**Cholla**: A NEW MASSIVELY-PARALLEL HYDRODYNAMICS CODE FOR ASTROPHYSICAL SIMULATION

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**Draft version October 17, 2014**

**ABSTRACT**

We present *Cholla* (Computational Hydrodynamics On Parallel Architectures), a new three-dimensional hydrodynamics code that harnesses the power of graphics processing units (GPUs) to accelerate astrophysical simulations. *Cholla* models the Euler equations on a static mesh using state-of-the-art techniques, including the unsplit Corner Transport Upwind (CTU) algorithm, a variety of exact and approximate Riemann solvers, and multiple spatial reconstruction techniques including the piecewise parabolic method (PPM). *Cholla* performs all hydrodynamical calculations in a massively-parallel manner, using GPUs to evolve the fluid properties of thousands of cells simultaneously while leaving the power of central processing units (CPUs) available for modeling additional physics. On current hardware, *Cholla* can update more than ten million cells per GPU-second while using an exact Riemann solver and PPM reconstruction with the CTU algorithm. Owing to the massively-parallel architecture of GPUs and the design of the *Cholla* code, astrophysical simulations with physically interesting grid resolutions (≥ 256³) can easily be computed on a single device. *Cholla* utilizes the Message Passing Interface library to extend calculations onto multiple devices, and exhibits nearly ideal scaling beyond 100,000 GPU cores. The excellent performance of *Cholla* is demonstrated on a suite of test problems that highlights the physical accuracy of our modeling and provides a useful comparison to other codes. We also provide a set of Appendices that uniformly documents all of the reconstruction methods and Riemann solvers implemented in *Cholla*, and discusses strengths and weaknesses of the various methods.

1. **INTRODUCTION**

Over the past fifty years, the field of computational hydrodynamics has grown to incorporate a wide array of numerical schemes that attempt to model a large range of astrophysical phenomena. From the pioneering work of Godunov (1959) and Courant et al. (1967), the sophistication of hydrodynamics solvers has steadily improved. Many astrophysical simulation codes now use high order reconstruction methods, implement very accurate or exact Riemann solvers, and model additional physics including gravity, cooling, magnetohydrodynamics, radiative transfer, and more (e.g., Kravtsov 1999; Knebe et al. 2001; Fryxell et al. 2000; Teyssier 2002; Hayes et al. 2006; Stone et al. 2008; Bryan et al. 2014). While these advanced techniques result in simulations of unprecedented physical accuracy, they can also be extremely computationally expensive. Given the detailed and expensive physical processes currently being modeled, new numerical approaches to modeling Eulerian hydrodynamics should be considered. This work presents the new, massively-parallel hydrodynamics code *Cholla* (Computational Hydrodynamics On Parallel Architectures) that leverages Graphics Processing Units (GPUs) to accelerate astrophysical simulations.

Historically, our ability to model larger and more complex systems has benefitted from improvements in computer technology, especially increased storage and faster clock speeds for central processing units (CPUs). Algorithmic improvements such as adaptive mesh refinement (e.g., Berger & Olliger 1984; Berger & Colella 1989) have had a major impact on the ability of codes to achieve higher resolution, but much of the basic structure of static mesh grid codes has remained. In the last decade, computer speed has improved significantly as a result of increased parallelization, and the fastest supercomputers now rely on hardware accelerators like GPUs or Intel Xeon Phi coprocessors to provide the bulk of their computational power. To leverage the full capabilities of these systems, multi-core CPU chips and accelerators must be used simultaneously in the context of a single hydrodynamics code. While similar parallelization and vectorization techniques apply to a variety of hardware accelerators, *Cholla* utilizes GPUs to perform all its hydrodynamical calculations.

Originally designed for on-screen pixel rendering, the GPU was quickly recognized as a potentially useful tool for scientific calculations. Companies such as NVIDIA and ATI produce general purpose GPUs for scientific computing, and have developed software like the CUDA (Compute Unified Device Architecture) extension to the C language used to program NVIDIA GPUs. Engineering *Cholla* to run natively on NVIDIA GPUs allows us to take advantage of the inherently parallel structure of grid-based Eulerian hydrodynamics schemes, and enables the substantial computational performance gain demonstrated in this paper.

Accelerators and other special purpose hardware have been used in astrophysical simulations for many years (e.g., Sugimoto et al. 1990; Aarseth 1999; Spurzem 1999; Portegies Zwart & Bedorf 2014). Several GPU-based hydrodynamical simulation methods have been recently presented, including AMR techniques (Schive et al. 2010; Wang et al. 2010), two dimensional Galerkin approaches (Chan et al. 2012), Smoothed Particle Hydrodynamics (SPH) codes (Sandalski 2012; Domínguez et al. 2013), and...
and hybrid schemes \cite{Kulikov2014}. Nonetheless, the field of massively-parallel astrophysical hydrodynamics is still relatively new. As is evidenced by the number of CPU codes presented in the literature, room exists for many different approaches optimized for different purposes. With Cholla, we have built a fast, GPU-accelerated static mesh hydrodynamics module that can be used efficiently in conjunction with a variety of additional physics. Beyond accelerating the hydrodynamics calculation, offloading the work onto the GPU frees the CPU to perform other tasks. This excess computational capacity makes Cholla an excellent bedrock for developing complex physical models that require hydrodynamics.

In the following sections, we fully describe Cholla. The code models solutions to the equations of hydrodynamics using the Corner Transport Upwind (CTU) algorithm \cite{Colella1990, Gardiner2008}, and includes multiple choices for both interface reconstruction and Riemann solvers. The CTU algorithm is presented in Section 2 along with brief descriptions of the reconstruction methods and Riemann solvers, which are fully documented in the Appendices. The code structure, including the simulation setup, CUDA functions, optimization strategies necessary to take advantage of the GPU architecture, and Message Passing Interface (MPI, Forum 1994) implementation and scalability, is described in Section 3. We then demonstrate the excellent performance of Cholla on a suite of canonical hydrodynamics tests in Section 4 and conclude in Section 5.

2. HYDRODYNAMICS

Hydrodynamics is relevant to many astrophysical processes and represents one of the most computationally demanding parts of numerical simulations. Creating a fast hydrodynamics solver is therefore an important step in increasing the resolution and speed with which astrophysical calculations can be performed. In this section, we present the equations modeled by Cholla, and then describe the numerical algorithms used to model them. Cholla includes a variety of reconstruction techniques and Riemann solvers, each of which is described below:

In differential conservation law form (see, e.g., \cite{Toro2000}), the multi-dimensional Euler equations can be written:

\begin{equation}
\frac{\delta \rho}{\delta t} + \frac{\delta (\rho u)}{\delta x} + \frac{\delta (\rho v)}{\delta y} + \frac{\delta (\rho w)}{\delta z} = 0,
\end{equation}

\begin{equation}
\frac{\delta (\rho u)}{\delta t} + \frac{\delta (\rho u^2 + p)}{\delta x} + \frac{\delta (\rho uv)}{\delta y} + \frac{\delta (\rho uw)}{\delta z} = 0,
\end{equation}

\begin{equation}
\frac{\delta (\rho v)}{\delta t} + \frac{\delta (\rho uv)}{\delta x} + \frac{\delta (\rho v^2 + p)}{\delta y} + \frac{\delta (\rho vw)}{\delta z} = 0,
\end{equation}

\begin{equation}
\frac{\delta (\rho w)}{\delta t} + \frac{\delta (\rho uw)}{\delta x} + \frac{\delta (\rho vw)}{\delta y} + \frac{\delta (\rho w^2 + p)}{\delta z} = 0,
\end{equation}

\begin{equation}
\frac{\delta E}{\delta t} + \frac{\delta [u(E + p)]}{\delta x} + \frac{\delta [v(E + p)]}{\delta y} + \frac{\delta [w(E + p)]}{\delta z} = 0.
\end{equation}

Here \( \rho \) is the mass density, \( u, v, \) and \( w \) are the \( x, y, \) and \( z \)-components of velocity, \( p \) is the pressure, and \( E \) is the total energy per unit volume,

\begin{equation}
E = \rho \left( \frac{1}{2} V^2 + e \right),
\end{equation}

where \( V = [u, v, w]^T \) is the three-component velocity vector. The total energy includes the specific internal energy, \( e \), and the specific kinetic energy,

\begin{equation}
\frac{1}{2} V^2 = \frac{1}{2} V \cdot V = \frac{1}{2} (u^2 + v^2 + w^2). \end{equation}

Equation 1 describes the conservation of mass, Equations 2-4 the conservation of momentum, and Equation 5 the conservation of energy. To model solutions to this system of conservation laws, an equation of state is also necessary. We use the equation of state for an ideal gas,

\begin{equation}
p = (\gamma - 1) \rho e,
\end{equation}

where \( \gamma \) is the ratio of specific heats. Incorporating a real gas equation-of-state model \cite{Colella1985} would not be incompatible with the structure of Cholla, though it is beyond the scope of our current work.

The Euler equations can also be written in vector notation. We define the vector of conserved quantities with components in three Cartesian dimensions,

\begin{equation}
u = [\rho, \rho u, \rho v, \rho w, \rho e]^T
\end{equation}

including density, the three components of velocity, and total energy. We will also refer to the vector of primitive variables,

\begin{equation}
w = [\rho, u, v, w, p]^T,
\end{equation}

that includes density, the three components of velocity, and pressure. We define three flux vectors

\begin{equation}
\mathbf{f} = \begin{bmatrix} \rho u \\ \rho u^2 + p \\ \rho u v \\ \rho u w \\ (E + p) u \end{bmatrix},
\end{equation}

\begin{equation}
\mathbf{g} = \begin{bmatrix} \rho v \\ \rho v w \\ \rho v^2 + p \\ (E + p) v \end{bmatrix},
\end{equation}

\begin{equation}
\mathbf{h} = \begin{bmatrix} \rho w \\ \rho w v \\ \rho w w \\ \rho w^2 + p \\ (E + p) w \end{bmatrix}.
\end{equation}

Using these definitions, we can compactly write the three dimensional Euler equations in the conservative form:

\begin{equation}
\frac{\delta \mathbf{u}}{\delta t} + \frac{\delta \mathbf{f}}{\delta x} + \frac{\delta \mathbf{g}}{\delta y} + \frac{\delta \mathbf{h}}{\delta z} = 0.
\end{equation}

The Euler equations can also be written using the primitive variables. In one dimension, the equations can be written as a set of linear hyperbolic equations of the form

\begin{equation}
\frac{\delta \mathbf{w}}{\delta t} + \mathbf{A}(\mathbf{w}) \frac{\delta \mathbf{w}}{\delta x} = 0.
\end{equation}
The matrix $A(w)$ is diagonalizable and can be written

$$A = RL\tilde{L},$$

(16)

where $R$ is a matrix of right eigenvectors, $A$ is a diagonal matrix of eigenvalues, and $L = R^{-1}$ is a matrix of left eigenvectors. The eigenvalues of $A$ are real and correspond to the speeds at which information propagates for the fluid equations. If we further define the characteristic variables, $\xi$, according to

$$d\xi = Ldw,$$

(17)

we can write the system a third way:

$$\frac{\delta\xi}{\delta t} + \Lambda \frac{\delta\xi}{\delta x} = 0.$$

(18)

This description is called the characteristic form of the Euler equations. The characteristic variables are sometimes called wave strengths, because they describe the magnitude of the jump in the primitive variables across an associated wave. For an extensive treatment of the Euler equations and related subjects see, e.g., [Laney (1998), LeVeque (2002), and Toro (2009)].

2.1. The CTU Algorithm

Cholla models the Euler equations using a three-dimensional implementation of the Corner Transport Upwind (CTU) algorithm optimized for magnetohydrodynamics (MHD; Colella 1990; Saltzman 1994; Gardiner & Stone 2008). The six-solve version of CTU used in Cholla is fully adapted for MHD and documented in both Gardiner & Stone (2008) and Stone et al. (2008). We will describe here an abbreviated version including only the steps relevant for hydrodynamics calculations.

The CTU algorithm is a Godunov-based method. A Godunov scheme uses a finite-volume approximation to model the Euler equations, evolving average values of the conserved variables $u$ in each cell using fluxes calculated at cell interfaces (Godunov 1959). In three dimensions, the calculation to update the conserved quantities $u^n$ at timestep $n$ can be written as

$$u^{n+1}_{(i,j,k)} = u^n_{(i,j,k)} + \frac{\Delta t}{\Delta x} [F^{n+\frac{1}{2}}_{(i-\frac{1}{2},j,k)} - F^{n+\frac{1}{2}}_{(i+\frac{1}{2},j,k)}]$$

$$+ \frac{\Delta t}{\Delta y} [G^{n+\frac{1}{2}}_{(i,j-\frac{1}{2},k)} - G^{n+\frac{1}{2}}_{(i,j+\frac{1}{2},k)}]$$

$$+ \frac{\Delta t}{\Delta z} [H^{n+\frac{1}{2}}_{(i,j,k-\frac{1}{2})} - H^{n+\frac{1}{2}}_{(i,j,k+\frac{1}{2})}].$$

(19)

Here the superscript $n+1$ refers to the next time step, and $u^{n+1}_{(i,j,k)}$ are the updated values of the conserved variables. The subscript $(i,j,k)$ refers to the three-dimensional Cartesian index of the cell. Indices that are displaced by half an integer refer to interfaces. For example, $(i-\frac{1}{2},j,k)$ is the interface between cell $(i-1,j,k)$ and cell $(i,j,k)$. The simulation time step is $\Delta t$, and $\Delta x$, $\Delta y$, and $\Delta z$ refer to the cell widths in each dimension. We use lower case versions of $u$ and $w$ when referring to a cell-averaged quantity, and uppercase versions when referring to an estimated value at a cell edge (e.g., $U$ and $W$). The fluxes in Equation (19) are averages in both space and time. Table 1 summarizes our notation.

When applied to every cell in the grid, Equation (19) is exact and conserves each of the quantities in $u$. However, the physical accuracy of a method based solely on Equation (19) is limited. Many hydrodynamics codes calculate the fluxes using only one-dimensional information as outlined in the method known as Strang (1968) splitting. While an elegant technique, Strang splitting may lead to asymmetries in hydrodynamics calculations and is not advantageous for some formulations of MHD (see, e.g., the discussion in [Balsara 2004]). Therefore, we employ instead the unsplit CTU algorithm to improve on the one-dimensional calculation of fluxes by taking into account transverse fluxes that can cross cell interfaces in multi-dimensional simulations.

Before beginning a simulation the computational domain and boundaries must be initialized, as described in Section 3. Once the fluid properties on the grid have been initialized, the first simulation time step, $\Delta t$, is calculated using the equation

$$\Delta t = C_0 \frac{\Delta x}{|u| + a},$$

(20)

where $C_0$ is the Courant-Friedrichs-Lewy (CFL) number and $a$ is the average sound speed in the cell. In adiabatic hydrodynamics, the sound speed is a function of pressure and density

$$a = \sqrt{\gamma p/\rho},$$

(21)

and can be calculated for each cell using the average values of the primitive variables. Thus, $|u| + a$ is the maximum wave speed in a cell with respect to the grid.

The minimum value of $\Delta t$ across the entire grid is determined and used in the CTU calculation, constraining the time step for every cell to be equal. In two or three dimensions, Equation (20) is modified such that the minimum required timestep is computed for each direction in the cell. In three dimensions, this minimization is computed as

$$\Delta t = C_0 \min \left( \frac{\Delta x}{|u| + a}, \frac{\Delta y}{|v| + a}, \frac{\Delta z}{|w| + a} \right).$$

(22)

With a suitable choice of $C_0$, Equation (22) ensures that the Courant condition is satisfied in all three dimensions. Note that for the six-solve CTU algorithm the CFL number must be below 0.5 for the solution to remain stable [Gardiner & Stone 2008].
Once we have calculated the timestep \( \Delta t \) we carry out the following procedure:

1. Reconstruct values of the conserved variables on both sides of every interface using the average value of the conserved quantities in adjacent cells. These reconstructed boundary values, denoted \( U_L \) and \( U_R \), represent an approximation to the true value of each conserved variable at the interface. For a multi-dimensional simulation, the reconstruction must be carried out in each dimension separately. We use additional subscripts to indicate which interface values we are calculating. For example, the left-hand reconstructed boundary value between cell \((i, j, k)\) and cell \((i+1, j, k)\) is denoted \( U_{L,(i+\frac{1}{2},j,k)} \). Cholla includes several different options for the interface reconstruction, which we describe in Section 2.2. In order for the CTU algorithm to be second-order accurate in time, the reconstructed boundary values must be evolved half a time step before using them to calculate fluxes. The initial time evolution is considered part of the reconstruction, and is also inherently one-dimensional. We label the initial time-evolved boundary values \( U^*_L \) and \( U^*_R \). An example of a piecewise-linear reconstruction in the \( x \)-direction is shown in Figure 1.

2. Using the initial time-evolved boundary values as inputs, solve a Riemann problem at each cell interface in each direction. The solution to the Riemann problems yields a set of one-dimensional fluxes, \( F^* \), \( G^* \), and \( H^* \), corresponding to the \( x \)-, \( y \)-, and \( z \)-interfaces, respectively. Like the boundary value arrays, the flux arrays contain five conserved value fluxes for each direction and interface. The Riemann solvers implemented in Cholla are described in Section 2.3.

