PATCHWORK: A Multipatch Infrastructure for Multiphysics/Multiscale/Multiframe Fluid Simulations

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Abstract

We present a “multipatch” infrastructure for the numerical simulation of fluid problems in which subregions require different grid scales, different grid geometries, different physical equations, or different reference frames. Its key element is a sophisticated client–router–server framework for efficiently linking processors supporting different regions (“patches”) that must exchange boundary data. This infrastructure may be used with a wide variety of fluid dynamics codes; the only requirement is that their primary dependent variables be the same in all patches, e.g., fluid mass density, internal energy density, and velocity. Its structure can accommodate either Newtonian or relativistic dynamics. The overhead imposed by this system is both problem and computer cluster architecture dependent. Compared to a conventional simulation using the same number of cells and processors employed on a problem not requiring multipatch methods, the cell update per processor rate decreases by an amount that can range from negligible to a factor of a few; however, even in these problems, the infrastructure can permit substantial decreases in the total number of cell updates required.

Key words: hydrodynamics – methods: numerical

1. Introduction

1.1. Importance of Multiphysics/Multiscale/Multiframe Capability

Many important physical processes involve heterogeneous systems in which the nature of the matter in different regions exhibits strong contrasts. The material may vary in its characteristic internal length or timescales, or in its local geometric symmetry. There may even be contrasts in which the physical mechanisms of importance differ between regions: for example, chemical reactions or self-gravity may be significant in some, but not all, locations. These regions may also move with respect to one another, perhaps changing shape as they do. When the regions have relative motion, the fact that the physics is often most concisely described in a system’s mean rest frame means that no single rest frame is appropriate for the entire problem. At the same time, interactions between these regions may nonetheless demand simulation methods allowing data from one region to inform the behavior of another.

Problems exhibiting strong contrasts in length or timescales are called “multiscale problems.” We will also use this term to include contrasts in grid symmetry. In multiscale problems, numerical methods work best with different grid systems in different regions, perhaps contrasting in resolution, perhaps in symmetry, e.g., polar versus Cartesian. Those involving disparities in mechanisms are called “multiphysics problems.” In these problems, one must solve entirely different equations: those of magnetohydrodynamics (MHD) rather than those of hydrodynamics, or with or without transport processes such as viscosity or diffusion. We dub problems with internal frame shifts “multiframe problems.” For these, it would be desirable to translate the equations from one frame to another in different portions of the calculation. Astrophysics is rich in problems to which at least one, and sometimes all, of these labels apply, and therefore at least one, and sometimes all, of the difficulties, both technical and conceptual, that they pose.

To illustrate their significance, consider a few examples. The topic that initially motivated our work is the mechanics of accretion around a binary system. For us, the partners in the binary are supermassive black holes (see, e.g., Schnittman 2013), but little about this problem changes whether the binary comprises a pair of protostars (e.g., as imaged and analyzed by Mayama et al. 2010) or a pair of black holes. In this situation, there are widely disparate scales because the structure of the circumbinary disk varies on the scale a of the orbital separation, whereas most of the accretion power emerges at the inner edges of the disks orbiting around the individual masses (often called “mini disks”), which could be a great deal smaller. In addition, throughout these disks, even to define the saturation level of the MHD turbulence supplying the accretion torques requires treatment of length scales that are small compared to disk scale heights, which could be considerably smaller than the radial scale. There are also differences in the symmetry of well-designed local grids. Because angular momentum transport in accretion disks is slow compared to the orbital time, it is very important that there be little numerical momentum diffusion; this fact demands a grid mimicking the symmetry of their nearly circular flow (see, e.g., Sorathia et al. 2013). However, such a grid would be polar and centered on the binary center of mass for the circumbinary disk, whereas for each mini disk, it would be polar and centered on the object whose gravity is most important for that disk. A single Cartesian grid for the entire system would likely produce an intolerable level of numerical diffusion. This binary accretion problem is also one that demands multiple reference frames for much the same reason it requires multiple subgrids.

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with different symmetries. The physics of the circumbinary flow is easiest to grasp in the center-of-mass frame, which is that of the individual mini disks in the frame of each member of the binary. Thus, one would like to be able to divide this calculation into at least three different zones, each with its own grid and reference frame.

Another example may be found in the tidal disruption of stars by supermassive black holes, which has become a subject of great interest in recent years as numerous examples have been found (in both optical/UV, e.g., Gezari et al. 2009 and Arcavi et al. 2014, and in X-rays: Auchettl et al. 2017). This is a multiscale problem because it is necessary both to resolve the dynamics within the star as it is broken apart and to follow the fluid dynamics of the debris as it gradually accretes onto the black hole. Measured in terms of gravitational radii $r_g$ defined relative to the mass $M$ of the black hole ($r_g \equiv GM/c^2$), main-sequence stars are $\sim 1M_\odot r_g$ in diameter, where $M_\odot$ is the black hole in units of $10^6 M_\odot$. Thus, to follow their breakup requires cells $\ll 0.1M_\odot r_g$ in size. On the other hand, the debris orbits have semimajor axes $\sim 10^3M_\odot r_g$, so that the fluid motion after stellar breakup takes place on a much larger scale. Nonetheless, despite this dramatic scale contrast, the breakup of the star is inextricably tied to the much larger-scale debris motion. It is also a multiphysics problem because stellar self-gravity, not surprisingly, is of the essence so long as the star stays in one piece, but after its matter is spread sufficiently widely, it becomes inconsequential. It is also a multiflare problem because the mechanics of a nearly hydrostatic star are definitely best viewed in the star’s frame where the fluid velocities are small, whereas the mechanics of an accretion flow are far more easily understood in the black hole frame. Its multiflare nature also creates a contrast in grid symmetry because coherent stars are most naturally treated in a spherical coordinate system whose origin is the center of the star, whereas flow around a black hole is best described in a cylindrical or spherical coordinate system whose origin is the center of the black hole. Thus, this problem, too, involves all of these categories of complication. In fact, we have chosen it as the subject of our first test problem for our new infrastructure. We will quote a few technical results from this test problem here; a full analysis will be published separately.

