Computing Interface Motion in Compressible Gas Dynamics

W. Mulder
S. Osher
J.A. Sethian

February 1990
CAM Report 90-03
COMPUTING INTERFACE MOTION IN COMPRESSIBLE GAS DYNAMICS

W. Mulder 1
Koninklijke/Shell Exploratien en Produktie Laboratorium
Rijswijk (Z-H), The Netherlands

S. Osher 2
Dept. of Mathematics, Univ. of California
Los Angeles, California 90024

and

J.A. Sethian 3
Dept. of Mathematics, Univ. of California
Berkeley, California 94720

Abstract

A "Hamilton-Jacobi" level set formulation of the equations of motion for propagating interfaces has been introduced recently by Osher and Sethian. This formulation allows fronts to self-intersect, develop singularities, and change topology. The numerical algorithms based on this approach handle topological merging and breaking naturally, work in any number of space dimensions, and do not require that the moving front be written as a function. Instead, the moving front is embedded as a particular level set in a fixed domain partial differential equation. Numerical techniques borrowed from hyperbolic conservation laws are then used to accurately capture the complicated surface motion that satisfies the global entropy condition for propagating fronts given by Sethian. In this paper, we analyze the coupling of this level set formulation to a system of conservation laws for compressible gas dynamics. We study both conservative and non-conservative differencing of the level set function, and compare the two approaches. As applications, we study the incompressible Rayleigh-Taylor and Kelvin-Helmholtz instabilities for air-air and air-helium boundaries. We perform numerical convergence studies of the method over a range of parameters, and analyze the accuracy of this approach applied to these problems.

1 Research supported in part by ONR Grant N-00014-86-K-0691, NSF Grant DMS88-11863 when this author was in residence at the University of California, Los Angeles

2 Research supported in part by ONR Grant N-00014-86-K-0691, NSF Grant DMS88-11863, DARPA Grant in the ACMP Program, and NASA Langley Grant NAG1-270.

3 Research supported in part by the Applied Mathematics Subprogram of the Office of Energy Research under contract DE-AC03-76SF00098.
COMPUTING INTERFACE MOTION IN COMPRESSIONABLE GAS DYNAMICS

A variety of physical phenomena involve propagating interfaces. The interface (or interfaces) separate regions which may differ according to their density, viscosity, or chemical type. The complexity of the motion of the interface can range from the particularly simple case of passive advection of two different colors, to problems in flame propagation and dendritic solidification, in which there is an intricate feedback mechanism between the local properties of the front and the physics on either side of it.

Recently, a new set of algorithms for following propagating interfaces has been developed. In [46], a Hamilton-Jacobi level set formulation for moving interfaces was introduced. These algorithms handle topological merging and breaking naturally, work in any number of space dimensions, and do not require that the moving front be written as a function. Instead, the moving front is embedded as a particular level set in a fixed domain partial differential equation. Numerical techniques borrowed from hyperbolic conservation laws are then used to accurately calculate the correct solution which satisfies the global entropy condition for propagating fronts given in [58].

These schemes have been used to model a variety of problems in front motion, flame propagation, and the geometry of moving surfaces, see [46,58,59]. This level set formulation of the moving fronts has been used as the basis for theoretical analysis of motion by mean curvature in [9,17].

In this paper, we analyze the coupling of this level set formulation to a system of conservation laws for compressible gas dynamics. We consider two different approaches. In one approach, the level set function is solved in non-conservative form, using the velocity obtained from conservative differencing of the standard hyperbolic system. In another approach, we directly incorporate the level set formulation into a system of five conservation laws, in which the moving front becomes one extra variable in the flow solver. In both the conservative and non-conservative settings, we also analyze a degenerate initialization of our level set approach, known as the color function. We then compare the various approaches, and discuss how the physics of the problem suggest the appropriate approach.
As application, we study the incompressible Rayleigh-Taylor and Kelvin-Helmholtz instabilities for air-air and air-helium boundaries. We compute the position of the moving interface, showing the development of plumes and rolls in the Rayleigh-Taylor instability and the rolling up of vortex structures in the Kelvin-Helmholtz instability. We perform numerical convergence studies of the method over a range of parameters, and analyze the accuracy of this approach applied to these problems.
I. Physical Problems

In this section, we discuss the two physical problems under investigation.

A. Physical Problems

The Rayleigh-Taylor instability occurs when a light fluid pushes a heavier one. Imagine a horizontal interface, in which a fluid with density \( \rho_1 \) lies above a fluid with density \( \rho_2 \). Here we assume that gravity is pointing downwards. If \( \rho_1 < \rho_2 \), the interface is stable, and the two fluids remain motionless. Small perturbations in the initial shape of the interface remain bounded. On the other hand, if \( \rho_1 > \rho_2 \), the interface is unstable. Small perturbations in the initial shape grow as the heavier fluid on the top pushes through these perturbations, and long fingers of the heavier fluid reach down into the lighter fluid. At the same time, plumes of the lighter fluid grow upward. The initial growth rate of the perturbations is exponential. Experimental observations indicate that the heavier fluid forms long "spikes" as it reaches into the lighter fluid, while the rising light fluid forms rounded tops, or "bubbles". The length of the interface increases dramatically, and can break into several parts, developing bubbles. Some examples where this instability can occur are in the collapse of a massive star, the laser implosion of deuterium-tritium fusion targets, and the electromagnetic implosion of a metal liner. One of the most straightforward examples is the novelty-store toy in which fluids of differing densities are trapped between two glass plates. By upending the apparatus, the lighter fluid rises to the top by forming long spikes in the interface. Bubbles can break off from the interface and later merge with other bubbles. The interface between the two fluids becomes highly complex, breaking into numerous different parts with wildly varying shapes.

In their most complicated form, the equations of motion are the equations of full viscous, compressible flow plus interface effects. Some important factors controlling the growth of instability are (1) the density ratio, which governs the growth of small amplitude perturbation (2) surface tension, which stabilizes wavelengths shorter than a critical wavelength (3) the viscosity, which reduces growth rate and regularizes the flow (4) compressibility, which reduces growth rate, and
(5) heterogeneity, which can excite instabilities of various wavelengths.

The Kelvin-Helmholtz instability occurs when one fluid is moving at a different rate relative to another. Imagine one fluid atop another, moving at different speeds initially parallel to the interface. The initial horizontal interface rolls up into large vortical structures, which serve to entrain the fluid. In compressible gas flow, the Kelvin-Helmholtz instability can be seen when a jet of fluid is injected into another, producing large vortical structures which roll up the interface between the two fluids. Another example is provided by parallel shear flow for incompressible fluids, which can be modeled through the study of vortex sheets. Here, the vorticity is zero everywhere except along an infinitely thin line or curve. A good example is flow around as trailing edge of a wing, which forms a vortex sheet whose strength depends on the given wing design. The ensuing motion and rollup of the vortex sheet affects both the drag on the wing and the flight of following aircraft.

For some experimental studies of these phenomena, we refer the interested reader to [12,16,28,29,36,37,50,52,54]. In addition, we draw the interested reader’s attention to the recent experiment on the three-dimensional Rayleigh-Taylor instabilities described in [29]. This paper contains some fascinating photographs of three-dimensional instabilities in circular tubes, and direct comparison with solutions from linear and non-linear theory developed in [28]. For studies of the theoretical aspects of Rayleigh-Taylor and Kelvin-Helmholtz instabilities, a possible starting point may be found in [4,5,6,8,20,27,34,39,40,43,44,49,53,55,57,63].

B. Numerical Studies

Two different types of numerical methods are often employed for computing interface problems in fluid mechanics. The first, or “Eulerian” type, compute the full Navier-Stokes equations in both fluids. In these techniques, the finite difference approximations are typically employed across the entire domain. The second, or “Lagrangian” type, reduce the equations of motion to equations for the interface itself. Here, one often uses markers to track the interface. One example in this category are vortex methods, which rely on a discrete approximation to a boundary integral along
the interface, see [3,11,18,30,31,32,33,39,51,64]. An excellent overview of some work on the Rayleigh-Taylor instability is due to Sharp [61]. Other calculations include [1,2,15,19,26,41,42,45,67]. Some particular beautiful calculations of compressible jets may be found in [7,66].

Hybrid "Eulerian-Lagrange" methods have also been employed. These methods are used in some of the earliest numerical calculations of the Rayleigh-Taylor instability, which were performed by Harlow and Welch [23]. In these calculations, the marker-and-cell method was introduced, in which a finite difference scheme is used to solve the full Navier-Stokes equations. One of the two fluids, say Type 1, is tracked by placing marker points at the centers of cells initially containing the chosen fluid. These markers are then advected with the computed fluid velocity. At subsequent times, cells are divided into three types: (a) those containing marker particles and whose neighboring cells contain marker particles (Type 1 fluid), (b) those not containing marker particles and whose neighboring cells also do not contain marker particles (Type 2 fluid), and (c) surface cells which must contain the boundary. Using this technique, a moving fluid interface was tracked. An extension of this technique was used in [14] to track the growth of a single mode of the Rayleigh-Taylor instability, showing the development of a large bubble and accompanying spike.

The most involved calculations using a combination Eulerian-Lagrangian scheme which couples the Navier-Stokes equations to a method for tracking fronts is the front tracking technology due to Glimm et al. [21,22]. In this work, the compressible Navier-Stokes equations are solved in the whole domain, and the interface is tracked through a set of marker particles on the moving interface. A variety of calculations of bubble and spike development for the Rayleigh-Taylor problem may be found in [21,22,61].
II. Equations of Motion for Propagating Interfaces

A. Statement of Problem

In the most general form, consider a propagating hypersurface $S(t)$ (that is, a curve in two space dimensions or a surface in three space dimensions) separating two regions in the domain. Here, $t$ is time, and $S(t): [0,\infty) \rightarrow \mathbb{R}^N$, $N=2,3$. Suppose that $S(t)$ propagates normal to itself with speed $F$. $F$ may vary along the interface $S(t)$, and depend on such factors as the position of the front $S(t)$, the direction of the normal $\vec{n}(t)$, the local curvature $K(t)$, as well as the time $t$. Note that the dependence of $F$ on the position $S(t)$ can generate tremendous complexity, since the physics on both sides of the interface may enter into the determination of $F$. Our goal is a numerical algorithm that follows the motion of $S(t)$.

It might seem most natural to formulate equations of motion by parameterizing the hypersurface and describing the evolution of the interface in terms of coordinate-free "Lagrangian" front properties, such as the local normal $\vec{n}$ and curvature $K$. Indeed, a standard numerical method for tracking moving fronts relies on discretizing such a parameterization with marker particles whose motion is determined by a discrete approximation to the appropriate equations of motion, see [68]. As shown in [58,59], such techniques can encounter considerable difficulties when sharp corners develop in the propagating interfaces or when the interface changes topology. A rigorous explanation of the inherent instability of this approach is given in the appendix of [46]. Instead, we consider an "Eulerian" formulation of the equations of motion which is more amenable to numerical approximation. The details of this formulation were first presented in [46].

