computed table of values, since kernel evaluations are computed frequently. The computational cost involved in calculating the kernel function is therefore the same whatever the functional form. In the calculations given in this thesis, the kernel is tabulated as $W(q)$ and $\partial W/\partial q$, where the table is evenly spaced in $q^2$ to give a better interpolation in the outer edges.

### 3.2.6 A general class of kernels

In this section we consider the possibility of constructing kernels based on smoother splines than the cubic but which retain compact support of size $2h$. A general class of such kernels may be derived by considering kernels of the form

$$f(q) = \sigma \begin{cases} 
(r-q)^n + A(\alpha-q)^n + B(\beta-q)^n, & 0 \leq q < \beta; \\
(r-q)^n + A(\alpha-q)^n, & \beta \leq q < \alpha; \\
(r-q)^n, & \alpha \leq q < r; \\
0, & q \geq r
\end{cases}$$

where $n$ is the order, $r$ is the compact support size (in this case $r = 2$), $A$ and $B$ are parameters to be determined and $\alpha$ and $\beta$ are the two matching points (with $0 < \beta < \alpha < r$), although an arbitrary number of matching points could be added. The formulation given above guarantees that the kernel and its derivatives are continuous at the matching points and zero at the compact support radius $W(r) = dW/dq(r) = 0$. To determine the parameters $A$ and $B$ we require two further constraints on the form of the kernel. For the kernels to resemble the Gaussian, we constrain the kernel gradient to be zero at the origin and also that the second derivative be minimum at the origin (this also constrains $n \geq 3$), ie.

$$W'(0) = 0, \quad W''(0) = 0. \quad (3.38)$$

For the moment we leave the matching points as free parameters. From the conditions (3.38), the parameters $A$ and $B$ are given in terms of the matching points by

$$A = \frac{r^n - (r^2 - \beta^2)}{\alpha^n - (\alpha^2 - \beta^2)}, \quad B = -\frac{r^{n-1} + A\alpha^{n-1}}{\beta^{n-1}}. \quad (3.39)$$

In one dimension the normalisation constant is given by

$$\sigma = \frac{n + 1}{2(A\alpha^{n+1} + B\beta^{n+1} + r^{n+1})}. \quad (3.40)$$

As an example we can construct a quintic ($n = 5$) kernel that closely resembles the cubic spline kernel (3.33) in all but the continuity of the second derivative. An example of such a kernel is given by the choice $\beta = 0.85, \alpha = 1.87$. This was chosen by constraining the second derivative to be equal to that of the cubic spline at the origin (ie. $W''(0) = -2$) and the turning point in the second derivative to be located as close
Figure 3.2: One dimensional stability properties of the kernels shown in Figure 3.1 for isothermal SPH. The $y$-axis gives the smoothing length in units of the particle spacing $\Delta x$, whilst the $x$-axis corresponds to wavenumber in units of $1/\Delta x$ (such that $kx \to 0$ represents the limit of an infinite number of particles per wavelength and $h \to \infty$ represents the limit of an infinite number of neighbours). Contours show the (normalised) square of the numerical sound speed from the dispersion relation (3.41). The quintic spline (top, centre) and Gaussian kernels show improved accuracy over the standard cubic spline kernel although at a higher computational cost. The kernels derived in §3.2.6 (bottom row) appear to give an improvement in accuracy for $h \gtrsim 1.1$ although degrade rapidly for $h \lesssim 1.1$ where the cubic spline retains a reasonable accuracy as possible to the that of the cubic spline ($W''(q \approx 1) = 0$; note that an exact match is not possible under the constraints given). This kernel is shown in Figure 3.1 (‘cubic-like quintic’). The stability properties are discussed in §3.2.7.

However, it would be more interesting to investigate whether other kernels with even better stability properties can be constructed. To this end we have performed a survey of parameter space for quintic ($n = 5$) kernels, from which we find that the most stable kernels are those with matching points in the range $\beta \approx 0.5$ with $\alpha \approx 1.7$ or $\beta \approx 0.7$ with $\alpha \approx 1.5$. These two kernels (‘New Quintic(1)’ and ‘New Quintic (2)’) are shown in Figure 3.1. The stability properties are discussed below.
3.2.7 Kernel stability properties

The accuracy of the kernels given in §3.2.5 and §3.2.6 may be compared via a stability analysis of the SPH equations. Detailed investigations of the stability properties of SPH have been given elsewhere (e.g. Morris 1996) and for this reason we refer the details of the stability analysis to appendix B (although as for the fluid equations, the linearised form of the SPH equations are derived from a variational principle). The result for one-dimensional SPH (for any equation of state) is the dispersion relation

\[
\omega_a^2 = \frac{2mP_0}{\rho_0} \sum_b [1 - \cos k(x_a - x_b)] \frac{\partial^2 W}{\partial x^2}(x_a - x_b, h) \\
+ \frac{m^2}{\rho_0^2} \left( c_s^2 - \frac{2P_0}{\rho_0} \right) \left[ \sum_b \sin k(x_a - x_b) \frac{\partial W}{\partial x}(x_a - x_b, h) \right]^2,
\]

(3.41)

where \( c_s = \frac{\partial P}{\partial \rho} \) is the sound speed. Figure 3.2 shows contours of the (normalised) square of the numerical sound speed \( C_{num}^2 = \omega^2 / k^2 \) as a function of wavenumber and smoothing length (both in units of the average particle spacing). The sums in (3.41) are calculated numerically assuming an (isothermal) sound speed and particle spacing of unity (both wavelength and smoothing length are calculated in units of the particle spacing). The quintic spline (top, centre) and the Gaussian (top right) show increasingly better stability properties over the standard cubic spline (top left) although at increased computational expense.

The stability properties of the ‘cubic-like’ quintic kernel derived in §3.2.6 (bottom left) are very similar to that of the cubic spline, except that the ‘trough’ in the contours of \( C_{num}^2 \) observed at \( h = 1.5\Delta p \) (where the closest neighbour crosses the discontinuity in the second derivative) is much smoother. However, the accuracy of this kernel appears to degrade for small smoothing lengths (\( h \lesssim 1.1\Delta p \)) where the cubic spline retains a reasonable accuracy. Of the remaining two kernels derived in §3.2.6 (bottom centre and bottom right), the second example (‘New Quintic (2)’) in particular appears to give slightly better accuracy than the cubic spline over the range \( h \gtrsim 1.1\Delta p \) although both kernels show the rapid decline in accuracy for small smoothing lengths (\( h \lesssim 1.1\Delta p \)) observed in the cubic-like quintic. It is worth noting that most multidimensional calculations use smoothing lengths in the range \( h = 1.1 - 1.2\Delta p \).

In summary the new kernels appear to give a small gain in accuracy over the cubic spline kernel, provided \( h \gtrsim 1.1\Delta p \). However, the gain in accuracy from the use of these alternative kernels is very minor compared to the substantial improvements in accuracy gained by the incorporation of the variable smoothing length terms (§3.3.4), which effectively act as a normalisation of the kernel gradient.

3.3 Fluid Equations

3.3.1 Continuity equation

The summation interpolant (3.5) takes a particularly simple form for the evaluation of density, ie.

\[
\rho_a = \sum_b m_b W_{ab}.
\]

(3.42)
Taking the (Lagrangian) time derivative, we obtain
\[ \frac{d\rho_a}{dt} = \sum_b m_b (v_{ba} - v_b) \cdot \nabla_a W_{ab}, \] (3.43)

which may be translated back to continuum form via the summation interpolant (3.5) to give
\[ \frac{d\rho}{dt} = \mathbf{v} \cdot \nabla \rho - \nabla \cdot (\rho \mathbf{v}), \]
\[ = -\rho (\nabla \cdot \mathbf{v}). \] (3.44)

This reveals that (3.43) and therefore (3.42) are SPH expressions for the continuity equation. It is a remarkable fact that the entire SPH formalism can be self-consistently derived using only (3.42) in conjunction with the first law of thermodynamics via a Lagrangian variational principle. Such a derivation demonstrates that SPH has a robust Hamiltonian structure and ensures that the discrete equations reflect the symmetries inherent in the Lagrangian, leading to the exact conservation of momentum, angular momentum and energy.

### 3.3.2 Equations of motion

The Lagrangian for Hydrodynamics is given by (Eckart, 1960; Salmon, 1988; Morrison, 1998)
\[ L = \int \left( \frac{1}{2} \rho v^2 - \rho u \right) dV, \] (3.45)

where \( u \) is the internal energy per unit mass. In SPH form this becomes
\[ L = \sum_b m_b \left[ \frac{1}{2} v_{b}^2 - u_b(\rho_b, s_b) \right], \] (3.46)

where as previously we have replaced the volume element \( \rho dV \) with the mass per SPH particle \( m \). We regard the particle co-ordinates as the canonical variables. Being able to specify all of the terms in the Lagrangian directly in terms of these variables means that the conservation laws will be automatically satisfied, since the equations of motion then result from the Euler-Lagrange equations
\[ \frac{d}{dt} \left( \frac{\partial L}{\partial \mathbf{v}_a} \right) - \frac{\partial L}{\partial x_a} = 0. \] (3.47)

The internal energy is regarded as a function of the particle’s density, which in turn is specified as a function of the co-ordinates by (3.42). The terms in (3.47) are therefore given by
\[ \frac{\partial L}{\partial \mathbf{v}_a} = m_a \mathbf{v}_a, \] (3.48)
\[ \frac{\partial L}{\partial x_a} = \sum_b m_b \left. \frac{\partial u_b}{\partial \rho_b} \right|_s \frac{\partial \rho_b}{\partial x_a}. \] (3.49)
3.3 Fluid Equations

From the first law of thermodynamics in the absence of dissipation we have
\[
\frac{\partial u_b}{\partial \rho_b}\bigg|_s = \frac{P_b}{\rho_b^2},
\] (3.50)
and using (3.42) we have
\[
\frac{\partial \rho_b}{\partial \mathbf{r}_a} = \sum_c m_c \nabla_a W_{bc} (\delta_{ba} - \delta_{ca}),
\] (3.51)
such that
\[
\frac{\partial L}{\partial \mathbf{r}_a} = \sum_b m_b \frac{P_b}{\rho_b^2} \sum_c m_c \nabla_a W_{bc} (\delta_{ba} - \delta_{ca}),
\] (3.52)
\[
= m_a \sum_b m_b \left( \frac{P_a}{\rho_a^2} + \frac{P_b}{\rho_b^2} \right) \nabla_a W_{ab},
\] (3.53)
where we have used the fact that the gradient of the kernel is anti-symmetric (i.e. $\nabla_a W_{ac} = -\nabla_a W_{ca}$). The SPH equation of motion in the absence of dissipation is therefore given by
\[
\frac{d\mathbf{v}_a}{dt} = -\sum_b m_b \left( \frac{P_a}{\rho_a^2} + \frac{P_b}{\rho_b^2} \right) \nabla_a W_{ab},
\] (3.54)
which can be seen to explicitly conserve momentum since the contribution of the summation to the momentum of particle $a$ is equal and opposite to that given to particle $b$ (given the antisymmetry of the kernel gradient). Taking the time derivative of the total angular momentum, we have
\[
\frac{d}{dt} \sum_a \mathbf{r}_a \times m_a \mathbf{v}_a = \sum_a m_a \left( \mathbf{r}_a \times \frac{d\mathbf{v}_a}{dt} \right),
\] (3.55)
\[
= \sum_a \sum_b m_a m_b \left( \frac{P_a}{\rho_a^2} + \frac{P_b}{\rho_b^2} \right) \mathbf{r}_a \times (\mathbf{r}_a - \mathbf{r}_b) F_{ab},
\] (3.56)
where the kernel gradient has been written as $\nabla_a W_{ab} = \mathbf{r}_a F_{ab}$. This last expression is zero since the double summation is antisymmetric in $a$ and $b$ (this can be seen by swapping the summation indices $a$ and $b$ in the double sum and adding half of this expression to half of the original expression, giving zero). Angular momentum is therefore also explicitly conserved.