3. Evolve the initial one-dimensional time-evolved boundary values half a time step using the transverse fluxes. For example, at the interface between cell \((i, j, k)\) and cell \((i+1, j, k)\) the transverse-flux-evolved boundary values are calculated using transverse-flux-evolved boundary values, denoted \( F^{n+\frac{1}{2}} \), \( G^{n+\frac{1}{2}} \), and \( H^{n+\frac{1}{2}} \). Cholla includes several different options for the transverse-flux-evolved boundary values half a time step using the transverse fluxes.

4. Use the transverse-flux-evolved boundary values, \( U^{n+\frac{1}{2}}_L \) and \( U^{n+\frac{1}{2}}_R \), as inputs for a new set of Riemann problems. The solution to these Riemann problems yields the CTU fluxes \( F^{n+\frac{1}{2}} \), \( G^{n+\frac{1}{2}} \), and \( H^{n+\frac{1}{2}} \). Each flux is calculated using a one-dimensional Riemann problem at a single interface, but includes contributions from adjacent perpendicular interfaces as a result of the evolution in Equation 23. The CTU fluxes are second-order accurate in time (Colella 1990).

5. Use the CTU fluxes to update the conserved quantities in each cell as in Equation 19.

\[
\begin{align*}
\mathbf{u}^{n+1}_{(i,j,k)} = & \mathbf{u}^n_{(i,j,k)} \\
& + \frac{\Delta t}{\Delta x} \left[ F^{n+\frac{1}{2}}_{(i-\frac{1}{2},j,k)} - F^{n+\frac{1}{2}}_{(i+\frac{1}{2},j,k)} \right] \\
& + \frac{\Delta t}{\Delta y} \left[ G^{n+\frac{1}{2}}_{(i,j-\frac{1}{2},k)} - G^{n+\frac{1}{2}}_{(i,j+\frac{1}{2},k)} \right] \\
& + \frac{\Delta t}{\Delta z} \left[ H^{n+\frac{1}{2}}_{(i,j,k-\frac{1}{2})} - H^{n+\frac{1}{2}}_{(i,j,k+\frac{1}{2})} \right].
\end{align*}
\]

The updated conserved quantities are used to calculate the next time step \( \Delta t^{n+1} \).

6. Repeat the algorithm until the final simulation time is reached.

| Symbol | Definition |
|--------|------------|
| \( U^*_L, U^*_R \) | Reconstructed boundary values at the left and right of an interface |
| \( F^*, G^*, H^* \) | Initial one-dimensional fluxes |
| \( R_{L,R}^{n+\frac{1}{2}}, R_{R,L}^{n+\frac{1}{2}} \) | Transverse-flux-evolved boundary values |
| \( F^{n+\frac{1}{2}}, G^{n+\frac{1}{2}}, H^{n+\frac{1}{2}} \) | CTU fluxes |
| \( u^{n+1} \) | Updated conserved average variables at time \( n+1 \) |
2.2. Interface Reconstruction

Cholla currently implements five methods for cell interface reconstruction. These include the piecewise constant method (PCM), two versions of the piecewise linear method (PLM), and two versions of the piecewise parabolic method (PPM). Differences between the versions of piecewise linear and piecewise parabolic reconstruction are demonstrated in the tests presented in Section 4. Access to multiple reconstruction options often proves useful, since lower-order methods are faster but higher-order methods are typically more accurate. Employing different versions of cell reconstruction also enables the impact of reconstruction on the evolution of a simulation to be quantified. Here, we give a brief overview of each of the reconstruction techniques implemented in Cholla. Detailed descriptions of the piecewise linear and piecewise parabolic options can be found in Appendix A.

2.2.1. Piecewise Constant Reconstruction

The simplest reconstruction technique is the piecewise constant method (Godunov 1959; Courant et al. 1967). In PCM, the initial time-evolved boundary values \( U^*_L \) and \( U^*_R \) are set equal to the cell average quantities on either side of the interface, i.e.

\[
U^*_L(i-\frac{1}{2},j,k) = u^n_{(i,j,k)}, \quad (25)
\]

and

\[
U^*_R(i+\frac{1}{2},j,k) = u^n_{(i,j,k)}, \quad (26)
\]

Note that in this notation, the boundary value at the right of the interface is at the left side of the cell, and vice versa. While the piecewise constant method is generally too diffusive for practical applications, it has merit for code testing, and is useful as a comparison to higher order reconstruction techniques.

2.2.2. Piecewise Linear Reconstruction

The second and third reconstruction techniques implemented in Cholla are both forms of the piecewise linear method, a scheme that is second-order accurate in space and time (e.g. Toro 2009). The PLMP reconstruction method detailed below primarily involves the primitive variables, while the PLMC method subsequently explicated involves projecting the primitive variables onto wave characteristics.

PLMP follows the method outlined in Chapter 13.4 of Toro (2009). First, the cell-average values of the primitive variables are used to calculate slopes of each variable across the left and right interface of each cell. We use the van Leer (1979) limiter to monotonize the slopes, thereby reducing the likelihood of spurious oscillations in a numerical solution. The limited slopes are used to calculate reconstructed values of the primitive variables at the cell interfaces, \( W_L \) and \( W_R \). To convert these reconstructed boundary values into input states for the Riemann problem, we need to evolve them by half a time step. For PLMP, we do this by converting primitive quantities back into conserved variables and calculating the associated fluxes using Equations 11-13. We use these fluxes to evolve the reconstructed boundary values half a time step, generating the initial time-evolved boundary states for the first set of Riemann problems, \( U^*_L \) and \( U^*_R \).

The second linear reconstruction technique, PLMC, is based on the method outlined in Stone et al. (2008). This reconstruction also uses a linear approximation to model the distribution of the conserved quantities in each cell, but limits the slopes of the characteristic variables (rather than the primitive quantities) and evolves the reconstructed boundary values differently. Rather than simply evolving the boundary values using the associated fluxes as in PLMP, we employ the more sophisticated approach first described and termed “characteristic tracing” in Colella & Woodward (1984). We calculate a first approximation to the time-evolved boundary values by integrating under the linear interpolation function used to calculate \( W_L \) and \( W_R \). The domain of dependence of the reconstructed boundary value integral is defined by the minimum (for the left-hand interface) or maximum (for the right-hand interface) wave speed. This integration is then corrected by including the contribution from each of the other characteristics approaching the interface. Once the corrections have been made, the calculation provides the initial time-evolved boundary values \( U^*_L \) and \( U^*_R \) that act as input states for the first set of Riemann problems. This process is more fully described in Appendix A.

2.2.3. Piecewise Parabolic Reconstruction

The remaining two reconstruction techniques implemented in Cholla are both versions of the piecewise parabolic method (PPM) originally described in Colella & Woodward (1984). We call the first method PPMP as it performs the reconstruction using primitive variables. Our PPMP implementation closely follows the FLASH code documentation (Fryxell et al. 2000). The second method, abbreviated PPMC, uses an eigenvalue decomposition to project onto the characteristic variables and is based on the Athena code documentation (Stone et al. 2008). Each PPM reconstruction method is described in detail in Appendix A.

The approach to slope limiting differs slightly between the two parabolic reconstruction techniques. PPMP calculates slopes at each interface the same way as PLMP, using van Leer (1979) limiting in the primitive variables. The slopes are limited in the characteristic variables for PPMC. In the parabolic methods, slopes are calculated using a stencil of five cells (two on either side of the cell for which we are calculating boundary values), which allows us to create a parabolic reconstruction of the primitive variables. This parabolic reconstruction makes PPM third-order accurate in space, though it remains only second-order accurate in time (Colella & Woodward 1984).

Several differences between the two parabolic reconstruction methods warrant further discussion. PPMP identifies and steepens the slopes near contact discontinuities, which results in a method that is less diffusive for contact waves. Downsides to contact steepening include the necessity of empirically determined criteria for selecting contact discontinuities and increased oscillatory behavior in the solution near shocks. In Cholla the ability to turn off contact steepening is retained, allowing an explicit comparison of results obtained with and without the technique. PPMP also flattens shocks that have become too narrow to be treated accurately, which reduces the potential for severe post-shock oscillations.
2.3. Riemann Solvers

Much effort has been devoted to finding efficient numerical algorithms to solve the Riemann problem (e.g., Toro 2009), a special initial value problem consisting of two constant states separated by a jump discontinuity. The Riemann problem has an implicit analytic solution that usefully enables a numerical model for Eulerian hydrodynamics, as it allows for the calculation of the flux across an interface separating two initially discontinuous states. An example of a Riemann problem is shown in Figure 2, with the upper panel showing the initial states, the middle panel showing the solution at the cell interface, and the bottom panel showing the resulting density distribution.

Riemann solvers that calculate a numerical solution to the exact Riemann problem can be incorporated in hydrodynamics codes. The implicit nature of the Riemann solution requires an iterative step in the exact numerical solver. A large number of Riemann problems must be solved for every time step in a simulation, and the corresponding computational cost is substantial. As a result, a variety of approximate Riemann solvers have been engineered to quickly solve an approximation to the Euler equations at the expense of some physical accuracy.

Cholla computes numerical solutions to the Riemann problems on GPUs. Floating point operations are performed very efficiently on a GPU: additional factors like memory latency and data transfer contribute a larger share of the computational expense of the method. Thus, adding the extra operations needed for the iterative procedure in an exact solver versus an approximate one does not impact the performance speed of Cholla in the same way as for a CPU-based code. However, there are certain problems where the extra diffusion in an approximate solver is helpful, for example to deal with the well known carbuncle instability (Quirk 1994) that affects grid-aligned shocks. For this reason, Cholla includes both an exact solver and the linearized solver first described by Roe (1981) that gives an exact solution to a linear approximation of the Euler equations. Detailed descriptions of our implementation of both solvers can be found in Appendix A.

2.3.1. The Exact Solver

Cholla implements the exact Riemann solver presented in Toro (2009). The solver uses a Newton-Raphson iteration to calculate the pressure of the gas in the intermediate state $W^m$ of the Riemann solution that lies between the initial states on the left and right of the interface, as shown in Figure 2. Once the pressure in the intermediate state has been found, the exact solution for the primitive variables between the left and right initial states can be calculated explicitly at any later point in time. The pressure and velocity are used to determine the solution at the cell interface, and the values of the primitive variables at that point are used to calculate the fluxes of conserved variables at the interface according to Equations 11-13. Transverse velocities are passively advected as scalar quantities.

The Toro Riemann solver gives a numerically exact solution to the Riemann problem in one dimension, and will never return negative densities or pressures if the input states are physically self-consistent. However, the input states on either side of the cell interface are estimated quantities, and because of the extrapolation involved in the reconstruction techniques they could be physically invalid. In these situations, the solver may be presented with an initial value problem without a physically valid solution. To prevent artificial vacuum or negative pressure solutions owing to such a circumstance, a pressure floor of $10^{-20}$ in the adopted unit scheme is enforced in Cholla. In practice, when using an exact Riemann solver the pressure floor has proved necessary only when performing the Noh test described in Section 4.

2.3.2. The Roe Solver

The Roe Solver
One common alternative to calculating an exact solution to the Riemann problem is to linearize the non-linear conservation laws and solve the resulting approximate problem exactly. In one dimension, the non-linear Euler equations can be replaced with the following linearized equation

$$\frac{\delta u}{\delta t} + A(\tilde{u}) \frac{\delta u}{\delta x} = 0,$$  \hspace{1cm} (27)

where $A$ is a constant Jacobian evaluated at some average state $\tilde{u}$ that is a function of the initial states on either side of the cell interface. This method was employed by Roe (1981), and Cholla includes a linearized solver very similar to the original Roe solver.

The first step in the Roe solver is to calculate the average state $\tilde{u}$. This average state, along with the eigenvalues, $\lambda^\alpha$, and left and right eigenvectors of the Jacobian $A$, $L^\alpha$ and $R^\alpha$, can be used to calculate the Roe fluxes at the interface:

$$F_{Roe} = \frac{1}{2} \left( F_L + F_R + \sum_{\alpha=1}^m \lambda^{\alpha} |\alpha| R^\alpha \right).$$  \hspace{1cm} (28)

Here, $\alpha = 1, m$ are the $m$ characteristics of the solution, and

$$\lambda^\alpha = L^\alpha \cdot \delta U$$  \hspace{1cm} (29)

are the characteristic variables, determined by projecting the differences in the initial left and right states, $\delta U = U_R - U_L$, onto the left eigenvectors. $F_L$ and $F_R$ are fluxes calculated with the left and right input states using Equation [11]. Expressions for the average state, $\tilde{u}$, as well as the eigenvalues and eigenvectors are given in Roe (1981) and Appendix B. The matrix $A$ is not actually needed in the calculation.

As pointed out by Einfeldt et al. (1991), there are certain Riemann problems that will cause any linearized solver to fail. In these cases, the linearized solution to the Riemann problem results in negative densities or pressures in the intermediate state calculated between the left and right input states. Because this intermediate state is used to calculate the fluxes returned by the solver, these unphysical solutions may lead to numerical pathologies. A failsafe is needed to deal with the case where the Roe solver produces negative pressures or densities. Following the method of Stone et al. (2008), we check the intermediate densities and pressures before returning the fluxes calculated with the Roe solver. Should any of them be negative, we revert to using the simpler HLLE Riemann solver, described below.

2.3.3. The HLLE Solver

The HLLE solver is a modification of the HLL solver first described by Harten et al. (1983) and later modified by Einfeldt (1988). Although the method is extremely diffusive for contact discontinuities, as demonstrated by Einfeldt et al. (1991) the HLLE solver is guaranteed to be positively conservative (that is, the density and internal energy remain positive). The HLLE solver calculates the interface flux using an average of the left and right state fluxes, together with bounding speeds comprising the largest and smallest physical signal velocities in the solution to the exact Riemann problem. If the Roe solver produces negative densities or pressures, we replace the Roe fluxes with a new numerical flux

$$F_{HLLE} = \frac{b^p F_L - b^m F_R}{b^p - b^m} + \frac{b^p b^m}{b^p - b^m} \delta U.$$  \hspace{1cm} (30)

The fluxes $F_L$ and $F_R$, and the slopes $\delta U$ are calculated as in the Roe solver. The signal velocities $b^p$ and $b^m$ are calculated using the largest and smallest eigenvalues of the Roe matrix as described in Appendix B. Because the HLLE solver quickly allows contact discontinuities to diffuse, we do not use it as a standalone Riemann solver in Cholla.

3. Code Architecture

Cholla is a grid-based hydrodynamics code that takes advantage of the massively parallel computing power of GPUs. In order to harness this power, Cholla was designed with the operation of the GPU in mind. In this section, we describe the overall structure of Cholla, including optimization strategies necessary to benefit from the parallel architecture of GPUs. As is standard in GPU programming, we will use the term “host” to refer to the CPU, and “device” to refer to the GPU.

Cholla consists of a set of C/C++ routines that run on the host plus functions called kernels that execute on a device. The device kernels and the host functions that call them are written in CUDA C, an extension to the C language introduced by NVIDIA. All of the CUDA functions are contained in a separate hydro module so that they can be compiled independently with the NVIDIA nvcc compiler. In addition, we have written a C/C++ version of the hydro module that performs the same calculations as all of the GPU kernels, so it is possible to run Cholla without using graphics cards. We use this mode for testing, but it is not recommended for performance since the structure of the code is optimized for use with GPUs.

3.1. Simulation Overview

Before detailing each piece of the code, we give a general overview of the steps followed by Cholla when a simulation is run. Given the power of a single GPU, small problems can easily be run on a single host/device pair. For large problems, Cholla can be run using the MPI library, and we describe our MPI implementation in Section 3.7. If MPI is enabled, the simulation volume will be split into subvolumes according to the number of processes. Each subvolume will then be treated as a self-contained simulation volume for the duration of each simulation time step. The main difference between an MPI and non-MPI simulation is the method for applying boundary conditions at the end of each time step; we describe that method in Section 3.7.

Portions of our algorithm that require information from potentially distant cells in the global simulation volume must be carried out on the host. The main host functions set initial conditions, apply boundary conditions, and perform any interprocess communications. Parts of the calculation that only require information from nearby cells can be carried out on the device. Because the bulk of the computational work resides in the CTU calculation that requires a stencil containing only

\[ \text{http://developer.nvidia.com} \]
FIG. 3.— Algorithmic procedure of a Cholla simulation. The initialization and application of boundary conditions are done on the GPU. The initialized data is then transferred to the CPU, where the hydrodynamics calculations are performed. The updated conserved variables must then be passed back to the GPU after each time step so that boundary cell information can be exchanged and the data output written (if necessary).

The steps in the Cholla algorithm are listed below and illustrated in Figure 3.

1. Initialize the simulation by setting the values of the conserved fluid quantities for all cells in the simulation volume, and calculate the first time step.

2. Transfer the array \( u \) of conserved variables to the GPU. The conserved variable array contains the values of each conserved quantity for every cell in the simulation volume.

3. Perform the CTU calculation on the GPU, including updating the conserved variable array and computing the next time step.

4. Transfer the updated conserved variable array back to the CPU.

5. Apply the boundary conditions. When running an MPI simulation, this step may require interprocess communication to exchange information for cells at the edges of subvolumes.