B. Eulerian Formulation

Given a closed hypersurface $\Gamma(t)$, we wish to produce an Eulerian formulation for the motion of the hypersurface propagating along its normal direction with speed $F$. We motivate the Eulerian formulation by a simple example, taken from [59].
Let $\Gamma(t)$ be a unit circle in $\mathbb{R}^2$ propagating outward with constant speed $F=1$ (See Figure 1a). Obviously, the solution at any time $t$ is just a circle with radius $(t+1)$. (See Figure 1b). Rather than describe the motion of this circle, we consider the motion of a surface $z = \psi(x, y, t)$ in $\mathbb{R}^3$ (See Figure 1c). The level set $\psi = 0$ of this surface is just the set of points in the $x-y$ plane corresponding to the propagating curve $\Gamma(t)$. That is,

$$\Gamma(t) = \{(x, y) \mid \psi(x, y, t) = 0\}$$

(2.1)

Thus, we have matched the motion of the front $\Gamma(t)$ in $\mathbb{R}^2$ with the evolution of a function $z = \psi(x, y, t)$ in $\mathbb{R}^3$. At this point, we must describe how to

1. Construct the initial value $\psi(x, y, 0)$

2. Derive the equations of motion for the evolving surface

We shall do this in some generality, referring to an $(N-1)$-dimensional hypersurface with arbitrary speed function $F$.

C. Construction of the Initial Value for $\psi$

Suppose we are given a closed, propagating $(N-1)$-dimensional hypersurface $\Gamma(t)$, where $\Gamma(t) : [0, \infty) \rightarrow \mathbb{R}^N$. A straightforward technique for constructing the initial front $\psi(\mathbf{x}, t=0)$, where $\mathbf{x} \in \mathbb{R}^N$, is to let

$$\psi(\mathbf{x}, t=0) = \pm d$$

(2.2)

where $d$ is the distance from $\mathbf{x}$ to $\Gamma(t=0)$, and the plus (minus) sign is chosen if the point $\mathbf{x}$ is outside (inside) the initial hypersurface $\Gamma(t=0)$. Thus, we have an initial function $\psi(\mathbf{x}, t=0) : \mathbb{R}^N \rightarrow \mathbb{R}$ with the property that

$$\Gamma(t=0) = \{\mathbf{x} \mid \psi(\mathbf{x}, t=0) = 0\}$$

Our goal is to now produce an equation for the evolving function $\psi(\mathbf{x}, t)$ which contains the
embedded motion of $\Gamma(t)$ as the level set $\psi = 0$.

D. Derivation of the Evolution Equation for $\psi$

We are given a propagating hypersurface $\Gamma(t)$ and a speed function $F$ at each point of the propagating hypersurface. Let $\mathbf{X}(t)$, $t \in [0, \infty)$ be the path of a point on the propagating front. That is, $\mathbf{X}(t=0)$ is a point on the initial front $\Gamma(t=0)$, and $\mathbf{X}_t = F(\mathbf{X}(t))$ and the vector $\mathbf{X}_t$ is in the direction normal to the front at $\mathbf{X}(t)$. Since the evolving function $\psi$ is always zero on the propagating hypersurface, we must have

$$\psi(\mathbf{X}(t), t) = 0$$

(2.3)

By the chain rule,

$$\psi_t + \sum_{i=1}^{N} \psi_x \cdot x_i = 0$$

(2.4)

where $x_i$ is the $i^{th}$ component of $\mathbf{X}$. Let $(u_1, u_2, \ldots, u_N) = (x_1, x_2, \ldots, x_N)$. Since

$$\sum_{i=1}^{N} \psi_x \cdot x_i = (\psi_x, \psi_y, \ldots, \psi_w) \cdot (u_1, u_2, \ldots, u_N) = F(\mathbf{X}(t)) \cdot \nabla \psi,$$

(2.5)

we then have the evolution equation for $\psi$, namely

$$\psi_t + F \cdot \nabla \psi = 0$$

(2.6)

We refer to this as a Hamilton-Jacobi "type" equation because, for speed function $F$ identically constant, we obtain a standard Hamilton-Jacobi equation.

To repeat, the position of the propagating hypersurface $\Gamma(t)$ is given as the level set

$$\Gamma(t) = \{ \mathbf{X} | \psi(\mathbf{X}, t) = 0 \}$$

(2.7)

where $\psi(\mathbf{X}, t)$ is the solution to the Hamilton-Jacobi-type equation

$$\psi_t + F \cdot \nabla \psi = 0$$

(2.8)

$$\psi(\mathbf{X}, t=0) = \pm \text{ distance } \Gamma(t=0)$$
E. Advantages to the Eulerian Formulation

There are three major advantages to this Eulerian Hamilton-Jacobi formulation. First, the evolving function \( \psi(x,t) \) always remains a function for reasonable \( F \). However, the level surface \( \psi = 0 \), and hence the propagating hypersurface \( \Gamma(t) \) may change topology, break, merge, and form sharp corners as the function \( \psi \) evolves. As an example, consider two circles in \( \mathbb{R}^2 \) expanding outward. The initial function \( \psi(x,t=0) \) is a double-humped function which is Lipschitz continuous, but not everywhere differentiable. As this function evolves according to Eqns. (2.7-8) the topology of the level set \( \psi = 0 \) corresponding to the propagating hypersurface \( \Gamma(t) \) can change. For example, as the two circles expand, they meet and merge into a single closed curve with two corners. This is reflected in the change of topology of the level set \( \psi = 0 \) in the propagating function.

The second major advantage of this Eulerian formulation concerns numerical approximation. Because \( \psi(x,t) \) remains a function as it evolves, we may use a discrete grid in the domain of \( x \) and substitute finite difference approximations for the spatial and temporal derivatives.

Finally, the Eulerian Hamilton-Jacobi formulation extends in an obvious way to moving surfaces in three space dimensions. All of the numerical methodology described below is easily generalized, with none of the complicated bookkeeping that plagues marker particle technology and volume of fluid methods.

F. Extension of \( F \) off the level surface \( \psi=0 \)

As mentioned earlier, \( F \) may depend on such factors as the position of the front and the local curvature. We point out a somewhat subtle issue that results from our Eulerian formulation. We have formed an extension of \( F \) off the propagating hypersurface to all of space. That is, the equation

\[
\Psi_t + F |\nabla \Psi| = 0
\]

applies to each level set \( \Psi = C \), and thus we have implicitly assumed that \( F \) is a function in \( \mathbb{R}^N \times [0, \infty): F(x,t) \) such that
\[ F(\xi, t) = F(\Gamma(t)) \quad \text{for} \quad (\xi, t) \in \Gamma(t) \]

How does one extend \( F \) off the propagating hypersurface \( \Gamma(t) \) to the entire domain? In previous work, (see [46,59]), the function \( F \) depended on the local curvature of the propagating level set \( \psi = 0 \). In this case, since the local curvature could be calculated for the entire family of level sets covering the domain, it is straightforward to extend \( F \) by using the value of the curvature at a point \( \xi \) in the domain determined by the particular level set passing through that point.

In the Rayleigh-Taylor and Kelvin-Helmholtz problems considered here, the level set \( \psi = 0 \) is carried by the underlying fluid advection, and thus the speed function \( F \) depends only on the position of the level set \( \psi = 0 \). Thus, we may quite naturally extend the speed \( F \) to the entire domain by moving each level set by the underlying fluid.

In more complicated cases, the speed function can depend on such factors as the local normal, boundary integrals along the level set \( \psi = 0 \) and other factors. In such cases, the extension of \( F \) off the propagating hypersurface to the entire domain is not straightforward. The most complicated interface motion studied to date using this Hamilton-Jacobi approach is dendritic solidification, see [60]. In that work, the motion of the front and extension of \( F \) requires the global evaluation of a time history-dependent boundary integral along the boundary. For details, see [60].
III. Compressible Flow and Propagating Interfaces:

In this section, we discuss how to couple the level set formulation for a propagating interface to a system of conservation laws. To begin, consider the system of equations which describe compressible flow, namely

$$\bar{\mathbf{q}}_t + [\mathbf{F}(\bar{\mathbf{q}})]_x + [\mathbf{G}(\bar{\mathbf{q}})]_y = \mathbf{H}(\bar{\mathbf{q}})$$  \hspace{1cm} (3.1)

where the vector $\bar{\mathbf{q}}$ is defined by

$$\bar{\mathbf{q}} = \begin{pmatrix} \rho \\ \rho u \\ \rho v \\ \rho \varepsilon \end{pmatrix}$$  \hspace{1cm} (3.2)

Here, $\rho = \rho(x, y, t)$ is the density, $u = u(x, y, t)$ is the velocity in the $x$ direction, $v = v(x, y, t)$ is the velocity in the $y$ direction, and $\varepsilon = \varepsilon(x, y, t)$ is the internal energy of the system. The flux functions $\mathbf{F}(\bar{\mathbf{q}})$ and $\mathbf{G}(\bar{\mathbf{q}})$ are given by

$$\mathbf{F}(\bar{\mathbf{q}}) = \begin{pmatrix} \rho u \\ \rho u^2 + P \\ \rho uv \\ \rho \varepsilon + uP \end{pmatrix} \quad \mathbf{G}(\bar{\mathbf{q}}) = \begin{pmatrix} \rho v \\ \rho uv \\ \rho v^2 + P \\ \rho \varepsilon + vP \end{pmatrix}$$  \hspace{1cm} (3.3)

The forcing function $\mathbf{H}(\bar{\mathbf{q}})$ depends on the particular problem under study. For the Rayleigh-Taylor problem, we assume that gravity $g$ is pointing up (the positive $y$ direction), and thus have

$$\mathbf{H}(\bar{\mathbf{q}}) = \begin{pmatrix} 0 \\ 0 \\ \rho g \\ \rho v g \end{pmatrix}$$  \hspace{1cm} (3.4)

For the Kelvin-Helmholtz problem, we assume that

$$\mathbf{H}(\bar{\mathbf{q}}) = \begin{pmatrix} 0 \\ 0 \\ 0 \\ 0 \end{pmatrix}$$  \hspace{1cm} (3.5)
Finally, we use the equation of state to link the pressure $P$ and the density, namely

$$P = (\gamma - \rho)(e - 1/2(u^2 + v^2))$$  \(3.6\)

where here we have used the typical $\gamma$-gas law.

Our goal now is to incorporate interface motion in this setting. Let $\Omega_1$ and $\Omega_2$ be two regions in $R^2$ separated by a curve $\Gamma(t=0)$ which is a small perturbation of a horizontally straight line. Suppose $\Omega_1$ is above $\Omega_2$, and that the density in $\Omega_1$ is less than that in $\Omega_2$. The system of conservation laws described above apply in both $\Omega_1$ and $\Omega_2$, with possibly different $\gamma$-law equations of state. Suppose the location of the propagating interface $\Gamma(t)$ is given by the level set $\psi(x, y, t) = 0$. Then the full motion of the two regions can be viewed as a single system of conservation laws, which may be solved by appropriate finite difference approximations. What remains is to couple the equations of motion for $\psi$ to the system given in Eqns. (3.1).