3.3.3 Energy equation

The energy equation also follows naturally from the variational approach, where we may choose to integrate either the particle’s internal energy $u$, its specific energy $e$ or even its specific entropy $s$. Integrating the specific energy guarantees that the total energy is exactly conserved and it is common practice to use this quantity in finite difference schemes. However the usual argument against this (which applies equally to
finite difference schemes) is that in some circumstances (where the kinetic energy is much greater than
the thermal energy) the thermal energy can become negative by round-off error. Integration of the specific
entropy has some advantages and has been argued for in both SPH (Springel and Hernquist, 2002) and finite
difference schemes (e.g. Balsara and Spicer 1999).

**Internal energy**

The internal energy equation in the absence of dissipation follows from the use of the first law of thermody-
namics (3.50), giving

\[
\frac{du_a}{dt} = \frac{P_a}{\rho_a^2} \frac{d\rho_a}{dt}.
\]  

(3.57)

Using (3.43) therefore gives

\[
\frac{du_a}{dt} = \frac{P_a}{\rho_a^2} \sum_b m_b v_{ab} \cdot \nabla_a W_{ab}.
\]  

(3.58)

**Total energy**

The conserved (total) energy is found from the Lagrangian via the Hamiltonian

\[
H = \sum_a \mathbf{v}_a \cdot \frac{\partial L}{\partial \mathbf{v}_a} - L,
\]  

(3.59)

where using (3.48) and (3.46) we have

\[
H = \sum_a m_a \left( \frac{1}{2} \mathbf{v}_a^2 + u_a \right),
\]  

(3.60)

which is simply the total energy of the SPH particles \(E\) since the Lagrangian does not explicitly depend on
the time. Taking the (Lagrangian) time derivative of (3.60), we have

\[
\frac{dE}{dt} = \sum_a m_a \left( \mathbf{v}_a \cdot \frac{d\mathbf{v}_a}{dt} + \frac{du_a}{dt} \right).
\]  

(3.61)

Substituting (3.54) and (3.58) and rearranging we find

\[
\frac{dE}{dt} = \sum_a m_a \frac{de_a}{dt} = \sum_a \sum_b m_a m_b \left( \frac{P_a}{\rho_a^2} \mathbf{v}_b + \frac{P_b}{\rho_b^2} \mathbf{v}_a \right) \cdot \nabla_a W_{ab},
\]  

(3.62)

and thus the specific energy equation (in the absence of dissipation) is given by

\[
\frac{de_a}{dt} = \sum_b m_b \left( \frac{P_a}{\rho_a^2} \mathbf{v}_b + \frac{P_b}{\rho_b^2} \mathbf{v}_a \right) \cdot \nabla_a W_{ab}.
\]  

(3.63)
Dissipative terms are discussed in §3.5.

**Entropy**

In the case of an ideal gas equation of state where

\[ P = A(s) \rho^\gamma, \]  
(3.64)

the function \( A(s) \) evolves according to

\[
\frac{dA}{dt} = \gamma - 1 \left( \frac{du}{dt} - \frac{P}{\rho^2} \frac{d\rho}{dt} \right). 
\]

This has the advantage of placing strict controls on sources of entropy, since \( A \) is constant in the absence of dissipative terms. The thermal energy is evaluated using

\[
u = \frac{A}{\gamma - 1} \rho^{\gamma - 1}. \]

(3.66)

This formulation of the energy equation has been advocated in an SPH context by Springel and Hernquist (2002).

### 3.3.4 Variable smoothing length terms

The smoothing length \( h \) determines the radius of interaction for each SPH particle. Early SPH simulations used a fixed smoothing length for all particles. However allowing each particle to have its own associated smoothing length which varies according to local conditions increases the spatial resolution substantially (Hernquist and Katz, 1989; Benz, 1990). The usual rule is to take

\[ h_a \propto \left( \frac{1}{\rho_a} \right)^{(1/\nu)}, \]

(3.67)

where \( \nu \) is the number of spatial dimensions, although others are possible (Monaghan, 2000). Implementing this rule self-consistently is more complicated in SPH since the density \( \rho_a \) is itself a function of the smoothing length \( h_a \) via the relation (3.42). A simple approach is to use the time derivative of (3.67), (Benz, 1990), ie.

\[
\frac{dh_a}{dt} = - \frac{h_a}{\nu \rho_a} \frac{d\rho}{dt}, 
\]

(3.68)

which can then be evolved alongside the other particle quantities. This rule works well for most practical purposes, and maintains the relation (3.67) particularly well when the density is updated using the continuity equation (3.43). However, it has been known for some time that, in order to be fully self-consistent, extra terms involving the derivative of \( h \) should be included in the momentum and energy equations (e.g. Nelson
Attempts to do this were, however, complicated to implement (Nelson and Papaloizou, 1994) and therefore not generally adopted by the SPH community. Recently Springel and Hernquist (2002) have shown that the so-called $\nabla h$ terms can be self-consistently included in the equations of motion and energy using a variational approach. Springel and Hernquist (2002) included the variation of the smoothing length in their variational principle by use of Lagrange multipliers, however, in the context of the discussion given in §3.3.2 we note that by expressing the smoothing length as a function of $\rho$ we can therefore specify $h$ as a function of the particle co-ordinates (Monaghan, 2002). That is we have $h = h(\rho)$ where $\rho$ is given by

$$\rho_a = \sum_b m_b W(r_{ab}, h_a).$$

(3.69)

Taking the time derivative, we obtain

$$\frac{d\rho_a}{dt} = \frac{1}{\Omega_a} \sum_b m_b v_{ab} \cdot \nabla_a W_{ab}(h_a),$$

(3.70)

where

$$\Omega_a = \left[1 - \frac{\partial h_a}{\partial \rho_a} \sum_c m_c \frac{\partial W_{ab}(h_a)}{\partial h_a}\right].$$

(3.71)

A simple evaluation of $\Omega$ for the kernel in the form (3.30) shows that this term differs from unity even in the case of an initially uniform density particle distribution (i.e. with constant smoothing length). The effects of this correction term even in this simple case are investigated in the sound wave tests described in §3.7.2.

The equations of motion in the hydrodynamic case may then be found using the Euler-Lagrange equations (3.47) and will therefore automatically conserve linear and angular momentum. The resulting equations are given by (Springel and Hernquist, 2002; Monaghan, 2002)

$$\frac{d\mathbf{v}_a}{dt} = -\sum_b m_b \left[ \frac{p_a}{\Omega_a \rho_a^2} \nabla_a W_{ab}(h_a) + \frac{p_b}{\Omega_b \rho_b^2} \nabla_a W_{ab}(h_b) \right].$$

(3.72)

Calculation of the quantities $\Omega$ involve a summation over the particles and can be computed alongside the density summation (3.69). To be fully self-consistent we solve (3.69) iteratively to determine both $h$ and $\rho$ self-consistently. We do this as follows: Using the predicted smoothing length from (3.68), the density is initially calculated by a summation over the particles. A new value of smoothing length $h_{\text{new}}$ is then computed from this density using (3.67). Convergence is determined according to the criterion

$$\frac{|h_{\text{new}} - h|}{h} < 1.0 \times 10^{-2}.$$  

(3.73)

For particles which are not converged, the density of (only) those particles are recalculated (using $h_{\text{new}}$). This process is then repeated until all particles are converged. Note that a particle’s smoothing length is only set equal to $h_{\text{new}}$ if the density is to be recalculated (this is to ensure that the same smoothing length that was
used to calculate the density is used to compute the terms in the other SPH equations). Also, the density only needs to be recalculated on those particles which have not converged, since each particle’s density is independent of the smoothing length of neighbouring particles. This requires a small adjustment to the density calculation routine (such that the density can be calculated only for a selected list of particles, rather than for all), but is relatively simple to implement and means that the additional computational cost involved is negligible (at least for the problems considered in this thesis). Note that in principle the calculated gradient terms (3.71) may also be used to implement an iteration scheme such as the Newton-Raphson method which converges faster than our simple fixed point iteration.

Where the variable smoothing length terms are not explicitly calculated, we use a simple averaging of the kernels and kernel gradients to maintain the symmetry in the momentum and energy equations (Hernquist and Katz, 1989; Monaghan, 1992), i.e.

\[
W_{ab} = \frac{1}{2} \left[ W_{ab}(h_a) + W_{ab}(h_b) \right],
\]

(3.74)

and correspondingly

\[
\nabla_a W_{ab} = \frac{1}{2} \left[ \nabla_a W_{ab}(h_a) + \nabla_a W_{ab}(h_b) \right].
\]

(3.75)

Many of the test problems in this thesis are performed using this simple formulation. This is in order to show (particularly in the MHD case) that satisfactory results on the test problems are not dependent on the variable smoothing length formulation. In almost every case, however, self-consistent implementation of the variable smoothing length terms as described above leads to a substantial improvement in accuracy (demonstrated, for example, in §3.7 and in the MHD case in §4.6). Perhaps the only disadvantage to the full implementation of the variable smoothing length terms is that the iterations of \( h \) with \( \rho \) mean that small density fluctuations are resolved by the method rather than being smoothed out, which may be disadvantageous under some circumstances (e.g. where the fluctuations are unphysical). One possible remedy for this might be to use a slightly different relationship between \( h \) and \( \rho \) than is given by (3.67).

### 3.4 Alternative formulations of SPH

In §3.3 the SPH equations of motion and energy were derived from a variational principle using only the density summation (3.42) and the first law of thermodynamics (3.50), leading to the equations of motion in the form (3.54) and the energy equation (3.58) or (3.63). However many alternative formulations of the SPH equations are possible and have been used in various contexts. In this section we demonstrate how such alternative formulations may also be derived self-consistently using a variational principle.

For example, a general form of the momentum equation in SPH is given by (Monaghan, 1992)

\[
\frac{d\nabla_a}{dt} = - \sum_b m_b \left( \frac{P_a}{\rho_a^{\sigma} \rho_b^{2-\sigma}} + \frac{P_b}{\rho_b^{\sigma} \rho_a^{2-\sigma}} \right) \nabla_a W_{ab},
\]

(3.76)
which is symmetric between particle pairs for all choices of the parameter $\sigma$ and therefore explicitly conserves momentum. Ritchie and Thomas (2001) use this form of the momentum equation with $\sigma = 1$ in their SPH formalism, finding that it gives slightly better results for problems involving large density contrasts (they also use a slightly different procedure for evaluating the density). Marri and White (2003), for similar reasons, use this equation with $\sigma = 3/2$, citing a reduction in the relative error in the force calculation on particle $a$ due to the influence of particle $b$ which is desirable in the case of particles with large density differences. However, it is apparent from the derivation given in §3.3.2 that forms of this equation other than the standard $\sigma = 2$ case cannot be derived consistently using the density summation (3.42) and correspondingly the continuity equation in the form (3.43). We are therefore led to the conclusion that a consistent formulation of the SPH equations using the general form of the momentum equation given above must involve modification of the continuity equation in some way. We show below that the general form of the continuity equation which is consistent with (3.76) is derived from the continuum equation

$$\frac{d\rho}{dt} = -\rho \nabla \cdot \mathbf{v}, \quad (3.77)$$

expressed in the form

$$\frac{d\rho}{dt} = \rho^{2-\sigma} \left[ \mathbf{v} \cdot \nabla (\rho^{\sigma-1}) - \nabla \cdot (\rho \mathbf{v}^{\sigma-1}) \right], \quad (3.78)$$

with SPH equivalent

$$\frac{d\rho_a}{dt} = \rho_a^{2-\sigma} \sum_b m_b \frac{(\mathbf{v}_a - \mathbf{v}_b)}{\rho_b^{2-\sigma}} \cdot \nabla a W_{ab}. \quad (3.79)$$

In order to demonstrate that this is so, we use this expression for the density to derive the equations of motion and energy via a variational principle.

