# Multimaterial Simulations using Ghost Fluid Methods

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# Abstract

The unsteady, compressible Euler equations for multimaterial flow in one dimension have been solved numerically by employing level-set based tracking of fronts and two versions of the Ghost Fluid Method. Equations of state for stiffened ideal gases are supported, to allow simulations with water. An exact Riemann solver is demonstrated, in addition to usage of both centered and upwind higher-order time-marching schemes, namely MUSCL-Hancock with HLLC and SLIC. Results from several varied test cases show that the implementation works in accordance with that presented in previous literature.

### 1. Introduction

Leonhard Euler first presented the momentum and continuity equations in  $1757^{[1]}$ , which were completed by the adiabatic condition presented by Laplace in 1816<sup>[2]</sup>. The energy balance equation, which is the last of what is now called the Euler equations, was not properly incorporated until the late nineteenth century [3], and although the much more general Navier-Stokes equations have been developed [4,5], the continued interest for and usefulness of the Euler equations is undisputable. They provide a robust framework for analyzing ideal fluids when viscuous effects are negligible, but cannot generally be solved analytically. Applications of the Euler equations include aerodynamics [6-8], atmospheric modelling and weather forecasts [9,10], astrophysics [11] and detonations and explosives [12–14], to name a few. Accordingly, precise and efficient methods for solving these non-linear, hyperbolic partial differential equations on arbitrary domains has been, and still is, a major field in modern computational fluid dynamics.

The Euler equations govern adiabatic and inviscid flow of a fluid. In the Froude limit (no external body forces) in one dimension, with density  $\rho$ , velocity u, total energy E and pressure p, they are given by

$$\partial_t \mathbf{U} + \partial_x \mathbf{F}(\mathbf{U}) = 0, \qquad (1)$$

where the vector of conserved quantities  $\mathbf{U}$  and their fluxes  $\mathbf{F}(\mathbf{U})$  are given by

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$$\mathbf{U} = \begin{bmatrix} \rho \\ \rho u \\ E \end{bmatrix} , \quad \mathbf{F} = \begin{bmatrix} \rho u \\ \rho u^2 + p \\ u(E+p) \end{bmatrix} .$$

It is sometimes convenient to work in terms of the primitive variables  $\mathbf{W} = \begin{pmatrix} \rho, & u, & p \end{pmatrix}^T$ . The total energy is the sum of the kinetic and potential energy of the system, i.e.

$$E = \frac{1}{2}\rho u^2 + \rho e\,, (2)$$

where e is the internal energy, related to the other variables through the equation of state. For an ideal gas, the equation of state is

$$e = \frac{p}{(\gamma - 1)\rho}, \tag{3}$$

where  $\gamma$  denotes the ratio of specific heats for the gas. Several other fluids can be approximated by a so-called stiffened ideal gas equation of state,

$$e = \frac{p + \gamma p_{\infty}}{(\gamma - 1)\rho} \,. \tag{4}$$

Here, a material-dependent stiffening parameter  $p_{\infty}$  has been introduced. Note that Eq. (4) reduces to Eq. (3) for materials with  $p_{\infty} = 0$ .

Many important milestones have led us to the current state of the art of solving the Euler equations numerically. Riemann identified and worked on the initial value problem for Eq. (1) with discontinuous initial conditions as early as  $1860^{[15]}$ ,

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but the first exact solution did not arrive until Godunov proposed an iterative scheme a century later [16]. Today, efficient approximate solvers such as HLLC (Harten-Lax-van Leer-Contact) [17] and Rotated-hybrid Riemann solvers [18] are readily available. The development of numerical analysis methods for partial differential equations got its first proper boost after the famous paper by Courant, Friedrichs and Lewis [19], but local Riemann problems<sup>[16]</sup>, conservative methods<sup>[20]</sup> and finite volume formulations<sup>[21]</sup> were necessary before the first proper, three-dimensional simulations of the Euler equations could be performed in the 1980s<sup>[22]</sup>. After the Monotonic Upstream-centered Scheme for Conservation Laws (MUSCL)<sup>[23]</sup> was introduced by van Leer as the first higher-order Total Variation Diminishing (TVD) scheme in 1979, several others have been developed, notably Toro's Weighted Average Flux (WAF) [24] and Flux/Slope LImiter Centred (SLIC/FLIC)<sup>[25]</sup> schemes.