1.2. State of the Art and Its Limitations

The desirability of overcoming these challenges has not gone entirely unnoticed by the computational community, and a number of partial solutions have been developed. Adaptive mesh refinement (AMR) methods can dynamically adjust spatial resolution to follow local length scales (Berger & Oliger 1984; Berger & Colella 1989); a closely related scheme, overlapping moving grids, can be used to follow a coherent region with a distinct spatial scale or symmetry. MultiProgram/ MultiData (MPMD) methods (Barney 2017) offer a convenient way to evolve different regions according to the different mechanisms acting in them. We will call these separate regions “patches,” hence the name “multipatch” for our general approach. Because each patch is run by an independent program under MPMD, their only interaction is through the exchange of boundary conditions.

For some of these methods, professionally supported implementations are available. Chombo, for example, is a particularly well-developed AMR package (Adams et al. 2014). It permits the use of two different methods to divide up regions into separate grids, embedded boundaries, and mapped multiblocks. There are also relativistic versions of these fixed multiblock methods (Schnetter et al. 2014; Clough et al. 2015). General relativistic hydrodynamics has been treated (Blakely et al. 2015) using the techniques of Overture (Brown et al. 1997), which offers users the option of moving overlapping grids. In this approach, a composite grid is formed in order to bridge two contrasting overlapping grids, with data from each interpolated onto the composite grid (Chesshire & Henshaw 1990). Numerical relativity calculations can use multiblock infrastructure with AMR, but all dependent variables must be defined in terms of a global Cartesian tensor basis (Pollney et al. 2017). Similarly, there are numerous MPMD systems permitting the computation of different physics in different parts of a global system. These range in their applications from linking multiscale fluid simulation to molecular dynamics (Nie et al. 2006) to modeling blood flow through the brain (Grinberg et al. 2013).

Another approach to solving the problems of multiple scales, but not multiple physics, is the use of moving unstructured grids (e.g., the codes AREPO, Springel 2010, and TESS, Duffell & MacFadyen 2011). Schemes like these very flexibly place resolution where it is required for the hydrodynamics. It has also recently become possible to extend them from hydrodynamics to magnetohydrodynamics (Duffell 2016; Mocz et al. 2016). They do not, however, naturally retain the virtues of conforming to natural symmetries of the problem (e.g., suppressing numerical diffusion by aligning cell axes with the fluid velocity), nor do they readily permit the use of contrasting physics in different regions. With significant effort, it is possible to avoid the first drawback (Duffell 2016), but a new solution must be created for each new problem.

Despite the real successes of all these different schemes, there remain significant barriers to their employment in many kinds of problems. Multiblocks must fit smoothly against one another in a fixed configuration, while embedded boundaries require Cartesian grids. Neither of these allows the relative motion of the cell blocks. Most importantly, none of the methods introduced so far achieves the simplification and efficiency gains that arise from following the physics of moving regions in their own reference frames.

The advantages of working in the most suitable reference frame can be substantial. Consider, for example, a hydrodynamics problem in which structure A, containing only motions subsonic relative to its own center of mass and varying on short length scales, moves supersonically within a larger background fluid B with longer gradient scales. If such a problem were treated with a moving grid scheme, the time step within region A would be severely limited by its supersonic velocity and its small cell sizes; transformation to the moving frame could reduce the number of time steps required by a large factor. Numerical accuracy would also be substantially improved as there would be no need to perform numerous
close subtractions of velocities in order to find the relative velocities between cells.

Analogous advantages can stem from equation simplification. Suppose, for example, that in the moving region there is a diffusive transport process that is unimportant in the background. Treating this transport process in the moving frame eliminates what would otherwise be a large, unnecessary, advective flux. If the velocity contrast between the regions approaches the relativistic level, treating everything in the background frame introduces serious conceptual problems: because classical diffusion causes instantaneous transmission of information (Morse & Feshbach 1953; Narayan 1992), such problems cannot be formulated covariantly. Transformation to a frame in which local motions are slow permits a clean use of the local Newtonian limit in which diffusive transport is mathematically consistent.

The limitations of existing methods severely crimp the study of many interesting problems, including the two we mentioned as motivating our work, the dynamics of accretion flow in binary systems and tidal disruptions. In the former case, none of the existing systems supports the optimal grid geometries, a pair of small spherical grids moving with respect to a larger (and coarser) spherical grid whose origin (the system’s center of mass) is stationary. Instead, one would be forced to cover a large region with small cells, none of them aligned with fluid motions and therefore incurring large numerical diffusion. Likewise, in none of them does one gain the advantage in numerical accuracy afforded by working in a locally co-moving reference frame. It is also difficult in this framework to avoid much of the simulation being burdened by a very small time step relevant in only a subregion. Simulations have been attempted, but they have been either restricted to 2D (Bowen et al. 2017; Ryan & MacFadyen 2017) or, if 3D, limited to very short duration (Bowen et al. 2018). The restrictions are even more severe for the tidal disruption problem. A number of simulations have been carried out whose grid origins follow the center of mass of the star (Cheng & Evans 2013; Guillochon & Ramirez-Ruiz 2013; Cheng & Bogdanović 2014; Guillochon et al. 2014), but they cannot continue to follow the career of the tidal debris during the much longer time that it orbits the black hole because their high-resolution grids are extremely inefficient for that problem. It is also important to compute the star’s self-gravity while it survives and before its tidal debris disperses, but then turn it off afterward for efficiency, yet no existing system provides the flexibility to ignore the regions where self-gravity is unimportant. These two astrophysical problems demonstrate the need for greater flexibility, but it is easy to imagine many other problems for which current methods are inadequate.

1.3. Our Innovation

The system we present here, which we call PATCHWORK, is designed to eliminate the limitations due to the use of a single reference frame for all patches while also maximizing the ability to simultaneously deal with issues of multiple scales, multiple grid symmetries, and multiple varieties of local physics. Exploiting the flexibility of the MPMD approach, it utilizes well-defined coordinate transformations informed by relativistic methods (but not restricted to relativistic problems) to simulate heterogeneous systems in which regions requiring independent treatment are regarded as independent processes operating in independent reference frames. Each patch has its own grid, with its own resolution scale and symmetry; among the many benefits offered by independent local coordinate systems, patches give an easy solution to the problems raised by coordinate singularities (e.g., at the origin of a polar system). Each patch also solves its own equations, whatever is necessary to do the job in that region.