A. Non-Conservative Differencing for $\psi$

For Eqn. (2.4), we have

$$\psi_t + u\psi_x + v\psi_y = 0$$ \(3.7\)

where $u = u_1, \ v = u_2$, and $\psi$ is the evolving function $\psi(x, y, t)$ such that

$$\Gamma(t) = \{(x, y) | \psi(x, y, t) = 0\}$$ \(3.8\)

Then one approach is to solve Eqn. (3.7), which is in non-conservative form, using the velocities $(u, v)$ obtained from the hyperbolic system given in Eqn. (3.1).

B. Conservative Differencing for $\psi$

Alternatively, we may put the equation of motion for the evolving function $\psi$ in conservation form. We have

$$(\rho\psi)_t + (\rho u\psi)_x + (\rho v\psi)_y = [\rho_t + (\rho u)_x + (\rho v)_y]\psi + \rho [\psi_t + u\psi_x + v\psi_y] = 0 + 0 = 0$$ \(3.9\)

For continuous $\psi$, the Rankine-Hugoniot jump conditions for this equation are the same as for the
conservation of mass equation (Eqn. 3.1). Thus, we may write a single system of conservation laws for the motion of the fluid in each region and the level set function \( \psi \), namely

\[
q_x + [F(q)]_x + [G(q)]_y = H(q)
\]

(3.10)

where

\[
q = \begin{pmatrix}
\rho \\
\rho u \\
\rho v \\
\rho \psi
\end{pmatrix}
\]

(3.11)

\[
F(q) = \begin{pmatrix}
\rho u \\
\rho u^2 + P \\
\rho u v \\
\rho u \psi + \mu P \\
\rho u \psi
\end{pmatrix}
\]

\[
G(q) = \begin{pmatrix}
\rho v \\
\rho v u \\
\rho v^2 + P \\
\rho v \psi + \nu P \\
\rho v \psi
\end{pmatrix}
\]

(3.12)

\[
H(q) = \begin{pmatrix}
0 \\
0 \\
\rho g \\
\rho v g \\
0
\end{pmatrix}
\text{Rayleigh–Taylor}
\]

\[
H(q) = \begin{pmatrix}
0 \\
0 \\
0 \\
0
\end{pmatrix}
\text{Kelvin–Helmholtz}
\]

(3.13)

All that remains is to formulate the equation of state. We define the pressure \( P \) by

\[
P = (\gamma(\psi) - 1) \rho(\epsilon - 1/2(u^2 + v^2))
\]

(3.14)

where

\[
\gamma(\psi) = \begin{cases}
\gamma_1 & \psi > 0 \\
\gamma_2 & \psi < 0 \\
? & \psi = 0
\end{cases}
\]

(3.15)

Away from \( \psi = 0 \), this is a conventional hyperbolic system of conservation laws with the standard propagation velocities of gas dynamics, and a triple linear degeneracy corresponding to the particle velocity. At \( \psi = 0 \), the fluxes are discontinuous, and it is not obvious what the correct conditions should be: this is reflected in the question mark "?" in Eqn. (3.15). In Section 5 we derive the appropriate condition at \( \psi = 0 \) and its numerical implementation.
We point out here that the extra work in computing the front is rather small. To solve the fundamental system of equations involves the use of a good numerical approximation to the system of conservation laws in two space dimensions. Computing the interface motion via the level set function requires either adding one more unknown to the system, namely \((\rho \psi)\), in a way that preserves the hyperbolic structure, or solving a separate equation in non-conservative form. In either case, the same finite difference grid lattice is used, and requires only one more array in the above data structures.
IV. Solving Hyperbolic Systems

A. General Outline

In this section, we lay the groundwork for our numerical methods. The field of hyperbolic solvers has grown rapidly in the past ten years, and good overviews of the material may be found in the review articles [48,56] and the references therein. Here, we give a brief flavor of the basic idea for those unfamiliar with the field.

The basic idea behind these methods is as follows. Consider, as a simple example, the \( n \) component linear hyperbolic system in one space variable, namely

\[
 u_t + [a(u)]_x = 0
\]  
(4.1)

Performing the differentiation, we then have

\[
 u_t + \left[ \frac{\partial a(u)}{\partial u} \right] u_x = u_t + A \overline{u}_x = 0
\]  
(4.2)

where \( A = \left[ \frac{\partial a}{\partial u} \right] \) is the (constant) \( mxm \) Jacobian matrix. Suppose \( T \) diagonalizes \( A \). Then

\[
 T A T^{-1} = \Lambda
\]  
(4.3)

where \( \Lambda \) is diagonal. Then if we define the vector

\[
 \overline{u} = Tu ,
\]  
(4.4)

we can premultiply Eqn. (4.2) by \( T \) and postmultiply by \( T^{-1} \) to obtain the decoupled diagonal system

\[
 \overline{u}_t + A \overline{u}_x = 0
\]  
(4.5)

Consider now the \( i^{th} \) component of the above diagonal system, namely

\[
 (\overline{u}_i)_t + A_k (\overline{u}_k)_x = 0
\]  
(4.6)
We may solve this equation exactly, since $A_u$ is a constant, and retrieve the solution $u$ by letting

$$u = T^{-1} \mathbf{u} \quad (4.6)$$

The strategy behind numerical algorithms for more general non-linear hyperbolic conservation laws is a time and space discretization of a non-linear version of the above process. Consider a lattice of points $x_i = ih, i = -3,-2,-1,0,1,2,3,...$ and the general system of the form

$$u_t + \left[ \frac{\partial A(u)}{\partial u} \right] u_x = u_t + A u_x = 0 \quad (4.7)$$

where now $A(u)$ is nonlinear. Let $u^n_i$ denote the approximate solution at time $n \Delta t$ at point $x_i$. In order to go from the solution at time $n \Delta t$ to the solution at time $(n+1) \Delta t$, at each point $x_i$ we compute the eigenvectors of the Jacobian matrix $A(u)$ to construct the diagonalizing matrices $T$ and $T^{-1}$.

The matrices $A$ and $T, T^{-1}$ are functions of $u$. Their values at an intermediate state between $u_i$ and $u_{i+1}$, denoted as $u_{i+1/2}$ at $x_{i+1/2}$, are approximated in Section 5 and used in the numerical procedure to update $u$ as follows. At each point $x_{i+1/2}$, we have a \textit{local Riemann problem}, which assumes a constant left initial state and a constant right initial state. Imagine then, that at each point $x_{i+1/2}$ at time $n \Delta t$, we consider the local Riemann problem which has initial state $u^n_{i+1/2}$ on the right and $u^n_{i-1/2}$ on the left (we postpone until later the calculation of these intermediate mesh values). Using approximate Riemann solvers, we solve this initial value problem for time step $\Delta t$, where $\Delta t$ is chosen small enough that waves traveling from neighboring Riemann problems do not interact. The matrices $A(u)$ and $T(u)$ play a key role in the approximate solutions to the Riemann problem. Details of these ideas may be found in [48,56].

\section*{B. The Equations of Motion for Gas Flow and Interface Motion}
Following the above outline, the first task is to compute the eigenvalues and eigenvectors of $A$, which is the Jacobian matrix of $F(q)$. There is a similarity between our set of five conservation laws (Eqns. 3.10-11) and the equations of two component inviscid gas flow studied by [35]. In those equations, the level set function $\psi$ is replaced by the mass fraction $Y$ of species one. In addition, the quantity $\gamma$ defined in Eqn. (3.15) is no longer a piecewise constant function of $\psi$, but instead for the two component mixture case is given by (see [35])

$$\gamma = \frac{Y c_n \gamma_1 + (1-Y)c_n \gamma_2}{Y c_n + (1-Y)c_n}$$  \hspace{1cm} (4.8)

where $c_n$ is the specific heat at constant volume of species $i$.

For simplicity of exposition we compute the one-space dimensional Jacobian, where we set $v=0$, and neglect the $\rho v$ equation in [Eqn. 3.10-11]. We note that $\gamma$ is a function of $Y$ for our problem and a function of $\psi$ for the two component problem. Using conserved variables, we may view

$$\gamma = \gamma \left( \frac{\rho Y}{\rho} \right) \quad \text{(two component gases)}$$  \hspace{1cm} (4.9)

$$\gamma = \gamma \left( \frac{\rho \psi}{\rho} \right) \quad \text{(level set, immiscible problem)}$$  \hspace{1cm} (4.10)

We let $\phi$ denote either $Y$ or $\psi$ for the two problems and obtain the Jacobian as in [35]

$$\frac{\partial F}{\partial q} = \begin{pmatrix}
0 & 1 & 0 & 0 \\
\frac{Y^2-3}{2} \mu^2 - \phi X & (3-\gamma)\mu & (\gamma-1) & X \\
-\frac{Y^2-1}{2} \mu^2 - \phi H - \phi \mu & \mu & \mu X \\
-\phi & \phi & 0 & \mu
\end{pmatrix}$$  \hspace{1cm} (4.12)

Here, the enthalpy $H$ is defined by

$$H = \frac{\rho e + p}{\rho}$$  \hspace{1cm} (4.13)

and
\[ X = \frac{p}{\gamma - 1} \frac{\gamma}{\rho}. \quad (4.14) \]

The eigenvalues of \( A \) are

\[ \lambda_1 = u - c, \quad \lambda_2 = u = \lambda_3, \quad \lambda_4 = u + c \quad (4.15) \]

where

\[ c = \sqrt{\frac{IP}{\rho}}. \quad (4.16) \]

A set of right eigenvectors is

\[ \begin{align*}
   r_1 &= \begin{pmatrix}
   1 \\
   u-c \\
   H-uc \\
   \phi
   \end{pmatrix}, \\
   r_2 &= \begin{pmatrix}
   1 \\
   u \\
   \frac{u^2}{2} \\
   \phi
   \end{pmatrix}, \\
   r_3 &= \begin{pmatrix}
   0 \\
   0 \\
   \frac{X}{\gamma-1} \\
   1
   \end{pmatrix}, \\
   r_4 &= \begin{pmatrix}
   1 \\
   u+c \\
   H+uc \\
   \phi
   \end{pmatrix}
\end{align*} \quad (4.17) \]

For our definition of \( \gamma(\psi) \), \( \phi X = 0 \) and

\[ X = \frac{p}{(\gamma-1)p} \delta(\psi-0)[\gamma_1 - \gamma]. \quad (4.18) \]

Obviously \( \delta(\psi-0) \) must be approximated numerically. We shall describe this in the next section.