### 3.4.1 Variational principle

In the derivation given in §3.3.2, the variables in the Lagrangian were explicitly written as a function of the particle co-ordinates (via the identity 3.42), guaranteeing the exact conservation of linear and angular momentum in the equations of motion via the use of the Euler-Lagrange equations. Using a more general form of the continuity equation, however, means that the density can no longer be expressed directly as a function of the particle co-ordinates and therefore that the derivation given in the previous section cannot be applied in this case. However we may still use the Lagrangian to derive the equations of motion by introducing constraints on $\rho$ in a manner similar to that of Bonet and Lok (1999). In this case conservation of momentum and energy can be shown to depend on the formulation of the velocity terms in the continuity equation (in particular that the term should be expressed as a velocity difference). Clearly the major disadvantage of using a continuity equation of any form rather than the SPH summation is that mass is no longer conserved exactly. It is shown in §4.3.2 that the kind of variational principle given below may also be used to derive the equations of motion and energy in the MHD case.
For stationary action we require

\[ \delta \int L \, dt = \int \delta L \, dt = 0, \tag{3.80} \]

where we consider variations with respect to a small change in the particle co-ordinates \( \delta \mathbf{r}_a \). We therefore have

\[ \delta L = m_a \mathbf{v}_a \cdot \delta \mathbf{v}_a - \sum_b m_b \frac{\partial u_b}{\partial \rho_b} \bigg|_s \delta \rho_b. \tag{3.81} \]

The Lagrangian variation in density is given, from (3.79), by

\[ \delta \rho_b = \rho_b^{2-\sigma} \sum_c \frac{m_c}{\rho_c^{2-\sigma}} (\delta \mathbf{r}_b - \delta \mathbf{r}_c) \cdot \nabla_b W_{bc}. \tag{3.82} \]

Using (3.82) and the first law of thermodynamics (3.50) in (3.81) and rearranging, we find

\[ \frac{\delta L}{\delta \mathbf{r}_a} = - \sum_b m_b \frac{P_b}{\rho_b^\sigma} \sum_c \frac{m_c}{\rho_c^{2-\sigma}} \nabla_b W_{bc} (\delta \rho_b - \delta \rho_a). \tag{3.83} \]

Putting this back into (3.80), integrating the velocity term by parts and simplifying (using \( \nabla_a W_{ab} = -\nabla_b W_{ba} \)), we obtain

\[ \int \left[ -m_a \frac{d}{dt} \mathbf{v}_a - \sum_b m_b \left( \frac{P_b}{\rho_b^{\sigma}} \frac{\rho_a^{2-\sigma}}{\rho_b^{2-\sigma}} + \frac{P_a}{\rho_a^{\sigma}} \frac{\rho_b^{2-\sigma}}{\rho_a^{2-\sigma}} \right) \nabla_a W_{ab} \right] \delta \mathbf{r}_a \, dt = 0, \tag{3.84} \]

from which we obtain the momentum equation in the form (3.76). This equation is therefore consistent with the continuity equation in the form (3.79). In the particular case considered by Marri and White (2003) (\( \sigma = 3/2 \)) this would imply a discrete form of the continuity equation given by

\[ \frac{d \rho_a}{dt} = \sqrt{\rho_a} \sum_b m_b \frac{\mathbf{v}_{ab}}{\sqrt{\rho_b}} \cdot \nabla_a W_{ab}. \tag{3.85} \]

Marri and White (2003) choose to retain the use of the usual SPH summation (3.42) to determine the density. In the case considered by Ritchie and Thomas (2001) (\( \sigma = 1 \)), the continuity equation becomes

\[ \frac{d \rho_a}{dt} = \rho_a \sum_b m_b \frac{\mathbf{v}_{ab}}{\rho_b} \cdot \nabla_a W_{ab}, \tag{3.86} \]

which is again somewhat different to the density estimation used in their paper. The continuity equation (3.86), when used in conjunction with the appropriate formulation of the momentum equation, has some advantages in the case of fluids with large density differences (e.g. at a water/air interface) since the term inside the summation involves only the particle volumes \( m/\rho \) rather than their mass, with the effect that large mass differences between individual particles have less influence on the calculation of the velocity divergence (Monaghan, private communication). An alternative is the formalism proposed by Ott and Schnetter (2003).
which we discuss in §3.4.3.

The internal energy equation consistent with the general momentum equation (3.76) is given by

$$\frac{du_a}{dt} = \frac{P_a}{\rho_a^\sigma} \sum_b m_b \frac{v_{ab}}{p_b^{2-\sigma}} \cdot \nabla W_{ab},$$

(3.87)

which is indeed the formalism used by Marri and White (2003) (with $\sigma = 3/2$) since it was found, unsurprisingly in this context, that integration of this equation resulted in much less numerical noise than using other formalisms of the internal energy equation (in conjunction with their use of (3.76) with $\sigma = 3/2$ as the momentum equation). The form of the total energy equation consistent with (3.76) and (3.79) is given by

$$\frac{de_a}{dt} = -\sum_b m_b \left( \frac{P_a}{\rho_a^\sigma} \frac{2-\sigma}{\rho_b^{2-\sigma}} v_{ab} + \frac{P_b}{\rho_b^\sigma} \frac{2-\sigma}{\rho_a^{2-\sigma}} v_{ba} \right) \cdot \nabla W_{ab}.$$  

(3.88)

We note the energy equation used by Ritchie and Thomas (2001) is different to the formulation given above (with $\sigma = 1$) and therefore variationally inconsistent with their implementation of the momentum equation. Hernquist and Katz (1989) point out that inconsistencies between the forms of the energy and momentum equations result in errors of $O(h^2)$ in the energy conservation. In this sense the difference between a consistent and inconsistent formalism is fairly minor, although a consistent formulation between the momentum and energy equations in general appears to lead to slightly improved results (as found by Marri and White). In practice we find that using alternative formulations of the continuity equation generally gives slightly worse results than (even inconsistent) use of the density summation.

### 3.4.2 General alternative formulation

The momentum equation (3.76) can be generalised still further by noting that the continuity equation (3.44) can be written as

$$\frac{d\rho}{dt} = \phi \left[ \mathbf{v} \cdot \nabla \left( \frac{\rho}{\phi} \right) - \nabla \cdot \left( \frac{\rho \mathbf{v}}{\phi} \right) \right],$$

(3.89)

with SPH equivalent

$$\frac{d\rho_a}{dt} = \phi_a \sum_b m_b \frac{v_{ab}}{\phi_b} \cdot \nabla W_{ab},$$

(3.90)

where $\phi$ is any scalar variable defined on the particles. Deriving the momentum equation consistent with this equation in the manner given above we find

$$\frac{d\mathbf{v}_a}{dt} = -\sum_b m_b \left( \frac{P_a \phi_a}{\rho_a^\sigma \phi_b} + \frac{P_b \phi_b}{\rho_b^\sigma \phi_a} \right) \nabla W_{ab},$$

(3.91)
which conserves momentum for any choice of $\phi$. In the case given in the previous section we would have $\phi = \rho^{2-\sigma}$. Choosing $\phi = \rho / \sqrt{P}$ gives

$$
\frac{dv_a}{dt} = - \sum_b m_b \left( \frac{\sqrt{T_a T_b}}{\rho_a \rho_b} \right) \nabla_a W_{ab}.
$$

(3.92)

which is the momentum equation used by Hernquist and Katz (1989). The continuity equation consistent with this form is therefore

$$
\frac{d\rho_a}{dt} = \frac{\rho_a}{\sqrt{T_a}} \sum_b m_b \frac{\sqrt{T_b}}{\rho_b} v_{ab} \cdot \nabla_a W_{ab},
$$

(3.93)

which at first sight appears somewhat bizarre, although it is certainly a valid expression of the continuity equation in SPH form. It is unclear whether using such alternative formulations of the continuity equation, in the name of consistency, has any advantages over the usual density summation. We leave it as an exercise for the reader to amuse themselves by exploring various other combinations of variables, noting that the forms of the internal and total energy equations consistent with (3.90) and (3.91) are given by

$$
\frac{du_a}{dt} = \frac{P_a}{\rho_a^2} \sum_b m_b \frac{\phi_a}{\phi_b} v_{ab} \cdot \nabla_a W_{ab},
$$

(3.94)

and

$$
\frac{de_a}{dt} = - \sum_b m_b \left( \frac{P_a \phi_a}{\rho_a^2} v_{ab} + \frac{P_b \phi_b}{\rho_b^2} v_{ab} \right) \cdot \nabla_a W_{ab}.
$$

(3.95)

### 3.4.3 Ott and Schnetter formulation

Other formulations of the SPH equations have also been proposed to deal with the problem of large density gradients. For example Ott and Schnetter (2003) propose modifying the SPH summation to give

$$
n_a = \sum_b W_{ab},
$$

$$
\rho_a = m_a n_a,
$$

(3.96)

that is where the number density of particles $n$ is calculated by summation rather than the mass density $\rho$. This is to improve the interpolation when particles of large mass differences interact. Taking the time derivative of (3.96), the continuity equation is given by (as in Ott and Schnetter 2003)

$$
\frac{d\rho_a}{dt} = m_a \sum_b v_{ab} \cdot \nabla_a W_{ab}.
$$

(3.97)

For equal mass particles this formalism is exactly the same as the usual summation (3.42). The formulation (3.96) enables the density to be expressed as a function of the particle co-ordinates and thus the derivation of the equations of motion and energy can be done in a straightforward manner using the Euler-Lagrange
equations, as in §3.3.2. The resulting equation of motion is given by

\[ m_a \frac{d\mathbf{v}_a}{dt} = -\sum_b \left( \frac{P_a}{n_a^2} + \frac{P_b}{n_b^2} \right) \nabla_a W_{ab}, \quad (3.98) \]

which is somewhat different to the equation of motion used in Ott and Schnetter (2003) (they use the form 3.76 with \( \sigma = 1 \)). The internal energy equation follows from the continuity equation (3.97) and the first law of thermodynamics (3.50). We find

\[ m_a \frac{du_a}{dt} = \frac{P_a}{n_a^2} \sum_b \mathbf{v}_{ab} \cdot \nabla_a W_{ab}. \quad (3.99) \]

Ott and Schnetter (2003) use a formulation of the internal energy equation where the pressure term is symmetrised, which is inconsistent with their use of (3.96). The total energy equation consistent with their formalism can also be derived using the Hamiltonian (§3.3.3) and is given by

\[ m_a \frac{de_a}{dt} = -\sum_b \left( \frac{P_a}{n_a^2} \mathbf{v}_b + \frac{P_b}{n_b^2} \mathbf{v}_a \right) \cdot \nabla_a W_{ab}. \quad (3.100) \]

In this case use of the self-consistent formalism presented above should lead to slightly improved results over the momentum and energy equations employed by Ott and Schnetter (2003), since the density is still calculated via a direct summation over the particles.