The relationships between these frames are defined by coordinate transformations with the ability to eliminate local mean velocities, so as to reap the benefits just described, while retaining a common overall time so that the entire simulation can advance together. This approach creates an important simplification—the same coordinate transformation that relates the array index space to spatial coordinates can also be used to eliminate bulk velocities. This single coordinate transformation also applies to the metric with respect to the coordinates. In addition, provided that the physical quantities entering into the boundary conditions are scalars, vectors, or rank-2 tensors, their transformation from the coordinate system of one region to that of another is well defined and straightforward. In this way, the free use of arbitrarily curvilinear and arbitrarily discretized coordinates can be combined with the virtues of treating the local physics in its most natural reference frame.

However, in order for the patches to exchange boundary conditions at simultaneous times, the time coordinate in all of the patches must be the same. In relativistic terms, this means that the coordinate transformations relating patches with relative motion are not Lorentz transformations; as a result, the reference frames of moving patches are in general non-inertial. This policy may be somewhat unfamiliar, but because the equations of physics can all be written in completely covariant fashion, their form under this sort of transformation is well defined.

Our version of the multipatch system also offers several additional features. Because the patches interact only through boundary condition exchange, they can have independent time steps; because long time-step patches need many fewer updates to traverse the same physical time, when parallelized, those patches can be assigned many more cells per processor to achieve better load balancing. In addition, patches can be added or removed from time to time as conditions change and different demands arise.

To mitigate the complexity and overhead created by interpatch communications when the computation is parallelized, we have created a client–router–server system that efficiently links the correct processors in each patch to their boundary condition partners in other patches.

Lastly, our package is structured as a “wrapper” to fit around a user-supplied hydrodynamic or magnetohydrodynamic simulation code. The only requirements placed upon these codes are that they should have fixed grids and their primary dependent variables (the ones whose boundary conditions are exchanged between patches) are all the same. In intrinsically conservative codes (e.g., HARM3D, our test code), boundary conditions are generally fixed through the “primitive variables” (density, velocity, internal energy) because the Riemann problem is defined in terms of them. Other algorithms may lead to other choices, and PATCHWORK should accommodate them. Here, we illustrate its performance only through HARM3D.

1.4. Outline of the Paper

In Section 2, we set out the principal features of our method. This presentation begins with an overview (Section 2.1) in
which we define what we mean by a “patch” and describe how different patches are related to one another. The next subsection discusses the principal operation of the multipatch system, the boundary condition exchange between neighboring patches. Following that, in Section 2.3, we detail how boundary data are interpolated from one patch’s grid to another’s. Section 2.4 briefly discusses how users can add new patches and remove old ones. Then, Section 2.5 presents the method’s core: the client-router-server architecture we created so that, in a parallelized environment, a processor in one patch is linked to the correct partner in a different patch in order to exchange boundary data. The final subsection, Section 2.6, explains how different patches can advance with different time steps yet remain synchronized. We lay out in the Appendix an overview of how these operations are organized into discrete routines and fluid codes can, with a small number of additional lines of code, be made compatible with the PATCHWORK system.

Section 3 presents a variety of tests of our multipatch implementation. In its first subsection, we demonstrate that a shock can pass smoothly from one patch to another without alteration. In the second, we show that even when patch symmetries contrast strongly, a blast wave can travel from one to the other and remain close to the Sedov–Taylor similarity solution. In the third, we examine the degree to which interpolation of data from one patch to another may lead to departures from rigorous conservation of mass, momentum, and energy, and discuss how such departures can be kept small.

Section 4 discusses computational efficiency by presenting benchmarking tests and scaling data for the overhead imposed by the multipatch system.

The last section summarizes the paper.

2. Method

We have designed our multipatch software to be compatible with any numerical simulation code in which the computational domain is discretized into a fixed set of discrete grid cells and the primary dependent variables, the ones exchanged between patches as boundary conditions, are the same in every patch. It also requires the geometric factors relevant to the operators of vector calculus to be defined in terms of metric elements; this is done as a matter of course in relativistic codes, but it is also a feature of a number of contemporary Newtonian codes such as Athena++ and the most recent version of Zeus (Sorathia et al. 2013; White et al. 2016). For development and testing purposes, we have used it with the finite volume general relativistic hydrodynamics code HARM3D (Noble et al. 2009) running in every patch. In the near future, we plan to port it to other codes to demonstrate its flexibility. We expect that, subject to the stipulation about variable consistency, it will be possible for different codes to run in different patches.

2.1. Overview

PATCHWORK’s structure is based on the concept of “patches.” A patch is a region of space defined by the user. Locations within it are described by its particular coordinate system and discretized according to its own particular grid. The time evolution of its fluid’s physical properties is governed by a particular set of equations, always including the Euler fluid equations, but potentially extensible to the Navier–Stokes equations or the MHD equations, and potentially supplementable by chemical or nuclear reaction networks, a Poisson solver for self-gravity, or other sorts of equations. Evolving an individual patch is the responsibility of an individual process within the MPMD environment. Although it is possible to run the simulation as a single program, using one program for each patch keeps the code simple and conceptually clear. As a result, the method is intrinsically parallelized: there must be at least one processor for each patch.

PATCHWORK coordinates a number of different individual patch processes through the incorporation of several specific routines into the fluid simulation code chosen by the user. Some are problem independent, but others are problem specific and therefore need to be written by the user. The most important functions of the PATCHWORK routines are to define the trajectory of each patch in terms of the “background coordinates” (see below for definition), calculate the coordinate transformation matrices necessary to translate physical quantities and locations between each patch and the background coordinates, control boundary condition exchange between different patches while maintaining “situational awareness” about which portions of a patch’s boundary adjoin other patches and which are on the edge of the physical problem volume, and synchronize the time steps in the different patches. In addition, there are several other optional multipatch-specific routines that will be described later.

A system of “background coordinates” underlies the entire region being simulated. The locations and boundaries of all the patches are defined in terms of this system. It is always Cartesian, and its time coordinate is the universal time for all of the patches’ coordinate systems. Its purpose is both to serve as a reference for positions and to serve as a “common language” for all patches to describe the locations of exchanged data.