C. Approximate Riemann Solvers

We must now solve the Riemann problem that occurs in the decoupled diagonalized system. We use second order TVD schemes, which can be based on either the true solution to the Riemann problem (Godunov's scheme), or, more likely, an approximate Riemann solver, e.g. Roe's [56]; Osher's [48] or van Leer's [65].
For our problem, the simplest scheme is van Leer’s, since it is based on a flux splitting

\[ f_{VL}(q_L, q_R) = f^+(q_L) + f^-(q_R) \]  

(4.19)

The eigenvalues of \( \frac{\partial f^+}{\partial q} \) (\( \frac{\partial f^-}{\partial q} \)) are all nonnegative (nonpositive). Each typically has one zero and two non-zero eigenvalues. However, there is no “switching” across the point \( \mu=0 \), thus the scheme is relatively viscous near stagnation points. This will be important in the solution of the Rayleigh-Taylor and Kelvin-Helmholtz problems.

For Osher’s scheme

\[ \hat{f}_O(q_L, q_R) = \frac{1}{2} [f(q_L) + f(q_R)] - \frac{1}{2} \int_{q_L}^{q_R} \frac{\partial f}{\partial q} (q) \, dq \]  

(4.20)

where the integral is taken along successive paths parallel to the right eigenvectors of \( \frac{\partial f}{\partial q} \).

The construction here is simplified by requiring that the Riemann invariants be constant along each path. This leads to a single equation for a single unknown for an intersection point. It can be shown that Newton’s method globally converges for this, (details elsewhere).

Roe’s scheme can be written as

\[ \hat{f}_R(q_L, q_R) = \frac{1}{2} [f(q_L) + f(q_R)] - \frac{1}{2} |A_{LR}|(q_R - q_L) \]  

(4.21)

where \( A_{LR} = A(q_L, q_R) \) is a matrix satisfying

\[ f(q_L) - f(q_R) = A_{LR}(q_L - q_R) \]  

(4.22)

It turns out that for \( \gamma \) law gas dynamics, \( A_{LR} \) can be chosen to be the Jacobian matrix evaluated at some intermediate state \( q_{LR} \), known as the “Roe average of \( q_L, q_R \).” This cannot be done for the system here. However, in [35] a Roe matrix for two component flow was constructed which is very close to the Jacobian at the Roe average. The expression is
\[ \frac{\partial F}{\partial q} = \begin{pmatrix} 0 & 1 & 0 & 0 \\ \frac{(\gamma-3)}{2} \hat{u}^2 - \hat{\gamma} \hat{X} & (3-\gamma) \hat{u} & \gamma - 1 & \hat{X} \\ -\hat{u} \hat{H} + \frac{(\gamma-1)}{2} \hat{u} \hat{Y} & \hat{H} - (\gamma-1) \hat{u}^2 & \gamma \hat{u} & \hat{u} \hat{X} \\ -\hat{u} \hat{Y} & \hat{Y} & 0 & \hat{u} \end{pmatrix} \]

(4.23)

where the averaged state \( \hat{q} = (\hat{\beta}, \hat{u}, \hat{\gamma}, \hat{\gamma} \hat{Y})^T \) is defined by:

\[ \hat{\beta} = \frac{\rho_L \sqrt{\rho_L} + \rho_R \sqrt{\rho_R}}{\sqrt{\rho_L} + \sqrt{\rho_R}} \]

(4.24)

\[ \hat{u} = \frac{u_L \sqrt{\rho_L} + u_R \sqrt{\rho_R}}{\sqrt{\rho_L} + \sqrt{\rho_R}} \]

(4.25)

\[ \hat{H} = \frac{H_L \sqrt{\rho_L} + H_R \sqrt{\rho_R}}{\sqrt{\rho_L} + \sqrt{\rho_R}} \]

(4.26)

\[ \hat{Y} = \frac{Y_L \sqrt{\rho_L} + Y_R \sqrt{\rho_R}}{\sqrt{\rho_L} + \sqrt{\rho_R}} \]

(4.27)

where \( \hat{\gamma} = \gamma(\hat{q}) \) and

\[ \hat{X} = \frac{C_1 C_2 (\gamma_1 - \gamma_2) \hat{Y}}{\hat{Y} C_1 + (1 - \hat{Y}) C_2} \]

(4.28)

with

\[ \hat{Y} = \frac{T_L \sqrt{\rho_L} + T_R \sqrt{\rho_R}}{\sqrt{\rho_L} + \sqrt{\rho_R}} \]

(4.29)

The matrix \( \hat{A} \) defined by the above is then diagonalizable: its eigenvalues are \( \hat{\gamma} - \hat{\gamma}, \hat{u}, \hat{u} - \hat{\gamma}, \)

where \( \hat{\gamma}^2 = (\gamma-1)(\hat{H} - \hat{u}^2/2), \) and its eigenvectors are given by expressions which are analogous to Eqn. (4.17)

This expression has an analogue in our immiscible case. We describe this and our numerical method in the next section.
V. Approximation to Equations of Motion

The system is discretized in space by a second order TVD (or ENO) scheme and in time by a two-stage TVD Runge-Kutta scheme which is second order accurate in time. We follow the approach described in [62] and stop at the second order accurate level.

Briefly, we set up a semi-discrete method of lines approximation to the system written as

\[ q_t = L(q) \]

(5.1)

The TVD operator \( L(q) \) approximates \( L(q) \) to 2nd order

\[ L(q) = L(q) + O(h^2) \]

(5.2)

for smooth \( q \), where \( h \) is the maximum mesh size. The Euler forward version

\[ q^{n+1} = q^n + \Delta t \cdot L(q^n) \]

(5.3)

is assumed to be total variation stable for

\[ \Delta t \leq \frac{2}{3} \left( \max \left( |u|/\Delta x + |v|/\Delta y + c\sqrt{1/\Delta x^2 + 1/\Delta y^2} \right) \right)^{-1} \]

(5.4)

A second order TVD Runge-Kutta time discretization is just Heun's method:

\[ q^* = q^n + \Delta t \cdot L[q^n] \]

(5.5)

\[ q^{n+1} = \frac{1}{2} (q^n + q^*) + \frac{\Delta t}{2} L[q^*] \]

which is stable under the same CFL condition as the Euler forward version.

Next we describe the space discretization

\[ L(q) = L^x[q] + L^y[q] + H(q) \]

(5.6)

Here, of course \( L^x \) approximates \(-F_x\), \( L_y \) approximates \(-G_y \), and \( H(q) \) is the exact value of \( H(q) \) at the grid point.
The most intricate part of the discretization involves \( L^x \) and \( L^y \). We describe \( L^x \) here; \( L^y \) is defined analogously.

\( L^x \) will be a conservation form finite difference scheme

\[
L^x = -\frac{1}{\Delta x} \left( \hat{f}_{j+1/2} - \hat{f}_{j-1/2} \right) 
\] (5.7)

where the numerical flux, \( \hat{f}_{j+1/2} \) is a second order accurate approximation to \( F(q) \) at the end point of a cell

\[
I_j = \left\{ x \mid x_{j-1/2} \leq x \leq x_{j+1/2} \right\} 
\] (5.8)

and \( q(x_j, t^n) \) is obtained at all time levels, for \( x_j = \frac{1}{2}(x_{j+1/2} + x_{j-1/2}) \), the cell center.

First we determine the Roe decomposition. The average Jacobian \( A_{j+1/2} \) for our system is analogous to the two component flow Roe matrix in [35]:

\[
A_{LR} = \begin{pmatrix}
0 & 1 & 0 & 0 & 0 \\
\frac{1}{2}(\dot{\gamma}-1)(\dot{u}^2+\dot{v}^2) - \dot{u}^2 - \dot{\psi} \ddot{X} & (\dot{\gamma}-1)\dot{u} & -\dot{\gamma} & \ddot{X} \\
-\dot{\gamma} & \ddot{\nu} & \ddot{\nu} & 0 & 0 \\
-\dot{\psi} & \ddot{\psi} & 0 & 0 & \ddot{\psi} \\
\end{pmatrix}
\] (5.9)

The averaged states are defined by

\[
\rho = \sqrt{\rho_L} \sqrt{\rho_R} 
\] (5.10)

\[
\ddot{u} = \frac{\sqrt{\rho_L u_L} \sqrt{\rho_R u_R}}{\sqrt{\rho_L} + \sqrt{\rho_R}} 
\] (5.11)

with \( \ddot{\nu}, \ddot{\psi} \), and \( \ddot{\psi} \) defined in the same way as \( \ddot{u} \). The condition that remains is:

\[
\ddot{\dot{X}} = \frac{(\rho_R - \rho_L)(\dot{\gamma}-1)[\rho_L(\rho - 1) - \rho_R(\rho - 1)]}{\rho(\rho_R - \rho_L)} 
\] (5.12)

Note that \( \dot{\rho} = \dot{\rho} \ddot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\dot{\do
by $a_0=\Delta p/(\rho \Delta \psi)$, and $a_1$ by $a_1=\Delta (\rho/(\gamma-1))/\rho \Delta \psi)$. Then Eqn. 5.11 becomes

$$\frac{d(\hat{\gamma}-1)}{d\hat{\psi}} = a_0(\hat{\gamma}-1) - a_1(\hat{\gamma}-1)^2$$

It is not possible to find a function $\gamma(\psi)$ satisfying Eqn. (5.13) with boundary conditions $\gamma(\psi_L) = \psi_L$. Therefore we choose

$$\gamma(\psi) = \frac{(\psi-\psi_L)\gamma_L + (\psi_A-\psi)\gamma_L}{\psi_A - \psi_L} \quad (5.14)$$

and let $\hat{\gamma}=\gamma(\hat{\psi})$. Now the condition given in Eqn. (4.22) is not satisfied consistently, but can be used to compute $\hat{\gamma}(\psi)$ or $\hat{X}$. Thus, we have obtained a matrix $\hat{A}$ which is almost equal to $A(\hat{\psi})$, except for $\hat{X}$.

$$\hat{A}_{LR} = \begin{pmatrix}
1 & 1 & 0 & 0 & 1 \\
\hat{\psi} - \hat{\psi}_e & \hat{\psi} & 0 & 0 & \hat{\psi}_e + \hat{\psi}_e \\
\hat{\psi} & \hat{\psi} & 1 & 0 & \hat{\psi}_e \\
\hat{H} - \hat{H}_e & \frac{1}{2}(\hat{\psi}_e^2 + \hat{\psi}^2) & \hat{\psi} & -\hat{X} + (\hat{\psi}-1) \hat{H} \hat{\psi}_e & \hat{H} + \hat{H}_e \\
\hat{\psi} & \hat{\psi} & 0 & 1 & \hat{\psi} \\
\end{pmatrix} \quad (5.15)$$

whereas the left eigenvectors $\mathbf{p}^I$ can be taken as the rows of $\mathbf{T}^{-1}$. The left eigenvectors obey the following relations:

$$\mathbf{p}^2 = \begin{pmatrix} 1 - 1/2(\hat{\psi} - 1)(\hat{\psi}_e^2 + \hat{\psi}^2)/2 + \psi \hat{X}/\hat{\psi}_e^2, (\hat{\psi} - 1) \hat{\psi}_e^2, (\hat{\psi} - 1) \hat{\psi}/\hat{\psi}_e^2, -\hat{X}/\hat{\psi}_e^2 \end{pmatrix} \quad (5.16)$$

$$\mathbf{p}^3 = (-\hat{\psi}, 0, 1, 0, 0)$$

$$\mathbf{p}^4 = (-\hat{\psi}, 0, 0, 0, 1)$$

$$\mathbf{p}^1 + \mathbf{p}^2 = (1, 0, 0, 0, 0) - \mathbf{p}^2$$

$$\mathbf{p}^1 - \mathbf{p}^2 = (\hat{\psi}/\hat{\psi}_e, 1/\hat{\psi}_e, 0, 0, 0)$$

The eigenvalues are the diagonal elements of

$$A = \text{diag}(\hat{\psi} - \hat{\psi}_e, \hat{\psi}, \hat{\psi}, \hat{\psi}_e, \hat{\psi}_e + \hat{\psi}_e) \quad (5.17)$$
The Roe-type matrix above has terms which are almost infinite (that is, behave like $\frac{1}{\Delta x}$), because of Eqn. (5.13). This mirrors the delta function occurring in the Jacobian when $\psi$ changes sign. In spite of this, no stability problems were found in our calculations below.