Considerations of multimaterial interactions and flow are among the more recent (less than twenty year old) developments within numerical solutions for the Euler equations. It is the goal of this report to demonstrate some techniques available for multimaterial simulations of the Euler equations in one dimension. Different fluids are characterized by their material properties, such as  $\gamma$ , and thus special care must be taken in the treatment of boundaries separating different fluids.

Firstly, the location of the boundary must be tracked as time evolves. The most common way of doing this is by the use of level set methods, as proposed by Osher and Sethian<sup>[26]</sup>. The level set function for a region is initiated such that it is negative inside the region, positive outside it and zero on the boundary. By taking into consideration the convection of the fluids under consideration, the level set function is then evolved so that its zeros move with the domain boundary. Osher and Fedkiw<sup>[27]</sup> provide an excellent overview of level set methods and their applications.

Secondly, the interaction between different fluids across the material interfaces must be modelled accurately. In order to do so, Fedkiw et al. developed the Original Ghost Fluid Method (OGFM)<sup>[28]</sup>. In ghost fluid methods, each material has ghost cells on the side of the boundary where the other material exists. In the OGFM, the pressures and velocities at these ghost cells are similar to that of the other material, and the densities are calculated by enforcing constant entropy across

the interface. This worked well for cases except when there were strong shock waves, since the entropy condition does not apply. As a response, the Modified Ghost Fluid Method (MGFM) was developed [29,30]. It is based on the realization that Riemann problems do not in general require the states on each side of the discontinuity to correspond to the same material. The cells in the ghost fluid are therefore updated by solving a local Riemann problem. A further development is the real Ghost Fluid Method (rGFM) [31], in which a more accurate interfacial boundary condition is applied. The implementation of the rGFM is outside the scope of this report.

The numerical methods which have been employed are explained in section 2, before several test cases and their results are discussed in section 3. Section 4 concludes the report.

#### 2. Numerical methods

### 2.1. Riemann solvers

Given a conservation equation and two sets of piecewise constant states separated by a single discontinuity, the initial value problem of evolving this system in time is called a Riemann problem. It is very useful in the study of the Euler equations for two reasons. Firstly, it allows for exact (up to an arbitrary accuracy) solutions for systems obeying Eq. (1) which have a single contact discontinuity. The other benefit of Riemann solvers is that the discretization of space which is inevitable in computational schemes for solving differential equations, allows for precise solvers of conservation equations based on the solutions of many local Riemann problems.

For the Euler equations and initial conditions

$$\mathbf{W}(x,t=0) = \begin{cases} \mathbf{W}_L, & x \le 0 \\ \mathbf{W}_R, & x > 0 \end{cases},$$

typical states the system can have are shown in Figure 1, in addition to characteristics for waves propagating in space-time. There are two types of resultant waves that propagate through space, in addition to the contact wave (dashed). The first is a shock wave, depicted as a thick line on the left, while the second is a rarefaction wave, shown as several gradually decaying lines on the right. Four different combinations of shock and rarefaction waves can occur on the left and right sides of the contact

discontinuity, and the result is only dependent on  $\mathbf{W}(x,0)$ .

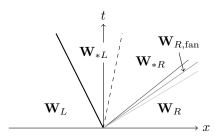


Figure 1: Possible wave configurations for the Riemann problem for Euler's equations in one dimension.

As seen from Figure 1, there are (up to) four unknown states for the Riemann problem: the left and right star states  $\mathbf{W}_{*K}$  located between the left and right waves, and additionally the states inside rarefaction waves  $\mathbf{W}_{K,\mathrm{fan}}$ . Note that the density in the K star state depends on whether the corresponding wave is of shock of rarefaction type. One must therefore be able to compute all these states, and also determine which of them is the correct one based on the speeds of the waves.