6. Output simulation data if desired.

The initialization of the simulation is carried out on the host. The initialization includes setting the values of the conserved variables for both the real and the ghost cells according to the conditions specified in a text input file. Ghost cells are a buffer of cells added to the boundaries of a simulation volume to calculate fluxes for real cells near the edges. The number of ghost cells reflects the size of the local stencil used to perform fluid reconstruction. Because updating the ghost cells at each time step may require information from cells that are not local in memory, the values of the ghost cells are set on the host before transferring data to the GPU.

Once the simulation volume has been initialized on the CPU, the hydrodynamical calculation begins. The host copies the conserved variable array onto the device. Because the GPU has less memory than the CPU, the conserved variable array associated with a single CPU may be too large to fit into the GPU memory at once. If so, Cholla uses a series of splitting routines described in Section 3.6 to copy smaller pieces of the simulation onto the GPU and carries out the hydrodynamics calculations on each subvolume. At the end of the hydro calculation the next time step is calculated on the device using a GPU-accelerated parallel reduction. The updated conserved variables and new time step are then transferred back to the host. The host updates the values of the ghost cells using the newly calculated values of the real cells, and Steps 2 - 5 repeat until the desired final simulation time is reached.

After each time step the values of the ghost cells are reset using the newly updated values of the conserved variables. Cholla includes three standard boundary conditions: periodic, reflective, and transmissive. These can be set in any combination on any of the borders of the simulation volume. For periodic and transmissive boundaries, the conserved variable values of each ghost cell are copied from the appropriate real cell. For reflective boundaries we follow the same process but reverse the sign of the perpendicular component of momentum. Cholla also includes the capability to define custom boundary conditions, such as the analytic boundaries specified in the Noh Strong Shock test (see Section 4.3.1). In a simulation performed using MPI communication, any necessary boundary regions are exchanged between relevant processes as described in Section 3.7.

3.2. Memory Structure

The data for a simulation in Cholla are contained in two structures. A header stores information about the size and shape of the domain, as well as global variables including the simulation time. A second structure contains the values of the conserved variables for each cell in the simulation. In an object oriented programming model, these values would often be stored in memory as an array of structures,

\[
\text{Cell}[0].\{\rho, \rho u, \rho v, \rho w, E\},
\]

\[
\vdots
\]

\[
\text{Cell}[N-1].\{\rho, \rho u, \rho v, \rho w, E\},
\]

where \( N \) is the total number of cells in the grid. In a CPU-based simulation code, this configuration can improve the performance of memory accesses.

While the object oriented model is intuitive, a different memory structure is more efficient for implementation on the GPU. On a GPU, calculations are done simultaneously by thousands of individual computational ele-
ments called cores, analogous to but individually much less powerful than a typical CPU core. The set of instructions carried out on a single GPU core is called a thread. The efficiency of the GPU comes in part from its ability to efficiently schedule the execution of millions of threads. The scheduling is done by streaming multiprocessors on the device that schedule threads for execution in groups called warps. Each thread warp performs a given set of operations simultaneously in the execution model often referred to as Single Instruction Multiple Data (SIMD). Given that data operations across cores on the GPU are rapidly executed in a massively parallel manner via the SIMD approach, hardware timescales such as the GPU global memory access time can represent a considerable fraction of the total computational expense of a calculation. Techniques to reduce the expense of global memory accesses include the organization of data needed by each thread warp into adjacent regions in physical memory. To facilitate this advantageous data locality, Cholla organizes conserved variables into a structure of arrays:

\[
\{ \rho_0 \ldots \rho_{N-1} \}, \\
\{ p\rho_0 \ldots p\rho_{N-1} \}, \\
\{ \rho u_0 \ldots \rho u_{N-1} \}, \\
\{ p\rho v_0 \ldots p\rho v_{N-1} \}, \\
\{ E_0 \ldots E_{N-1} \}.
\]

The thread warps can retrieve the conserved quantities within this structure of arrays in global memory with unit stride in memory accesses, reducing collisions in the access pattern.

The process of initiating a data transfer from the host to the device involves an associated computational overhead. Limiting the number of transfers required by the algorithm mitigates this overhead, as hardware latency may cause many small transfers to take longer than one large transfer. The allocation of a single structure containing the conserved variable arrays ensures a contiguous data layout in memory, and limits the required data transfers to two (single transfers to and from the device).

3.3. The GPU Grid

Once the simulation volume has been initialized on the host and the values of the ghost cells have been set, the array of conserved variables is transferred to global memory on the GPU. When kernels are then executed on the device, the GPU launches a grid of thread blocks. The GPU grid and thread block dimensions are set by the programmer and are application dependent. Cholla typically uses one or two dimensional grids of one dimensional thread blocks; the latter arrangement is illustrated in Figure [2]. We emphasize that the dimensions of the GPU grid are not constrained to match the dimensions of the simulation, as the location of a cell in the simulation volume can always be mapped to a unique index within the GPU grid of thread blocks.

The dimensions of the GPU grid can affect the efficiency with which the device performs calculations and dictate the mapping from a real-space cell index to a thread index. To define the thread index, CUDA C includes built-in data elements that return specific values for each thread, as shown in the following pseudo-code:

\[
\text{blockId} = \text{blockIdx.x} + \text{blockIdx.y}\cdot\text{gridDim.x} \\
tid = \text{threadIdx.x} + \text{blockId} \cdot \text{blockDim.x}.
\]

Here, blockIdx returns the ID of the thread block within the grid, blockDim returns the dimensions of the grid, threadIdx returns the ID of the thread within the block, and blockDim returns the dimensions of the block. By combining these pre-defined quantities, a unique global index can be calculated for each thread. We use the thread index to assign each thread the work of computing the conserved variable update for a single cell in the simulation. For Cholla, we choose a one dimensional block of threads because most of the kernels are one dimensional in nature. The PPM reconstruction, for example, requires only a one dimensional stencil and is carried out separately for the x, y, and z interfaces. Because a new GPU grid with different dimensions can be initiated every time a device function is called, the thread index calculation and subsequent mapping to a real cell index must be performed within each GPU kernel. No data needs to be transferred back to the CPU between kernels, as all information needed between kernels is stored in the GPU global memory.

3.4. The GPU Kernels

Cholla leverages a modular design that enables an easy selection of the reconstruction method or Riemann solver, and facilitates the incorporation of new features. Each reconstruction method and Riemann solver is performed through an associated kernel executed by the GPU. A routine implementing the CTU algorithm calls these kernels through a wrapper function that segregates CUDA calls from the rest of the code. This organization allows for a flexible compilation structure in which non-CUDA code (including MPI calls) can be compiled with standard C compilers. Within the CUDA wrapper for the CTU algorithm, the following steps are followed:
1. Allocate arrays within GPU global memory to hold the conserved variables $\mathbf{u}$, the initial time-evolved boundary values $\mathbf{U}^*$, the initial one-dimensional fluxes $\mathbf{F}^*$, the transverse-flux-evolved boundary values $\mathbf{U}^{n+\frac{1}{2}}$, and the CTU fluxes $\mathbf{F}^{n+\frac{1}{2}}$.

2. Transfer the conserved variable data from the host to the device and store in the newly allocated arrays in GPU global memory.

3. Call the reconstruction kernel for each dimension.

4. Call the Riemann solver kernel for each dimension.

5. Call the kernel to perform the transverse flux update (Equation 23).

6. Call the Riemann solver kernel for each dimension again.

7. Call the kernel to update the conserved variables and calculate the next time step (Equation 24).

8. Transfer the conserved variable arrays back to the CPU.

Step 1 involves the allocation of memory on the GPU, and it should be noted that the global GPU memory available is typically small compared with the CPU memory. Depending on the device, the simulation size, the number of MPI processes, and the domain decomposition, each process’s conserved variable data may exceed the available GPU memory. An excess may occur even if the local grid governed by each process is small (e.g., $128^3$). When necessary, Cholla uses a set of splitting routines to divide the simulation volume into more manageable subvolumes that are then treated according to the steps listed above. Section 3.6 describes these splitting routines in more detail.

The GPU kernel calls in Steps 3-7 resemble traditional C function calls, but kernels are implemented with additional variables that establish the dimensions of the grid of thread blocks launched by the GPU. For example, the syntax for calling Cholla’s PPM reconstruction function is:

```
PPM_reconstruction<<<BlocksPerGrid,
ThreadsPerBlock>>>(<function_parameters>)),
```

where the triple chevron syntax, `<<<,>>>`, informs the CUDA-enabled compiler that this function should be executed on the GPU device. Since the amount of data processed at once by the GPU is limited by its available global memory, `BlocksPerGrid` can always be set large enough to assign a thread to each cell.

Separate kernels carry out different parts of the CTU algorithm, but each kernel shares common elements. Every kernel must begin with a calculation of the index of each thread, as described in Section 3.3. Using the appropriate mapping, the index of each thread of the kernel can be translated to a unique real-space cell index in the simulation volume. The threads within the kernel then retrieve necessary cell data from the GPU global memory. For the reconstruction function, these data would include the values of the conserved variables for the cell assigned to that thread, as well as those of the nearby cells within the reconstruction stencil. Once the data have been retrieved, the threads carry out any relevant calculations, and load the result into the relevant GPU global memory array. Once all of the threads have finished their calculations the kernel returns, and the process continues through each of the steps listed above.

### 3.5. Time Step Calculation

The implementation of most kernels described in the previous section (reconstruction, Riemann solver, and transverse flux update) follows closely the descriptions of the CTU calculation given in Section 2. However, the final conserved variable update kernel in each iteration of the algorithm is extended to include the calculation of the next simulation time step $\Delta t^{n+1}$ via a parallel reduction operation performed on the GPU. A brief explanation of this parallel reduction operation follows.

Thread blocks on the GPU have a limited amount of “shared memory” that each thread in the block can access rapidly, as illustrated in Figure 4. At the end of the conserved variable computation, the updated conserved variable data for each cell are stored in the private register memory assigned to each thread. The updated values of the conserved variables are used by each thread to calculate the minimum time step associated with its cell, according to Equation 22. Each thread then loads the time step it calculated into an array the size of the thread block in shared memory - note that each thread block has its own array. The threads in the block then perform a tree-based parallel reduction on the time step array, finding the minimum time step for the entire block. This minimum value is uploaded into an array in the GPU global memory, and is then passed back to the CPU. The final reduction is performed on the CPU, looping through the array of calculated time steps. This array only has `BlocksPerGrid` elements, reducing the required calculations on the CPU for the time step computation approximately by the ratio of the number of cells in the computational volume to `BlocksPerGrid`.

Calculating the time step on the GPU achieves a performance gain relative to a CPU since executing a large number of floating point operations is extremely efficient on the GPU. The shared memory reduction on the GPU reduces the number of loops needed on the CPU by a factor of `ThreadsPerBlock` (typically set to 128 for an NVIDIA Kepler K20X GPU). For reference, we find the parallel GPU reduction time step calculation for a $1920 \times 1080$ simulation can achieve a $100 \times$ performance gain relative to a single CPU core depending on the architecture ($\sim 1$ms vs. $\sim 100$ms).

### 3.6. Subgrid Splitting

As mentioned in previous sections, the total amount of memory on a single device may be quite limited when compared to the memory available on the host. At present, a typical GPU has only a few gigabytes of global memory, though this number has been increasing with each new generation of devices. Current devices can typically only hold the information for a 3D hydrodynamical simulation of size $\sim 128^3$, including conserved variables and fluxes. In Cholla, slightly more cells can fit for 1D and 2D simulations owing to the reduced number of transverse interface states and fluxes. Often, the limited
memory resources mean that the simulation volume associated with a local process may need to be successively subdivided to fit on a GPU. We term this subdivision process “subgrid splitting”.

In practice, subgrid splitting is typically only needed for multidimensional simulations or 1D simulations with millions of cells. A description of the 1D subgrid splitting is provided below as a straightforward example. First, the size of the simulation volume that will fit on the GPU at once given the global memory available is calculated and stored in a variable, e.g. \( \text{MaxVol} \). The local volume governed by each local process is further split into subvolumes of size less than or equal to \( \text{MaxVol} \). We refer to these subvolumes as “subgrid blocks”. The CTU calculations for each subgrid block are performed on the GPU sequentially, including any necessary ghost cells from nearby subvolumes. Memory buffers on the CPU are used to avoid overwriting grid cells that act as ghost cells for neighboring subgrid blocks. The procedure of copying data to the GPU, calculating, and transferring data back to the CPU is repeated until the hydro step for the entire simulation volume has completed. Because the conserved variables for the simulation are contiguous in memory on the host, copying them into buffers via \text{memcpy} contributes a negligible amount to the total simulation run time.

To illustrate the subgrid blocking method, Figure 5 displays a two-dimensional grid with an example subgrid blocking by a factor of four. Each of the four subgrid blocks (red, green, blue, purple) require real cells from adjacent subgrid blocks to act as subgrid ghost cells to form the full computational stencil for the fluid reconstruction and CTU calculations (indicated by colored dashed lines). Since these subgrid blocks also abut either local simulation boundaries between local processes or global boundaries at the edges of the illustrated region, they also require standard ghost cells (gray regions) to complete their computational stencils. The subgrid block regions outlined by the dashed lines are transferred sequentially to the GPU for the CTU calculation, taking care to preserve in memory the subgrid ghost cell boundary regions between subgrid blocks until the entire local volume has been processed. The standard ghost cells are updated after the CTU calculation, since they depend either on communication between MPI processes or the global boundary conditions of the simulation. Note that Figure 5 illustrates a very small grid for convenience, and the actual subgrid regions of a 2D simulation would be orders of magnitude larger.

We performed a variety of tests using subgrid blocks of different shapes in order to determine a method of division that helps minimize GPU communication overhead. For 2D simulations, splitting the volume into approximately square regions works well. For 3D simulations, we find that maintaining an aspect ratio that is approximately even in the \( y \) and \( z \) directions with a longer aspect ratio in the \( x \)-direction works well. In practice, we keep the aspect ratio even in \( y \) and \( z \) while maintaining an aspect ratio in \( x \) that is roughly 5 times the geometric mean of the \( y \)- and \( z \)-aspect ratios. Memory access overheads can be reduced by first copying multiple subgrid blocks into buffers on the CPU, and then transferring subarrays containing individual subgrid blocks to the GPU for computation. Even in simulations where local volumes must be subdivided into subgrid blocks dozens of times the overhead associated with copying the conserved variables into buffers on the CPU is insignificant, typically limited to 5% of the total time taken for the CTU calculation.

3.7. MPI Implementation and Scaling

The massively parallel algorithm implemented by \textit{Cholla} can be adapted to execute on multiple GPUs simultaneously. \textit{Cholla} can thereby gain a multiplex advantage beyond the significant computation power afforded by a single GPU. The parallelization is implemented using the MPI library. The global simulation volume is decomposed into subvolumes, and the subvolumes are each assigned a single MPI process. In \textit{Cholla}, each MPI process runs on a single CPU that has a single associated GPU, such that the number of MPI processes, CPUs, and GPUs are always equal. When the simulation volume is initialized, each process is assigned its simulation subvolume and surrounding ghost cells. Since the hydrodynamical calculation for every cell is localized to a finite stencil, only the ghost cells on the boundary of the volume may require updating from other processes via MPI communication every timestep. Compared with a simulation done on a single CPU/GPU pair, additional overheads for a multi-process simulation can therefore include MPI communications needed to exchange information at boundaries and potential inefficiencies in the GPU computation introduced by the domain decomposition. While domain decomposition influences communi-
cations overheads in all MPI-parallelized codes by changing the surface area-to-volume ratio of computational subvolumes, domain decomposition additionally affects the performance of a GPU-accelerated code by changing the ratio of ghost to real cells in memory that must be transferred to the GPU. Since memory transfers to the GPU are a significant overhead, often of order 30% of the entire time taken to complete the hydro module on the GPU, domain decompositions that limit the required amount of ghost cells on a local process are favorable. Cholla therefore allows for two different domain decompositions, described below.

3.7.1. Slab Decomposition

Following the domain decomposition utilized by the Fastest Fourier Transform in the West discrete Fourier transform library (FFTW; Frigo & Johnson 2005), Cholla can use a slab-based decomposition in which the simulation volume is sliced only in one dimension. In the slab decomposition a maximum of two boundaries may be shared between processes, and because there are limited communications the slab decomposition proves efficient for simulations run with a small number of processes. With the addition of more processes the slabs grow narrower, the ratio of boundary ghost cells to real cells for each subvolume increases rapidly, and the time required to exchange boundary cells between processes remains nearly constant. Although these features cause a computational inefficiency that continues to degrade with increasing numbers of processes, Cholla nonetheless includes an optional slab decomposition for use with limited processes and in conjunction with FFTW.