Algorithm (TVD-Roe).

Given the states $q_j = q(x_j)$, where the $x_j$ are grid points, we compute the fluxes $f_j = f(q_j)$. To determine the numerical flux $\hat{f}_{j+1/2}$, we transform to characteristic (Riemann invariant-like) variables. We denote the left eigenvectors (computed in Eqn. (5.15)), the right eigenvectors, and the eigenvalues of $A_{j+1/2} = A_{j+1}$ (see Eqn. 5.15), (left state $q_j = q_j$, right state $q_{j+1}$) by $l_{\mu j+1/2}^{(\nu)}$, $r_{\nu j+1/2}^{(\mu)}$, $\lambda_{\nu j+1/2}^{(\mu)}$, $\nu = 1, 2, 3, 4, 5$. Computation shows that

$$\lambda_{\mu j+1/2}^{(1)} = u_{j+1/2} - c_{j+1/2} = u_{j+1/2} - \sqrt{\frac{\gamma_{j+1/2} P_{j+1/2}}{\rho_{j+1/2}}}$$  \quad (5.18)

$$\lambda_{\mu j+1/2}^{(2)} = \lambda_{\mu j+1/2}^{(3)} = \lambda_{\mu j+1/2}^{(4)} = u_{j+1/2}$$ \quad (5.19)

$$\lambda_{\mu j+1/2}^{(5)} = u_{j+1/2} + c_{j+1/2}$$ \quad (5.20)

Also,

$$l_{\mu j+1/2}^{(\nu)} \cdot r_{\nu j+1/2}^{(\mu)} = \delta_{\mu \nu} = \begin{cases} 1 & \text{if } \nu = \mu \\ 0 & \text{if } \nu \neq \mu \end{cases}$$ \quad (5.21)

We may decompose

$$f_k = \sum_{j=1}^{N} r_{\nu j+1/2}^{(\mu)} \cdot G_k^{(\nu)}, \quad k = j-1, j, j+2$$ \quad (5.22)

where

$$G_k^{(\nu)} = l_{\mu j+1/2}^{(\nu)} \cdot f_k$$ \quad (5.23)

The next step is just second order accurate ENO integration on $G_k^{(\nu)}$, namely
\[ G_L^{(v)} = G_j^{(v)} + 1/2 \Delta_j^{(v)} \]  
(5.24)

\[ G_H^{(v)} = G_j^{(v)} - 1/2 \Delta_j^{(v)} \]  
(5.25)

If

\[ a_j^{(v)} = G_j^{(v)} - G_{j+1}^{(v)} \quad b_j^{(v)} = G_{j+1}^{(v)} - G_j^{(v)} \]  
(5.26)

then

\[ \Delta_j^{(v)} = \begin{cases} 
  a_j^{(v)} & \text{if } |a_j^{(v)}| \leq |b_j^{(v)}| \\
  b_j^{(v)} & \text{otherwise}
\end{cases} \]  
(5.27)

Upwind differencing is now applied to the field

\[ G_{j+1/2}^{(v)} = \begin{cases} 
  G_L^{(v)} & \text{if } \lambda_{j+1/2}^{(v)} \geq 0 \\
  G_H^{(v)} & \text{otherwise}
\end{cases} \]  
(5.28)

Finally, we transform back by letting

\[ f_{j+1/2} = \sum_{\nu=1}^{5} r_{j+1/2}^{(\nu)} \cdot G_{j+1/2}^{(\nu)} \]  
(5.29)

There is a (slightly nonstandard) version of a second order accurate TVD Roe-based scheme, see [62]. As such, it is known to admit stationary expansion shocks. An entropy fix due to Harten [24] is obtained through

\[ G_{j+1/2} = 1/2 \left[ G_L^{(v)} + G_H^{(v)} - g(\lambda_{j+1/2}) \left( G_H^{(v)} - G_L^{(v)} \right) \right] \]  
(5.30)

where
\[ g(x) = \begin{cases} 
\text{sign}(x) & \text{if } |x| \geq \epsilon_f \\
\frac{x^2 + \epsilon_f^2}{2 \epsilon_f} & \text{if } \epsilon_f^2 < |x| < \epsilon_f \\
\frac{\epsilon_f^4 + \epsilon_f^2}{2 \epsilon_f^2} & \text{if } |x| < \epsilon_f^2 
\end{cases} \quad (5.31) \]

We choose \( \epsilon_f = 0.1 c_{j+1/2} \). This entropy fix suppresses unphysical expansion shocks, and is only applied for the characteristic field \( v=1 \) if \( (u-c)_j < 0 < (u-c)_{j+1} \), and for field \( v=5 \) if \( (u+c)_j < 0 < (u+c)_{j+1} \).

Finally, we consider the case for computations where \( \psi \) is not in conservative form. The term \( uv_x \) is approximated via

\[ (\Delta x)uv_x = u_j^+ \left( [\psi_j + 1/2(\Delta \psi)_j] - [\psi_{j-1} + 1/2(\Delta \psi)_{j-1}] \right) + \quad (5.32a) \]

\[ u_j^- \left( [\psi_{j+1} - 1/2(\Delta \psi)_{j+1}] - [\psi_j - 1/2(\Delta \psi)_j] \right) \]

where

\[ u_j^+ = \max(u_j, 0) \quad u_j^- = \min(u_j, 0) \quad (5.32b) \]

and

\[ (\Delta \psi)_j = \text{smaller in absolute value of } (\psi_{j+1} - \psi_j, \psi_j - \psi_{j-1}) \quad (5.32c) \]

The other term \( vw_y \) is approximated analogously.

The spatial discretization of the remaining four equations is as described above, and the time discretization is just Heun's method, once again.
VI. Results

A. Rayleigh-Taylor Instability

The numerical experiments were performed on a rectangular domain with walls on the lower and upper side, and periodic boundaries in the horizontal direction. Gravity acts in the upward direction. The horizontal size of the domain is chosen as unit length. An initial sine perturbation has a wavelength $\lambda$ of the same size. As initial conditions, we use the solution of the linearized equations given in [21]. This solution refers to the air-air case. The relevant parameters are the initial density ratio $D=\rho_a/\rho_s$ and $M^2=g\lambda^2/c_s^2$. The subscript $a$ refers to the gas just above the interface, the subscript $b$ to gas just below the interface. The sound speed just below the interface $c_b$ is set to 1, as is the density ($c_a=1$, $\rho_a=1$). The constant of gravity follows from $M^2$. The adiabatic exponent $\gamma_a=\gamma_b=1.40$.

Figure 1 shows contours of $\psi$ at values of $-1/32$, 0, and 1/32, for times 0, 1, ..., 6. The grids have size $64\times128$ and $192\times384$, respectively. Symmetry is forced: the computations have been carried out on a grid with half the size in the horizontal direction. The initial density ratio $D$ is 2, the amplitude of the initial perturbation is 0.015, and $M^2=0.5$.

In Figure 1a, we see that a small sinusoidal perturbation grows into the expected mushroom-shaped object and develops side rolls. However, tripling the mesh size, shown in Figure 1b, does not produce a refined picture. Instead, pronounced oscillations develop, and smaller rolls appear on the surface of the basic structure. This suggests that the solution does not converge under refinement. As a test, we compute the relative error in $\psi$ defined by

$$E^2_h(\psi) = \| \psi^{2h} - I_h^{2h} \psi_h \|$$

(6.1)

where the superscript $h$ denotes the cell size of the uniform grid, and $2h$ the grid size after coarsening. In order to compare the two, we apply the restriction operator $I_h^{2h}$ to the solution of the fine mesh and volume average to produce values for comparison with the coarse mesh size. The initial
data obey \( E^2_\psi = O(h^2) \), implying second-order accuracy. For short time, the error decreases as \( h \) is refined. However, for larger times \( T > 4 \), the error increases as the mesh is refined. In Table 1, we show these results for the inviscid \((\mu=0)\) case.

This indicates that the problem is physically unstable. To obtain a stable solution, we add physical viscosity. The Navier-Stokes equations without heat conduction are used. Following Stokes hypothesis, the second coefficient of viscosity \( \lambda = \frac{2}{3} \mu \). The first coefficient of viscosity \( \mu \) is chosen to be constant. The spatial discretization is based on the usual central differences. The timestep is chosen as

\[
\Delta t = \left( \frac{\lambda_1}{CFL_1} + \frac{\lambda_2}{CFL_2} \right)^{-1} \tag{6.2}
\]

where

\[
\lambda_1 = \max (|u| + |v| + c \sqrt{2})/h, \quad \lambda_2 = \frac{14}{3} \frac{\mu}{h^2} \frac{1}{\min(p)} \tag{6.3}
\]

Here the maximum and minimum are computed over the grid. We use \( CFL_1 = 0.3 \) and \( CFL_2 = 1 \).

The addition of physical viscosity stabilizes the problem. We performed runs with grid size of 32, 48, 64, 96, 128 and 192 mesh points across the horizontal width, with twice as many points in the vertical direction. The results indicate that convergence improves with larger values of the viscosity \( \mu \). Table 1a shows the relative errors \( E^2_\psi \) in \( \psi \), computed from a grid refinement sequence with \( h^{-1} \) equal to 32, 48, 64, 96, 128, and 192. The number of points in the vertical direction is twice that amount.

Table 1b shows the order of accuracy \( p \), estimated by a weighted least-squares fit to \( \log E^2_\psi(\psi) = b_0 + p \log h \), using \( h^{-2} \) as weight. The grids used have 64, 96, 128, 192, 256, and 384 points in the vertical direction, and half that number in the horizontal direction (actually 1/4, with the forced symmetry). This provides 4 data points for each least-squares fit. It is clear that without viscosity, the error increases under grid refinement. In the viscous case, convergence improves with
larger values of the viscosity $\mu$.