### 3.5 Shocks

In any high-order numerical scheme, the simulation of shocks is accompanied by unphysical oscillations behind the shock front. This occurs because in discretising the continuum equations (in the SPH case using 3.5) we assume that the fluid quantities are smoothly varying on the smallest length scale (in SPH this is the smoothing length \( h \)). This means that discontinuities on such scales are not resolved by the numerical method. The simplest approach to this problem is to introduce a small amount of viscosity into the simulation which acts to spread out the shock front so that it can be sufficiently resolved (von Neumann and Richtmyer, 1950; Richtmyer and Morton, 1967). This is similar to the way in which shock fronts are smoothed out by nature, although in the latter case the effect occurs at a much finer level. The disadvantage of using such an ‘artificial’ viscosity is that it can produce excess heating elsewhere in the simulation. As such the use of artificial viscosity is regarded by many numerical practitioners as outdated since most finite difference schemes now rely on methods which either restrict the magnitude of the numerical flux across a shock front in order to prevent unphysical oscillations (such as total variation diminishing (TVD) schemes) or by limiting the jump in the basic variables across the shock front using the exact solution to the Riemann problem (Godunov-type schemes). There remain, however, distinct advantages to the use of an artificial viscosity, primarily that, unlike the Godunov-type schemes, it is easily applied where new physics is introduced (such as a more complicated equation of state than the ideal gas law) and the complexity of the algorithm does not
increase with the number of spatial dimensions. In the case of magnetohydrodynamics, artificial viscosity is commonly used even in standard finite-difference codes\(^3\) since the Riemann problem is difficult to solve and computationally expensive. Furthermore, dissipative terms are often still used even when a Riemann solver has been implemented (e.g. Balsara 1998). For these reasons artificial viscosity methods continue to find widespread usage, particularly in simulations using unstructured or Lagrangian meshes (Caramana et al., 1998).

In recent years it has been shown that Godunov-type schemes can in fact be used in conjunction with SPH by regarding interacting particle pairs as left and right states of the Riemann problem (Cha and Whitworth, 2003; Inutsuka, 2002; Parshikov and Medin, 2002; Monaghan, 1997). In this manner the implementation of Godunov-type schemes to multidimensional problems is greatly simplified in SPH because the one-dimensional Riemann problem is solved between particle pairs, removing the need for complicated operator splitting procedures in higher dimensions. The formalism presented by Cha and Whitworth (2003) is remarkably simple to incorporate into any standard SPH code. A Godunov-type scheme for MHD in SPH would be extremely useful (although not widely applicable), but it is well beyond the scope of this thesis. We therefore formulate artificial dissipation terms using the formulation of Monaghan (1997) which is generalised to the MHD case in §4.5. The problem of excess heating is addressed by the implementation of switches to turn off the dissipative terms away from shock fronts, described in §3.5.2.

### 3.5.1 Artificial viscosity and thermal conductivity

A variety of different formulations of artificial viscosity in SPH have been used, however the most common implementation is that given by Monaghan (1992), where the term in equation (3.54) is given by

\[
\left( \frac{d\mathbf{v}^a}{dt} \right)_{\text{diss}} = \sum_b m_b \frac{\alpha \bar{c}_{ab} \mu_{ab} + \beta \mu_{ab}^2}{\bar{\rho}_{ab}} \nabla_a W_{ab}, \quad \mu_{ab} = \frac{h \mathbf{v}_{ab} \cdot \mathbf{r}_{ab}}{r_{ab}^2 + 0.01h^2}, \tag{3.101}
\]

where \(\mathbf{v}_{ab} = \mathbf{v}_a - \mathbf{v}_b\) (similarly for \(\mathbf{r}_{ab}\)), barred quantities refer to averages between particles \(a\) and \(b\), and \(c\) refers to the sound speed. This viscosity is applied only when the particles are in compression (i.e. \(\mathbf{v}_{ab} \cdot \mathbf{r}_{ab} < 0\)), is Galilean invariant, conserves total linear and angular momentum and vanishes for rigid body rotation. The \(\beta\) term (quadratic in \(\mathbf{v}_{ab}\)) represents a form of viscosity similar to the original formulation of von Neumann and Richtmyer (1950) and becomes dominant in the limit of large velocity differences (i.e. high Mach number shocks). The \(\alpha\) term is linear in \(\mathbf{v}_{ab}\) and is dominant for small velocity differences\(^4\). Most astrophysical SPH implementations follow Monaghan (1992) in setting \(\alpha = 1\) and \(\beta = 2\) which provides the necessary dissipation near a shock front.

The term given by equation (3.101) was constructed to have the properties described above, however in the relativistic case it was unclear as to what form such an artificial viscosity should take. Chow and Monaghan (1997) thus formulated an artificial viscosity for ultra-relativistic shocks in SPH by analogy with

---

\(^3\)for example in the widely used ZEUS code for astrophysical fluid dynamics (Stone and Norman, 1992)

\(^4\)The introduction of such a term into artificial viscosity methods is generally attributed to Landshoff (1955) (see, e.g. Caramana et al. 1998)
Riemann solvers. This is outlined by Monaghan (1997) in a discussion of SPH and Riemann solvers. The essential idea is to regard the interacting particles as left and right Riemann states and to construct a dissipation which involves jumps in the physical variables. The dissipation term in the force (giving artificial viscosity) therefore involves a jump in the velocity variable and is similar to (3.101), taking the form (for $v_{ab} \cdot r_{ab} < 0$)

$$\left( \frac{d\mathbf{v}_a}{dt} \right)_{diss} = -\sum_b m_b \frac{\alpha v_{sig}(\mathbf{v}_a - \mathbf{v}_b) \cdot \hat{r}_{ab}}{2\bar{\rho}_{ab}} \nabla_a W_{ab},$$

(3.102)

where $v_{sig}$ is a signal velocity and $\hat{r}_{ab} \equiv (\mathbf{r}_a - \mathbf{r}_b)/|\mathbf{r}_a - \mathbf{r}_b|$ is a unit vector along the line joining the particles. Note that this formalism differs from (3.101) in that a factor of $h/|\mathbf{r}_{ab}|$ has been removed. Also the $0.01h^2$ term has been removed from the denominator since for variable smoothing lengths it is unnecessary. The jump in velocity involves only the component along the line of sight since this is the only component expected to change at a shock front. In a similar manner, the dissipative term in the specific energy equation (3.63) is given by

$$\left( \frac{de_a}{dt} \right)_{diss} = -\sum_b m_b v_{sig}(e_a^* - e_b^*) \frac{\hat{r}_{ab} \cdot \nabla_a W_{ab}}{2\bar{\rho}_{ab}},$$

(3.103)

where $(e_a^* - e_b^*)$ is the jump in specific energy. The specific energy used in this term is given by

$$e_a^* = \begin{cases} \frac{1}{2}\alpha(\mathbf{v}_a \cdot \hat{r}_{ab})^2 + \alpha_a u_a, & \mathbf{v}_a \cdot \mathbf{r}_{ab} < 0; \\ \alpha_a u_a, & \mathbf{v}_a \cdot \mathbf{r}_{ab} \geq 0; \end{cases}$$

(3.104)

that is, where the specific kinetic energy has been projected along the line joining the particles, since only the component of velocity parallel to this vector is expected to jump at a shock front. Note that in general we use a different parameter $\alpha_a$ to control the thermal energy term and that this term is applied to particles in both compression and rarefaction.

The signal velocity represents the maximum speed of signal propagation along the line of sight between the two particles. Whilst many formulations could be devised, it turns out that the results are not sensitive to the particular choice made. A simple estimate of the signal velocity is given by

$$v_{sig} = c_a + c_b - \beta \mathbf{v}_{ab} \cdot \hat{r}_{ab}$$

(3.105)

where $c_a$ denotes the speed of sound of particle $a$ and $\beta \sim 1$, such that $v_{sig}/2$ is an estimate of the maximum speed for linear wave propagation between the particles. The $\beta$ term, which acts as a von Neumann and Richtmyer viscosity as in equation (3.101), arises naturally in this formulation. Practical experience suggests, however, that $\beta = 2$ is a better choice. For a more general discussion of signal velocities we refer the reader to Monaghan (1997) and Chow and Monaghan (1997).

The contribution to the thermal energy from the dissipative terms is found using

$$\frac{du_a}{dt} = \frac{de_a}{dt} - \mathbf{v}_a \cdot \frac{d\mathbf{v}_a}{dt}.$$

(3.106)
In this case we obtain

\[
\left( \frac{dt_u}{dt} \right)_{diss} = \sum \frac{m_b}{2 \rho_{ab}} \left\{ -\frac{1}{2} \alpha \left[ (v_a - v_b) \cdot \hat{r}_{ab} \right]^2 + \alpha_u (u_a - u_b) \right\} \hat{r}_{ab} \cdot \nabla_a W_{ab}
\]

(3.107)

which is added to the non-dissipative term (3.58). The first term is the positive definite contribution to the thermal energy from the artificial viscosity (since the kernel gradient is always negative). The second term (involving a jump in thermal energy) provides an artificial thermal conductivity. Physically this means that discontinuities in the thermal energy are also smoothed.

The artificial dissipation given by (3.102)-(3.107) is used as a basis for constructing an appropriate dissipation for the MHD case in §4.5.

### 3.5.2 Artificial dissipation switches

#### Artificial viscosity

In both (3.101) and (3.102) the artificial viscosity is applied universally across the particles despite only being needed when and where shocks actually occur. This results in SPH simulations being much more dissipative than is necessary and can cause problematic effects where this dissipation is unwanted (such as in the presence of shear flows). A switch to reduce the artificial viscosity away from shocks is given by Morris and Monaghan (1997). Using this switch in multi-dimensional simulations substantially reduces the problematic effects of using an artificial viscosity in SPH.

The key idea is to regard the dissipation parameter \( \alpha \) (c.f. equation 3.102) as a particle property. This can then be evolved along with the fluid equations according to

\[
\frac{d\alpha_a}{dt} = -\frac{\alpha_a - \alpha_{\text{min}}}{\tau_a} + \mathcal{S}_a,
\]

(3.108)

such that in the absence of sources \( \mathcal{S} \), \( \alpha \) decays to a value \( \alpha_{\text{min}} \) over a timescale \( \tau \). The timescale \( \tau \) is calculated according to

\[
\tau = \frac{h}{\mathcal{C} v_{\text{sig}}},
\]

(3.109)

where \( h \) is the particle’s smoothing length, \( v_{\text{sig}} \) is the maximum signal propagation speed at the particle location and \( \mathcal{C} \) is a dimensionless parameter with value \( 0.1 < \mathcal{C} < 0.2 \). We conservatively use \( \mathcal{C} = 0.1 \) which means that the value of \( \alpha \) decays to \( \alpha_{\text{min}} \) over \( \sim 5 \) smoothing lengths.