The division of the simulation volume for Cholla’s slab decomposition is straightforward. When the FFTW library slab decomposition is used, the slab width on each process is optimized for accelerating discrete Fourier transform computations. Otherwise, the number of cells in the \( x \)-dimension spanning the total simulation volume is divided evenly across the number of processes, and any remaining cells are split as evenly among the processes as possible. Once the domains have been assigned, each process initializes the real cells associated with its volume and exchanges boundary cells. First, each process posts a receive request for each MPI boundary. If the process has a global simulation boundary along the \( x \)-direction, it posts either one receive request in the case of reflective, transmissive, or analytic global boundary conditions, or two in the case of global periodic boundary conditions. Processes that are surrounded by other processes will always have two MPI boundaries. The processes then send the initialized values of the real cells from their subvolume that are needed by other processes. While waiting for the cell exchange communications to complete, each process computes the cell values on its non-MPI boundaries (typically the \( y \) - and \( z \)-boundaries). This asynchronous ordering of communication and boundary computation minimizes the amount of time the CPU must sit idle while waiting to receive boundary cell information. Once all the receives have completed, each process proceeds through the CTU step as though it were an independent simulation volume. At the end of each time step, boundary cells must again be exchanged along with the information from each local subvolume required to determine the global simulation time step.

3.7.2. Block Decomposition

To improve upon the computational inefficiencies of the slab decomposition, Cholla includes a block decomposition that seeks to minimize the aspect ratios of the simulation subvolume evolved by each process. For a block decomposition, up to six MPI communications for cell exchanges may be required per timestep. Despite this increased number of communications, the reduction of the surface area-to-volume ratio of the block decomposition improves its efficiency beyond that achieved by a slab decomposition for large numbers of processes.

In the case of a block decomposition the \( x \)-, \( y \)-, and \( z \)-dimensions for each subvolume are kept as even as possible. Once a simulation volume is appropriately divided and local real cells initialized by each process, boundary cells between subvolumes must be exchanged. To keep the number of MPI communications to a maximum of six, the processes exchange boundary cells in a specific order. First, each process posts a receive request for any MPI boundaries on an \( x \)-face. While waiting for those data to arrive, the processes set \( x \)-ghost cells on any non-MPI \( x \) faces. Once the \( x \)-boundaries arrive, the processes post receive requests for MPI boundaries on \( y \)-faces, including the corner regions just transferred in the exchange along the \( x \)-direction. While waiting for the \( y \)-boundaries to arrive, each process computes the ghost cell values along non-MPI \( y \)-boundaries. By first sending the \( x \)-boundaries, processes can receive information needed for the \( y \)-boundary exchange from diagonally-adjacent processes without directly exchanging information with those processes. The same procedure is followed for the \( z \)-boundaries.

Figure 6 illustrates this process for a 2D simulation with periodic boundaries and a four-process decomposition. Each process first initializes its real cells, represented by the large colored squares. To perform the hydrodynamical simulation timestep in parallel across separate processes, each process must receive boundary ghost cells from the real cells hosted by surrounding processes. The boundary regions for each process are outlined in Figure 6 and labelled 1 and 2. The processes first exchange \( x \)-boundary information (region 1) via two MPI communications. Once the \( x \)-boundary exchange is complete, \( y \)-boundary information (region 2) is exchanged, including the corner regions received from other processes. The same procedure is repeated for \( z \)-boundaries in a 3D simulation. Following this pattern keeps the required number of MPI communications to a maximum of six, instead of the potential twenty-six communications that would be required to separately exchange each face, edge, and corner boundary region for an interior subvolume in a 3D simulation.

We note briefly that the block decomposition implemented in Cholla may also be adapted to enable the use of Fast Fourier Transform libraries that use a block decomposition, such as the Parallel FFT package written by Steve Plimpton.

3.7.3. Scaling

The scalability of the Cholla MPI implementation to more than one GPU warrants a discussion. To study the

\[ \text{http://www.sandia.gov/~sjplimp/docs/fft/README.html} \]
scaling of the code, the GPU-accelerated International Business Machines iDataplex cluster El Gato at the University of Arizona was used. Using El Gato, we have tested both the strong and weak scaling of Cholla using up to 64 GPUs (∼184,000 GPU cores). The results are shown in Figures 7 and 8. For both scaling tests, a three-dimensional sound wave perturbation problem with periodic boundary conditions and a block decomposition is used to maximize the number of MPI communications required per timestep. In both the strong and weak scaling tests, Cholla updates an average of $6.3 \times 10^6$ cells per second using the NVIDIA Kepler K20X GPUs available on El Gato. The 3D sound wave perturbation requires work to be done by each cell and uses third-order spatial reconstruction and an exact Riemann solver. The test is therefore relatively inefficient. By contrast, the 2D Kelvin-Helmholtz test described in Section 4 Cholla updates an average of $1.2 \times 10^7$ cells per second.

For the strong scaling test, a $512^3$ grid is evolved for 10 time steps. The timing results for the total test runtime, the CTU algorithm (performed on the GPU), and the boundary computation including ghost cell exchange communication are tracked separately. We exclude the simulation initialization from the runtime, as it comprises a significant fraction of the runtime for these short tests and obscures the results. As Figure 7 shows, the overall scaling of Cholla is close to ideal, with the CTU step scaling slightly better than ideal. That the boundary condition computation does not scale as well primarily owes to the reduced number of MPI communications needed in runs with small numbers of processes compared with tests utilizing large numbers of processes where every subvolume boundary requires an MPI communication per timestep. The scaling between an 8 process run and a 64 process run, both of which require MPI communications for all boundaries, is close to ideal. All of the GPU calculations contained within the CTU step scale better than ideal at 64 processes. The increased efficiency at 64 processes owes to the decomposition decreasing the cells per process below the number that necessitates subgrid splitting, thereby reducing the CPU-GPU communications overhead. The dip at 14 processes reflects an unavoidable inefficient $2 \times 7$ process domain decomposition.

The weak scaling performance of Cholla is shown in Figure 8. A sound wave perturbation test is again used, but in this test the size of the total computational volume is rescaled with the number of processes to keep the number of cells per process approximately constant at $\approx 322^3$ (note that each process uses its own distinct GPU during the simulation). The total runtime efficiency as a function of the number of processes remains roughly constant beyond a single process. The CTU algorithm comprises the majority of the computational cost of each timestep, and exhibits nearly perfect weak scaling. The boundary condition calculation for the serial case does not involve MPI communications and is correspondingly inexpensive when using a single process. With two or more processes, MPI communications induce an additional overhead beyond the single process case. However, the weak scaling of the boundary conditions is reasonably maintained to 64 processes.
Fig. 8.— Weak scaling performance of Cholla. A sound wave perturbation test is used, and the total number of cells calculated by each process is scaled to maintain \( \approx 322^3 \) cell for a single process. The total runtime (blue points) remains roughly constant up to 64 processes, as does the time taken for the GPU portion of the code (red points).

4. TESTS

A large variety of hydrodynamics tests exist in the literature, some of which have been used for several decades (e.g. Sod 1978). Their ubiquity makes these canonical tests an excellent way to compare the performance of Cholla with other codes. In addition, many tests have been designed to explicitly show the failings of hydrodynamics solvers in various environments or highlight the circumstances where they perform exceptionally well. In choosing the tests shown below, we attempt to demonstrate the breadth of problems Cholla can simulate. We also demonstrate the effects of changing reconstruction methods or Riemann solvers, and show differences in the outcomes of tests where they are relevant. If not otherwise specified, the following tests were performed using piecewise parabolic reconstruction with the characteristic variables (PPMC) and an exact Riemann solver.

Before delving into the specifics of each test, we make a note about the convergence rate of Cholla. Many shock-capturing methods revert to first order at shocks (see, e.g. Laney 1998), so to test the convergence rate of Cholla the smooth perturbation test described in Stone et al. (2008) is employed. Both the PPMP and PPMC implementations in Cholla demonstrate second-order convergence in the L1 error norm out to grid resolutions of 1024 cells.

4.1. 1D Hydrodynamics

4.1.1. Sod Shock Tube

The Sod problem (Sod 1978) is often the first test performed by hydrodynamics codes, and we do not diverge from precedent here. Though the Sod problem is not a difficult test to run, the solution contains several im-

Fig. 9.— The solution to the Sod shock tube test using PPMP with a resolution of 100 cells. The exact solution is shown as a line with points from the Cholla simulation over plotted. Features seen in the density plot include a rarefaction wave expanding from the initial discontinuity at \( x = 0.5 \), a rightward moving contact discontinuity at \( x \approx 0.7 \), and a rightward moving shock at \( x \approx 0.85 \).
Fig. 10.— The solution to the Sod shock tube test using PPMC with a resolution of 100 cells. The exact solution is shown as a line with points from the Cholla simulation over plotted. The same fluid features are seen as in Figure 9, with the contact discontinuity slightly less narrowly resolved.
The strong rarefaction test, or 123 problem, was originally used by Einfeldt et al. [1991] to illustrate a scenario that causes a subset of approximate Riemann solvers to fail. Because the solution contains a region where the energy is largely kinetic and the pressure is close to vacuum, the Roe solver (or any other linearized solver) will produce negative densities or pressures in the numerical solution (Einfeldt et al. [1991]). The initial conditions consist of a fluid with constant density and pressure but opposite receding velocity at the center. Specifically, we set \( \rho_L = \rho_R = 1.0, P_L = P_R = 0.4, u_L = 2.0, \) and \( u_R = 2.0. \) In Figure 12 we show the results of this test on a grid of 128 cells at time \( t = 0.15 \) using PPMP reconstruction and the exact solver. This test is not a challenge using the exact solver, but without modification the Roe solver would fail on this problem. For this reason, we test the density and pressure produced in the solution by the Roe solver and revert to the HLLE solver if necessary. With that fix we can run the problem with either solver, and in fact the HLLE fluxes are only needed on the first step of the simulation.

4.1.4. Shu and Osher Shocktube

The Shu-Osher shocktube test shows the tendency of PPM to cut off maxima in smoothly varying problems as a result of the slope limiters imposed in the reconstruction method (Shu & Osher [1989]). The test consists of a strong shockwave propagating into a region with a sinusoidally varying density. The initial conditions are \( \rho_L = 3.857143, u_L = 2.629369, \) and \( p_L = 10.3333; \) \( \rho_R = 1 + 0.2 \sin(3\pi x), u_R = 0, \) and \( p_R = 1.0. \) We run the problem on the domain \( x = [-1, 1] \) with the initial discontinuity at \( x = -0.8. \) The results of the test using both 200 cells and 800 cells are shown in Figure 13. As can be seen, the low resolution solution does lose some of the amplitude of the peaks. Using newer versions of the limiting functions can help alleviate this problem (Colella & Sekora [2008] Stone et al. [2008]), although we have not yet implemented these limiters in Cholla.

4.1.5. Interacting Blast Waves

Originally described in Colella & Woodward [1984], the interacting blast wave test helps quantify the behavior of a code near strong shocks and contact discontinuities. The test consists of a medium with initially constant density \( p = 1.0, \) with \( \gamma = 1.4 \) on the domain \( x = [0, 1]. \) Reflecting boundary conditions are used. Two shocks are initialized on either side of the domain, with \( p = 1000 \) for \( x < 0.1, p = 100 \) for \( x > 0.9, \) and \( p = 0.01 \) in between. The problem is run until time \( t = 0.038, \) at which point the shocks and rarefactions in the initial solution have interacted multiple times.

We show plots of the density computed with both PPMP and PPMC in Figure 14. We plot a low resolution solution with 400 grid cells over a high resolution reference solution computed with 9600 cells. As can be seen in the figure, PPMP does an excellent job keeping the contact discontinuities at \( x = 0.6 \) and \( x = 0.8 \) contained within just two zones, as compared to the solution computed with PPMC in which the contacts are smeared over many cells. In addition, PPMC tends to more severely cut off the maximum at \( x = 0.75, \) while PPMP does a decent job of keeping the full height although the peak is slightly offset. Both reconstruction...
4.2.1. Implosion Test

The implosion test is a converging shock problem first presented in Hui et al. (1999). The version presented here is described in Liska & Wendroff (2003), and begins with a square region of high density and pressure containing a diamond-shaped region of low density and pressure. These initial conditions evoke the traditional Sod shock tube problem extended to two dimensions, but inclined to the grid by 45 degrees rather than aligned as in the one dimensional case. As the test begins material moves inward rapidly toward the center, leading to an implosion. When run for a short amount of time, this test demonstrates the ability of a code to resolve contact discontinuities and other fluid features for a non-grid aligned shock tube. When run for enough time to evolve well past the initial shock tube solution, the test illustrates the symmetry (or lack thereof) of a code.

Figure 15 shows the results of the implosion test run with PPMC and an exact solver at an early time $t = 0.045$ and a later time $t = 2.5$. The problem was run on a $400 \times 400$ grid with a domain $x = [0, 0.3]$, $y = [0, 0.3]$ and reflecting boundary conditions at every boundary, comprising the upper right quadrant of the axisymmetric test described above. The initial density and pressure within the diamond-shaped region are $\rho = 0.125$ and $p = 0.14$, while outside the density and pressure are $\rho = 1.0$ and $p = 1.0$. Initial velocities are zero, as in the Sod test. A discontinuous interface is located along the diagonal running from $(0.15, 0)$ to $(0, 0.15)$. In the upper panel of Figure 15 a rarefaction fan can be seen expanding outward from this interface. As the upper panel shows, Cholla does an excellent job resolving the contact at early
Fig. 14.— Numerical solution for the interacting blast wave test using PPMP (top) and PPMC (bottom) with a resolution of 400 cells, plotted over a reference solution with 9600 cells. This solution is shown at $t = 0.038$, when the original shocks and rarefactions have interacted several times. The test was designed to capture a code’s ability to maintain narrow features.

At the later time a jet has appeared in the solution. The production of the jet is a direct result of ability to preserve symmetry in the Cholla solution to numerical accuracy. Liska & Wendroff (2003) demonstrated that codes that employ non-symmetry preserving methods like Strang splitting may fail to produce the jet-like feature. The fact that this test is so sensitive to the symmetry of the problem makes it useful for diagnosing potential coding errors, but the test also demonstrates the extent to which a non-symmetric algorithm can impact the physical accuracy of the result. As this test

Fig. 15.— Numerical solutions for an implosion test. Top: The $400 \times 400$ implosion test at $t = 0.045$; 31 density contours from 0.35 to 1.1 are overlaid on a color-scale pressure map, with only the inner region of the computational domain shown, $x = [0, 0.22]$ and $y = [0, 0.22]$. Middle: The same test at $t = 2.5$; 36 density contours from 0.125 to 1 are overlaid on a color-scale pressure map. Bottom: A $4096 \times 4096$ version of the implosion test at $t = 2.5$. The same contour levels are drawn.
shows, relatively large-scale features in the solution can be completely lost if a code fails to maintain a sufficient level of symmetric accuracy.

The bottom panel of Figure 15 shows the results of this test recomputed at a much higher resolution of 4096 × 4096. The same large-scale features are apparent in the solution, but as expected the small-scale density perturbations and shape of the jet have clearly not converged. However, this high resolution test serves as further evidence of the ability of Cholla to preserve axisymmetry even in a very difficult problem. At this extreme resolution, the code must perform over 200,000 time steps and more than $3 \times 10^{12}$ cell updates to reach time $t = 2.5$. At that point, the results are still exactly symmetric (to floating-point precision), demonstrating that symmetry preservation in Cholla is a robust feature of the code.

4.2.2. Explosion Test

The explosion test, also from Liska & Wendroff (2003), is designed to test the evolution of an unstable contact discontinuity and is highly sensitive to numerical noise in the initial conditions. This noise seeds an instability that grows as the solution evolves. The test starts with a domain $x = [0, 1.5], y = [0, 1.5]$ that contains a circularly symmetric region of high density and pressure, with $\rho = 1$ and $p = 1$ inside a circle with radius $r = 0.4$. Reflecting inner boundaries and transmissive outer boundaries are used. Outside the circle the density and pressure are set to $\rho = 0.125$ and $p = 0.1$. The initial velocities are zero. Because the problem is sensitive to initial perturbations at the interface, the density and pressure for cells are area-weighted at the boundary. For each cell on the boundary of the circle the percentage of the area inside the radial boundary is computed, and the initial cell data weighted appropriately.

The test problem is performed on a grid of $400 \times 400$ cells, and Figure 16 shows the result of the calculation using PLMP and PPMP at $t = 3.2$. As expected, the higher order reconstruction method does a better job preserving the narrow structure of the contact, but is also more susceptible to structure along the interface as the instability develops. Thus, as the problem progresses, the lower order more diffusive method may result in a cleaner solution. We note that both methods preserve the exact symmetry of the problem, provided the initial conditions are symmetric.

4.2.3. Kelvin-Helmholtz Instability

A Kelvin-Helmholtz instability test demonstrates the extent to which a hydrodynamics code resolves mixing caused by shear flows. In this test, two fluids at different densities flow past each other and characteristic eddies appear and grow at the interface between the fluids. The growth of the eddies in the linear regime can be analytically described (Chandrasekhar 1961) and depends on properties of the fluid and the interface itself. At a discontinuous interface, small eddies will develop first at the grid scale of the simulation, and these will gradually grow and combine into larger eddies as shown in Figure 17.

The exact nature of the instability depends sensitively on the resolution and initial conditions of the test (e.g., Robertson et al. 2010). The test shown in Figure 17 was run on a $1920 \times 1080$ grid, with a domain $x = [0, 1], y = [0, 0.5625]$ in order to maintain square cells. The simulation initial conditions include a dense fluid with density $\rho = 2.0$ in the middle third of the box, surrounded by a less dense fluid with density $\rho = 1.0$ in the outer thirds. The denser fluid has a velocity $u = -0.5$, and the less dense fluid has a velocity $u = 0.5$; the $y$-velocities are initially $v = 0$. The entire simulation volume is initialized in pressure equilibrium with $p = 2.5$. A small-amplitude perturbation is added to the $x$- and $y$-velocities of every cell in the grid, in proportion to the $x$-position following $u = u + 0.01\sin(2\pi x)$ and $v = v + 0.01\sin(2\pi x)$. The simulation is evolved to $t = 1.0$, by which time the growth of the eddies has entered the non-linear regime.