Figure 2a shows $\psi$ at time 6, for various choices of $\mu$. Figure 2b shows the density $\rho$ for the same parameters. Since there is no feedback mechanism from the front to the fluid, the density of the fluid is a good indicator of the front position. Comparison of the two figures reveals the cosmetic character of $\psi$.

The above calculations consider a conservative differencing of $\psi$, initialized as the signed distance to the initial front, as given in Eqn. (2.2). We now consider alternatives to this approach. To begin, other researchers have tracked fronts by following the evolution of a "color" function, which is $-1$ on one side of the front and $+1$ on the other. A sophisticated variant of this idea using a version of SLIC to gain subcell resolution was employed in [13] to performed detailed calculations and comparison with experiment of a shock wave hitting a gas interface. We may incorporate a color function into our code by initializing $\psi$ to $\pm 1$. Figure 3a displays the result of a computation identical to the one in Fig. 1a, but using the color function instead of $\psi$. Comparison shows that the color function suggests a faster evolution of the instability than $\psi$. This is highlighted in Fig. 3b, which shows part of vertical cross-section through the middle of Fig. 3a.

Next, we consider non-conservative differencing, as discussed in Section III.A. Here, the standard four-component hyperbolic system is solved, and those velocities are then used in a second order accurate upwind fashion to advect $\psi$ using Eqn. (3.7), as described in the text. In Figure 3b, we compare the results of conservative and non-conservative differencing of both the initialized distance function for $\psi$ and the color function. Our results here seem to indicate that the non-conservative differencing of $\psi$ using the level set initial distance function is most desirable.

The motion of the front becomes significantly more complicated when we allow feedback between the front location and the fluid mechanics. Consider an air-helium boundary. Here, the bottom gas is air with $\gamma_1=1.40$ (Air), and the top gas is helium with $\gamma_2=1.63$ (He). As initial condition, we again use the linearized solution. Using the molecular weights $\mu_1=28.964$ and $\mu_2=4.00260$, we set $c_1=1$ and $\rho_1=1$, and find a density ratio by assuming constant temperature and pressure across
the interface. This implies that the density ratio $D = \rho_1 / \rho_2 = \mu_1 / \mu_2$ and that $c_s^2 = c_2^2(\gamma / \gamma_2)$. 

In Figure 4a, we model this problem using conservative differencing of the level set/gas dynamic five component hyperbolic system and the original distance function $\psi$ initialization. Calculations are performed on a 32x56 grid. The small initial bubble grows upwards into a long plume. We show contours of $\psi$ at -1/32, 0, 1/32. In Figure 4b, we perform the same calculation using non-conservative differencing for $\psi$. For comparison, in Figure 4c we show results using $\psi$ initialized as a piecewise constant color function, namely -1 on one side and +1 on the other side of the interface. Finally, the results of a different computation with the effective adiabatic exponent based on the concentration $Y$, as in [35], is shown in Fig. 4d. We have included $\psi$ in this computation as well, as a passive scalar. Contours of $\psi$ are presented in Fig. 4e.

Computations based on the concentration $Y$ model different physics. Still, the plot of the passive $\psi$ corresponds fairly closely to the one in Fig. 4a. A comparison between Figs. 4c and 4d shows that it is not so easy to determine the position of the (smeared) front from the concentration.

B. Kelvin-Helmholtz Instability

Next, we perform calculations of the Kelvin-Helmholtz instability. As initial conditions, we take constant pressure and temperature above and below the interface, with zero vertical velocity. The initial shape is a sine perturbation. Above the interface, the gas moves towards the left with velocity $u = -u_0$, below the interface, the horizontal velocity is $u = u_0$. For the air-air case, we set the density and the sound speed to 1 everywhere. Again we have periodic boundaries in the horizontal direction, and walls at the bottom and top. Gravity is not included.

In Figure 5a, we show the evolution of an initial perturbation with amplitude $a = 0.1$ and $a = 0.25$. We use a 128x256 grid. We study an air-air interaction, so that the level function $\psi$ is passively advected. We plot values of $\psi$ at -1/32, 0, 1/32. The results show the rollup of a vortex structure as it progresses through several turns. The small oscillations in the shape seem to indicate, once
again, that the problem is physically unstable. We check this by analyzing the computed solution at
time $t=6$ for various values of $\alpha$. In Figure 5b, we show the results of a calculation on a 32x32,
64x64, 96x96, and 128x128 grid. The refinement in mesh size at fixed time indicates that the results
are not stable.

Next, we add physical viscosity to the system. In Table 2a and 2b, we show the error $E_1^2(\psi)$
measured in the $l_1$ norm as a function of the grid size and viscosity $\mu$ at various times. The intro-
duction of physical viscosity stabilizes the problem. In Figure 5c, we show zero contours of $\psi$
at time $t=6$, with viscosity $\mu = 0, 10^{-4}, 10^{-3}, 5 \times 10^{-3}$ going from left to right. As expected, the intro-
duction of physical viscosity slows the rollup.

Figure 6a shows the evolution for the air-helium case. The gas below the interface is helium.
The initial conditions are: $c_a=1, \rho_a=1, D=\rho_b/\rho_a=\mu_b/\mu_a, \quad c_0^2=\gamma \rho_a/\gamma_b, \quad u=\pm u_0, \quad v=0$. This
corresponds to constant initial pressure and temperature. We let $u_0=0.5$. The initial sine perturbation
has an amplitude $a=0.1$. The roll-up can no longer be resolved after a time between 4 and 5.
Discussion

In this paper, we have discussed the coupling of the level set formulation of interface motion to the equations of compressible gas dynamics. We have considered two approaches. In one approach, the level set equation is posed in non-conservative form, and coupled to the four-component system. Alternatively, we have shown that a conservative version of the level set function \( \psi \) can be directly incorporated as a five-component system of hyperbolic conservation laws using standard shock technology. In both conservative and non-conservative settings, we have examined the distance function initialization of the level set function \( \psi \) and a degenerate initialization using the color function.

The efficiency of these various techniques depends on the particular problem under study. In the Rayleigh-Taylor problem we considered, the normal velocity varies continuously across the interface, unlike the density \( \rho \), which undergoes a jump. In this case, the non-conservative formulation for \( \psi \) uses a smooth \( \mathbf{u} \), and our results indicate that this approach is preferable to direct incorporation of \( \psi \) into the conservative system because of the discontinuity in \( \rho \psi \). It seems reasonable to expect that for problems in which \( \mathbf{u} \) jumps across the interface, the conservative approach will be preferable.

In the problems considered here, the front velocity does not depend on the geometry of the interface. All that is needed is a rough location of the front to determine the selected region for the gas constant. Thus, the ability of the Hamilton-Jacobi level set formulation to accurately calculate curvature and normal direction is untapped in this simple calculation. For such simple problems, the color function is an adequate initialization, and leads to only slightly worse performance; however, we point that it is no cheaper than our original level set approach. Furthermore, in more sophisticated problems, see, for example, [60], the color function idea is insufficient and the full capabilities of the level set approach are utilized.
Finally, we have computed the solution to two complex physical phenomena. To what degree are these solutions accurate? First, we point out that in the zero viscosity limit, both of the problems are physically unstable. Our calculations in this case do not converge with respect to mesh refinement. We believe the following is a plausible scenario. Our schemes introduce artificial viscosity which decreases with decreasing mesh size. For a coarse enough mesh, the numerical viscosity stabilizes instabilities that occur in the zero viscosity limit, and the solution is smooth. This can be seen in the calculations with coarse grids given in Figure 1c. As the mesh size is refined, and the artificial viscosity lessens, small physical instabilities are not suppressed and instead grow, as seen in the finer grid calculations of Figure 1c.

In order to justify this hypothesis, we should be able to demonstrate that, given some amount of physical viscosity, we can compute on a fine enough grid so that the physical viscosity dominates the numerical viscosity on the results are unchanged with respect to further grid refinement. This is the experiment indicated in Tables 1 and 2. On the basis of this, we believe that our technique is capturing a reasonable portrait of the solution in the viscous cases, and reflects the physical instability of the problem in the zero viscous limit case. Of course, the particular unstable solution shown in the case $\mu = 0$ means little; only the gross features are of significance.

In future work, we hope to use the notion of subcell resolution [25] together with the level set formulation to account more accurately for the small scale geometry of the front.
Appendix: Color Versus Smooth \( \psi \)

Consider the one-dimensional motion of a contact discontinuity. Let its speed be \( u_0 \). Then our system reduces to

\[
\frac{\partial w}{\partial t} + u_0 \frac{\partial w}{\partial x} = 0 \quad w = \begin{pmatrix} \rho \\ \rho \psi \end{pmatrix}
\]  

(A1)

A first-order discretization of this system introduces numerical viscosity, which can be modeled by the equivalent equation

\[
\frac{\partial w}{\partial t} + u_0 \frac{\partial w}{\partial x} = \epsilon \frac{\partial^2 w}{\partial x^2}
\]

(A2)

Transforming to moving coordinates \( x' = x - u_0 t, t' = t \) produces the heat equation

\[ w_t = \epsilon w_{xx} \]  

(A3)

where the primes have been dropped. The solution is

\[ w(x,t) = \int K(x,y)w(y,0)dy \]  

(A4)

where the kernel

\[ K(x,y) = \frac{1}{2} (\rho \epsilon t)^{-\frac{1}{2}} \exp \left( -\frac{(x-y)^2}{4\epsilon t} \right) \]  

(A5)

For initial data

\[ w(x,0) = \begin{cases} \psi_L & \text{if } x < 0 \\ \psi_R & \text{if } x > 0 \end{cases} \]  

(A6)

the solution is

\[ w(x,t) = \psi_L + (\psi_R - \psi_L)S(x,t) \quad S(x,t) = \frac{1}{2} \left[ 1 + \text{erf} \left( \frac{x}{2\sqrt{\epsilon t}} \right) \right] \]  

(A7)

Let the color function be denoted by \( \psi^c \), and the smooth version by \( \psi^S \). Their initial data are
\[
\psi^c(x,t) = \begin{cases} 
-1 & x<0 \\
1 & x>0 
\end{cases} 
\]  
(A8)

and

\[
\psi^s(x,t) = x 
\]  
(A9)

respectively. The initial density distribution is \( \rho(x,0) = \rho_L \) for negative and \( \rho(x,0) = \rho_R \) for positive \( x \). The solutions are

\[
\psi^c(x,t) = \frac{-\rho_L + (\rho_R + \rho_L)S(x,t)}{\rho_L + (\rho_R - \rho_L)S(x,t)} 
\]  
(A10)

and

\[
\psi^s(x,t) = x + \frac{(\rho_R - \rho_L)}{\rho_L + (\rho_R - \rho_L)S(x,t)} \left( \frac{\epsilon_l}{\pi} \right)^{1/2} \exp \left( -\frac{x^2}{4\epsilon_l} \right) 
\]  
(A11)

At \( x=0 \), we find

\[
\psi^s(0,t) = \frac{\rho_R - \rho_L}{\rho_R + \rho_L} 
\]  
(A12)

\[
\psi^c(0,t) = \frac{\rho_R - \rho_L}{\rho_R + \rho_L} \left( \frac{\epsilon_l}{\pi} \right)^{1/2} 
\]  
(A13)

Let \( A = (\rho_R - \rho_L)/(\rho_R + \rho_L) \). Then the point where \( \psi^c(x,t) = 0 \) is, for small \( A \),

\[
x_f^c = A \sqrt{\epsilon_l} 
\]  
(A14)

whereas for the initially smooth \( \psi^s(x,t) \) we find

\[
x_f^s = A \sqrt{\epsilon_l/\kappa} 
\]  
(A15)

Both are wrong: we should have \( x_f = 0 \). The smooth function \( \psi^s(x,t) \) is better than the color function \( \psi^c(x,t) \) in monitoring the position of the front, but only by a factor \( 2\kappa/\kappa = 0.64 \).