Individual patches can have any shape or size, provided only that they fit within the background coordinate grid. They can be stationary relative to the background coordinates or move. Their internal coordinate systems and grids are entirely independent of all other patches’ spatial coordinate systems and grids. It is convenient in many problems to divide the patches into two categories, “global” and “local.” Frequently, one patch provides the great majority of boundary condition data for the other patches and occupies all or a large part of the problem volume. When that is the case, that patch is deemed “global,” and its reference frame is tied to the frame of the background coordinates. Its internal spatial coordinate system, however, can still be defined independently of the background coordinates. Although it is often convenient to have a global patch, it is not a requirement of the system. Any patch not designated as “global” is considered to be “local.”

When two or more patches overlap in their spatial coverage, only one of them governs the dynamics within the overlap volume. We then speak of the “active patch” updating the properties of the “uncovered cells” and the “inactive patch” containing the “covered cells.” If one patch is a local patch and the other is a global patch, the local patch is always the active one. If two or more local patches overlap, the user designates the hierarchy of activity in advance. As covered cells approach the patch boundary of the active patch, they become ghost cells for other cells in their patch that are already uncovered. At that point, they are filled by interpolation from the active patch cells covering them.

One spacetime is specified for the entire problem volume with a metric defined on the background coordinates. This spacetime can be described in any of the patches by means of
the appropriate coordinate transformation from the background system to the patch coordinate system. In some instances, the fluid mass in one or more of the patches may be important to gravity throughout the problem volume. The best way to account for this contribution to gravity depends on circumstances. Relativistic velocities or strong gravity demands solution of the Einstein field equations. Because these equations, like the hydrodynamics equations, are hyperbolic, they can also be solved within the multipatch framework. On the other hand, if the fluid moves more slowly and the gravitational field is weak, the Poisson equation, which is elliptic, is appropriate. This case requires the global patch physics repertoire to include Newtonian self-gravity, and the density distribution from any local patch with significant mass must be interpolated to the global grid, although possibly with crude resolution.

Physical consistency likewise demands that all patches are updated according to the same time coordinate and must reach a given value of this time coordinate together. Such synchronization is achieved automatically if all advance with the same time step. However, as we discuss below (Section 2.6), this is not necessary. If some patches can be evolved stably and accurately with a longer time step than others, it is necessary only for the patches to all be synchronized after one time step of the patch with the longest step. Note that because there is a single time coordinate for all patches, the coordinate transformations between them are not Lorentz transformations unless the relative velocity between the two patches being linked is zero and both patches are inertial.

In the course of each update, patches bordering on one another must exchange boundary condition data. Accomplishing this step is the core of our system.

2.2. Boundary Condition Exchange between Patches

Within any particular patch, we distinguish three types of boundary zones for individual processors. Ghost zones covered by other processors in the patch are in the first category. The second category comprises ghost zones lying on the physical boundary of the problem. The third category, those ghost zones covered by processors assigned to other patches, is of greatest interest to the multipatch scheme.

The first two can be handled by the standard devices found in existing fluid codes. Here we describe how the boundary information is obtained for ghost zones in the third category. We begin by displaying an example so that readers can easily visualize the issues involved (Figure 1). In this figure, the fluid’s internal energy density is represented by color contours, and grid cells are delineated by black lines. We have chosen to follow the fluid mechanics in this example by means of two patches, a finely resolved Cartesian local patch and a more coarsely resolved polar global patch whose radial grid is logarithmically spaced. In the upper panel of the figure, the physical boundary of the Cartesian patch is shown by the inner white box; the area covered by its “ghost zones,” the cells needed to establish boundary conditions for the physical region, lies between the two white boxes. The lower panel shows the converse situation: the jagged white contour shows the boundary of territory in the global patch not covered by physical cells of the local patch; the cells between that jagged contour and the white cells are where the global patch needs boundary data.

![Figure 1](image.png)

The first step is to discover which processors in which patches have the information. To minimize interpatch communication time, we organize this process to avoid exchanging unused data. Because this procedure is almost independent of whether the patch needing boundary data is a global or a local patch, for this part of the discussion, we call them “patch A” and “patch B.” We begin by labeling all the zones in patch A (here this happens to be the global patch) with an integer array, illustrated in Figure 2. The values in this “flag array” denote...
whether a zone is a ghost zone, and if so, what type of ghost zone. This array must be updated at each time step if any of the relevant patches move (at each synchronization time step in the case of heterogeneous time steps; Section 2.6). In the figure, the white zones are in the interior of patch A and have nothing to do with boundary conditions. Gray zones are the zones in patch A completely covered by patch B; they, too, are irrelevant to boundary conditions. A zone in the global patch is considered to be covered by the local patch if its center falls within the local patch’s physical region. The red and blue zones are the covered cells in patch A that act as ghost cells for the uncovered cells in patch A. The red cells directly touch uncovered patch A cells. Fluxes across their inner (in a topological sense) faces are used in the updates of the uncovered cells they touch. Blue zones are the outer layer of ghost cells needed for the updates of the uncovered cells in patch A; in HARM3D, the ghost-cell zone is three cells wide, so the blue cells are either the second or third ghost cell from the last uncovered patch A cell along at least one dimension. They are used in the internal reconstruction by which cell-center values of fluid quantities are extrapolated to the face touching the physical boundary. As long as the number of ghost cells is adjusted appropriately, any reconstruction method should in principle work with PATCHWORK. In the tests presented here, we used piecewise parabolic reconstruction (Colella & Woodward 1984) with an MC (monotonized central-differenced) slope limiter.

It is important to note that this system is thoroughly agnostic about many of the possible choices made in different codes. Because the coordinates at which the boundary data are needed are determined by the fluid code operating in the requesting patch, it does not matter whether that code defines the variables at cell centers or at the centers of cell faces or anywhere else; it knows the locations at which it needs the information, and it is the job of the responding code (which may be an entirely different one) simply to interpolate its data, no matter how defined in terms of location within cells, to the proper point. The system is even capable of accommodating codes with different numbers of ghost cells. HARM3D, for example, requires three layers of ghost cells, but PATCHWORK contains a parameter that can be set to whatever number of layers the user’s code needs.