As the eddies grow, more mixing between the high density and low density material occurs. Resolving this mixing is an important task for a hydrodynamics code, as the amount of mixing can have a significant impact on broad features in the simulation outcome. The level of mixing tends to increase with resolution as well.
as with higher order reconstruction techniques. Therefore, Kelvin-Helmholtz instabilities highlight the importance of having an efficient high order reconstruction method and a fast code. As expected for a high resolution grid code with a high order reconstruction method, Cholla does an excellent job of resolving the shear mixing.

4.3. 3D Hydrodynamics

4.3.1. Noh’s Strong Shock

The Noh strong shock test, originally described in one dimension by Noh (1987), demonstrates how well a code can track a strong, high mach number shock. This test is considered difficult to perform in either two or three dimensions, as many hydrodynamics codes cannot run the test accurately and some fail completely (Liska & Wendroff 2003). The test starts with a constant density of $\rho_0 = 1.0$ throughout the grid, with zero pressure and constant velocity $|V| = 1.0$ toward the origin. For this test, the adiabatic index is set to $\gamma = \frac{5}{3}$. These initial conditions result in a formally infinite strength shock reflecting outward from the origin with spherical symmetry. Cholla cannot be run with zero pressure, so we set the initial pressure to a low number, $p_0 = 10^{-6}$, but we note that the results are relatively insensitive to the initial pressures below $p_0 \sim 10^{-3}$.

The Noh test is initialized in an octant on the domain $[0, 1]$ with reflecting inner boundaries. The outer boundaries are set according to the analytic solution for the density and energy, which in two or three dimensions is

$$\rho(t) = \rho_0 \left(1 + \frac{t}{r} \right)^{n-1},$$

where $r$ is the radius in polar or spherical coordinates, and $n$ is the dimensionality of the problem. The momentum follows from the velocity and the solution for the density, and the total energy is set to

$$E(t) = \frac{\rho_0}{\gamma - 1} + 0.5 \rho(t).$$
We evolve the solution to \( t = 2.0 \), by which time the shock has propagated through more than half of the computational domain. The density immediately in front of the shock as well as the density of the post shock gas can also be calculated analytically. In the 3D case, the gas immediately before the shock has a density of \( \rho = 16 \), and the post-shock gas has a corresponding density of \( \rho = 64 \).

Running the Noh test on a Cartesian grid creates strong, grid-aligned shocks that provoke a behavior in the numerical solution known as the carbuncle instability. The carbuncle instability arises as a result of oscillatory crossflow solutions to the Riemann problem near such shocks (Quirk 1994). This problem is addressed by implementing a form of the H correction, as described in Sanders et al. (1998) and detailed in Appendix C. By incorporating information about the fastest transverse wave speeds, the H correction adds dissipation to the 1D fluxes calculated by the Roe Riemann solver that reduces the carbuncle strength.

The result of the Noh test using PPMC with and without the H correction can be seen in Figure 13. Without the H correction, the solution suffers from strong oscillatory behavior, particularly along the edges where the shock is aligned with the grid. In the version with the H correction applied, the unstable behavior along the axes is effectively absent. The region near the origin where the density dips down is not related to the carbuncle instability, and is a density error known as “wall heating”. This feature that the 1D Noh test was originally designed to demonstrate. The slight noise along the shock front is a result of the strength of the shock, and is similar to the minor oscillations seen in the 1D strong shock test. We note that implementing the H correction increases the stencil required for CTU. Because the current version of Cholla is designed to accommodate a maximum of four ghost cells, we currently implement the H correction only with PPMC.

5. CONCLUSIONS

In this work we have presented Cholla, a new, massively-parallel, three-dimensional hydrodynamics code optimized for Graphics Processor Units (GPUs). Cholla uses the unsplit Corner Transport Upwind algorithm (Colella 1990; Gardiner & Stone 2008), multiple Riemann solvers, and a variety of reconstruction methods to model numerical solutions to the Euler equations on a static mesh.

In writing the code, we have maintained a modular structure that allows for the implementation of different hydrodynamical schemes. Cholla features five methods for interface reconstruction, including the first-order piecewise constant method, second-order linear reconstruction methods, and two third-order methods based on the original piecewise parabolic method developed by Colella & Woodward (1984). There are multiple Riemann solvers, including the exact solver from Toro (2009) and a linear solver based on the method of Roe (1981). Incorporating multiple reconstruction and Riemann solver methods provides the ability to test results for a dependence on the particular numerical techniques used, and the strengths and weaknesses of the different methods are discussed. Cholla also implements an optional diffusive correction called the H correction (Sanders et al. 1998) to suppress instabilities along grid-aligned shocks. The H correction adds additional diffusion to the Roe fluxes based on the fastest transverse velocities. The Appendices of this paper detail all of the equations used in the code, and supplement the discussion of each method presented in the main text.

The strategies employed in designing Cholla to run natively on GPUs are extensively detailed. The necessity of transferring data to and from the GPU with every time step requires a specific memory layout to improve efficiency. Once information has been transferred to the GPU, the CTU integration algorithm can be effectively divided into kernel functions that execute on the device, similar to functions in a traditional CPU code. Each of these kernels is self-contained and contributes to the modularity of the code. Because the GPU has limited global memory storage, the simulation volume must often be subdivided to optimize performance and compensate for GPU memory constraints. The strategy we employ to execute this “subgrid splitting” efficiently is also presented.

The architectural differences between CPUs and massively-parallel GPUs require the elucidation of several key concepts in GPU programming. When a device kernel is called, the GPU launches a large set of individual computational elements called threads. A single kernel call can launch millions of threads. In Cholla, each thread is assigned the work of computing the updated hydrodynamic conserved variables for a single real cell in the simulation volume. The streaming multiprocessors on the GPUs handle the work of assigning threads to GPU cores, which means that Cholla will be easily transportable to newer generations of GPU hardware or other coprocessors. Thousands of cores operating simultaneously on each device results in thousands of simultaneous cell updates, making the hydrodynamics solver in Cholla very fast. GPUs are designed to optimize for throughput as opposed to latency. As demonstrated in the GPU-based time step calculation presented in Section 3.5 and adding additional floating point calculations to GPU kernels is relatively inexpensive. This feature can be exploited to incorporate more physics on the GPU, such as cooling, at relatively little computational cost.

The scalability of Cholla and its performance on a range of hydrodynamics problems was also documented. Using the Message Passing Interface (MPI) library (Ferry 1994), Cholla can be run across many GPUs in parallel. The code incorporates both slab-based and block-based domain decompositions and exhibits excellent strong and weak scaling in block-decomposed tests beyond 64 GPUs and 100,000 GPU cores. We present the results of a suite of canonical hydrodynamics tests in one, two, and three dimensions. The state-of-the-art art hydrodynamics algorithms employed enables Cholla to perform accurately on a wide variety of tests, and the GPU architecture makes the computations fast without sacrificing physical accuracy. Since a single GPU can compute a simulation with nontrivial resolution, using Cholla with a cluster presents the option of running many problems to explore large parameter spaces rapidly. Further, the excellent weak scaling of Cholla suggests that very large problem sizes can be tackled on large GPU-enabled supercomputers.

Lastly, we note that we have designed Cholla to per-
form almost all of the hydrodynamics calculations on the 
GPU, freeing the CPU for other tasks (see e.g. Figure 3). By performing calculations on the CPU and GPU 
simultaneously, additional physics could be modeled on 
the GPU. Logical extensions to Cholla include using Fourier 
transforms to solve gravity or drive turbulence. The ad-
dition of a magnetohydrodynamics module on the GPU
is also an attractive possibility, as Cholla uses an unsplit 
integration algorithm that is optimized for MHD [Gar-
diner & Stone 2008].

We are grateful to Jordan Stone for creating the 
acronym Cholla. This material is based upon work sup-
ported by the National Science Foundation Graduate Re-
search Fellowship under Grant No. DGE-1143953, as well 
as a National Science Foundation Grant No. 1228509.

APPENDIX

A. RECONSTRUCTION METHODS

To calculate the input states for the Riemann solvers used in Cholla, appropriate values of the conserved variables at 
each interface must be reconstructed using the cell-averaged quantities. Each Riemann problem requires input states at 
either side of a cell interface, referred to as \( U_L^* \) and \( U_R^* \) in the context of the CTU algorithm presented in Section 2.1. 
While previously \( L \) and \( R \) indicated the left and right of the interface, in this Appendix a cell-centered labeling is used to 
document the procedure for computing the input states at the left and right boundaries of a single cell.

The input states calculated at the left and right sides of the cell will be labeled \( U_L^* \) and \( U_R^* \) in the conserved variables, or \( W_L^* \) and \( W_R^* \) in the primitive variables. As described in Table 1, the asterisk indicates the time-evolved input state. The boundary values reconstructed before time evolution will be labeled \( W_L \) and \( W_R \). For each method described below, only the steps involved in the reconstruction for the \( x \)-interfaces are shown. The \( y \)- and \( z \)-reconstructions proceed in the same manner but with an appropriate change of stencil. The notation will drop the \( i \), \( j \), and \( k \) subscripts unless they are needed for clarification.

A.1. PLMP

The simplest practical reconstruction method implemented in Cholla is PLMP, a piecewise linear reconstruction with slope limiting applied in the primitive variables. The stencil to calculate the boundary values for cell \( i \) contains cells \( i - 1 \) to the left and \( i + 1 \) to the right. The first step in the method converts the cell-averaged values of the 
conserved variables into the primitive variables, \( w = [\rho, u, v, w, p]^T \). The cell-averaged primitive values are then used to reconstruct boundary values in the primitive variables, \( W_L \) and \( W_R \) at the left and right sides of cell \( i \) using a local, piece-wise linear reconstruction [Toro 2009]:

\[
W_L = w_i - \frac{1}{2} \delta w_i, \\
W_R = w_i + \frac{1}{2} \delta w_i,
\]

where \( \delta w_i \) is a vector containing the slope of each primitive variable across cell \( i \). To compute \( \delta w_i \), we first calculate the left, right, and centered differences in the primitive variables across each of the cell interfaces:

\[
\delta w_L = w_i - w_{i-1}, \quad \delta w_R = w_{i+1} - w_i, \quad \delta w_C = 0.5(w_{i+1} - w_{i-1}).
\]

A monotonized central-difference limiter [van Leer 1977] is then used to compute \( \delta w_i \):

\[
\delta w_i = \begin{cases} 
\text{sgn} (\delta w_C) \min(|\delta w_C|, 2|\delta w_L|, 2|\delta w_R|), & \delta w_L \delta w_R > 0 \\
0, & \text{otherwise},
\end{cases}
\]

where \( \text{sgn} \) is defined as

\[
\text{sgn}(x) = \begin{cases} 
-1, & x < 0 \\
1, & \text{otherwise}.
\end{cases}
\]

Each of the primitive variables is treated independently in the limiting process, so the vector of primitive variable 
slopes (for extrapolations to the left cell face) can be simply written as

\[
\delta w_L = \{\delta \rho_L, \delta u_L, \delta v_L, \delta w_L, \delta p_L\}^T.
\]

The primitive variable slopes for extrapolating to the right cell face can be similarly defined.

The last step in computing input states for the Riemann problem is to evolve the reconstructed boundary values by 
half a time step \( \Delta t/2 \). The time evolution is modeled using the conserved form of the Euler equations, and \( W_L \) and 
\( W_R \) are therefore converted back into conserved variables, \( U_L \) and \( U_R \) and used to calculate the associated fluxes via 
Equation 11. These fluxes are used to evolve the reconstructed boundary values and obtain input states appropriate 
for the Riemann problem:

\[
U_i^* = U_L + 0.5 \frac{\Delta t}{\Delta x} [F(U_L) - F(U_R)]
\]
\[ \mathbf{U}_R^+ = \mathbf{U}_R + 0.5 \frac{\Delta t}{\Delta x} [\mathbf{F}(\mathbf{U}_L) - \mathbf{F}(\mathbf{U}_R)]. \] (A7)

### A.2. PLMC

The second reconstruction method, PLMC, is also based on a piecewise linear reconstruction but with the slope limiting computed using the characteristic variables. The stencil again contains one cell to the left and right of cell \( i \). After converting the cell-averaged quantities from conserved to primitive variables, an eigenvector decomposition of the Euler equations is performed using the characteristic variables as described in Section 2. First, the eigenvalues of the linear system of equations for cell \( i \) are calculated. For adiabatic hydrodynamics, the eigenvalues correspond to the three wave speeds,

\[ \lambda^m = u_i - a_i, \quad \lambda^0 = u_i, \quad \lambda^p = u_i + a_i, \] (A8)

where \( a_i \) is the average sound speed in cell \( i \). The quantities \( \lambda^m \) and \( \lambda^p \) are speeds of the acoustic waves and \( \lambda^0 \) is the speed of the contact wave. The corresponding eigenvalue for any advected scalar quantity (such as the transverse velocities in multidimensional problems) is simply the speed of the fluid in the normal direction \( u_i \).

The left (\( \delta \mathbf{w}_L \)), right (\( \delta \mathbf{w}_R \)), and centered (\( \delta \mathbf{w}_C \)) differences in the primitive variables shown in Equation A2 are then calculated. These differences are projected onto the characteristic variables, \( \delta \xi \), using the left eigenvectors given in Appendix A of Stone et al. (2008). Rather than reproduce the the expressions for each eigenvector, equations describing the final projections are shown since they are actually used in the GPU kernel. The projection of the left difference is

\[
\delta \xi_L = \begin{bmatrix}
-0.5(\rho_1 \delta u_L/a_i + \delta p_L/a_i^2)
\delta p_L - \delta p_L/a_i^2
\delta u_L
0.5(\rho_1 \delta u_L/a_i + \delta p_L/a_i^2)
\end{bmatrix},
\] (A9)

where \( \delta \rho_L, \delta u_L, \delta v_L, \delta w_L \), and \( \delta p_L \) are the components of the primitive variable difference vector \( \delta \mathbf{w}_L \). The projections for the right and central differences are calculated in the same manner, yielding \( \delta \xi_R \) and \( \delta \xi_C \).

The characteristic differences are then monotonized using the van Leer (1977) limiter, computed as

\[
\delta \xi = \begin{cases} 
\text{sgn}(\delta \xi_C) \min( |\delta \xi_C|, 2|\delta \xi_L|, 2|\delta \xi_R|), & \delta \xi_L, \delta \xi_R > 0 \\
0, & \text{otherwise}.
\end{cases}
\] (A10)

We project the monotonized differences in the characteristic variables back onto the primitive variables, providing slopes in each variable that are analogous to the limited slopes described in PLMP:

\[
\delta \mathbf{w}_i = \begin{bmatrix}
\delta \xi_0 + \delta \xi_1 + \delta \xi_4 \\
(\frac{1}{\rho_1})(-\delta \xi_0 + \delta \xi_1) \\
\delta \xi_2 \\
\delta \xi_3 \\
\frac{1}{a_i^2}(\delta \xi_0 + \delta \xi_4)
\end{bmatrix},
\] (A11)

Here, the numeric subscripts to refer to the components of the vector \( \xi \).