Figure 7 illustrates what happens for the second-order ENO/ROE scheme. Two contacts are moving on a one-dimensional periodic grid. The density and color function are smeared, due to
numerical viscosity. Although the numerical viscosity is smaller than for the first-order scheme, the zero-crossings of $\psi^C$ and $\psi^S$ appear to display the effect described above. Non-conservative differencing for both $\psi^S$ and $\psi^C$ are also shown. The non-conservative scheme for $\psi^S$ seems to be the best choice.
REFERENCES


| \((2h)^{-1}, (h)^{-1}\)| time | \(\mu = 0\) | \(5 \times 10^{-4}\) | \(1 \times 10^{-3}\) | \(5 \times 10^{-3}\) |
|---|---|---|---|---|
| 32,64 | 0 | 1.15 - 5 | 1.15 - 5 | 1.15 - 5 | 1.15 - 5 |
| | 2 | 4.03 - 4 | 3.22 - 4 | 5.13 - 4 | 3.46 - 4 |
| | 4 | 7.37 - 3 | 4.79 - 3 | 3.70 - 3 | 1.08 - 3 |
| | 6 | 2.79 - 2 | 2.47 - 2 | 2.02 - 2 | 5.67 - 3 |
| 48,96 | 0 | 5.11 - 6 | 5.11 - 6 | 5.11 - 6 | 5.11 - 6 |
| | 2 | 2.79 - 4 | 1.80 - 4 | 2.46 - 4 | 1.62 - 4 |
| | 4 | 6.40 - 3 | 3.42 - 3 | 2.27 - 3 | 5.48 - 4 |
| | 6 | 2.88 - 2 | 1.80 - 2 | 1.39 - 2 | 3.19 - 3 |
| 64,128 | 0 | 2.87 - 6 | 2.87 - 6 | 2.87 - 6 | 2.87 - 6 |
| | 2 | 2.34 - 4 | 1.21 - 4 | 1.55 - 4 | 9.71 - 5 |
| | 4 | 5.90 - 3 | 2.58 - 3 | 1.58 - 3 | 3.46 - 4 |
| | 6 | 3.57 - 2 | 1.22 - 2 | 9.59 - 3 | 2.10 - 3 |
| 96,192 | 0 | 1.28 - 6 | 1.28 - 6 | 1.28 - 6 | 1.28 - 6 |
| | 2 | 1.71 - 4 | 6.85 - 5 | 8.04 - 5 | 5.05 - 5 |
| | 4 | 5.71 - 3 | 1.52 - 3 | 8.42 - 4 | 1.92 - 4 |
| | 6 | 6.20 - 2 | 6.64 - 3 | 5.14 - 3 | 1.16 - 3 |

Table 1a. Relative error \(E_h^2(\psi)\) measured in the \(\ell_1\) norm, as a function of grid size and viscosity \(\mu\) at times 0, 2, 4, and 6, for the Rayleigh Taylor problem.

<table>
<thead>
<tr>
<th>(\mu)</th>
<th>(\text{norm})</th>
<th>(t = 0)</th>
<th>(t = 2)</th>
<th>(t = 4)</th>
<th>(t = 6)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>(\ell_1)</td>
<td>2.00</td>
<td>1.04</td>
<td>0.16</td>
<td>-0.93</td>
</tr>
<tr>
<td></td>
<td>(\ell_\infty)</td>
<td>1.95</td>
<td>0.01</td>
<td>-0.41</td>
<td>-0.81</td>
</tr>
<tr>
<td>(5 \times 10^{-4})</td>
<td>(\ell_1)</td>
<td>2.00</td>
<td>1.62</td>
<td>1.22</td>
<td>1.42</td>
</tr>
<tr>
<td></td>
<td>(\ell_\infty)</td>
<td>1.95</td>
<td>0.83</td>
<td>0.98</td>
<td>0.78</td>
</tr>
<tr>
<td>(1 \times 10^{-3})</td>
<td>(\ell_1)</td>
<td>2.00</td>
<td>1.63</td>
<td>1.46</td>
<td>1.43</td>
</tr>
<tr>
<td></td>
<td>(\ell_\infty)</td>
<td>1.95</td>
<td>0.78</td>
<td>1.29</td>
<td>0.97</td>
</tr>
<tr>
<td>(5 \times 10^{-3})</td>
<td>(\ell_1)</td>
<td>2.00</td>
<td>1.67</td>
<td>1.51</td>
<td>1.46</td>
</tr>
<tr>
<td></td>
<td>(\ell_\infty)</td>
<td>1.95</td>
<td>0.61</td>
<td>0.84</td>
<td>1.32</td>
</tr>
</tbody>
</table>

Table 1b. Order of accuracy, estimated from the relative error in \(\psi\) measured in the \(\ell_1\) and \(\ell_\infty\) norm, as a function of the viscosity \(\mu\) at various time, for the Rayleigh Taylor problem.
### Table 2a. Relative error $E_h^2(\psi)$ measured in the $l_1$ norm, as a function of grid size and viscosity $\mu$ at times 0, 2, 4, and 6, for the Kelvin-Helmholtz problem.

<table>
<thead>
<tr>
<th>$(2h)^{-1}, h^{-1}$</th>
<th>time</th>
<th>$\mu = 0$</th>
<th>$1 \times 10^{-4}$</th>
<th>$5 \times 10^{-4}$</th>
<th>$1 \times 10^{-3}$</th>
<th>$5 \times 10^{-3}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>32, 64</td>
<td>0</td>
<td>3.94 - 5</td>
<td>3.94 - 5</td>
<td>3.94 - 5</td>
<td>3.94 - 5</td>
<td>3.94 - 5</td>
</tr>
<tr>
<td></td>
<td>2</td>
<td>6.64 - 3</td>
<td>6.47 - 3</td>
<td>5.45 - 3</td>
<td>4.73 - 3</td>
<td>2.79 - 3</td>
</tr>
<tr>
<td></td>
<td>4</td>
<td>1.28 - 2</td>
<td>1.20 - 2</td>
<td>0.96 - 3</td>
<td>0.80 - 3</td>
<td>0.44 - 3</td>
</tr>
<tr>
<td></td>
<td>6</td>
<td>1.64 - 2</td>
<td>1.64 - 2</td>
<td>1.25 - 2</td>
<td>1.04 - 2</td>
<td>0.53 - 3</td>
</tr>
<tr>
<td></td>
<td>8</td>
<td>1.04 - 2</td>
<td>1.21 - 2</td>
<td>1.36 - 2</td>
<td>1.14 - 2</td>
<td>0.55 - 3</td>
</tr>
<tr>
<td>48, 96</td>
<td>0</td>
<td>1.76 - 5</td>
<td>1.76 - 5</td>
<td>1.76 - 5</td>
<td>1.76 - 5</td>
<td>1.76 - 5</td>
</tr>
<tr>
<td></td>
<td>2</td>
<td>5.64 - 3</td>
<td>5.18 - 3</td>
<td>3.71 - 3</td>
<td>2.89 - 3</td>
<td>1.57 - 3</td>
</tr>
<tr>
<td></td>
<td>4</td>
<td>1.34 - 2</td>
<td>1.24 - 2</td>
<td>0.74 - 3</td>
<td>0.59 - 3</td>
<td>0.25 - 3</td>
</tr>
<tr>
<td></td>
<td>6</td>
<td>1.77 - 2</td>
<td>1.69 - 2</td>
<td>1.19 - 2</td>
<td>0.87 - 3</td>
<td>0.32 - 3</td>
</tr>
<tr>
<td></td>
<td>8</td>
<td>1.86 - 2</td>
<td>1.62 - 2</td>
<td>0.96 - 3</td>
<td>0.92 - 3</td>
<td>0.34 - 3</td>
</tr>
<tr>
<td>64, 128</td>
<td>0</td>
<td>9.93 - 6</td>
<td>9.93 - 6</td>
<td>9.93 - 6</td>
<td>9.93 - 6</td>
<td>9.93 - 6</td>
</tr>
<tr>
<td></td>
<td>2</td>
<td>5.11 - 3</td>
<td>4.33 - 3</td>
<td>2.68 - 3</td>
<td>1.95 - 3</td>
<td>1.02 - 3</td>
</tr>
<tr>
<td></td>
<td>4</td>
<td>1.29 - 2</td>
<td>1.13 - 2</td>
<td>0.68 - 3</td>
<td>0.49 - 3</td>
<td>0.20 - 3</td>
</tr>
<tr>
<td></td>
<td>6</td>
<td>1.89 - 2</td>
<td>1.74 - 2</td>
<td>1.03 - 2</td>
<td>0.68 - 3</td>
<td>0.21 - 3</td>
</tr>
<tr>
<td></td>
<td>8</td>
<td>1.85 - 2</td>
<td>1.82 - 2</td>
<td>0.81 - 3</td>
<td>0.56 - 3</td>
<td>0.23 - 3</td>
</tr>
</tbody>
</table>

### Table 2b. Order of accuracy, estimated from the relative error in $\psi$ measured in the $l_1$ and $l_\infty$ norm, as a function of the viscosity $\mu$ at various time, for the Kelvin-Helmholtz instability.