Once patch A determines the locations of its ghost zones’ centers, a list of the background coordinates for these locations is sent to all other potential patch B’s along with a request for interpolated values of the fluid variables at the coordinate locations. Patch B interpolates within its grid in order to find the values at the locations desired by patch A. It then transforms the data from its coordinate system to the background coordinate system, using the coordinate transformation Jacobian linking patch B to the background system. Only then are the boundary data transmitted back to patch A, which transforms it from the background system to its own coordinates. This procedure enables every patch to deal with the incoming coordinate list independently without knowing anything about other patches. Doing things this way is especially important when patch B moves.

Note that if patch A is the global patch, the first step is done differently. At initialization, the local patches are informed of the cell locations at which the global patch dependent variable data are defined. Because the local patches know their own positions in terms of background coordinates, they can determine on their own what data the global patch needs. This alternate procedure has the virtue of diminishing interpatch data transmission.

### 2.3. Interpolation

Although remarked on only briefly in the preceding subsection, there are a number of subtleties to data interpolation, and multiple mechanisms may be used. In the current version of our system, we use a comparatively simple method, but this could readily be upgraded to something more sophisticated for problems requiring it.

In principle, an arbitrary number of zones could be used to support interpolation to a single point. However, it is generally best for the interpolation stencil to extend away from the point by a number of zones that is no more than the number of ghost-zone layers (usually two to three), so that the stencil does not extend into another processor’s domain.

For our current method, we employ trilinear interpolation. We locate the grid corner closest to the interpolation point and define the stencil in 3D to be the centers of all eight cells touching that corner. This method works quite well when the dimensions of the cells in patch A and patch B are comparable (see Section 3.3), but can lead to errors when they are not. In some sense, this is unsurprising: if there is structure on the finest scale supported by one of the patches, it cannot be well-represented by a much coarser grid in the other. However, the trouble can also move in the opposite direction because the eight cells in the finer grid nearest the interpolation point may together cover only a small part of the volume of the ghost zone in question if its grid is much coarser. Sometimes errors of this latter variety can be substantially reduced by replacing the values in the inner layer of ghost cells with a wider average.

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8 If patch A were a local patch, it would have gray zones only if patch B were another local patch and patch B took priority over patch A; we have not yet implemented “local–local” boundary data exchange, but plan to do so soon.
over nearby cells. Such an operation effectively magnifies the volume of the finer-scale grid contributing to the coarser-grid ghost cells.

Without special methods, interpolation does not necessarily conserve quantities. To achieve strict mass (or momentum or energy) conservation in our data interpolation could require identifying all cells that fall within the ghost cell and summing their contributions. If the ghost-cell boundaries cut obliquely (or even worse, in a curve) across some of the interpolation cells, one would need to adjust their volumes accordingly. Although this is possible if both global and local patches are in Cartesian coordinates, it becomes a nontrivial mathematical problem once any of the patches are in curvilinear coordinates. In Section 3.3, we test quantitatively how closely our interpolation method comes to conserving mass and momentum.

2.4. Adding and Removing Patches

Stationary or moving patches can be added or removed throughout the simulation anywhere within the physical problem volume. This is done using the flag array for the ghost zones discussed in Section 2.2. Although these flags are most often used to signal the need for data interpolation from overlapping patches, they can also be used to signal the need to interpolate data for other reasons as well—such as removing or introducing a new patch. To remove a local patch, one temporarily changes the flags on all the zones in the global patch covered by the local patch to "ghost zones" so that all of them are filled with interpolated data provided by the local patch. Once these zones are filled with data, one changes the flags back to their normal state. To add a local patch, one creates a new patch process, and to define its initial condition sets all its cell flags to indicate they are ghost zones. Just as in the patch removal operation, these cells are then filled with the data they need, and the flags can be reset to normal as soon as that is done. However, the simulation must be stopped immediately after a patch removal or immediately prior to a patch addition because either one demands a new domain decomposition for processor assignment.

In principle, patches could be added or removed while running. To do so, however, requires having a clear criterion for when to make the change, a specific plan for the reallocation of processors because MPMD does not permit any change in the total number of processors while running, and synchronization of the resumption of fluid updates between all processors and patches. For the time being, we have not implemented such a scheme.

2.5. Parallelization and Interpatch Communication

One of the most difficult tasks in developing a multipatch code is its parallelization. It requires a sophisticated infrastructure combining two levels of data communication. In one level, boundary data exchange within a patch, a single executable exchanges information between its multiple processors exactly in the way made familiar by non-multipatch parallelized methods. In the other level, boundary data exchange between patches, it is necessary to enable effective data communication when the pairing of processors with overlapping boundaries evolves dynamically, and two independent executables, both running within the MPMD environment, must be coordinated.

To describe how we achieve this, we first define a notation. We label each CPU in a simulation by \( C_{ij} \), where \( i \) is a patch ID and \( j \) denotes the local CPU rank within that patch. We set the patch ID of the global patch to \( i = 0 \) and that of local patches to \( i = 1, 2, \ldots, N - 1 \), where \( N \) is the total number of patches in the simulation, including the global patch. The index \( j \) runs from 0 to \( n_i - 1 \), where \( n_i \) is the number of CPUs used for patch \( i \).

Consider a CPU at the edge of a patch, designated \( C_{i0} \). This CPU possesses boundary zones that need to be filled with interpolated values. It needs to know which CPUs \( C_{jk} \) in other patches handle cells lying under these zones (there could be multiple values of \( j \) satisfying this criterion, and sometimes multiple values of \( k \)). It must also contact them to request interpolation values. The partner CPUs \( C_{jk} \), on the other hand, need to know in advance that other CPUs may be contacting them. Because these relationships constantly change if the patches move relative to one another, this information must somehow be updated dynamically, even though the patches may have differing time steps.