### 2.1.1. Exact solver

The exact solver, as first introduced by Godunov<sup>[16]</sup>, has been implemented following Chapter 4 of Toro's comprehensive overview of Riemann solvers and their uses<sup>[32]</sup>, but extended to take into consideration that different (stiffened) ideal gases may be present on each side of the discontinuity.

It can be proven that the pressure in the star states  $p_*$  is given by the root of the nonlinear function

$$f(p) = f_L(p) + f_R(p) - (u_L - u_R).$$
 (5)

Here,

$$f_K(p) = \begin{cases} \frac{2a_K}{\gamma_K - 1} \left[ \left( \frac{p + p_{\infty, K}}{p_K + p_{\infty, K}} \right)^{\frac{\gamma_K - 1}{2\gamma_K}} - 1 \right], & p \le p_K \\ \left( p - p_K \right) \left( \frac{A_K}{B_K + p} \right)^{\frac{1}{2}}, & p > p_K \end{cases},$$

where  $a_K = \sqrt{\frac{\gamma(p_K + p_{\infty,K})}{\rho_K}}$  is the speed of sound and

$$A_K = \frac{2}{(\gamma_K+1)\rho_K} \,,\, B_K = \frac{(\gamma_K-1)p_K+2\gamma_K p_{\infty,K}}{\gamma_K+1} \,. \label{eq:AK}$$

In the exact solver, the pressure in the star region is found by applying an iterative scheme to Eq. (5). Due to the nature of f(p), it is well-suited for iterative schemes, but it is important to ensure that the pressure stays positive. For this project, Newton-Raphson iterations were implemented, taking the initial guess as  $\frac{1}{2}(p_L + p_R)$ . After finding  $p_*$ , the velocity in the star region is computed as

$$u_* = \frac{u_L + u_R - [f_L(p_*) - f_R(p_*)]}{2}.$$
 (6)

When the pressure and velocity in the star region have been computed, one can calculate the wave speeds (one per shock wave, two per rarefaction wave and one for the contact discontinuity) and use this to sample the state at a given point S=x/t in space-time. Pseudocode for this procedure is given in Algorithm 1. Formulas for wave speeds and densities in different states have been left out for brevity, but they in general depend on the initial conditions (including material properties).

```
Data: S, \mathbf{W}_L, \mathbf{W}_R, p_*, u_*, material properties Result: \mathbf{W}(S)
```

```
if S \leq speed of contact discontinuity then
    if left wave is rarefaction then
        if S \leq speed of rarefaction head then
            return \mathbf{W}_L;
        else
            if S > speed of rarefaction tail then
               return \mathbf{W}_{*L,\mathrm{fan}};
             return \mathbf{W}_{L,\mathrm{fan}};
            end
        end
    else
        if S \leq speed of shock wave then
            return \mathbf{W}_L;
        else
            return \mathbf{W}_{*L};
        end
    end
else
    perform similar analysis for sampling point
    on right side of contact discontinuity;
```

**Algorithm 1:** Sample exact solution of Riemann problem given pressure and velocity in star states.

#### 2.1.2. HLLC

By far the most computationally expensive part of the exact solver is the iterative root finding procedure for finding  $p_*$ . Since most modern schemes require the solutions of local Riemann problems between all points in the computational domain, high gains in terms of computational efficiency can be achieved by employing exact solvers instead. One such solver is the HLLC solver [17], which has been implemented for this project.

When compared to the exact solver, the main simplifications in HLLC are that the pressure in the star region is approximated instead of found by iterative schemes; that states inside rarefactions are not taken into consideration, and that wave speeds are estimations, either direct or pressure-based. In this contribution, the pressure is estimated as

$$p_* \approx \frac{1}{2}(p_L + p_R) + \frac{1}{8}(u_L - u_R)(\rho_L + \rho_R)(a_L + a_R),$$
(7)

and set equal to zero if the approximation is negative. Based on the approximation for  $p_*$ , the wave speeds  $S_K$  of the left and right travelling waves are computed, with the type of wave taken into consideration. The speed of the contact discontinuity is then approximated as

$$S_* \approx \frac{\rho_L u_L (S_L - u_L) - \rho_R u_R (S_R - u_R) - (p_L - p_R)}{\rho_L (S_L - u_L) - \rho_R (S_R - u_R)}.$$
(8)

There are four possible states, separated by the three (approximated) wave speeds. Contrary to the exact solver, where the output is the state  $\mathbf{W}(S)$ , the HLLC solver outputs a flux  $\mathbf{F}^{\text{HLLC}}(\mathbf{W}_L, \mathbf{W}_R)$  based on the state corresponding to S=0. This is because the flux at the intermediate point between cells is the quantity employed in the update formula as introduced in Subsection 2.2. For an overview of the approximate states and the flux calculation employed in the HLLC solver, see Chapter 9 of Toro's book [32].