As in PLMP, these slopes are subsequently used to create a linear interpolation for reconstructing boundary values of the primitive variables:

\[
\mathbf{W}_{L,A} = \mathbf{w}_i - \frac{1}{2} \delta \mathbf{w}_i,
\]

\[
\mathbf{W}_{R,A} = \mathbf{w}_i + \frac{1}{2} \delta \mathbf{w}_i.
\] (A12)

The primitive variable boundary values are further monotonized to ensure that they are numerically bounded by the neighboring cell values:

\[
\mathbf{W}_{L,B} = \max[\min(\mathbf{w}_i, \mathbf{w}_{i-1}), \mathbf{W}_{L,A}]
\]

\[
\mathbf{W}_L = \min[\max(\mathbf{w}_i, \mathbf{w}_{i-1}), \mathbf{W}_{L,B}]
\]

\[
\mathbf{W}_{R,B} = \max[\min(\mathbf{w}_i, \mathbf{w}_{i+1}), \mathbf{W}_{R,A}]
\]

\[
\mathbf{W}_R = \min[\max(\mathbf{w}_i, \mathbf{w}_{i+1}), \mathbf{W}_{R,B}].
\] (A13)

enabling a slope vector to be computed from these adjusted boundary values as

\[
\delta \mathbf{w}_i = \mathbf{W}_R - \mathbf{W}_L.
\] (A14)

The reconstructed boundary values must be evolved in time to calculate appropriate input states for the Riemann problem. Instead of simply evolving the reconstructed values using associated fluxes as in Equations A6 and A7, the characteristic tracing method of Colella & Woodward (1984) is employed. To obtain a first approximation for the input states, an integration under the linear interpolation function is performed using the minimum wave speed to
define the domain of dependence for the left side of the cell, \( \lambda^m \), and the maximum wave speed for the right side of the cell, \( \lambda^p \):

\[
\begin{align*}
\tilde{W}_L^* &= W_L - 0.5 \frac{\Delta t}{\Delta x} \min(\lambda^m, 0) \delta w_i, \\
\tilde{W}_R^* &= W_R - 0.5 \frac{\Delta t}{\Delta x} \max(\lambda^p, 0) \delta w_i.
\end{align*}
\] (A15)

The input states are then corrected by accounting for the portion of each wave that does not reach the interface over a time \( \Delta t/2 \) as a result of the presence of the other waves. Correction terms are only needed for characteristics propagating toward each interface. The eigenvector projection and correction for each element of \( W^* \) is shown below, tracking the correction terms in the vectors \( s_L \) and \( s_R \). For the left side of cell \( i \),

\[
W_L^* = \tilde{W}_L^* + 0.5 \frac{\Delta t}{\Delta x} s_L
\] (A16)

with

\[
s_L = \begin{bmatrix}
(\lambda^m - \lambda^0)(\delta p_i - \delta p_i/a_i^2) \\
0 \\
(\lambda^m - \lambda^0)\delta v_i \\
(\lambda^m - \lambda^0)\delta w_i
\end{bmatrix}
+ \begin{bmatrix}
0.5(\lambda^m - \lambda^0)(\delta u_i/a_i + \delta p_i/a_i^2) \\
0.5(\lambda^m - \lambda^0)(\delta u_i + \delta p_i/(a_i p_i)) \\
0 \\
0.5(\lambda^m - \lambda^0)(\rho_i \delta u_i a_i + \delta p_i)
\end{bmatrix}.
\] (A17)

The first term is associated with the contact wave and is added only if \( \lambda^0 < 0 \). The second term is associated with the right acoustic wave and is added only if \( \lambda^p < 0 \). If both \( \lambda^0 \) and \( \lambda^p \) are greater than 0, there is no need for a correction because those waves cannot affect the left interface of the cell. For the right side of cell \( i \),

\[
W_R^* = \tilde{W}_R^* + 0.5 \frac{\Delta t}{\Delta x} s_R
\] (A18)

with

\[
s_R = \begin{bmatrix}
0.5(\lambda^p - \lambda^0)(-\rho_i \delta u_i/a_i + \delta p_i/a_i^2) \\
0.5(\lambda^p - \lambda^0)(\delta u_i - \delta p_i/(a_i p_i)) \\
0 \\
0.5(\lambda^p - \lambda^0)(-\rho_i \delta u_i a_i + \delta p_i)
\end{bmatrix}
+ \begin{bmatrix}
(\lambda^p - \lambda^0)(\delta p_i - \delta p_i/a_i^2) \\
(\lambda^p - \lambda^0)(\rho_i \delta u_i a_i + \delta p_i) \\
(\lambda^p - \lambda^0)\delta v_i \\
(\lambda^p - \lambda^0)\delta w_i
\end{bmatrix}.
\] (A19)

Here the first term is associated with the left acoustic wave and is added only if \( \lambda^m > 0 \), while the second term is associated with the contact wave and is added only if \( \lambda^0 > 0 \). As with the left interface, the corrections only apply if the waves are moving toward the interface. Once the corrections have been made, \( W_L^* \) and \( W_R^* \) can be used as inputs to the Riemann problem.

### A.3. PPMC

**Cholla** also includes implementations of third-order spatial reconstruction techniques, including two varieties of the piecewise parabolic method developed by [Colella & Woodward] (1984). The piecewise parabolic method with slope limiting applied in the characteristic variables (PPMC) is presented first as it shares several steps with PLMC. Our implementation of this method closely follows that outlined in [Stone et al.] (2008).

The first step in PPMC is to calculate monotonized slopes for cells \( i-1 \), \( i \), and \( i+1 \). These slopes are labeled \( \delta w_{i-1} \), \( \delta w_i \), and \( \delta w_{i+1} \). The limited slopes for each cell are calculated in a manner identical to that described in PLMC, following Equations [A8] - [A11]. Since slopes must be calculated for all three cells, the stencil for PPMC contains two cells to the left and right of cell \( i \). Once the limited slope vectors for all three cells have been calculated, the algorithm proceeds as follows.

Using the monotonized linear slopes, a parabolic interpolation is computed and used to calculate the reconstructed boundary values:

\[
W_{L,A} = \frac{1}{2}(w_i + w_{i-1}) - \frac{1}{6}(\delta w_i - \delta w_{i-1}),
\]
\[
W_{R,A} = \frac{1}{2}(w_{i+1} + w_i) - \frac{1}{6}(\delta w_{i+1} - \delta w_i).
\] (A20)

Monotonicity constraints are applied to the reconstructed boundary values to ensure that they lie between the average values in neighboring cells. If the cell contains a local minimum or maximum, both interface values are set equal to the cell average:

\[
W_{L,A} = W_{R,A} = w_i, \quad \text{if } (W_{R,A} - w_i)(w_i - W_{L,A}) \leq 0
\] (A21)

If the parabolic interpolation violates monotonicity as a result of a steep gradient, the interface values are modified as:

\[
W_{L,A} = 3w_i - 2W_{R,A}, \quad \text{if } (W_{R,A} - W_{L,A})(W_{L,A} - (3w_i - 2W_{R,A})) < 0
\]
\[
W_{R,A} = 3w_i - 2W_{L,A}, \quad \text{if } (W_{R,A} - W_{L,A})[(3w_i - 2W_{L,A}) - W_{R,A}] < 0
\] (A22)
These reconstructed boundary values are further adjusted using the minmod limiter operation:

\[
\begin{align*}
W_{L,B} &= \max(\min(w_i, w_{i-1}), W_{L,A}) \\
W_L &= \min(\max(w_i, w_{i-1}), W_{L,B}) \\
W_{R,B} &= \max(\min(w_i, w_{i+1}), W_{R,A}) \\
W_R &= \min(\max(w_i, w_{i+1}), W_{R,B})
\end{align*}
\] (A23)

At this point, a monotonized parabolic interpolation can be reconstructed. New slopes are computed that account for the adjusted boundary values:

\[
\delta w_i = W_R - W_L.
\] (A24)

These slopes are used to compute the time-evolved left and right boundary values by integrating under a parabolic interpolation function:

\[
\begin{align*}
\tilde{W}_L &= W_L - \frac{1}{2} \alpha^m \left[ \delta w_i + w_6(1 + \frac{2}{3} \alpha^m) \right], \\
\tilde{W}_R &= W_R - \frac{1}{2} \beta^p \left[ \delta w_i - w_6(1 - \frac{2}{3} \beta^p) \right],
\end{align*}
\] (A25)

\[
\begin{cases}
\alpha^m = \frac{\Delta t}{\Delta x} \min(\lambda^m, 0), & \beta^m = \frac{\Delta t}{\Delta x} \max(\lambda^m, 0), \\
\alpha^0 = \frac{\Delta t}{\Delta x} \min(\lambda^0, 0), & \beta^0 = \frac{\Delta t}{\Delta x} \max(\lambda^0, 0), \\
\alpha^p = \frac{\Delta t}{\Delta x} \min(\lambda^p, 0), & \beta^p = \frac{\Delta t}{\Delta x} \max(\lambda^p, 0).
\end{cases}
\] (A27)

Here we have borrowed from the notation of Colella & Woodward (1984) to define \(\alpha^m\) and \(\beta^p\), unit-free variables associated with the characteristic speeds

As in PLMC, the minimum characteristic speed, \(\lambda^m\), is used to define the domain of dependence for the left interface, and the maximum characteristic speed, \(\lambda^p\), is used for the right interface. The primitive variable time-evolved boundary values \(\tilde{W}_L^*\) and \(\tilde{W}_R^*\) are first approximations to the input states for the Riemann problem.

The input states must now be corrected by accounting for the other characteristics propagating toward the interface. At the left side of the cell, we compute

\[
s_L = \begin{bmatrix}
E_0 - E_4/a_i^2 \\
E_2 \\
E_3
\end{bmatrix} + \begin{bmatrix}
0.5(\rho_i B_1/a_i + B_1/a_i^2) \\
0.5[B_1 + B_4/(a_i \rho_i)] \\
0.5(\rho_i B_1 a_i + B_4)
\end{bmatrix},
\] (A28)

where the first term is added only if \(\lambda^0 < 0\), and the second only if \(\lambda^p < 0\) - otherwise there is no correction. In the above,

\[
E = \frac{1}{2} \alpha^m \left[ \delta w_i + w_6(1 + \frac{2}{3} \alpha^m) \right] - \frac{1}{2} \alpha^0 \left[ \delta w_i + w_6(1 + \frac{2}{3} \alpha^0) \right],
\] (A29)

is associated with the contact wave, and

\[
B = \frac{1}{2} \alpha^m \left[ \delta w_i + w_6(1 + \frac{2}{3} \alpha^p) \right] - \frac{1}{2} \alpha^p \left[ \delta w_i + w_6(1 + \frac{2}{3} \alpha^p) \right].
\] (A30)

is associated with the right-most acoustic wave. Subscripts denote the elements of \(E\) and \(B\). Similarly, for the right side of the cell,

\[
s_R = \begin{bmatrix}
0.5(-\rho_i C_1 a_i + C_4/a_i^2) \\
0.5[C_1 - C_4/(a_i \rho_i)] \\
0.5(-\rho_i C_1 a_i + C_4)
\end{bmatrix} + \begin{bmatrix}
(D_0 - D_4/a_i^2) \\
D_2 \\
D_3
\end{bmatrix}.
\] (A31)

The first term accounts for the correction owing to the left-most acoustic wave and is added only if \(\lambda^m > 0\), and the
second term accounts for the correction from the contact wave and is added only if $\lambda^0 > 0$. In this case,

$$C = \frac{1}{2} \beta^p \left[ \delta w_i - w_0 (1 - \frac{2}{3} \beta^p) \right] - \frac{1}{2} \beta^m \left[ \delta w_i - w_0 (1 - \frac{2}{3} \beta^m) \right],$$

$$D = \frac{1}{2} \beta^p \left[ \delta w_i - w_0 (1 - \frac{2}{3} \beta^p) \right] - \frac{1}{2} \beta^p \left[ \delta w_i + w_0 (1 + \frac{2}{3} \beta^p) \right].$$

(A32)

With these correction terms input states for the Riemann problem can be calculated as

$$W_L^* = \tilde{W}_L^* + s_L \quad \text{and} \quad W_R^* = \tilde{W}_R^* + s_R.$$

(A33)

A.4. PPMP

The *Cholla* implementation of the piecewise parabolic method computed in the primitive variables closely follows the original description in *Colella & Woodward* (1984), with some additional notation adapted from *Fryxell et al.* (2000). For convenience, the following description assumes a uniform cell size. PPMP is the most complicated of the reconstruction methods implemented in *Cholla*, and the algorithm follows this brief outline:

1. Reconstruct boundary values using parabolic interpolation.
2. Steepen contact discontinuities, if necessary.
3. Flatten shocks, if necessary.
4. Ensure that the parabolic distribution is monotonic.
5. Integrate under the parabolic interpolation to determine input states for the Riemann problem.
6. Use characteristic tracing to correct the input states.

A.4.1. Parabolic Interpolation

The first step in PPMP is to reconstruct the boundary values using a parabolic interpolation with limited slopes. The interpolation is identical to that shown in Equation (A20)

$$W_L = \frac{1}{2} (w_i + w_{i-1}) - \frac{1}{6} (\delta w_i - \delta w_{i-1}),$$

$$W_R = \frac{1}{2} (w_{i+1} + w_i) - \frac{1}{6} (\delta w_{i+1} - \delta w_i).$$

(A34)

However, in PPMP the slopes are limited in the primitive variables using the *van Leer* (1977) limiter. The slopes $\delta w_{i-1}$, $\delta w_i$, and $\delta w_{i+1}$ are all calculated following Equations (A2) and (A3).

A.4.2. Contact Discontinuity Steepening

Once the interface values have been reconstructed, contact discontinuity steepening is applied to the interface values for the density, $W_L[0] = \rho_L$ and $W_R[0] = \rho_R$. Whether steepening is applied depends on a number of necessary criteria. First, to avoid steepening density jumps owing to numerical noise, steepening is only applied if the density difference between cells exceeds a minimum relative size:

$$\frac{|\rho_{i+1} - \rho_{i-1}|}{\min(\rho_{i+1}, \rho_{i-1})} > 0.01.$$  

(A35)

If the density jump is large enough, we further require that the pressure jump across the cell be sufficiently small:

$$\frac{|p_{i+1} - p_{i-1}|}{\min(p_{i+1}, p_{i-1})} < 0.1 \gamma \frac{|\rho_{i+1} - \rho_{i-1}|}{\min(\rho_{i+1}, \rho_{i-1})}.$$  

(A36)

Next, the second derivative of the density distribution across cells $i - 1$ and $i + 1$ is estimated as

$$\delta^2 \rho_i = \frac{\rho_{i+1} - 2 \rho_i + \rho_{i-1}}{6 \Delta x^2}.$$  

(A37)

The product of the second derivatives then determines whether the local density profile on either side of the cell $i$ has the same concavity by requiring

$$\delta^2 \rho_{i-1} \delta^2 \rho_{i+1} > 0.$$  

(A38)

Assuming the three conditions listed in Equations (A35), (A36), and (A38) are satisfied, a dimensionless quantity involving the first and third derivatives of density is calculated as

$$\eta_i = \frac{(\delta^2 \rho_{i+1} - \delta^2 \rho_{i-1}) \Delta x^2}{\rho_{i+1} - \rho_{i-1}}.$$  

(A39)
This quantity is used to compute the steepening coefficient $\eta_i$ from the parameters determined heuristically in Colella & Woodward (1984):

$$\eta_i = \max[0, \min(20\delta \eta_i - 1, 1)]. \quad (A40)$$

The steepening coefficient $\eta_i$ and the monotonized slopes $\delta w_{i-1}[0] = \delta \rho_{i-1}$ and $\delta w_{i+1}[0] = \delta \rho_{i+1}$ are then used to steepen the left and right interface density values, providing

$$\begin{align*}
\rho_L &= \rho_L(1 - \eta_i) + (\rho_{i-1} + 0.5\delta \rho_{i-1})\eta_i \\
\rho_R &= \rho_R(1 - \eta_i) + (\rho_{i-1} - 0.5\delta \rho_{i+1})\eta_i
\end{align*} \quad \text{(A41)}$$

**A.4.3. Shock Flattening**

Because of their self-steepening property, shocks in PPMP can become under-resolved, i.e. narrow enough to be contained within a single cell. Tests have demonstrated that shocks contained within a single cell tend to lead to severe oscillations near the shock front, while those spread over two or more cells do not pose a problem (Colella & Woodward 1984; Fryxell et al. 2000). A solution is to flatten numerically the interpolation near problematic shocks, reverting to a first-order reconstruction in circumstances where the shock is empirically deemed too narrow. To determine whether a shock needs flattening, a shock steepness parameter $S$ is calculated to compare the pressure gradient across two and four cells:

$$S_i = \frac{\rho_{i+1} - \rho_{i-1}}{\rho_{i+2} - \rho_{i-2}} \quad \text{(A42)}$$

The steepness parameter is used to construct a dimensionless coefficient

$$\tilde{F}_i = \max(0, \min[1, 10(S_i - 0.75)]) \quad \text{(A43)}$$

that may cover the range $\tilde{F}_i = [0, 1]$. This formulation is designed to ensure that only shocks contained within fewer than two cells are steepened. Further, we set $\tilde{F}_i = 0$ if the relative pressure jump is not large and the shock is not steep, when

$$\frac{|\rho_{i+1} - \rho_{i-1}|}{\min(\rho_{i+1}, \rho_{i-1})} < \frac{1}{3} \quad \text{(A44)}$$

or if the velocity gradient is positive (indicating that the fluid is not being compressed in the direction along which we are reconstructing boundary values), when

$$u_{i+1} - u_{i-1} > 0 \quad \text{(A45)}$$

The same rules are applied to the dimensionless parameters $\tilde{F}_{i-1}$ and $\tilde{F}_{i+1}$. This procedure means that PPMP requires a stencil with three cells on both sides of cell $i$. Here we are calculating shocks in the $x$-direction; in the $y$- and $z$-directions, the components $v$ and $w$ are used to test the velocity gradient. The final flattening coefficient for cell $i$ is set as

$$F_i = \begin{cases} 
\max(\tilde{F}_i, \tilde{F}_{i+1}), & \text{if } \rho_{i+1} - \rho_{i-1} < 0, \\
\max(\tilde{F}_i, \tilde{F}_{i-1}), & \text{otherwise.} \end{cases} \quad \text{(A46)}$$

We use this value of $F_i$ to modify the interface values. Unlike in discontinuity steeping, for shock flattening every primitive variable is modified:

$$\begin{align*}
W_L &= F_i w_i + (1 - F_i) W_L \\
W_R &= F_i w_i + (1 - F_i) W_R
\end{align*} \quad \text{(A47)}$$

If $F_i = 0$, the expression has no effect on the interface values. If $F_i = 1$ the zone average values are used for the interface values, effectively replacing the original limited slope for cell $i$ with a flatter slope.