<table>
<thead>
<tr>
<th>$\mu$</th>
<th>norm</th>
<th>$t = 0$</th>
<th>$t = 2$</th>
<th>$t = 4$</th>
<th>$t = 6$</th>
<th>$t = 8$</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>$l_1$</td>
<td>1.99</td>
<td>0.36</td>
<td>0.04</td>
<td>-0.21</td>
<td>-0.48</td>
</tr>
<tr>
<td></td>
<td>$l_\infty$</td>
<td>1.31</td>
<td>-0.19</td>
<td>-0.58</td>
<td>-1.02</td>
<td>-1.11</td>
</tr>
<tr>
<td>$1 \times 10^{-4}$</td>
<td>$l_1$</td>
<td>1.99</td>
<td>0.60</td>
<td>0.18</td>
<td>-0.08</td>
<td>-0.51</td>
</tr>
<tr>
<td></td>
<td>$l_\infty$</td>
<td>1.31</td>
<td>0.02</td>
<td>-0.51</td>
<td>-0.90</td>
<td>-1.33</td>
</tr>
<tr>
<td>$5 \times 10^{-4}$</td>
<td>$l_1$</td>
<td>1.99</td>
<td>1.07</td>
<td>0.60</td>
<td>0.36</td>
<td>0.58</td>
</tr>
<tr>
<td></td>
<td>$l_\infty$</td>
<td>1.31</td>
<td>0.81</td>
<td>0.46</td>
<td>-0.07</td>
<td>0.33</td>
</tr>
<tr>
<td>$1 \times 10^{-3}$</td>
<td>$l_1$</td>
<td>1.99</td>
<td>1.31</td>
<td>1.14</td>
<td>0.72</td>
<td>0.49</td>
</tr>
<tr>
<td></td>
<td>$l_\infty$</td>
<td>1.31</td>
<td>1.24</td>
<td>0.92</td>
<td>0.29</td>
<td>0.18</td>
</tr>
<tr>
<td>$5 \times 10^{-3}$</td>
<td>$l_1$</td>
<td>1.99</td>
<td>1.47</td>
<td>1.41</td>
<td>1.30</td>
<td>1.23</td>
</tr>
<tr>
<td></td>
<td>$l_\infty$</td>
<td>1.31</td>
<td>1.00</td>
<td>1.60</td>
<td>1.00</td>
<td>0.82</td>
</tr>
</tbody>
</table>
Figure captions

Fig. 1a. Contours of $\psi$ at values of $-1/32$, 0, and 1/32, for a 64$\times$128 grid. The computation has been carried out on a 32$\times$128 grid, with forced symmetry.

Fig. 1b. As Fig. 1a, but for a 192$\times$384 grid.

Fig. 1c. Grid refinement sequence at time 6. We have $h^{-1}=32$, 48, 64, 96, 128, and 192, from left to right, top to bottom.

Fig. 2a. Contours of $\psi$ at values of $-1/32$, 0, and 1/32, for the computations described in Table 1 at time 6. The viscosity $\mu$ has values $0, 5\times10^{-4}, 1\times10^{-3}, 5\times10^{-3}$, and increases from left to right.

Fig. 2b. As Fig. 2a, but now the density is plotted. Contours are 0.1 apart.

Fig. 3a. Contours of the color function at $-0.5$, 0.0, and 0.5 for a 64$\times$128 grid. The computation has been carried out on a 32$\times$128 grid, with forced symmetry.

Fig. 3b. Vertical cross section halfway Fig. 3a at time 6. Shown are $\rho$ (drawn line), $\psi$ (dashed), and the color function (dots). Also shown are runs for non-conservative differencing.

Fig. 4a. Contours of $\psi$ at values of $-1/32$, 0, and 1/32, for a 32$\times$256 grid, in the Helium-Air case.

Fig. 4b. Same parameters and initialization with non-conservative differencing.

Fig. 4c. As Fig. 4a, but now $\psi$ has been initialized as -1 and +1, representing a color function. Contours are at -0.5, 0.0, and 0.5.

Fig. 4d. As Fig. 4a, but using the concentration $Y$ to determine the effective value of $\gamma$. Contours are drawn at 0.25, 0.50, and 0.75.

Fig. 4e. The (passive) function $\psi$ at values $-1/32$, 0, and 1/32, for the same computation as in Fig. 4d.

Fig. 5a. Kelvin-Helmholtz instability. Shown are contours of the (passive) function $\psi$ at values $-1/32$, 0, and 1/32. The grid has 128$\times$256 points.

Fig. 5b. Grid refinement sequence at time 6, for $h^{-1}=32$, 48, 64, 96, and 128.

Fig. 5c. Zero contours of $\psi$ for the computations described in Table 2 at time 6. The viscosity $\mu$ has values $0, 1\times10^{-4}, 5\times10^{-4}, 1\times10^{-3}, 5\times10^{-3}$, and increases from left to right.

Fig. 6. Kelvin-Helmholtz instability for air (above the interface) and helium (below the interface), on a 128$\times$256 grid.

Fig. 7. Propagation of two contact discontinuities on a periodic grid. The velocity $u=0.5$, the initial density is 1.0 or 0.2. Shown is the result at time 2, using second-order ENO/ROE. In the absence of numerical viscosity, the results would be identical to the initial data. The jumps in density (drawn line) occur at $x=0$ and $x=0.5$. The dashed line represents the function $\psi^{(a)}$, initialized with $-\sin(2\pi x)$, whereas the dotted line displays the color function $\psi^{C(a)}$, initialized with $-1$ if $\rho=1.0$ and +1 if $\rho=0.2$. Also shown are the cases with non-conservative differencing.
Fig. 1a. Contours of $y$ at values of $-1/32$, 0, and $1/32$, for a 64x128 grid. The computation has been carried out on a 32x128 grid, with forced symmetry.

X002: $D = 2.000$, $M^2 = 0.500$, $a = 0.015000 [64x128](2,2) \, i\text{grid}=2, \, i\text{curv}=1, \, K = 0.00000, \, d = 0.03125$
Fig. 1b. As Fig. 1a, but for a 192x384 grid.

X005: D = 2.000, M**2 = 0.500, a = 0.015000 [192x384](2,2) igrid=2, icurv=1, K = 0.00000, d = 0.03125
Fig. 1c. Grid refinement sequence at time 6. We have \( n = 32, 48, 64, 96, 128, \) and 192, from left to right, top to bottom.

W001: \( D = 2.000, M = 0.500, a = 0.015000 \) [32x 64](2,2) igrid = 2, icurv = 1, K = 0.00000, d = 0.03125
Fig. 2a. Contours of $\psi$ at values of $-1/32$, 0, and $1/32$, for the computations described in Table 1 at time 6. The viscosity $\mu$ has values $0, 5 \cdot 10^{-4}, 1 \cdot 10^{-3}, 5 \cdot 10^{-3}$, and increases from left to right.

$\sqrt[001]{D} = 2.000, M^* = 0.500, a = 0.015\text{000}[, 192 \times 384](2,2) \text{grid} = 2, \text{icurv} = 1, K = 0.00\text{000}, d = 0.03\text{125}$
Fig. 2b. As Fig. 2a, but now the density is plotted. Contours are 0.1 apart.

V001: D= 2.000, M**2= 0.500, a= 0.015000[ 192x 364](2,2) igrid=2, icurv=1, K= 0.00000, r, 0.10000
Fig. 3a. Contours of the color function at -0.5, 0.0, and 0.5 for a 64x128 grid. The computation has been carried out on a 32x128 grid, with forced symmetry.

XC02: D = 2.000, M**2 = 0.500, a = 0.015000[ 64x128](2,2) igrid=2, icurv=3, K = 0.00000, d = 0.50000
Fig. 3b. Vertical cross section halfway Fig. 3a at time 6. Shown are $\rho$ (drawn line), $\psi$ (dashed), and the color function (dots). Also shown are runs for non-conservative differencing.
Fig. 4a. Contours of $\psi$ at values of $-1/32$, $0$, and $1/32$, for a $32 \times 256$ grid, in the Helium-Air case.

Y002: $D = 7.236$, $M^2 = 1.000$, $a = 0.015000[32 \times 256](2,2)\text{igrid}=2$, $\text{icurv}=1$, $K = 0.00000$, $d = 0.03125$
Fig. 4b Same parameters and initialization with non-conservative differencing.

Tn0: D = 7.236, M**2 = 1.000, a = 0.015000! 32x 256(2.2) or.d,crv,psi:2 1 1, K = 0.00000, d, 0.03125
Fig. 4c. As Fig. 4a, but now \( y \) has been initialized as -1 and +1, representing a color function. Contours are at -0.5, 0.0, and 0.5.

C002: \( D = 7.236, M^2 = 1.000, a = 0.015000 \), \( 32 \times 256 \)(2,2) \( igrid=2, icurv=3, K = 0.00000, d, 0.50000 \)
Fig. 4d. As Fig. 4a, but using the concentration $Y$ to determine the effective value of $\gamma$. Contours are drawn at 0.25, 0.50, and 0.75.

Parameter values: $D = 7.236$, $M^2 = 1.000$, $a = 0.015000 \times 32 \times 256$, $i\text{grid}=2$, $i\text{curv}=1$, $K = 0.00000$, $y = 0.25000$. 
Fig. 4e. The (passive) function \( \psi \) at values -1/32, 0, and 1/32, for the same computation as in Fig. 4d.

MO02: \( D = 7.236, M^{**2} = 1.000, a = 0.015000, 32x256(2,2) igrid=2, icurv=1, K = 0.00000, d, 0.03125 \)
Fig. 6a. Kelvin-Helmholtz instability. Shown are contours of the (passive) function $\psi$ at values $-1/32, 0$, and $1/32$. The grid has $128 \times 256$ points.

KHA4: $D = 1.000, M^2 = 0.250, a = 0.100000[128 \times 256](2,2)$ igrid = 1, icurv = 1, $K = 0.00000$, d, 0.03125
Fig. 9b. Grid refinement sequence at time 6, for $h^{-1} = 32, 48, 64, 96, \text{ and } 128$. 

$\text{GHG: } D = 1.000, M^2 = 0.250, a = 0.100000[32 \times 64](2,2) \text{ grid=1, icurv=1, } K = 0.00000, d, 0.00000$
Fig. 5c. Zero contours of $\psi$ for the computations described in Table 2 at time 6. The viscosity $\mu$ has values $0, 1 \times 10^{-4}, 5 \times 10^{-4}, 1 \times 10^{-3}, 5 \times 10^{-3}$, and increases from left to right.

KHV0: $D = 1.000, M_{ms} = 0.250, a = 0.100000[128x256](2.2)\ i_{grid}=1, i_{curv}=1, K = 0.00000, d = 0.00000$
Fig. 6. Kelvin-Helmholtz instability for air (above the interface) and helium (below the interface), on a 128x256 grid.

KD04: D= 7.236, \(M^2 = 0.500\), \(a=0.100000\) [128x256] (2,2) igrid=1, icurv=1, K=0.00000, d, 0.00000
Fig. 7. Propagation of two contact discontinuities on a periodic grid. The velocity $u_x=0.5$, the initial density is 1.0 or 0.2. Shown is the result at time 2, using second-order ENO/ROE. In the absence of numerical viscosity, the results would be identical to the initial data. The jumps in density (drawn line) occur at $x=0$ and $x=0.5$. The dashed line represents the function $\psi^s(x)$, initialized with $-\sin(2\pi x)$, whereas the dotted line displays the color function $\psi^c(x)$, initialized with $-1$ if $\rho=1.0$ and $+1$ if $\rho=0.2$. Also shown are the cases with non-conservative differencing.