The source term \( \mathcal{S} \) is chosen such that the artificial dissipation grows as the particle approaches a shock front. We use (Rosswog et al., 2000)

\[
\mathcal{S} = \max(-\nabla \cdot \v, 0)(2.0 - \alpha),
\]

(3.110)

such that the dissipation grows in regions of strong compression. Following Morris and Monaghan (1997)
where the ratio of specific heats $\gamma$ differs from 5/3 (but not for the isothermal case), we multiply $S$ by a factor

$$\left[ \ln \left( \frac{5/3+1}{5/3-1} \right) \right] / \left[ \ln \left( \frac{\gamma+1}{\gamma-1} \right) \right]$$

(3.111)

The source term is multiplied by a factor $(2.0 - \alpha)$ as the standard source term given by Morris and Monaghan (1997) was found to produce insufficient damping at shock fronts when used in conjunction with the Monaghan (1997) viscosity. The source term (3.110) is found to provide sufficient damping on the Sod (1978) hydrodynamic shock tube problem and in the MHD shock tube tests we describe in chapter §4.6 (ie. $\alpha_{\text{max}} \sim 1$ for these problems). In order to conserve momentum the average value $\bar{\alpha} = 0.5(\alpha_a + \alpha_b)$ is used in equations (3.102), (3.104) and (3.107). A lower limit of $\alpha_{\text{min}} = 0.1$ is used to preserve order away from shocks (note that this is an order of magnitude reduction from the usual value of $\alpha = 1.0$ everywhere).

The numerical tests in §4.6 demonstrate that use of this switch gives a significant reduction in dissipation away from shocks whilst preserving the shock-capturing ability of the code.

**Artificial thermal conductivity**

A similar switch to that used in the artificial viscosity may therefore be devised for the artificial thermal conductivity term, with the parameter $\alpha_u$ evolved according to

$$\frac{d \alpha_{u,a}}{dt} = -\frac{\alpha_{u,a} - \alpha_{u,\text{min}}}{\tau_a} + S_a,$$

(3.112)

where the decay timescale $\tau$ is the same as that used in (3.108) and in this case we use $\alpha_{u,\text{min}} = 0$. The corresponding source term is given by

$$S = |\nabla \sqrt{u}|,$$

(3.113)

which is constructed to have dimensions of inverse time. The gradient term is computed according to

$$\nabla \sqrt{u} = \frac{1}{2} u^{-1/2} \nabla u,$$

(3.114)

where

$$\nabla u_a = \frac{1}{\rho_a} \sum_b m_b (u_a - u_b) \nabla a W_{ab}(h_a).$$

(3.115)

Use of this switch ensures that artificial thermal conductivity is only applied at large gradients in the thermal energy. The need to do so in dissipation-based shock capturing schemes is often concealed by smoothing of the initial conditions in shock tube tests (§3.7.3). From the first law of thermodynamics (3.50) we infer that gradients in the thermal energy correspond to large gradients in the density. In a hydrodynamic shock these occur either at the shock front or at the contact discontinuity. Artificial viscosity is not required at
3.6 Timestepping

the contact discontinuity because the pressure is constant across it. Using unsmoothed initial conditions and in the absence of artificial thermal conductivity, a significant overshoot in thermal energy occurs at the contact discontinuity (this phenomenon is known as ‘wall heating’ and is illustrated in Figure 3.9). The resulting glitch in the pressure is often ascribed to ‘starting errors’ due to the unsmoothed initial conditions. However, applying smoothing to the initial conditions of a shock-tube test means that gradients across the contact discontinuity remain smoothed throughout the evolution (see e.g. Figure 3.8), removing the need for artificial thermal conductivity which acts to spread gradients in the thermal energy. Whilst there is also a gradient in thermal energy at a shock front, this is smoothed out by the application of artificial viscosity there and so the need for artificial thermal conductivity can go unnoticed. In §3.7.3 we present results of the standard Sod (1978) shock tube test, showing the effectiveness of the switch discussed above in applying the requisite amount of smoothing at the contact discontinuity.

3.6 Timestepping

3.6.1 Predictor-corrector scheme

We integrate the SPH equations in this thesis using a slight modification of the standard predictor-corrector (Modified Euler) method which is second order accuracy in time (Monaghan, 1989). The predictor step is given by

\[
v^{1/2} = v^0 + \frac{\Delta t}{2} f^0, \]
(3.116)

\[
r^{1/2} = r^0 + \frac{\Delta t}{2} v^{1/2}, \]
(3.117)

\[
e^{1/2} = e^0 + \frac{\Delta t}{2} \dot{e}^0, \]
(3.118)

where in practice we use \(f^0 \approx f^{1/2}\) and \(\dot{e}^0 \approx \dot{e}^{1/2}\) to give a one-step method. The rates of change of these quantities are then computed via the SPH summations using the predicted values at the half step, ie.

\[
f^{1/2} = f(r^{1/2}, v^{1/2}) \quad \dot{e}^{1/2} = \dot{e}(r^{1/2}, v^{1/2}) \]
(3.119)

The corrector step is given by

\[
v^* = v^0 + \frac{\Delta t}{2} f^{1/2}, \]
(3.120)

\[
r^* = r^0 + \frac{\Delta t}{2} v^*, \]
(3.121)

\[
e^* = e^0 + \frac{\Delta t}{2} \dot{e}^{1/2}, \]
(3.122)

and finally

\[
v^1 = 2v^* - v^0, \]
(3.123)
\[ r^1 = 2r^* - r^0, \quad (3.124) \]
\[ e^1 = 2e^* - e^0. \quad (3.125) \]

Note that in this scheme the position updates in both the predictor and corrector steps use the updated value of velocity. This effectively means that the position is updated using both the first and second derivatives. From numerical experiments we find that this scheme gives much better stability properties. Where evolved, density, smoothing length, magnetic field and the dissipation parameters follow the energy evolution. The total energy \( e \) is interchangeable for the thermal energy \( u \).

### 3.6.2 Reversible integrators

The simple predictor-corrector method given above is adequate for all the problems considered in this thesis since the integration time is quite short. For large simulations over long timescales, however, the accuracy and stability of the integration method needs more careful attention. In the past decade or so a substantial research effort has been devoted to the development of high accuracy so-called ‘geometric’ integrators for Hamiltonian systems (e.g. Hut et al., 1995; Stoffer, 1995; Huang and Leimkuhler, 1997; Holder et al., 2001; Hairer et al., 2002). Since SPH in the absence of dissipative terms can derived from a Hamiltonian variational principle, much of this work is applicable in the SPH context. The primary condition for the construction of a geometric integrator is time-reversibility (that is, particle quantities should return to their original values upon reversing the direction of time integration). It is fairly straightforward to construct a reversible integrator for the SPH equations in the case of a constant smoothing length, where the density summation is used and where the pressure is calculated directly from the density (such that the force evaluation uses only the particle coordinates). The standard leapfrog algorithm is one such example. In general, however, the construction of a reversible scheme is complicated by several factors. The first is the use of a variable timestep (which immediately destroys the time-symmetry in the leapfrog scheme, although see Holder et al. (2001) for recent progress on this). The second complicating factor is that the reversibility condition becomes more difficult when equations with rates of change involving the particle velocity are used (such as the thermal or total energy equation or the continuity equation for the density). In this case the construction of a reversible integrator for SPH necessarily involves the calculation of derivatives involving the velocity in separate step to the force evaluation, leading to additional computational expense. A third complicating factor is the use of individual particle timesteps in large SPH codes, although symplectic methods have also been constructed for this case (Hairer et al., 2002).

### 3.6.3 Courant condition

The timestep is determined by the Courant condition

\[ dt_c = C_{\text{cour}} \min \left( \frac{h}{v_{\text{sig}}} \right) \quad (3.126) \]
where \( h = \min(h_a, h_b) \) and \( v_{\text{sig}} \) is the maximum signal velocity between particle pairs. This signal velocity is similar to that used in the artificial dissipation terms (§3.5), except that we use

\[
v_{\text{sig}} = \frac{1}{2} (v_a + v_b + \beta |v_{ab} \cdot \mathbf{j}|)
\]

(3.127)

with \( \beta = 1 \) when \( v_{ab} \cdot \mathbf{j} > 0 \) (ie. where the dissipation terms are not applied). The minimum in (3.126) is taken over all particle interactions and typically we use \( C_{\text{coar}} = 0.4 \).

Although this condition is sufficient for all of the simulations described here, in general it is necessary to pose the additional constraint from the forces

\[
dt_f = C_f \min \left( \frac{h_a}{|a_a|} \right)^{1/2},
\]

(3.128)

where \( a_a \) is the acceleration on particle \( a \) and typically \( C_f = 0.25 \).

### 3.7 Numerical tests

#### 3.7.1 Implementation

Unless otherwise indicated the simulations use the density summation (3.42), the momentum equation (3.54) and the energy equation in the form (3.63). The numerical tests presented throughout this thesis were implemented using a code written by the author as a testbed for MHD algorithms.

**Neighbour finding**

Since the code has been designed for flexibility rather than performance, we take a simplified approach to neighbour finding using linked lists. The particles are binned into grid cells of size \( 2h \) where \( h \) is the maximum value of smoothing length over the particles. Particles in a given cell then search only the adjoining cells for contributing neighbours. This approach becomes very inefficient for a large range in smoothing lengths such that for large simulations it is essential to use a more effective algorithm. A natural choice is to use the tree code used in the computation of the gravitational force (Hernquist and Katz, 1989; Benz et al., 1990).

**Boundary conditions**

Boundary conditions are implemented using either ghost or fixed particles. For reflecting boundaries, ghost particles are created which mirror the SPH particles across the boundary. These particles are exact copies of the SPH particles in all respects except for the velocity, which is of opposite sign on the ghost particle, producing a repulsive force at the boundary. For periodic boundary conditions the ghosts are exact copies of the particles at the opposite boundary. In the MHD shock tube tests considered in §4.6 involving non-zero velocities at the boundaries, boundary conditions are implemented in one dimension by simply fixing the
properties of the 6 particles closest to each boundary. Where the initial velocities of these particles are non-zero their positions are evolved accordingly and a particle is removed from the domain once it has crossed the boundary. Where the distance between the closest particle and the boundary is more than the initial particle spacing a new particle is introduced to the domain. Hence for inflow or outflow boundary conditions the resolution changes throughout the simulation.

3.7.2 Propagation and steepening of sound waves

We initially consider the propagation of linear sound waves in SPH. This test is particularly important in the MHD case (§4.6.4) since it highlights the instability in the momentum-conserving formalism of SPMHD. In this case we investigate the dependence of sound speed on smoothing length and the damping due to artificial viscosity.

Particle setup

The particles are initially setup at equal separations in the domain $x = [0, 1]$ using ghost particles (§3.7.1) to create periodic boundary conditions. The linear solution for a travelling sound wave in the $x$-direction is given by

$$
\rho(x,t) = \rho_0(1 + Asin(kx_0 - \omega t)),
$$

(3.129)

$$
v_x(x,t) = C_s Asin(kx_0 - \omega t),
$$

(3.130)

where $\omega = 2\pi C_s / \lambda$ is the angular frequency, $C_s$ is the sound speed in the undisturbed medium and $k = 2\pi / \lambda$ is the wavenumber. The initial conditions therefore correspond to $t = 0$ in the above. The perturbation in density is applied by perturbing the particles from an initially uniform setup. We consider the one dimensional perturbation

$$
\rho = \rho_0[1 + Asin(kx)],
$$

(3.131)

where $A = D / \rho_0$ is the perturbation amplitude. The cumulative total mass in the $x$ direction is given by

$$
M(x) = \rho_0 \int [1 + Asin(kx)]dx
= \rho_0[x - A\cos(kx)]_0^x,
$$

(3.132)

such that the cumulative mass at any given point as a fraction of the total mass is given by

$$
\frac{M(x)}{M(x_{max})},
$$

(3.133)
3.7 Numerical tests

Figure 3.3: Representative results from the isothermal sound wave tests in one dimension using the standard cubic spline kernel with a fixed smoothing length. The figure on the left shows the results after 5 periods (corresponding to 5 crossings of the computational domain) using $h = 1.5 \Delta p$. The figure on the right shows the results using a fixed smoothing length but with the correction from the variable smoothing length terms.