## 2.2. Schemes for the Euler equations

The Euler equations as given in Eq. (1) are said to be a system of conservation equations in differential form. In integral form, on a spatial domain  $[x_L, x_R]$ , the same system of equations can be written

$$\int_{x_L}^{x_R} \mathbf{U}(x, t_2) dx = \int_{x_L}^{x_R} \mathbf{U}(x, t_1) dx$$
$$- \int_{t_1}^{t_2} \mathbf{F}(\mathbf{U}(x_R, t)) dt + \int_{t_1}^{t_2} \mathbf{F}(\mathbf{U}(x_L, t)) dt. \quad (9)$$

From Eq. (9), it is straightforward to discretize the Euler equations to produce a conservative time-marching scheme through the Finite Volume Method (FVM). Let the spatial domain  $x \in [0,1]$  be divided into N equal cells of width  $\Delta x = 1/N$ . We denote by  $x_i$  the center of the i-th cell, i.e.  $x_i = (i + 1/2)\Delta x$ . Temporal discretization is done with variable time step  $\Delta t^n$ , so that  $t^n = \sum_{i=1}^n \Delta t^i$ .

We write the discrete approximation of  $\mathbf{U}(x_i, t^n)$  as  $\mathbf{U}_i^n$ , and let it be the weighted average of  $\mathbf{U}(x, t^n)$  in the cell with boundaries at  $x_{i-1/2}$  and  $x_{i+1/2}$ :

$$\mathbf{U}_{i}^{n} \approx \frac{1}{\Delta x} \int_{x_{i-1/2}}^{x_{i+1/2}} \mathbf{U}(x, t^{n}) dx$$
 (10)

Similarly, the fluxes at each interface are approximated as

$$\mathbf{F}_{i+1/2}^{n} \approx \frac{1}{\Delta t^{n}} \int_{t_{n}}^{t_{n+1}} \mathbf{F}(\mathbf{U}(x_{i+1/2}, t)) dt. \quad (11)$$

By applying Eq. (9) on each cell  $[x_{i-1/2}, x_{i+1/2}]$ , and by inserting the discrete approximations in Eqns. (10) and (11), a conservative, time-marching scheme for evolving the Euler equations in time is

$$\mathbf{U}_{i}^{n+1} = \mathbf{U}_{i}^{n} - \frac{\Delta t^{n}}{\Delta x} \left( \mathbf{F}_{i+1/2} - \mathbf{F}_{i-1/2} \right) . \tag{12}$$

The Courant-Friedrichs-Lewis condition for convergence demands that

$$\Delta t^n \le \frac{\Delta x}{S_{\max}^n} \,, \tag{13}$$

so we let  $\Delta t^n = C_{\rm CFL} \Delta x/S_{\rm max}^n$ , where  $0 \le C_{\rm CFL} \le 1$ . In our implementation, we use  $C_{\rm CFL} = 0.9$ . Transmissive boundary conditions are implemented on both boundaries, so that

$$\mathbf{W}_{-i}^{n+1} = \mathbf{W}_{-(i-1)}^{n}, \quad \mathbf{W}_{N+i}^{n+1} = \mathbf{W}_{N+(i-1)}^{n},$$
(14)

for  $i = 1, ..., N_{GC}$  where the number of ghost cells  $N_{GC}$  equals the order of the scheme employed.