To solve this problem, we construct a client–router–server system, setting up interpatch communication relationships that can persist throughout the simulation. In its simplest form, one CPU in each patch is chosen to serve as the router, its liaison with all the other patches. Then, when client CPU \( C_{i'} \) needs information from beyond the boundary of patch \( i \), it transforms the coordinates of the cell centers in question to background coordinates and broadcasts that list to processors \( C_{k'j} \), where the \( r_k \) processor in patch \( k \) is the designated router for that patch. The router processor in the \( k \)th patch then transforms the list from background coordinates to patch \( k \) coordinates. If all of the cell centers on the list lie outside patch \( k \), the router replies accordingly. On the other hand, if some of them are inside patch \( k \), the router processor determines which of the other processors working on patch \( k \) have responsibility for those cells and distributes the request to those processors. These processors, the servers, interpolate their data to the correct positions, transform the results to background coordinates, and return the results to the router. Finally, the router transmits the information back to the client, CPU \( C_{i'} \).

This communication scheme is conceptually simple and easy to code. However, if only a single CPU is given router duties for an entire patch, the communication load is shared very unevenly and the great majority of processors sit idle while waiting for the routers to finish their work. To divide the workload more evenly, we regard all processors as potential routers for their patch and redefine the client–router relationship uniquely for each individual CPU (see Figure 3). These relationships are defined at the beginning of the simulation and remain unchanged unless patches are added or removed. For example, one may decide that \( C_{i0} \) always contacts \( C_{0j} \) for any information regarding patch 0, \( C_{i1} \) always contacts \( C_{0j} \), and so on. The function of the router is unchanged; it still determines which, if any, of the processors on its patch holds the information requested and acts as the go-between connecting clients and servers.

Although the varying numbers of processors per patch make an exactly even division of labor impossible, a simple assignment scheme can spread it in a reasonably even-handed manner. If \( C_{i'j} \) requires information regarding patch \( p_k \), it
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D. N. K. L. is the smallest radius in patch instance of data exchange, a client CPU in a local patch, and enlarged for visualization of the information exchange system. The patches reside inside the global patch, but they are placed outside the global requests are shown by red arrows, data returned by blue arrows. The local routers, and servers are marked with C (patch relevant server CPUs and transmits it back to the client. Even while these two CPUs on its patch to collect the requested data and returns the data to its client.

Simultaneously, CPU in patch 1 CPUs communicate with their partners in the global patch, it is possible CPU C1, also requests data from the global patch, working with its global patch router C10, which in turn collects the information from the relevant server CPUs and transmits it back to the client. Even while these two patch 1 CPUs communicate with their partners in the global patch, it is possible for a CPU in the global patch, for example, C10 (lower right in patch 0), to be a client, requesting data from other patches such as patch 2; in this case, the router is C2.

Contacts

\[
\text{router} - p_2(C^{p_1}) = C^{p_1 \mod n_{p_2}}.
\]

Note that CPUs on patch p2 could have 0 or multiple clients on patch p1, depending on their index, \(n_{p_1}\), and \(n_{p_2}\).

2.6. Heterogeneous Time Steps

One of the common problems in simulating multiscale systems with grid-based hydrodynamics codes is that the time step of the entire computational domain is limited to a small value by a few regions with small grid cells and high characteristic fluid velocity. As a result, the remainder of the simulation, where the intrinsic timescales can be much larger, is required to integrate with unnecessarily short time steps, leading to a large computational cost. However, the multiramp method, in which different regions are updated by independent processes, allows each patch to have its own time step while nonetheless evolving the system in a fashion synchronized across all patches. We call this mode of operation “heterogeneous time steps”, in contrast to the simpler “homogeneous time-step” mode in which all patches are forced to have the same time step.

Heterogeneous time steps can be managed with great flexibility. The only restriction placed on the time steps in different patches is that the update times should all be synchronized at intervals equal to the longest of the time steps, \(\Delta T \equiv \max_k(\Delta t_k)\), where, as before, \(k\) is an index labeling the different patches. To optimize computational resource use, before initiating a run, the user adjusts the number of CPUs assigned to each patch so that the wall-clock time to advance by a time \(\Delta T\) is approximately the same for all patches.

In practice, the coordination works as follows. At the nth synchronization time \(t^{\text{sync}}_n\), the different patches exchange boundary condition data. They then also exchange information about their time steps so that they can determine which is the longest and in which patch it is found (call that patch \(K\)). If all of the other patches receive their boundary data either from patch \(K\) or the problem boundary, the next synchronization time is set to be \(t^{\text{sync}}_{n+1} = t^{\text{sync}}_n + \Delta T\). If \(\Delta T\) is a factor \(Q_l(>1)\) by definition) larger than the time step \(\Delta t_l\) in some other patch \(l\), patch \(l\) performs \(\sim [\{Q_l\}]\) updates, where \([x]\) is the greatest integer \(\leq x\), while patch \(K\) works to advance to \(t^{\text{sync}}_{n+1}\). When patch \(l\) reaches a time \(t'\) such that \(\Delta t_{n+1} - t' < \Delta t_l\), \(\Delta t_l\) is reset to \(\Delta t_{n+1} - t'\) to achieve synchronization. Patches that have arrived at \(t^{\text{sync}}_{n+1}\) before the rest of the patches wait until all have reached it. When that has been achieved, the cycle is repeated. Because, by definition, conditions in patch \(K\) change by at most a modest amount over a time \(\Delta T\), it is unnecessary for the other patches to receive boundary condition information from it during their individual time steps within the interval \(\Delta T\). However, when there are more than two patches, it is possible that some patches may require boundary data from other patches whose time steps are shorter than \(\Delta T\). When that occurs, that pair must exchange boundary data at times determined by the longer of their two time steps. Note that processors within the same patch trade boundary data in the usual way at each of that patch’s internal time steps.

If all of the patches are solving the same equations, optimal load balancing can be achieved when patch \(K\) can be identified with reasonable reliability in advance, and the ratios \(Q_l\) can similarly be estimated. If those criteria are met, all that is necessary is to assign processors in patch \(l\) a number of cells \(N_l \approx N_k/Q_l\). Depending on system architecture, this simple load-balancing method may be constrained by the total memory available to processors supporting large numbers of cells.

In some situations, the \(Q_l\) might be essentially fixed throughout the simulation. For example, this would be the case in a simulation of gas dynamics in an isotropic gravitational potential in which the patches are nested spherical shells. In such a situation, the time step for each shell would always be \(\approx \frac{\Omega(k, r)}{r_{\text{min}}k} / 2 \pi \), where \(N_k\) is the number of azimuthal cells in patch \(k\), \(\Omega(r)\) is the orbital frequency as a function of radius, and \(r_{\text{min}}\) is the smallest radius in patch \(k\). In such a case, load balancing could be achieved fairly reliably and would need no adjustment during the simulation.