Figure 3.4: Representative results from the isothermal sound wave tests in one dimension using the standard cubic spline kernel with a variable smoothing length that varies with density. The figure on the left shows the results after 5 periods using a simple average of the kernel gradients, whilst the figure on the right shows the results using the consistent formulation of the variable smoothing length terms.

For equal mass particles distributed in $x = [0, x_{\text{max}}]$ the cumulative mass fraction at particle $a$ is given by $x_a / x_{\text{max}}$ such that the particle position may be calculated using

$$\frac{x_a}{x_{\text{max}}} = \frac{M(x_a)}{M(x_{\text{max}})}.$$  \hspace{1cm} (3.134)

Substituting the expression for $M(x)$ we have the following equation for the particle position

$$\frac{x_a}{x_{\text{max}}} - \frac{x_a - A \cos(kx_a)}{[x_{\text{max}} - A \cos(kx_{\text{max}})]} = 0,$$ \hspace{1cm} (3.135)

which we solve iteratively using a simple Newton-Raphson rootfinder. With the uniform particle distribution as the initial conditions this converges in one or two iterations.
One dimensional tests

Initially we consider one dimensional, isothermal simulations using a fixed smoothing length (for which the results of the stability analysis given in §3.2.7 hold). The cubic spline kernel is used with $h = 1.5\Delta p$ where $\Delta p$ is the initial particle spacing. This value of smoothing length was chosen because in Figure 3.2 the cubic spline is seen to significantly underestimate the sound speed at this value of $h$. The simulation is setup using 100 particles (corresponding to $k_x = 0.0628$ in Figure 3.2) and a wave amplitude of 0.005 to ensure that the wave remains essentially linear throughout the simulation. No artificial viscosity is used. For isothermal simulations, the pressure is calculated directly from the density using $P = \rho c_s^2$. The sound speed given by the SPH simulations is estimated from the temporal spacing of minima in the total kinetic energy of the particles.

A representative example of these simulations is given in the left hand side of Figure 3.3 after five crossings of the computational domain. The amplitude is well maintained by the SPH scheme, however the wave lags behind the exact solution, giving a significant phase error as expected from the stability analysis (Figure 3.2). The sound speed obtained from the numerical tests is plotted in Figure 3.5 for a range of smoothing length values (solid points). In this case the results show excellent agreement with the analytic results using the dispersion relation (3.41) given by the solid line (this line corresponds to $k_x \approx 0$ in Figure 3.2). We observe that, depending on the value of $h$ the numerical sound wave can both lag and lead the exact solution (in Figure 3.5 this corresponds to sound speeds less than or greater than unity).

In §3.3.4 it was noted that the variable smoothing length terms normalise the kernel even in the case of a fixed smoothing length. The results of the fixed smoothing length simulation with this correction term are shown by the dashed line in Figure 3.5, with a representative example given in the right hand side of Figure 3.3. The numerical wave speed appears much closer to the theoretical value of unity.

Results using a smoothing length which varies with density according to (3.68) are given by the dot-dashed line in Figure 3.5, with a representative example shown in Figure 3.4. The phase error is slightly lower than either of the fixed smoothing length cases. Including the normalisation of the kernel gradient from the variable smoothing lengths (§3.3.4) gives numerical sound speeds very close to unity (dotted line in Figure 3.5). A representative example of these simulations is given in the right hand panel of Figure 3.4 after 5 periods. The results in this case show excellent agreement with the exact (linear) solution, with a small amount of steepening due to nonlinear effects.

The results of this test indicate that, whilst alternative kernels can give slight improvements in accuracy over the standard cubic spline (§3.2.7), a substantial gain in accuracy can be gained firstly by the use of a variable smoothing length and secondly by self-consistently accounting for $\nabla h$ terms in the formulation of the SPH equations. These terms act as a normalisation of the kernel gradient which appear to effectively remove the dependence of the numerical sound speed on the smoothing length value.
3.7 Numerical tests

Figure 3.5: Summary of the isothermal sound wave tests using 100 particles. The numerical sound speed from the SPH simulations is shown plotted against the (mean) smoothing length in units of the average particle spacing. Results using the cubic spline kernel with a fixed smoothing length (solid points) may be compared with the analytic result (solid line, under points) from the dispersion relation (3.41) (this line corresponds to \( kx = 0 \) in Figure 3.2). The dashed line gives the numerical results using the cubic spline with a fixed smoothing length but incorporating the correction from the \( \nabla h \) terms, which show much lower phase errors. The dotted and dot-dashed lines give numerical results using the cubic spline with a variable smoothing length with and without the \( \nabla h \) terms respectively. In both cases the results show a substantial improvement over the fixed smoothing length case, much more so than from the use of alternative kernels (e.g. the New Quintic (2) from §3.2.6, given by the solid line).

Effects of artificial viscosity

In the absence of any switches, the artificial viscosity is specified according to (3.102) with \( \alpha = 1, \beta = 2 \) everywhere. The results of the sound wave propagation with artificial viscosity turned on are shown in the left panel of Figure 3.6. After 5 crossings of the computational domain the wave is severely damped by the artificial viscosity term. The effect is to reduce the order of the numerical scheme since convergence to the exact solution is much slower. The results using the artificial viscosity switch discussed in §3.5.2 are shown in the right panel of Figure 3.6. The results show good agreement with the linear solution, demonstrating that use of the artificial viscosity switch very effectively restores the numerical schemes ability to propagate small perturbations without excessive damping.

Finally, we demonstrate the usefulness of the artificial viscosity switch by considering the steepening of a nonlinear sound wave. In this case the initial amplitude is 0.05 and artificial viscosity is applied using the switch. The wave profile at \( t = 5 \) is shown in Figure 3.7 and is significantly steepened compared to the initial conditions (solid line). The use of the switch enables the steepening to be resolved, however some oscillations are found to occur ahead of the steepened wave.
Figure 3.6: (left) Isothermal sound wave with amplitude = 0.005 in one dimension with artificial viscosity applied uniformly to particles in compression (i.e. $\alpha = 1, \beta = 2$) and (right) applied using the viscosity switch with $\alpha_{\text{min}} = 0.1$.

Figure 3.7: Nonlinear isothermal sound wave in one dimension showing steepening to shock. The wave profile is shown after 5 crossings of the computational domain, corresponding to 5 periods. The initial conditions are a linear wave with amplitude 0.05 (solid line). With artificial viscosity applied using the switch the steepening is resolved, although some oscillations are observed to occur ahead of the steepened wave.

3.7.3 Sod shock tube

The standard shock tube test for any compressible fluid dynamics code is that of Sod (1978). The problem consists of dividing the domain into two halves, one consisting of high pressure, high density gas whilst the other is low pressure and low density. These two portions of gas are allowed to interact at $t = 0$, resulting in a shock and rarefaction wave which propagate through the gas. This test illustrates the shock capturing ability of the 1D code and thus provides a good test of the artificial viscosity formalism ($\S$3.5). It is also the basis for the MHD shock tube considered in $\S$4.6.3. We set up the problem using 450 SPH particles in the domain $x = [-0.5, 0.5]$. The particles are setup with uniform masses such that the density jump is modelled by a jump in particle separation. Initial conditions in the fluid to the left of the origin are given by $(\rho, P, v_x) = [1, 1, 0]$ whilst conditions to the right are given by $(\rho, P, v_x) = [0.125, 0.1, 0]$ with $\gamma = 1.4$. The particle separation to the left of the discontinuity is 0.01.

Figure 3.8 shows the results of this problem at $t = 0.2$. The exact solution, calculated using the exact
3.7 Numerical tests

Riemann solver given in Toro (1992) is given by the solid line. In this case artificial viscosity has been applied uniformly to particles in compression (ie. using $\alpha = 1$), whilst no artificial thermal conductivity has been used (ie. $\alpha_u = 0$). The results are generally good although there is significant deviation in the slope of the rarefaction wave. This can be traced largely to the smoothing applied to the initial conditions. Following Monaghan (1997) (although a similar procedure is applied in many published versions of this test), the initial discontinuities in density and pressure were smoothed over several particles according to the rule

$$ A = \frac{A_L + A_R e^{x/d}}{1 + e^{x/d}} $$

(3.136)

where $A_L$ and $A_R$ are the uniform left and right states with respect to the origin and $d$ is taken as half of the largest initial particle separation at the interface (ie. the particle separation on the low density side). Where the initial density is smoothed the particles are spaced according to the rule

$$ \rho_a(x_{a+1} - x_{a-1}) = 2\rho_R \Delta R $$

(3.137)

where $\Delta R$ is the particle spacing to the far right of the origin with density $\rho_R$. Note that initial smoothing lengths are set according to the rule $h \propto 1/\rho$ and are therefore also smoothed. Where the total energy $\hat{e}$ is integrated we smooth the basic variable $u$ construct the total energy from the sum of the kinetic and internal energies.

Such smoothing of the initial conditions can be avoided altogether if the density summation (3.42) is used, particularly if the smoothing length is updated self-consistently with the density. The results of this problem using unsmoothed initial conditions are shown in Figure 3.9. The artificial viscosity is applied uniformly whilst no artificial thermal conductivity has been used. In this case the rarefaction profile agrees extremely well with the exact solution (solid line). The unsmoothed initial conditions highlight the need for artificial thermal conductivity since the thermal energy overshoots at the contact discontinuity with a resulting glitch in the pressure profile. The gradient in thermal energy at the shock front does not show this effect due to the smoothing of the shock by the artificial viscosity term. The results of this test with a small amount of artificial thermal conductivity applied using the switch discussed in §3.5.2 are shown in Figure 3.10. The variable smoothing length terms have also been used in this case, although results are similar with a simple average of the kernel gradients in the force equation (3.54). The contact discontinuity is smoothed over several smoothing lengths by the thermal conductivity term, removing the overshoot in the thermal energy. The resulting profiles compare extremely well with the exact solution (solid line).

Finally, the results of this test where both the artificial viscosity and conductivity are controlled using the switches described in §3.5.2 are shown in Figure 3.11. The top row shows the velocity and thermal energy profiles compared with the exact solution (solid line), whilst the bottom row shows the time-varying co-efficients $\alpha$ and $\alpha_u$ of the viscosity and thermal conductivity respectively. With the unsmoothed initial conditions and the viscosity switch there is a slight oscillation in the velocity profile at the head of the rarefaction wave. The variable smoothing length terms have been used in this case involving the consistent update of the smoothing length with density (§3.3.4). If a simple average of the kernel gradients is used
Figure 3.8: Results of the Sod shock tube problem in one dimension. The simulation uses 450 particles with conditions in the fluid initially to the left of the origin given by \((\rho, P, v_x) = [1, 1, 0]\) whilst conditions to the right are given by \((\rho, P, v_x) = [0.125, 0.1, 0]\) with \(\gamma = 1.4\). Initial profiles of density and pressure have been smoothed and artificial viscosity is applied uniformly. Agreement with the exact solution (solid line) is generally good, but note the deviation from the exact solution in the rarefaction wave due to the initial smoothing.