All the most common schemes used for solving conservation laws are of the same form as Eq. (12), and the difference between them is in the evaluation of the fluxes  $\mathbf{F}_{i\pm1/2}$ . For this project, we have implemented so-called total variation diminishing (TVD) high resolution methods for this purpose. High resolution methods are accurate to at least second order; they avoid spurious oscillations in the solution, capture discontinuities and shock waves well and satisfy a discrete form of the entropy condition. There are two main classes of higher-order flux evaluation schemes; the upwind schemes are based on the solution of local Riemann problems, while the centered schemes are extensions of the First ORder CEntered (FORCE) flux:

$$\mathbf{F}^{\text{FORCE}}(\mathbf{U}_L, \mathbf{U}_R) = \frac{1}{2} \left[ \mathbf{F}_0 + \frac{1}{2} (\mathbf{F}_L + \mathbf{F}_R) \right] - \frac{1}{4} \frac{\Delta x}{\Delta t^n} (\mathbf{U}_R - \mathbf{U}_L) . \quad (15)$$

Here,  $\mathbf{F}_K = \mathbf{F}(\mathbf{U}_K)$  and

$$\mathbf{U}_0 = \frac{1}{2}(\mathbf{U}_L + \mathbf{U}_R) - \frac{1}{2}\frac{\Delta t}{\Delta x}(\mathbf{F}_R - \mathbf{F}_L).$$
 (16)

In general, upwind schemes offer higher accuracy than centered once, but they are computationally more demanding and difficult to implement.

The MUSCL-Hancock<sup>[23]</sup> and SLIC<sup>[25]</sup> schemes, as mentioned in the Introduction, exemplify upwind and centred schemes, respectively. They are similar in many regards, and both consist of three main steps.

First, the data cell averages  $\mathbf{U}_i^n$  are replaced locally by linear functions so that at the left and right cell interfaces, the boundary extrapolated values are

$$\mathbf{U}_{i}^{L} = \mathbf{U}_{i}^{n} - \frac{1}{2}\Delta_{i}, \quad \mathbf{U}_{i}^{R} = \mathbf{U}_{i}^{n} + \frac{1}{2}\Delta_{i}.$$
 (17)

Here, the slope vector  $\Delta_i$  has been introduced, which in our case is chosen such that the schemes are TVD. For the MUSCL-Hancock scheme, we have implemented limited slopes <sup>[7]</sup>, while SLIC uses slope limiters <sup>[25]</sup>.

After calculating the boundary extrapolated values for each cell, these are advanced to time  $t^n + \Delta t^n$  to obtain

$$\bar{\mathbf{U}}_{i}^{L,R} = \mathbf{U}_{i}^{L,R} - \frac{1}{2} \frac{\Delta t}{\Delta x} \left[ \mathbf{F}(\mathbf{U}_{i}^{R}) - \mathbf{F}(\mathbf{U}_{i}^{L}) \right]. \quad (18)$$

The third and final step is the actual evaluation of the intercell flux  $\mathbf{F}_{i+1/2}$  based on the partially evolved boundary extrapolated values. For MUSCL-Hancock, this is done by using the HLLC solver to solve the local Riemann problem across each cell interface, *i.e.* 

$$\mathbf{F}_{i+1/2}^{\text{MUSCL}} = \mathbf{F}^{\text{HLLC}}(\bar{\mathbf{W}}_{i}^{R}, \bar{\mathbf{W}}_{i+1}^{L}). \tag{19}$$

For SLIC, however, the intercell flux is directly evaluated as the FORCE flux with

$$\mathbf{F}_{i+1/2}^{\text{SLIC}} = \mathbf{F}^{\text{FORCE}}(\bar{\mathbf{U}}_i^R, \bar{\mathbf{U}}_{i+1}^L). \tag{20}$$

#### 2.3. Level-set method

In order to represent the interfaces between materials, we employ a level-set function  $\phi(x)$  whose sign is a boolean indicator of the material present. The interfaces between materials are thus located at the points where  $\phi=0$ . Figure 2 shows the simple case of a single material interface separating materials A and B; several interfaces have been incoorporated by letting  $\phi$  be piecewise linear throughout the domain in a zig-zag pattern.

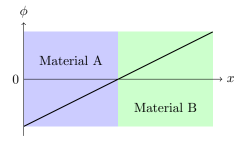


Figure 2: Simple example of level-set function for one material interface. In the left region, where  $\phi(x) < 0$ , material A is present, while material B is present in the right region.