**A.4.4. Monotonicity**

The next step in the reconstruction is to ensure that the parabolic distribution of each of the variables is monotonic by checking for local maxima and minima and modifying steep gradients, as in Equations (A21) and (A22)

$$\begin{align*}
W_L &= W_R = w_i, & \text{if } (W_R - w_i)(w_i - W_L) &\leq 0 \\
W_L &= 3w_i - 2W_R, & \text{if } (W_R - W_L)(W_L - (3w_i - 2W_R)) &< 0 \\
W_L &= 3w_i - 2W_L, & \text{if } (W_R - W_L)((3w_i - 2W_L) - W_R) &< 0.
\end{align*} \quad \text{(A48)}$$

**A.4.5. Calculation of the Input States**

By this stage, reconstruction of the boundary values has been completed and the input states for the Riemann problem can be calculated. Once again, the characteristic tracing method of Colella & Woodward (1984) is used. First, the speeds of the three characteristics are defined as in PLMC and PPCM:

$$\lambda^e = u_i - a_i, \quad \lambda^0 = u_i, \quad \lambda^p = u_i + a_i. \quad \text{(A49)}$$
Again, \( a_i \) is the sound speed in cell \( i \) calculated using average values of the density and pressure. Because we have adjusted the boundary values from the original parabolic interpolation, we must adjust the values of the slopes across the cell so that the parabolic interpolation retains the correct average value:

\[
\delta w_i = W_R - W_L, \quad w_6 = 6[w_i - 0.5(W_L + W_R)].
\]  

We now define \( \alpha \) and \( \beta \), two variables that are associated with the characteristic wave speeds approaching the left and right interfaces,

\[
\alpha^m = \frac{\Delta t}{\Delta x} \min(\lambda^m, 0), \quad \beta^m = \frac{\Delta t}{\Delta x} \max(\lambda^m, 0),
\]

\[
\alpha^0 = \frac{\Delta t}{\Delta x} \min(\lambda^0, 0), \quad \beta^0 = \frac{\Delta t}{\Delta x} \max(\lambda^0, 0),
\]

\[
\alpha^p = \frac{\Delta t}{\Delta x} \min(\lambda^p, 0), \quad \beta^p = \frac{\Delta t}{\Delta x} \max(\lambda^p, 0).
\]

We use these variables to calculate a time-evolved boundary value associated with each characteristic:

\[
W^m_L = W_L - \frac{1}{2} \alpha^m \left[ \delta w_i + w_6 (1 + \frac{2}{3} \alpha^m) \right]
\]

\[
W^0_L = W_L - \frac{1}{2} \alpha^0 \left[ \delta w_i + w_6 (1 + \frac{2}{3} \alpha^0) \right]
\]

\[
W^p_L = W_L - \frac{1}{2} \alpha^p \left[ \delta w_i + w_6 (1 + \frac{2}{3} \alpha^p) \right]
\]

\[
W^m_R = W_R - \frac{1}{2} \beta^m \left[ \delta w_i - w_6 (1 - \frac{2}{3} \beta^m) \right]
\]

\[
W^0_R = W_R - \frac{1}{2} \beta^0 \left[ \delta w_i - w_6 (1 - \frac{2}{3} \beta^0) \right]
\]

\[
W^p_R = W_R - \frac{1}{2} \beta^p \left[ \delta w_i - w_6 (1 - \frac{2}{3} \beta^p) \right]
\]

For example, \( W^m_L \) is the time-evolved boundary value at the left interface of the cell obtained by integrating under the characteristic associated with the left acoustic wave. If the fluid flow is supersonic toward the right, the left acoustic wave is not approaching the interface, and \( W^m_L \) is simply equal to the reconstructed boundary value. The same integration appeared in Equation (A25), though in this case we explicitly integrate under every characteristic and not just the characteristic approaching the interface with the greatest speed. For the density, normal velocity, and pressure, we refer to value calculated using the characteristic approaching the interface at the greatest speed as the “reference state”, e.g. \( W^m_L \) for the left interface and \( W^0_R \) for the right. As with PPMC, this reference state is our first guess at the input state for the Riemann problem. For the transverse velocities, we use the states associated with the advection speed, \( W^p_L \) and \( W^p_R \).

These reference states are only first-order accurate approximations and the input states can be further corrected to account for the presence of other characteristics approaching the interface. Following the notation in [Colella & Woodward (1984)], the description of the algorithm continues in terms of the primitive variables. The sound speeds for the reference states on the left and right of the cell are computed

\[
a_L = \sqrt{\frac{\gamma p^0_L}{\rho^m_L}} \quad \text{and} \quad a_R = \sqrt{\frac{\gamma p^0_R}{\rho^p_R}},
\]

along with correction terms that are added to the reference state,

\[
\chi^m_L = -\frac{1}{2 \rho^m_L a_L} \left( u^m_L - u^p_L - \frac{p^0_L + p^m_L}{\rho^m_L a_L} \right),
\]

\[
\chi^0_L = \frac{1}{2 \rho^0_L a_L} \left( u^0_L - u^m_L - \frac{p^0_R - p^m_L}{\rho^0_L a_L} \right),
\]

\[
\chi^p_L = \frac{p^m_L - p^0_L}{(\rho^m_L a_L)^2} + \frac{1}{\rho^m_L} - \frac{1}{\rho^0_L},
\]

\[
\chi^m_R = -\frac{1}{2 \rho^m_R a_R} \left( u^m_R - u^p_R - \frac{p^0_R + p^m_R}{\rho^m_R a_R} \right),
\]

\[
\chi^0_R = \frac{1}{2 \rho^0_R a_R} \left( u^0_R - u^m_R - \frac{p^0_R - p^m_R}{\rho^0_R a_R} \right),
\]

\[
\chi^p_R = \frac{p^m_R - p^0_R}{(\rho^m_R a_R)^2} + \frac{1}{\rho^m_R} - \frac{1}{\rho^0_R}.
\]
In the event that the characteristic is not traveling toward the interface these correction terms are set to zero,

\[
\begin{align*}
\chi_L^0 &= 0 \quad \text{if } \chi^0 > 0, \\
\chi_L^p &= 0 \quad \text{if } \chi^p > 0, \\
\chi_R^m &= 0 \quad \text{if } \chi^m < 0, \\
\chi_R^0 &= 0 \quad \text{if } \chi^0 < 0.
\end{align*}
\]  
(A55)

The correction terms are then used with the reference state integration to calculate the final input states for the Riemann problem on each side of the cell:

\[
\begin{align*}
\rho_L &= \left( \frac{1}{\rho_L^m} - \chi_L^0 - \chi_L^p \right)^{-1}, \\
\rho_R,i &= \left( \frac{1}{\rho_R^m} - \chi_R^m - \chi_R^0 \right)^{-1}, \\
u_L &= u_L^m + \rho_L^m a_L \chi_L^p, \\
u_R &= u_R^p - \rho_R^p a_R \chi_R^m, \\
p_L &= p_L^m + (\rho_L^m a_L)^2 \chi_L^p, \\
p_R &= p_R^p + (\rho_R^p a_R)^2 \chi_R^m, \\
v_L &= v_L^0, \\
v_R &= v_R^0, \\
w_L &= w_L^0, \\
w_R &= w_R^0.
\end{align*}
\]  
(A56)

B. RIEMANN SOLVERS

B.1. The Exact Solver

The exact solver used in Cholla follows the solver presented in Chapter 4 of [Toro (2009)], adapted for implementation in CUDA C. The algorithm to solve the Riemann problem is presented below, using an x-interface as an example. In the following section the CTU notation from Section 2.1 is used, where states are labeled at the left and right of the interface.

First, the input states at the left and right of the interface are converted to the primitive variables, \(W_L^p\) and \(W_R^p\). (Between GPU kernels like the interface reconstruction and the Riemann problem, calculated values are stored in the conserved form.) These vectors are used to compute the corresponding sound speed on either side of the interface

\[
\begin{align*}
\alpha_L &= \sqrt{\gamma p_L / \rho_L} \quad \text{and} \quad \alpha_R = \sqrt{\gamma p_R / \rho_R}. 
\end{align*}
\]  
(B1)

To determine the Riemann solution, the exact pressure \(p^m\) and velocity \(u^m\) in the intermediate state must be computed (see Figure 2). We use the [Toro (2009)] primitive variable Riemann solver to provide an initial approximation to the intermediate state pressure, given by

\[
\tilde{p} = 0.5(p_L + p_R) + 0.125(u_L - u_R)(p_L + p_R)(a_L + a_R). 
\]  
(B2)

Because \(\tilde{p}\) is an approximation and the solution for \(p\) cannot be negative, we set \(\tilde{p} = 0\) if the calculated pressure is below zero. A two-shock Riemann solver is then used to calculate a more accurate estimate,

\[
p_0 = \frac{9Lp_L + 9Rp_R - (u_R - u_L)}{9L + 9R},
\]  
(B3)

with

\[
g_k = \sqrt{\frac{A_k}{\rho + B_k}}, \quad A_k = \frac{2}{(\gamma + 1)\rho_k}, \quad B_k = \frac{\gamma - 1}{\gamma + 1}p_k, \quad \text{where } k = L, R.
\]  
(B4)

To maintain positivity, a pressure floor of \(p_0 \geq 10^{-6}\) is enforced in this estimate. The pressure \(p_0\) is then used as a starting point in a Newton-Raphson iteration to compute the exact solution for the pressure in the intermediate region. We define the pressure functions \(f_L\) and \(f_R\),

\[
f_k = \left\{ \begin{array}{ll}
\frac{2a_k}{\gamma - 1} & \text{if } p_0 \leq p_k \text{ (rarefaction)}, \\
(p_0 - p_k) \left( \frac{A_k}{p_0 + B_k} \right) \frac{2a_k}{\gamma - 1} & \text{if } p_0 > p_k \text{ (shock)},
\end{array} \right.
\]  
(B5)

and their first derivatives \(f'_L\) and \(f'_R\),

\[
f'_k = \left\{ \begin{array}{ll}
\frac{1}{\rho_k a_k} \left( \frac{p_0}{p_k} \right) \frac{2a_k}{\gamma - 1} & \text{if } p_0 \leq p_k \text{ (rarefaction)}, \\
\left[ \frac{A_k}{B_k + p_0} \right] \left( 1 - \frac{p_0 - p_k}{2(B_k + p_0)} \right) \frac{2a_k}{\gamma - 1} & \text{if } p_0 > p_k \text{ (shock)},
\end{array} \right.
\]  
(B6)
Again, $k = L$ or $R$, and $A_k$ and $B_k$ are as defined above. These quantities are used to calculate the pressure in the intermediate state, $p^m$:

$$p^m = p_0 - \frac{f_L + f_R + u_R - u_L}{f'_L + f'_R}. \tag{B7}$$

We then compare the newly computed pressure, $p^m$, to the previously computed pressure,

$$\Delta = 2\frac{|p^m - p_0|}{|p^m + p_0|}. \tag{B8}$$

If $\Delta$ is greater than a relative tolerance (e.g., $10^{-6}$), the values of $f_k$, $f'_k$, and $p^m$ are recomputed using the updated value of $p^m$ in place of $p_0$ in Equations \[B5\] and \[B6\]. When $\Delta$ is less than the tolerance, the procedure halts. Having calculated a suitably accurate pressure $p^m$, the pressure can then be used to compute the velocity in the intermediate state as

$$u^m = 0.5(u_L + u_R + f_R - f_L). \tag{B9}$$

Once the values of $p^m$ and $u^m$ in the intermediate state have been computed, the values for each of the primitive variables can be calculated at the cell interface. To do this, a number of conditions are tested to determine where in the Riemann solution the interface lies. For pure hydrodynamics there are ten possible outcomes.

If $u^m \geq 0$, the contact discontinuity is to the right of the cell interface. We then check to see if there is a rarefaction wave on the left, i.e. if $p^m \leq p_L$. If so, there are three possible solutions.

- If $u_L - a_L \geq 0$ the interface is in the left data state, and the solution is simply the input data on the left:

  $$\rho = \rho_L, \quad u = u_L, \quad p = p_L. \tag{B10}$$

- If $u^m - a_L \left(\frac{p^m}{p_L}\right)^{\frac{1}{\gamma-1}} < 0$ the interface is in the intermediate data state to the right of the fan, but left of the contact:

  $$\rho = \rho_L \left(\frac{p^m}{p_L}\right)^{\frac{1}{\gamma}}, \quad u = u^m, \quad p = p^m. \tag{B11}$$

- Otherwise, the interface is within the rarefaction fan:

  $$\rho = \rho_L \left(\frac{a}{a_L}\right)^{\frac{2}{\gamma-1}}, \quad u = a, \quad p = p_L \left(\frac{a}{a_L}\right)^{\frac{2\gamma}{\gamma-1}}, \quad \text{where} \quad a = 2\frac{\gamma + 1}{\gamma + 1}\left(a_L + \frac{\gamma - 1}{2}u_L\right). \tag{B12}$$

If there is a shock to the left, rather than a rarefaction, i.e. if $p^m > p_L$, the shock speed is calculated as

$$s_L = u_L - a_L \sqrt{\frac{(\gamma + 1) p^m}{2\gamma p_L}} + \frac{\gamma - 1}{2\gamma}. \tag{B13}$$

- If $s_L \geq 0$ the interface samples the left data state:

  $$\rho = \rho_L, \quad u = u_L, \quad p = p_L. \tag{B14}$$

- Otherwise, the interface samples the intermediate data state to the left of the contact:

  $$\rho = \rho_L \left(\frac{p^m}{p_L} + \frac{\gamma - 1}{\gamma + 1}\right) / \left(\frac{p^m}{p_L} \frac{\gamma - 1}{\gamma + 1} + 1\right), \quad u = u^m, \quad p = p^m. \tag{B15}$$

If instead $u^m < 0$, the contact discontinuity is to the left of the cell interface and there is a similar set of five possible outcomes. If there is a rarefaction wave on the right of the Riemann solution, i.e. if $p^m \leq p_R$ there are three possibilities:

- If $u_R + a_R \leq 0$ the interface samples the right data state:

  $$\rho = \rho_R, \quad u = u_R, \quad p = p_R. \tag{B16}$$

- If $u^m + a_R \left(\frac{p^m}{p_R}\right)^{\frac{1}{\gamma-1}} \geq 0$ the interface samples the intermediate state to the left of the fan, but right of the contact:

  $$\rho = \rho_R \left(\frac{p^m}{p_R}\right)^{\frac{1}{\gamma}}, \quad u = u^m, \quad p = p^m. \tag{B17}$$
is also needed. These Roe average states are used to calculate the eigenvalues of the Roe Jacobian, the cell interface are determined. The fluxes of the conserved variables can then be calculated following Equation 11:

\[
\rho = \rho_R \left( \frac{a}{a_R} \right)^{\frac{2\gamma}{\gamma+1}}, \quad u = -a, \quad p = p_R \left( \frac{a}{a_R} \right)^{\frac{2\gamma}{\gamma+1}}, \quad \text{where } a = \frac{2}{\gamma+1} \left( a_R + \frac{\gamma-1}{2} u_R \right).
\]  

(B18)

If \( p^m > p_R \) there is a shock to the right rather than a rarefaction, the shock speed is calculated as

\[
s_{LR} = u_R + a_R \sqrt{\left( \frac{\gamma+1}{2\gamma} \right) \frac{p^m}{p_R} + \frac{\gamma-1}{2\gamma}}.
\]  

(B19)

• If \( s_L \leq 0 \) the interface samples the right data state:

\[
\rho = \rho_R, \quad u = u_R, \quad p = p_R.
\]  

(B20)

• Otherwise, the interface samples the intermediate data state to the right of the contact:

\[
\rho = \rho_R \left( \frac{p^m}{p_R} + \frac{\gamma-1}{\gamma+1} \right) / \left( \frac{p^m}{p_R} \frac{\gamma-1}{\gamma+1} + 1 \right), \quad u = u^m, \quad p = p^m.
\]  

(B21)

After determining where in the Riemann solution the interface samples, the evolved primitive variables \( \rho, u, \) and \( p \) at the cell interface are determined. The fluxes of the conserved variables can then be calculated following Equation 11

\[
F = \begin{bmatrix} \rho u \\ \rho u^2 + p \\ \rho u w_k \\ \rho u w_k + (E + p)u \end{bmatrix},
\]  

(B22)

where \( k = L \) or \( R; \) \( L \) if \( u \geq 0, \) and \( R \) if \( u < 0. \) These conditions reflect the fact that transverse velocities are simply advected with the flow. Equation 12 is used to compute similar fluxes in the \( y \)-direction, and Equation 13 is used for fluxes in the \( z \)-direction.

B.2. The Roe Solver

Rather than using an expensive iterative procedure to calculate the exact solution to the Riemann problem, the Roe (1981) Riemann solver calculates an exact solution to a linearized version of the Euler equations. Below the procedure for calculating the Roe fluxes at an \( x \)-interface is detailed. The \( y \)- and \( z \)-interface calculations are identical modulo an appropriate change of variables. Should the approximate Roe solver fail, we include a failsafe based on the method of Stone et al. (2008) where HLLE fluxes (Harten et al. 1983; Einfeldt 1988) are substituted. The Roe and HLLE solvers are described below.