More often, however, the \(Q_l\) may vary as functions of time. Because the MPMD environment does not permit the dynamic reassignment of processors from one program to another, when this condition obtains perfect load balancing through the adroit assignment of processors to patches will nearly always be an unreachable goal. Nonetheless, as we show in Section 4, even approximate load balancing by combining an appropriately chosen number of processors per cell in each patch with heterogeneous time steps can lead to significant gains in computational efficiency relative to homogeneous time-step
operation. These gains can be sustained even if the ratios $Q_i$ change significantly through the simulation if the user periodically stops the simulation and restarts with an adjusted choice in numbers of processors per patch.

3. Physics Tests

In this section, we showcase the performance quality of the multipatch method. The tests appropriate to this system are different from those useful to verify fluid codes because the multipatch infrastructure does not directly update fluid quantities; rather, it transfers results from one region to an adjacent one. Consequently, the focus of our tests is PATCHWORK’s ability to bridge patches without undermining the quality of the underlying code’s solution of the fluid problem.

That the issue is enforcing consistency between patches rather than the quality of the solution within individual patches explains why we do not present special tests of the method’s multiphysics capability. Nearly all examples of local physics (e.g., viscosity, different equations of state) affect the way the fluid state variables (mass density, internal energy density, velocity/momentum density) behave, but do not change which variables are transmitted from patch to patch. Consequently, if our system works for a homogeneous physics example, it works just as well for a multiphysics example. Even if the magnetic field is important to the physics in one patch but not others, it must be the case that for some reason (e.g., high resistivity) magnetic fields weaken greatly near the borders of that single patch. There is then no need to transfer magnetic field data to the other patches because it is not relevant to them.

First we demonstrate that it accurately reproduces the analytic solutions to two classic hydrodynamic simulation test cases even when critical features of these solutions pass through patch boundaries, and the grid symmetries and resolutions of the patches differ sharply. We then explore how well non-conservative interpolation maintains conservation of mass, momentum, and energy, and identify the conditions in which it does not.

3.1. Sod Shock Tube

In this test, we demonstrate that patch boundaries create no significant artifacts when shocks and rarefaction waves travel from one patch to another. For this test, we created a square planar problem volume in which, following the Sod prescription (Sod 1978), the fluid is initially at rest everywhere, but there is a sharp pressure and density discontinuity at a specific value of $x$ within the volume. There is no initial variation as a function of $y$. Within this volume, we placed a local patch and gave it a constant velocity so that it moves diagonally in the $xy$ direction. We performed four runs to demonstrate the code’s performance in a variety of coordinate system configurations. In all, the background spacetime is taken to be Minkowski.

In three of these, the coordinates for both the global and local patches are Cartesian, while the fourth uses Cartesian coordinates for the global patch and cylindrical for the local patch. All Cartesian–Cartesian configurations have aligned local and global grids, and in all three, the global patch cells are eight times the size of the local patch cells. Because the cylindrical–Cartesian test is designed to explore sensitivity to grid symmetry contrast rather than grid-scale contrast, the uniform cylindrical grid of the local patch has cells comparable in size to the Cartesian cells of the global patch.

In two of the Cartesian–Cartesian tests, the local patch moves slowly relative to the global patch; in one of these tests, the local patch is placed so that the shock passes through it, while in the other the local patch is placed where the rarefaction wave runs across it. In the third Cartesian–Cartesian test, the discontinuity runs through the global patch at the start, but the local patch travels rapidly enough to run through the rarefaction wave, the contact discontinuity, and the shock, and then emerge on the far side. The problem solved in the cylindrical–Cartesian test is similar to the third Cartesian–Cartesian test in that the discontinuity starts within the local patch, while the local cylindrical patch moves at the slower velocity used in the first two Cartesian–Cartesian tests.

Because HARM3D is framed in terms of relativistic dynamics, it is convenient to choose $c$ as the unit of speed. Given arbitrary code-units of length $\ell_0$ and mass density $\rho_0$, the unit of time is $\ell_0/c$ and the unit of pressure is $\rho_0 c^2$. To ensure Newtonian flow (Hawley et al. 1984), it suffices to make $p/c^2 \ll 1$ when measured in code-units. We also chose an adiabatic index $\gamma = 1.4$.

Our problem volume was 40 code-units on a side. For the Cartesian–Cartesian tests, there were 400$^3$ cells in the global patch, each with dimension $0.1 \times 0.1$. The three Cartesian local patches had side lengths of 8 and were cut into 640$^3$ cells of dimension $0.0125 \times 0.0125$, so that each cell was $1/8$ the size (per dimension) of those in the global patch. For the cylindrical–Cartesian test, the global patch had 800$^3$ cells, each $0.05 \times 0.05$, while the local patch’s cylindrical grid consisted of 240 uniform cells over $2 \leq r < 8$ in cylindrical radius and 1000 uniform cells over the full azimuthal extent. The cylindrical local patch requires a cutout at its center so as to avoid the coordinate singularity at the origin associated with polar coordinates. In this cylindrical grid, the largest azimuthal cell size was approximately equal to the global patch’s cell size while the radial cell width was half the global patch’s cell size.

For the initial state, the gas was divided into left (L) and right (R) states, with density and pressure $\rho_L = 1.0 \times 10^{-4}$, $p_L = 1.0$ and $p_R = 1.25 \times 10^5$, $\rho_R = 0.1$. The sound speed was therefore $\approx3 \times 10^{-3}$ on both sides, clearly subrelativistic. Zero-gradient boundary conditions were used for the problem exterior. In the “shocked slow patch” test, the state divide was placed at $x = -6$, while it was located at $x = 0$ for the cylindrical–Cartesian case, at $x = 7$ for the “fast patch” run, and at $x = 6$ for the “rarefaction slow patch” case. The local patch’s origin initially coincided with the point ($-15, 15$) in the fast patch simulation, while all other Sod tests began with the local patch centered at the point (0, 10). When the patch moved slowly, its velocity was $V = 10^{-3}(\hat{x} - \hat{y})/\sqrt{2}$; i.e., it traveled subsonically and considerably slower than the shock front. When the local patch moved fast, its velocity was $V = 5 \times 10^{-2}(\hat{x} - \hat{y})/\sqrt{2}$, i.e., it traveled supersonically and $\approx6.2$ times faster in the $\hat{x}$ direction than the shock front.