Initializing the level-set function is trivial, but it is a good idea to make sure that the zeros end up between cell centers, so that each grid point has well-defined material properties. In order to capture the evolution of the system, the level-set function itself must be advanced in time for each time step. Since  $\phi$  is a consequence of the state of the fluid rather than a property, it is not coupled to the other variables in the system. Thus it is governed by the advection equation

$$\partial_t \phi + u \partial_x \phi = 0, \qquad (21)$$

Due to frequent discontinuities in density at material interfaces, conservative schemes lead to smearing of the locations of the zeros. Since the only

interesting information contained in  $\phi$  are these locations, non-conservative methods can be safely implemented as long as  $\phi$  remains smooth and not too sharp. In our implementation, we have implemented the first-order upwind scheme

$$\phi_i^{n+1} = \begin{cases} \phi_i^n - u_i^n \frac{\Delta t^n}{\Delta x} (\phi_i^n - \phi_{i-1}^n), & u_i^n > 0\\ \phi_i^n - u_i^n \frac{\Delta t^n}{\Delta x} (\phi_{i+1}^n - \phi_i^n), & u_i^n \le 0 \end{cases}$$
(22)

Note that  $u_i^n$  is the velocity of material A at  $x_i$  if  $\phi_i^n < 0$ , and the velocity of material B otherwise. In order to avoid the level set function becoming too sharp, we reinitialize it after every update by demanding  $|\phi_{i+1}^n - \phi_i^n| = \Delta x$ . In this procedure, we also make sure that the zeros of  $\phi$  lie between grid points.

## 2.4. Ghost fluid methods

Multimaterial simulations require one final consideration in addition to interface tracking; namely which boundary conditions to enforce across these These dynamic boundary conditions should be thermodynamically consistent in order to capture the physical interactions between materials. In the computational implementation, the Euler equations are solved on a separate domain for each material, where that material is present in all cells. For the regions where  $\phi(x) < 0$ , fluid A is present in domain A, while ghost fluid B is present in domain B. The opposite is true when  $\phi(x) > 0$ . The only interesting cells in the ghost fluids are the  $N_{GC}$  points located immediately adjacent to a material interface, since these affect the solution in the actual fluid. It is the goal of ghost fluid methods to update these ghost cells in a thermodynamically consistent manner.

## 2.4.1. Original ghost fluid method

The first successful attempt at implementing a ghost fluid method was performed by Fedkiw et al. in  $1998^{[28]}$ . In this approach, which we refer to as OGFM, the entropy S is held constant across the ghost fluid interface, in order to give conditions for thermodynamic consistency. The assumption of constant entropy is not entirely true in the case of shocks, so one does not expect excellent capturing of large shock waves. The pressure and velocity in the ghost cells are set equal to the pressure and velocity in the same cells of the other material domain. It is the density which is extrapolated by

enforcing constant entropy. To derive how, we note that the entropy of an ideal gas is given by

$$S = c_V \ln \left(\frac{p}{\rho^{\gamma}}\right) + \text{const},$$
 (23)

so if two cells i and j have the same entropy and contain the same material, they must satisfy

$$\rho_i = \left(\frac{p_i}{p_j}\right)^{1/\gamma} \rho_j \,. \tag{24}$$

In order to avoid "overheating" across the material boundary, an isobaric fix has been implemented for OGFM by enforcing constant entropy from the neighbour of the boundary cell to the boundary as well as from the boundary and into the ghost fluid. A schematic illustration, including the isobaric fix, is given in Figure 3.

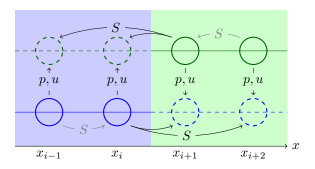


Figure 3: Updating of cells in ghost fluids as implemented in OGFM. The pressure and velocities are inherited from the other material, while entropy is kept constant across the ghost fluid interface. With the isobaric fix (grey), the entropy condition is enforced from further inside the actual fluid, to reduce overheating.