The Roe solver starts with the calculated input vectors of primitive variables at the left and right of an interface, \( W_L^* \) and \( W_R^* \). Fluxes corresponding to the left and right state, \( F_L \) and \( F_R \), are calculated following Equation 11. The Roe average state, \( \bar{u} = (\bar{\rho}, \bar{u}, \bar{v}, \bar{w}, \bar{H})^T \), is then computed:

\[
\begin{align*}
\bar{\rho} &= \sqrt{\rho_L \rho_R} \\
\bar{u} &= \frac{\sqrt{\rho_L \rho_R} u_L + \sqrt{\rho_R u_R}}{\sqrt{\rho_L} + \sqrt{\rho_R}} \\
\bar{v} &= \frac{\sqrt{\rho_L \rho_R} v_L + \sqrt{\rho_R v_R}}{\sqrt{\rho_L} + \sqrt{\rho_R}} \\
\bar{w} &= \frac{\sqrt{\rho_L \rho_R} w_L + \sqrt{\rho_R w_R}}{\sqrt{\rho_L} + \sqrt{\rho_R}} \\
\bar{H} &= \frac{\sqrt{\rho_L} H_L + \sqrt{\rho_R} H_R}{\sqrt{\rho_L} + \sqrt{\rho_R}},
\end{align*}
\]  

(B23)

with the enthalpy, \( H = (E + p)/\rho \), used instead of the pressure. The average sound speed

\[
\bar{a} = \sqrt{(\gamma - 1)(\bar{H} - 0.5 \bar{V}^2)}, \quad \text{where } \bar{V}^2 = \bar{u}u + \bar{v}v + \bar{w}w
\]  

(B24)

is also needed. These Roe average states are used to calculate the eigenvalues of the Roe Jacobian,

\[
\lambda^m = \bar{u} - \bar{a}, \quad \lambda^0 = \bar{a}, \quad \text{and } \lambda^p = \bar{u} + \bar{a}.
\]  

(B25)

If the flow is supersonic (\( \lambda^m \geq 0 \) to the right, or \( \lambda^m \leq 0 \) to the left), the solver returns the appropriate left or right state fluxes \( F_L \) or \( F_R \). If flow is subsonic, the calculation of the Roe fluxes proceeds.

Differences in the conserved variables between the left and right states are computed:

\[
\begin{align*}
\delta \rho &= \rho_R - \rho_L \\
\delta m_x &= m_{x,R} - m_{x,L} \\
\delta m_y &= m_{y,R} - m_{y,L} \\
\delta m_z &= m_{z,R} - m_{z,L} \\
\delta E &= E_R - E_L.
\end{align*}
\]  

(B26)
These differences are projected onto the characteristics by multiplying by the left eigenvector associated with each eigenvalue. The resulting characteristics are the wave strengths, \( \xi \), from Equation 29:

\[
\begin{align*}
\xi_0 &= \delta \rho N_a (0.5 \gamma' \tilde{V}^2 + \tilde{u} \tilde{a}) - \delta m_x N_a (\gamma' \tilde{u} + \tilde{a}) - \delta m_y N_a \gamma' \tilde{w} - \delta m_z N_a \gamma' \tilde{w} + \delta E N_a \gamma' \\
\xi_1 &= -\delta \rho \tilde{v} + \delta m_y \\
\xi_2 &= -\delta \rho \tilde{w} + \delta m_z \\
\xi_3 &= \delta \rho (1 - N_a \gamma' \tilde{V}^2) + \delta m_x \gamma' \tilde{u} / \tilde{a}^2 + \delta m_y \gamma' \tilde{v} / \tilde{a}^2 + \delta m_z \gamma' \tilde{w} / \tilde{a}^2 - \delta E \gamma' / \tilde{a}^2 \\
\xi_4 &= \delta \rho N_a (0.5 \gamma' \tilde{V}^2 - \tilde{u} \tilde{a}) - \delta m_x N_a (\gamma' \tilde{u} - \tilde{a}) - \delta m_y N_a \gamma' \tilde{v} - \delta m_z N_a \gamma' \tilde{w} + \delta E N_a \gamma'
\end{align*}
\]

where \( N_a = 1/(2 \tilde{a}^2) \) and \( \gamma' = \gamma - 1 \). Numeric subscripts denote the elements of \( \xi \). Each characteristic variable is multiplied by its associated eigenvalue, yielding a vector of coefficients, \( C \):

\[
C_0 = \xi_0 |\lambda^m|, \quad C_1 = \xi_1 |\lambda^0|, \quad C_2 = \xi_2 |\lambda^0|, \quad C_3 = \xi_3 |\lambda^0|, \quad C_4 = \xi_4 |\lambda^p|
\]

The product of these coefficients with the right eigenvectors are then summed, keeping track of the summation in the vector \( s \):

\[
\begin{align*}
s_0 &= C_0 + C_3 + C_4 \\
s_1 &= C_0 (\tilde{u} - \tilde{a}) + C_3 \tilde{u} + C_4 (\tilde{u} + \tilde{a}) \\
s_2 &= C_0 \tilde{v} + C_1 + C_3 \tilde{v} + C_4 \tilde{v} \\
s_3 &= C_0 \tilde{w} + C_2 + C_3 \tilde{w} + C_4 \tilde{w} \\
s_4 &= C_0 (\tilde{H} - \tilde{u} \tilde{a}) + C_1 \tilde{v} + C_2 \tilde{w} + 0.5 C_3 \tilde{V}^2 + C_4 (\tilde{H} + \tilde{u} \tilde{a})
\end{align*}
\]

By this stage, all information needed to compute the Roe fluxes has been obtained. The Roe fluxes are then computed as

\[
F_{Roe} = \frac{1}{2} (F_L + F_R - s)
\]

Before returning these fluxes, however, the intermediate states are examined for possible negative densities or pressures. The intermediate states, labeled \( U_{L}^m \) and \( U_{R}^m \) in analogy with the exact Riemann solver, are calculated by projecting the characteristic variables onto the conserved variables. Each characteristic variable is multiplied by its associated right eigenvector and the results summed, in turn, to the left state. The left intermediate state, \( U_{L}^m \), is calculated as

\[
\begin{align*}
\rho_{L}^m &= \rho_L + \xi_0 \\
\rho u_{L}^m &= \rho u_L + \xi_0 (\tilde{u} - \tilde{a}) \\
\rho v_{L}^m &= \rho v_L + \xi_0 \tilde{v} \\
\rho w_{L}^m &= \rho w_L + \xi_0 \tilde{w} \\
E_{L}^m &= E_L + \xi_0 (\tilde{H} - \tilde{u} \tilde{a})
\end{align*}
\]

If \( \lambda^0 > \lambda^m \), we check for negative density and pressure. We then move on to the right intermediate state, \( U_{R}^m \):

\[
\begin{align*}
\rho_{R}^m &= \rho_R^m + \xi_3 \\
\rho u_{R}^m &= \rho u_R + \xi_3 \tilde{u} \\
\rho v_{R}^m &= \rho v_R + \xi_1 + \xi_3 \tilde{v} \\
\rho w_{R}^m &= \rho w_R + \xi_2 + \xi_3 \tilde{w} \\
E_{R}^m &= E_R^m + \xi_1 \tilde{v} + \xi_2 \tilde{w} + 0.5 \xi_3 \tilde{V}^2.
\end{align*}
\]

If \( \lambda^p > \lambda^0 \) then negative densities and pressures are possible and, if present, the Roe fluxes are replaced with HLLE fluxes. Otherwise the Roe fluxes calculated in Equation B30 are returned.

Little additional work must be done to calculate the HLLE fluxes since many of the required quantities are computed for the Roe fluxes. The HLLE solver constructs a single average state between the right and left input states, neglecting the contact wave. Ignoring the contact wave means that density discontinuities diffuse quickly, but the HLLE solver has the advantage of always producing positive intermediate densities and pressures, as shown in Einfeldt et al. (1991).

The HLLE flux algorithm starts with the computation of the sound speed for the left and right input states, \( a_L \) and \( a_R \). Bounding speeds are then calculated, defined by the minimum and maximum Roe eigenvalues and the left and right acoustic waves:

\[
b^m = \min[\min(\lambda^m, u_L - a_L), 0], \quad b^p = \max[\max(\lambda^p, u_R + a_R), 0]
\]

These speeds are used to compute left and right fluxes:

\[
F_L = F(W_L) - b^m U_L \quad \text{and} \quad F_R = F(W_R) - b^p U_R,
\]
where $\mathbf{F}(\mathbf{W})$ is calculated using Equation [11] and $\mathbf{U}$ refers to the conserved variables on either side of the interface, i.e. the input states for the Riemann problem. These intermediate fluxes are then employed to compute the HLLE fluxes, given by

$$ F_{\text{HLLE}} = \frac{1}{2} \left[ (\mathbf{F}_L + \mathbf{F}_R) + (\mathbf{F}_L - \mathbf{F}_R) \left( \frac{\mathbf{b}^p + \mathbf{b}^m}{\mathbf{b}^p - \mathbf{b}^m} \right) \right]. $$

(B35)

If the Roe solver fails during the transverse flux or conserved variable update steps of the CTU algorithm, these HLLE fluxes may be used.

C. THE H CORRECTION

As mentioned in the main text, strictly upwind multidimensional integration algorithms like CTU are susceptible to a particular type of numerical pathology, referred to as the carbuncle instability owing to its appearance \cite{Quirk1994}. The carbuncle instability arises in problems containing strong, grid-aligned shocks as a result of insufficient dissipation when inserting a one dimensional Riemann flux into a multidimensional problem. In the scenario when a planar shock is present perpendicular to the direction of the fluid flow, crossflow oscillations arise that grow and lead to severe inaccuracies in the numerical model. To correct this phenomenon, Cholla implements a solution devised by Sanders et al. \cite{Sanders1998} that accounts for the perpendicular velocities when calculating the Riemann flux normal to an interface. The correction thereby adds sufficient dissipation to mitigate the carbuncle instability. This method for suppressing the carbuncle instability is called H correction in reference to the shape of the required stencil.

In Cholla, H Correction is implemented in conjunction with the Roe Riemann solver. The correction is only applied to the fluxes calculated during the second set of Riemann problems, between the transverse flux update and the final conserved variable update of the CTU algorithm. To apply the H correction, at every interface a quantity $\eta$ is computed that depends on the velocity normal to the interface and the sound speed. For the $x$-interface at $(i + \frac{1}{2},j)$ in a two dimensional simulation, this quantity can be computed as

$$ \eta_{(i+\frac{1}{2},j)} = \frac{1}{2}[(u_R + a_R) - (u_L + a_L)]. $$

(C1)

where $u_R$ and $u_L$ are the normal velocity at the interface calculated using the right and left input states, and $a_R$ and $a_L$ are the corresponding sound speeds. For $y$- and $z$-interfaces, the velocity components $v$ and $w$ would be used. Once a value of $\eta$ has been calculated for each interface, a further quantity $\eta^H$ is computed for each interface by finding the maximum value of $\eta$ for each of the surrounding interfaces:

$$ \eta_{(i+\frac{1}{2},j)}^H = \max[\eta_{(i-1,j+\frac{1}{2})}, \eta_{(i-1,j-\frac{1}{2})}, \eta_{(i+\frac{1}{2},j-1)}, \eta_{(i+\frac{1}{2},j+\frac{1}{2})}, \eta_{(i+1,j-\frac{1}{2})}]. $$

(C2)

For $x$-interfaces, information about the maximum wavespeeds at the adjacent $y$-interfaces that could affect the Riemann solution are thereby incorporated. In three dimensions, the calculation of $\eta^H$ uses an appropriate nine-point stencil. The $\eta^H$ quantity is used in conjunction with the Roe Riemann solver when calculating the CTU fluxes for an interface ($\mathbf{F}^{n+\frac{1}{2}}, \mathbf{G}^{n+\frac{1}{2}}, \mathbf{H}^{n+\frac{1}{2}}$) by replacing the eigenvalues $\lambda^a$ in Equation [B28] with

$$ |\lambda^a| = \max(|\lambda^a|, \eta^H). $$

(C3)

This substitution serves to add crossflow dissipation to the one-dimensional Roe fluxes and mitigates the carbuncle instability, as seen in Figure 18. Using the H correction comes at an additional cost, as its stencil requires adding an extra ghost cell on each side of the simulation in every dimension.

REFERENCES

Aarseth, S. J. 1999, PASP, 111, 1333
Balsara, D. S. 2004, ApJS, 151, 149
Berger, M. J., & Colella, P. 1989, Journal of Computational Physics, 82, 64
Berger, M. J., & Oliger, J. 1984, Journal of computational Physics, 53, 484
Bryan, G. L., et al. 2014, ApJS, 211, 19
Chan, C.-k., Mitra, D., & Brandenburg, A. 2012, Phys. Rev. E, 85, 036315
Chandrasekhar, S. 1961, Hydrodynamic and Hydromagnetic Stability (Oxford University Press)
Colella, P. 1990, Journal of Computational Physics, 87, 171
Colella, P., & Glaz, H. M. 1985, Journal of Computational Physics, 59, 264
Colella, P., & Sekora, M. D. 2008, Journal of Computational Physics, 227, 7069
Colella, P., & Woodward, P. R. 1984, Journal of Computational Physics, 54, 174
Courant, R., Friedrichs, K., & Lewy, H. 1967, IBM Journal of Research and Development, 11, 215
Domínguez, J. M., Crespo, A. J. C., Valdez-Balderas, D., Rogers, B. D., & Gómez-Gesteira, M. 2013, Computer Physics Communications, 184, 1848
Einfeldt, B. 1988, SIAM Journal of Numerical Analysis, 25, 294
Einfeldt, B., Roe, P. L., Munz, C. D., & Sjogreen, B. 1991, Journal of Computational Physics, 92, 273
Einfeldt, B., Roe, P. L., Munz, C. D., & Sjogreen, B. 1991, Journal of Computational Physics, 92, 273
Forum, M. P. 1994, MPI: A Message-Passing Interface Standard, Tech. rep., Knoxville, TN, USA
Frigo, M., & Johnson, S. G. 2005, Proceedings of the IEEE, 93, 216, special issue on “Program Generation, Optimization, and Platform Adaptation”
Fryxell, B., et al. 2000, ApJS, 131, 273
Gardiner, T. A., & Stone, J. M. 2008, Journal of Computational Physics, 227, 4123
Godunov, S. K. 1959, Math. Sbornik, 47, 271
Harfst, S., Guandalris, A., Merritt, D., Spurzem, R., Portegies Zwart, S., & Berckzil, P. 2007, New Astron., 12, 357
Harten, A., Lax, P. D., & Van Leer, B. 1983, SIAM Review, 25, 35
Hayes, J. C., Norman, M. L., Fiedler, R. A., Borden, J. O., Li, P. S., Clark, S. E., ud-Doula, A., & Mac Low, M.-M. 2006, ApJS, 165, 188
Hui, W. H., Li, P. Y., & Li, Z. W. 1999, Journal of Computational Physics, 153, 596
Knebe, A., Green, A., & Binney, J. 2001, MNRAS, 325, 845
Kravtsov, A. V. 1999, PhD thesis, NEW MEXICO STATE UNIVERSITY
Kulikov, I. 2014, ApJS, 214, 12
Laney, C. B. 1998, Computational Gasdynamics (Cambridge University Press)
LeVeque, R. J. 2002, Finite Volume Methods for Hyperbolic Problems (Cambridge University Press)
Liska, R., & Wendroff, B. 2003, SIAM Journal of Scientific Computing, 25, 995
Noh, W. F. 1987, Journal of Computational Physics, 72, 78
Portegies Zwart, S., & Bédorf, J. 2014, ArXiv e-prints
Portegies Zwart, S. F., Baunegardt, H., Hut, P., Makino, J., & McMillan, S. L. W. 2004, Nature, 428, 724
Quirk, J. J. 1994, International Journal for Numerical Methods in Fluids, 18, 555
Robertson, B. E., Kravtsov, A. V., Gnedin, N. Y., Abel, T., & Rudd, D. H. 2010, MNRAS, 401, 2463
Roe, P. L. 1981, Journal of Computational Physics, 43, 357
Saltzman, J. 1994, Journal of Computational Physics, 115, 153
Sandalski, S. 2012, PhD thesis, California State University, Long Beach
Sanders, R., Morano, E., & Druguet, M.-C. 1998, Journal of Computational Physics, 145, 511
Schive, H.-Y., Tsai, Y.-C., & Chiueh, T. 2010, ApJS, 186, 457
Shu, C.-W., & Osher, S. 1989, Journal of Computational Physics, 83, 32
Sod, G. A. 1978, Journal of Computational Physics, 27, 1
Spurzem, R. 1999, Journal of Computational and Applied Mathematics, 109, 407
Stone, J. M., Gardiner, T. A., Teuben, P., Hawley, J. F., & Simon, J. B. 2008, ApJS, 178, 137
Strang, G. 1968, SIAM Journal on Numerical Analysis, 5, 506
Sugimoto, D., Chikada, Y., Makino, J., Ito, T., Ebisuzaki, T., & Umemura, M. 1990, Nature, 345, 33
Teyssier, R. 2002, A&A, 385, 337
Toro, E. F. 2009, Riemann Solvers and Numerical Methods for Fluid Dynamics (Springer)
van Leer, B. 1977, Journal of Computational Physics, 23, 276
—. 1979, Journal of Computational Physics, 32, 101
Wang, P., Abel, T., & Kachler, R. 2010, New Astronomy, 15, 581