Figure 3.9: Results of the Sod shock tube problem using unsmoothed (purely discontinuous) initial conditions. Artificial viscosity has been applied uniformly whilst no artificial thermal conductivity has been used. In the absence of any smoothing of the initial conditions the rarefaction profile agrees well with the exact solution (solid line). The thermal energy is observed to overshoot at the contact discontinuity. There is also a small overshoot in velocity at the right end of the rarefaction wave.
3.7 Numerical tests

Figure 3.10: Results of the Sod shock tube problem using unsmoothed initial conditions and applying a small amount of artificial thermal conductivity using the switch described in §3.5.2. Artificial viscosity is applied uniformly. The overshoot in the thermal energy observed in Figure 3.9 is corrected for by the smoothening of the contact discontinuity produced by the thermal conductivity term. The variable smoothing length terms have also been used in this case, although results are similar with a simple average of the particle kernels.

Figure 3.11: Velocity and thermal energy profiles (top row) in the Sod shock tube problem using unsmoothed initial conditions and where both artificial viscosity and thermal conductivity are applied using the switches discussed in §3.5.2. The bottom row shows the time-varying co-efficients $\alpha$ and $\alpha_u$ of the viscosity and thermal conductivity respectively. With the unsmoothed initial conditions and the viscosity switch there is a slight oscillation in the velocity profile at the head of the rarefaction wave. The variable smoothing length terms have also been used in this case.
instead the oscillations in the rarefaction wave are still present but slightly less pronounced. In effect, the iterations of density and smoothing length make the scheme much more sensitive to small perturbations, since a small change in the smoothing length will be reflected in the density profile and vice-versa. This means that structures in the simulation are in general better resolved and is clearly advantageous. However also mean that small errors in the density evolution are amplified where they may otherwise have been smoothed out by the numerical scheme.

### 3.7.4 Blast wave

In this test we consider a more extreme version of the shock tube test considered previously. In this problem the initial conditions in the fluid to the left of the origin are given by \((\rho, P, v_x) = [1, 1000, 0]\) whilst conditions to the right are given by \((\rho, P, v_x) = [1, 0, 1, 0]\) with \(\gamma = 1.4\). The 10\(^4\) pressure ratio across the initial discontinuity results in a strong blast wave which propagates into the fluid to the right of the origin. The velocity of the contact discontinuity is very close to that of the shock, producing a sharp density spike behind the shock front. This test therefore presents a demanding benchmark for any numerical code.

The results of this test at \(t = 0.01\) are shown in Figure 3.12. The agreement with the exact solution (solid line) is excellent. The contact discontinuity is spread sufficiently by the artificial thermal conductivity to be resolved accurately. In this simulation the density summation and the average of the kernel gradients has been used.
used and the artificial viscosity is applied using the viscosity switch. The SPH results may be compared with those given in Monaghan (1997). Although we use the same formulation of the dissipative terms as in Monaghan (1997), in that paper the artificial thermal conductivity is applied only for particles in compression, resulting in a need to smooth the initial discontinuity in the pressure. With the thermal conductivity term applied using the switch the contact discontinuity is spread sufficiently in order to be resolved accurately and smoothing of the initial conditions is therefore unnecessary. In the SPH solution given by Monaghan (1997) the spike in density is observed to overshoot the exact solution, which is not observed in this case. This is due to the use of the density summation (3.42) rather than evolving the continuity equation (3.43) as in Monaghan (1997). Use of the continuity equation is more efficient since it does not require an extra pass over the particles in order to calculate the density. Using alternative formulations of the pressure term in the momentum equation (e.g. using equation (3.76) with $\sigma = 1$) gives similar results (although the Hernquist and Katz (1989) formulation (3.92) appears to produce negative pressures on this problem). Using the consistent alternative formulations of the continuity equation, however, appears to worsen the overshoot observed in the density spike compared to the usual continuity equation (for example in the $\sigma = 1$ case, the density spike overshoots to $\rho_{\text{max}} \approx 10$ when the continuity equation (3.86) is used).

### 3.7.5 Cartesian shear flows

In a recent paper Imaeda and Inutsuka (2002) (hereafter II02) have suggested that SPH gives particularly poor results on problems involving significant amounts of shear. The simplest test considered by II02 is a Cartesian shear flow. The setup is a two dimensional, uniform density $\rho = 1$ box in the domain $0 \leq x \leq 1$ and $0 \leq y \leq 1$ with a shear velocity field $v_x = 0, v_y = \sin(2\pi x)$ and periodic boundary conditions in the $x-$ and $y-$ directions. In general such flows are known (at least in the incompressible case) to be unstable to Kelvin-Helmholtz instabilities at the inflection point in the velocity profile (e.g. Drazin and Reid, 1981). However, a straightforward stability analysis of this flow demonstrates that it is indeed stable to small perturbations in the $x-$direction (note, however that the application of viscosity can significantly affect the stability properties for these types of problems).

We setup the problem using 2500 (50 x 50) particles initially arranged on a cubic lattice. The smoothing length we use is set according to

$$h = \eta \left( \frac{m}{\rho} \right)^{\frac{1}{2}},$$

where we use $\eta = 1.2$, resulting in an initially uniform value of $h = 0.024$. The smoothing length is allowed to change with density according to (3.68), although this has little effect since the density remains close uniform throughout the simulation. The equation of state is isothermal such that the pressure is given in terms of the density via $P = c_s^2 \rho$. As in II02, we consider both the pressure-free case ($c_s = 0$) and also using $c_s = 0.05$, in both cases using no artificial viscosity. The results for the pressure-free case are shown in Figure 3.13. After 50 dynamical times (defined as one crossing of the computational domain at the highest velocity, i.e. in this case $t_{\text{dyn}} = 1$) the density remains extremely close to uniform ($\Delta \rho \approx 10^{-3} \rho$) and the
particle positions remain ordered. Results in II02 show large errors ($\Delta \rho / \rho \gtrsim \rho$) in the density in less than 1 dynamical time. Similar results are obtained in the $c_s = 0.05$ case, shown after 20 dynamical times in Figure 3.14. Again, the amplitude of the density error is very small ($\Delta \rho \approx 10^{-2} \rho$). Some disruption in the particle distribution is observed to occur at later times, however in the absence of any artificial viscosity small compressible modes are not damped in any way and in the absence of a high accuracy timestepping algorithm such disorder might reasonably be expected. Also it is well known that the particles initially arranged on a cubic lattice will eventually move off the lattice and settle to a more isotropic close packed distribution (e.g. Morris 1996).

![Figure 3.13: Particle positions (left) and density evolution (right) in the pressure-free Cartesian shear flow test with shear velocity field $v_x = 0, v_y = \sin(2\pi x)$. The amplitude of the density error is extremely small ($\Delta \rho \approx 10^{-3} \rho$)](image)

![Figure 3.14: Particle positions (left) and density evolution (right) in the Cartesian shearing box test with sound speed $c_s = 0.05$ and shear velocity field $v_x = 0, v_y = \sin(2\pi x)$. The amplitude of the density error is very small ($\Delta \rho \approx 10^{-2} \rho$)](image)

The question is, therefore: Why do the results obtained in II02 show so much error in the density evolution? The major factor appears to be the particle setup. The details of the particle setup are not given in II02, however by inspection of their figures it appears that the particles are arranged in a quasi-random fashion. The density errors observed in their paper may therefore be an amplification (by the shear flow) of initial perturbations in the density distribution due to the particle setup. A second contributing factor is that the value of smoothing length used by II02 is very low (they use $\eta = 1$ in equation (3.138), whereas typical values for $\eta$ lie in the range $1.1 - 1.2$ in most multidimensional SPH implementations). However, even with their choice of smoothing length $h = 1.0(m/\rho)^{1/2}$, we still find that the density remains essentially constant.
3.7 Numerical tests

Figure 3.15: Toy star static structure. 200 SPH particles are set up in an initially uniform distribution along the x axis and allowed to evolve under the influence of the linear force. The SPH particles are shown by the solid points after damping to an equilibrium distribution. The agreement with the exact quadratic \( \rho = 1 - x^2 \) solution (solid line) is extremely good.

3.7.6 Toy stars

A disadvantage of many of the test problems found in the astrophysical fluid dynamics literature is that, being designed for grid-based codes, they all involve some kind of boundary condition. For codes designed ultimately to simulate self-gravitating gas it is useful to have benchmarks based on a finite system. Secondly, simple, exact, nonlinear solutions to the equations of hydrodynamics are few and far between, and this even more so in the case of magnetohydrodynamics. For this reason we investigate benchmarks based on a simple class of exact solutions which we call ‘Toy Stars’. The equations of hydrodynamics are modified by the addition of a linear force term which is proportional to the co-ordinates (which means that the particles move in a paraboloidal potential centred on the origin). The one dimensional equation of motion is given by

\[
\frac{dv}{dt} = -\frac{1}{\rho} \frac{\partial P}{\partial x} - \Omega^2 x, \tag{3.139}
\]

where \( \Omega \) is the angular frequency. In the following we rescale the equations in units such that \( \Omega^2 = 1 \). The toy star force has many interesting properties and was even considered by Newton as an example of the simplest many-body force. The toy star equations with \( \gamma = 2 \) are also identical in form to the shallow water equations.

Assuming a polytropic equation of state (ie. \( P = K\rho^\gamma \)) with constant of proportionality \( K = 1/4 \) and \( \gamma = 2 \), the Toy Star static structure at equilibrium is easily derived from (3.139) as

\[
\rho = \rho_0 (1 - x^2) \tag{3.140}
\]

In this thesis we will simply consider the most interesting toy star problem which is the calculation of the fundamental oscillatory mode since it turns out to be an exact, non-linear solution. However, a perturba-
tion analysis can be used to derive linear solutions to the Toy Star equations which also present interesting benchmarks for numerical codes. An investigation of the linear modes using SPH, together with a detailed comparison of the oscillation frequencies with the linear solution is given in Monaghan and Price (2004). The non-linear solution for arbitrary $\gamma$ may be derived by considering velocity perturbations in the form

$$v = A(t)x,$$  \hfill (3.141)

where the density is given by

$$\rho^{\gamma-1} = H(t) - C(t)x^2.$$  \hfill (3.142)

The exact solution (Monaghan and Price, 2004) for the parameters $A$, $H$ and $C$ is given in terms of the ordinary differential equations

$$\dot{H} = -AH(\gamma - 1),$$  \hfill (3.143)

$$\dot{A} = \frac{2K\gamma}{\gamma - 1}C - 1 - A^2,$$  \hfill (3.144)

$$\dot{C} = -AC(1 + \gamma).$$  \hfill (3.145)

which can be solved numerically with ease. The relation

$$A^2 = -1 - \frac{2\sigma C}{\gamma - 1} + kC^\frac{1}{\gamma-1},$$  \hfill (3.146)

where $k$ is a constant which is determined from the initial values of $A$ and $C$. The exact solution equations (3.143)-(3.145) take particularly simple forms for the case $\gamma = 2$.