# 2.4.2. Modified ghost fluid method

In order to be able to solve a wider range of problems, including those with strong shock waves or large density gradients, the MGFM was developed  $^{[29,30]}$ . In this approach, the intermaterial boundary conditions are enforced in a way which is based on solving a local Riemann problem across the interface. As explained in Subsection 2.1, it is unproblematic so solve a Riemann problem with different material properties on each side of the initial discontinuity. Given the two states immediately adjacent to the material interface,  $\mathbf{W}_L$  and  $\mathbf{W}_R$ , and the corresponding material properties, we calculate the star states  $\mathbf{W}_{*L}$  and  $\mathbf{W}_{*R}$ , and set the ghost fluid cells equal to them. Thus every extrapolated value in the ghost fluid corresponds to

a physical state of the system, and we have thermodynamical consistency. In order to compute the star states, the exact solver is employed.

### 2.5. Complete algorithm

A summary of all the steps made in the simulations is given in Algorithm 2.

```
Data: initial configuration of system,
  choice of scheme, N, t_{\rm end}
 Result: configuration of system at time t_{\text{end}}
 set t = 0;
 initiate computational domain;
 while t < t_{\text{end}} \text{ do}
     find locations of material interfaces;
     for each interface do
        update ghost fluid using GFM;
     \mathbf{end}
     calculate \Delta t^n from Eq. (13);
     if t + \Delta t^n > t_{\text{end}} then
      \Delta t^n \to t_{\rm end} - t;
     update boundaries according to Eq. (14);
     advance \phi according to Eq. (22);
     for each material do
         compute fluxes using Eq. (19) or (20);
         evolve system in time with Eq. (12);
     end
     t \to t + \Delta t^n;
 \quad \mathbf{end} \quad
 if initially there was one interface then
     compute exact solution for reference;
Algorithm 2: Complete structure for simulating
```

# 3. Results

multimaterial system

The different tests described in this section aim to show how the implemented methods cope under different circumstances. Since there are quite many different choices one can make for each method, we inform that unless otherwise specified:

- fluxes are computed using the MUSCL-Hancock scheme.
- slope limiters are of minbee type,
- OGFM is implemented with the isobaric fix.

#### 3.1. Moving contact discontinuity

To illustrate how the combination of a level set function and GFMs allow capture of a moving contact discontinuity, test A has a single discontinuity in density, and the system moves with constant speed to the right. Full test specifications are given in Table 1. Air occupies the entire domain, so both single-material solvers and multimaterial GFM solvers are employed.

Table 1: Initial conditions for test A.				
$x \in$	ho	u	p	$\gamma$
[0, 0.25]	1.000	0.5	1	1.4
(0.25, 1]	0.138	0.5	1	1.4

Results for test A are shown in Figure 4. Please note that only half of the spatial domain is included, since the discontinuity is located here. As is evident, both ghost fluid methods capture the discontinuity far better than the single-material solvers, in fact no error at all is apparent. Of the schemes used for the single-material solver, the MUSCL-Hancock scheme provides a higher accuracy around the discontinuity, as expected.

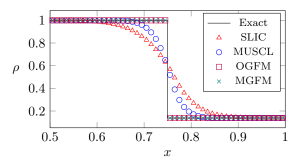


Figure 4: Results from test A with N = 100 and  $t_{end} = 1$ .

Test B is identical to test A except for the fact that the discontinuity now separates two different fluids; namely air and helium ( $\gamma=1.67$ ). As such, only the GFM solvers have been utilized.

Results for test B are shown in Figure 5, which demonstrates that the solvers work well for the multimaterial case as well.

- 3.2. Simple ghost fluid tests
- 3.3. Multimaterial shock tubes for gases
- 3.4. Water-gas shock tube test
- 3.5. More interfaces!

### 4. Conclusions

Everythings went better as expectance!

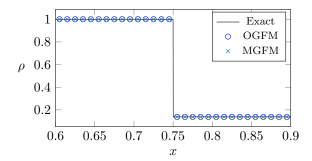


Figure 5: Results from test B with N = 100 and  $t_{end} = 1$ .

#### Acknowledgements

#### Thanks Steve.

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