**Static structure**

The simplest test with the toy star is to verify the static structure. We setup 200 SPH particles equally spaced along the x axis with $x = [-1, 1]$ with zero initial velocity and a total mass $M = 4/3$. The particles are then allowed to evolve under the influence of the linear force, with the velocities damped using the artificial viscosity. The particle distribution at equilibrium is shown in Figure 3.15 and shows extremely good agreement with the exact solution (eq. 3.140).

**Non linear test cases**

For the non-linear tests the one dimensional Toy star is initially set up using 200 equal mass particles distributed along the x axis. Although in principle we could use the particle distribution obtained in the previous test as the initial conditions, it is simpler just to space the particles according to the static density profile (3.140). The SPH equations are implemented using the summation over particles to calculate the density and the usual momentum equation with the linear force subtracted. The equation of state is specified by
3.7 Numerical tests

Figure 3.16: Results of the SPH non linear Toy star simulation with $\gamma = 2$ and initial conditions $v = x$, $\rho = 1 - x^2$ (ie. $A = C = H = 1$). Velocity and density profiles are shown after approximately one oscillation period, with the SPH particles indicated by the solid points and the exact solution by the solid line in each case. Equal mass particles are used with a variable initial separation.

Figure 3.17: Results of the SPH non linear Toy star simulation with $\gamma = 5/3$ and initial conditions $v = x$, $\rho = (1 - x^2)^{3/2}$ (ie. $A = C = H = 1$ with $\gamma = 5/3$). Velocity and density profiles are shown after approximately three oscillation periods and the exact solution is given by the solid line.

using $P = K \rho^\gamma$, where for the cases shown we set $K = 1/4$. The smoothing length is allowed to vary with the particle density, where we take simple averages of kernel quantities in the SPH equations in order to conserve momentum.

The exact (non-linear) solution is obtained by numerical integration of equations (3.143)-(3.145) using a simple improved Euler method. We use the condition (3.146) as a check on the quality of this integration by evaluating the constant $k$, which should remain close to its initial value.

Results for the case where initially $A = C = H = 1$ (and therefore $k = 4$) are shown in figure 3.16 at $t = 3.54$ (corresponding to approximately one oscillation period) alongside the exact solution shown by the solid lines. No artificial viscosity is applied in this case. The agreement with the exact solution is excellent.
Note that the sound speed in this case is $C_s = 1/\sqrt{2}$ such that using the parameter $A = 1$ results in supersonic velocities at the edges of the star (the solution is therefore highly non-linear).

Figure 3.17 shows the SPH results for a simulation with $\gamma = 5/3$ and the same initial parameters as Figure 3.16. Velocity and density profiles are shown at time $t = 11.23$ corresponding to approximately three oscillation periods. No artificial viscosity is used. The agreement with the exact solution (solid lines) is again extremely good.

Results of simulations with artificial viscosity turned on are similar, although with a small damping of the kinetic energy over time.

3.8 Summary

In this chapter we have thoroughly reviewed the SPH algorithm. Alternatives to the standard cubic spline kernel were investigated in §3.2.5 and §3.2.6, on the basis of their stability properties. Higher order spline kernels giving closer approximations to the Gaussian were found to give better stability properties although at the price of an increase in computational expense due to the greater number of contributing neighbours. The possibility of constructing kernels with better stability properties based on smoother splines but retaining compact support of size $2h$ was investigated, with good results for smoothing lengths $h \gtrsim 1.1$ (in units of the average particle spacing). However, the gain in accuracy from the use of these alternative kernels is very minor compared to the substantial improvements in accuracy gained by the incorporation of the variable smoothing length terms (§3.3.4)

The discrete equations of SPH were formulated self-consistently from a variational principle in §3.3, leading naturally to equations which explicitly conserve momentum, angular momentum and energy. Artificial dissipation terms used to capture shocks were then discussed, where in §3.5.2 a new switch to control the application of artificial thermal conductivity was considered (the importance of which is highlighted in the numerical tests described in §3.7). The consistent formulation of the SPH equations incorporating a variable smoothing length was discussed in §3.3.4, which are shown to lead to increased accuracy in a wide range of problems (including linear waves (§3.7.2), shock tubes (§3.7.3), Cartesian shear flows (§3.7.5) and toy stars (§3.7.6)). It was shown in §3.4 that consistent formulations of SPH when alternative formulations of the momentum equation are used can be derived from a variational principle by modifying the form of the continuity equation. Various timestepping algorithms were discussed in §3.6, particularly the need to perform a separate pass over the particles to compute derivatives involving the velocity for a reversible integration of the SPH equations. Finally several numerical tests were presented.

The linear sound wave tests (3.7.2) demonstrated a phase error in the SPH simulation of sound waves dependent on the value of the smoothing length and related to the use of kernels with compact support. This phase error was shown to be largely corrected for by allowing the smoothing length to vary with density and self-consistently accounting for the extra terms which arise in the SPH equations. Also the damping of small perturbations induced by the artificial viscosity term was found to be significantly reduced by use of the artificial viscosity switch described in §3.5.2. In the second test problem, the standard shock tube test of Sod (1978), the importance of applying a small amount of artificial thermal conductivity was highlighted, which
3.8 Summary

avoids the need to artificially smooth the initial conditions of such problems. The SPH algorithm was also shown to give good results on a more extreme version of this test (§3.7.4). Thirdly (§3.7.5), the Cartesian shear flow tests given by Imaeda and Inutsuka (2002) were examined, demonstrating that SPH gives good results on this problem for uniform particle setups and does not show the large errors encountered by these authors. Finally, the SPH algorithm was tested against several exact, non-linear solutions derived for systems of particles, known as ‘Toy Stars’ and was shown to give results in excellent agreement with theory.
Appendix B

SPH stability analysis

In this appendix we perform a stability analysis of the standard SPH formalism derived in §3.3. Since the SPH equations were derived directly from a variational principle, the linearised equations may be derived from a second order perturbation to the Lagrangian (3.46), given by

$$\delta L = \sum_b m_b \left[ \frac{1}{2} v_b^2 - \delta \rho_b \frac{d u_b}{d \rho_b} - \frac{(\delta \rho_b)^2}{2} \frac{d^2 u_b}{d \rho_b^2} \right]$$

(B.1)

where the perturbation to $\rho$ is to second order in the second term and to first order in the third term. The density perturbation is given by a perturbation of the SPH summation (3.42), which to second order is given by

$$\delta \rho_a = \sum_b m_b \delta x_{ab} \frac{\partial W_{ab}}{\partial x_a} + \sum_b m_b (\delta x_{ab})^2 \frac{\partial^2 W_{ab}}{\partial x_a^2}$$

(B.2)

The derivatives of the thermal energy with respect to density follow from the first law of thermodynamics, ie.

$$\frac{d u}{d \rho} = \frac{P}{\rho^2}, \quad \frac{d^2 u}{d \rho^2} = \frac{d}{d \rho} \left( \frac{P}{\rho^2} \right) = \frac{c_s^2 - 2P}{\rho^3}$$

The Lagrangian perturbed to second order is therefore

$$\delta L = \sum_b m_b \left[ \frac{1}{2} v_b^2 - \frac{P_b}{\rho_b^2} \sum_c m_c (\delta x_{bc})^2 \frac{\partial^2 W_{bc}}{\partial x_a^2} - \frac{(\delta \rho_b)^2}{2 \rho_b^2} \left( c_s^2 - \frac{2P_b}{\rho_b} \right) \right]$$

(B.3)

The perturbed momentum equation is given by using the perturbed Euler-Lagrange equation,

$$\frac{d}{dt} \left( \frac{\partial L}{\partial v_a} \right) - \frac{\partial L}{\partial (\delta x_a)} = 0.$$  

(B.4)

---

1Note that the first order term may be decoded into continuum form to give the usual expression

$$\delta \rho = -\rho_0 \nabla \cdot (\delta r)$$

where $\rho_0$ refers to the unperturbed quantity.
where
\[
\frac{\partial L}{\partial v_a} = m_a v_a \quad (B.5)
\]
\[
\frac{\partial L}{\partial (\delta x_a)} = -m_a \sum_b m_b \left( \frac{P_a}{\rho_a} + \frac{P_b}{\rho_b} \right) \delta x_{ab} \frac{\partial^2 W_{bc}}{\partial x_a} \nonumber
\]
\[
- \frac{m_a}{2} \sum_b m_b \left[ \delta \rho_a \left( \frac{c_s^2 - 2P_b}{\rho_b} \right) + \delta \rho_b \left( \frac{c_s^2 - 2P_a}{\rho_a} \right) \right] \frac{\partial W_{ab}}{\partial x_a} \quad (B.6)
\]
giving the SPH form of the linearised momentum equation
\[
\frac{d^2 \delta x_a}{dt^2} = - \sum_b m_b \left( \frac{P_a}{\rho_a} + \frac{P_b}{\rho_b} \right) \delta x_{ab} \frac{\partial^2 W_{bc}}{\partial x_a} \nonumber
\]
\[
- \sum_b m_b \left[ \left( \frac{c_s^2 - 2P_b}{\rho_b} \right) \delta \rho_a + \left( \frac{c_s^2 - 2P_a}{\rho_a} \right) \delta \rho_b \right] \frac{\partial W_{ab}}{\partial x_a} \quad (B.7)
\]
Equation (B.7) may also be obtained by a direct perturbation of the SPH equations of motion derived in §3.3.2. For linear waves the perturbations are assumed to be of the form
\[
x = x_0 + \delta x, \quad (B.8)
\]
\[
\rho = \rho_0 + \delta \rho, \quad (B.9)
\]
\[
P = P_0 + \delta P. \quad (B.10)
\]
where
\[
\delta x_a = X e^{ik_a \cdot x}, \quad (B.11)
\]
\[
\delta \rho_a = D e^{ik_a \cdot x}, \quad (B.12)
\]
\[
\delta P_a = c_s^2 \delta \rho_a. \quad (B.13)
\]
Assuming equal mass particles, the momentum equation (B.7) becomes
\[
- \omega^2 X = \frac{2mP_0}{\rho_0^2} X \sum_b \left[ 1 - e^{ik_a \cdot x_b} \right] \frac{\partial^2 W}{\partial x_a^2} \frac{m}{\rho_0} \left( \frac{c_s^2 - 2P_b}{\rho_b} \right) D \sum_b \left[ 1 + e^{ik_a \cdot x_b} \right] \frac{\partial W}{\partial x_a} \quad (B.14)
\]
From the continuity equation (3.43) the amplitude $D$ of the density perturbation is given in terms of the particle co-ordinates by
\[
D = X m \sum_b \left[ 1 - e^{ik_a \cdot x_b} \right] \frac{\partial W}{\partial x_a} \quad (B.15)
\]
Finally, plugging this into (B.14) and taking the real component, the SPH dispersion relation (for any equation of state) is given by
\[
\omega_a^2 = \frac{2mP_0}{\rho_0^2} X \sum_b \left[ 1 - \cos k(x_a - x_b) \right] \frac{\partial^2 W}{\partial x_a^2} (x_a - x_b, h)
\]
\[ \frac{m^2}{\rho_0^2} \left( c_s^2 - \frac{2P_0}{\rho_0} \right) \left[ \sum_b \sin k(x_a - x_b) \frac{\partial W}{\partial x}(x_a - x_b, h) \right]^2, \]  

(B.16)

For an isothermal equation of state this can be simplified further by setting \( c_s^2 = \frac{P_0}{\rho_0} \). An adiabatic equation of state corresponds to setting \( c_s^2 = \gamma P_0 / \rho_0 \).
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