The results of all four cases can be compared with exact analytic solutions (Laney 1998). In Figure 4, we show data from the “shocked slow patch” run. At $t = 900$ (the left column), the shock has just entered the local patch; at $t = 1600$ (middle column), both the shock and the contact discontinuity run through the local patch; at $t = 3000$ (right column), the shock has exited the local patch, but the contact discontinuity remains within it. In all stages, the multipatch solution closely follows the exact analytic solution. The only noticeable departure is a slight smoothing of the contact discontinuity,
visible in the density plot at $t = 1600$, due to the fact that the discontinuity formed in the coarser global patch. Although almost invisible in these plots, it is also worth pointing out an obvious consequence of the multipatch approach: the shock wave is always only two or three cells thick and is therefore considerably sharper in physical space where it runs through the local patch. We note that where the global and local patches overlap, the global patch values plotted are those interpolated from local patch values even though the values may not be used in the global patch’s evolution; all other global patch values shown are those resulting from the global patch’s update procedure.

The second test, the “rarefaction slow” case, is shown in Figure 5. In all three snapshots, part, but not all, of the rarefaction wave is contained in the local patch. Where the multipatch formalism has affected the results, agreement with the analytic solution is essentially perfect; the only departures are a very slight rounding of the trailing edge of the rarefaction wave apparent at the earlier times when the local patch has never been anywhere near this edge. These departures are the

Figure 4. Shock tube test problem for the Cartesian–Cartesian “shocked slow patch” case, observed at three times, $t = 900$ (left), $t = 1600$ (middle), and $t = 3000$ (right). Note that the horizontal axis scales are different for all three times in order to highlight different segments of the problem. Each column of three panels shows 1D cuts in density $\rho$, pressure $p$, and velocity $v_x$ as functions of $x$ at $y = 10$. Data from the global patch are shown with small green dots, data from the local patch with large cyan dots, and data from the analytic solution are shown with a black line.
The third Cartesian–Cartesian run tests whether supersonically moving local patches create any special problems. The results of this test are shown in Figure 6. As can be seen, small errors are induced in the fluid velocity between the shock front and the rarefaction wave. However, these are not due to any property of the multipatch method; similar errors are produced in conventional monopatch calculations whenever the Mach number of the reference frame’s velocity relative to the shock velocity is significant. We have compared the errors seen in this multipatch test to those seen in a simulation of an identical problem in which only the global patch is present, i.e., a conventional monopatch run, but with the gas initially given a bulk uniform velocity. In this comparison run, the errors are \( \approx 2\% \). The errors in the multipatch test are, depending on the location, generally smaller than, but in a few places comparable to, those in the monopatch run. We have also repeated the multipatch test illustrated in Figure 6 with a less extreme local patch velocity, an \( \hat{x} \) velocity relative to the shock front only 2× the shock speed rather than 6.2. The errors in this test are also typically smaller than, but in a few places...
comparable to, those in the figure. We therefore expect at most modest-amplitude errors when high-contrast local patches pass at high Mach numbers through shocks, and considerably smaller errors when the relative speed is small, as would often be a desirable choice. Indeed, one of the advantages of the multipatch method is the flexibility it offers to choose preferred reference frames in different portions of the problem.

In our last Sod test case, the cylindrical–Cartesian run, the shock traverses a different sequence of patches as a function of the \( y \)-coordinate. For instance, the shock wave traveling along \( y = 10 \) from \( x = 0 \) starts in the global patch, then enters the local patch through its inner radial boundary, and ultimately exits the local patch’s outer radial boundary as it re-enters the global patch. Along other constant-\( y \) trajectories, the waves may start in the local patch and emerge in the global patch or always reside in the global patch.

This test’s results are illustrated quantitatively in Figure 7. Unlike the situation in the first three tests, the local patch’s grid here no longer conforms to the symmetry of the initial data. We show 2D contours of the three fluid quantities from the run to demonstrate how well PATCHWORK maintains the problem’s linear symmetry despite the cylindrical local patch. While the

Figure 6. Shock tube test problem for the Cartesian–Cartesian “fast patch” case, observed at three times, \( t = 575 \) (left), \( t = 675 \) (middle), and \( t = 900 \) (right). Again, the horizontal axis scales are different for all three times in order to highlight different segments of the problem, and each column of three panels shows 1D cuts in density \( \rho \), pressure \( p \), and velocity \( v_x \) as functions of \( x \) at \( y = 3 \). Symbols are also as in Figure 4.
rarefaction and postshock states show no signs of $y$-asymmetry, we do find minor artifacts at the contact discontinuity (at $x \approx 8$ in the images) in $\rho$ and $v_x$. These artifacts are so small that they are difficult to see except in the line plot of $v_x$. The artifacts in $\rho$ take the form of a one- or two-cell displacement of the contact discontinuity. They originate from the two points where the newly formed contact discontinuity intersects the local patch’s inner boundary and then travel with the contact discontinuity. The artifact in $v_x$ also begins when the contact discontinuity crosses the local patch’s inner boundary and similarly travels with the contact discontinuity. It, however, takes the form of a $\sim 5\%$ error in $v_x$ along a half-circle “echo” of the patch boundary. The amplitude of the artifacts decreases with finer grids. Also, the artifacts are not cylindrically symmetric on the local patch because they are advected with the $x$-oriented velocity of the solution while the local patch moves diagonally in the $x$–$y$ plane.

3.2. Sedov–Taylor Blast Wave

The purpose of this test was to show the performance of the multipatch when at least one of the patches has a grid whose symmetry is a poor match to the natural symmetry of the problem and to demonstrate that crossing a patch boundary separating regions of different grid symmetry introduces no ill effects. To that end, we study a Sedov–Taylor 3D spherical blast wave (Sedov 1959) with a central local patch using Cartesian coordinates and a global patch using spherical coordinates. As a standard of comparison, we also contrast a monopatch simulation with entirely Cartesian coordinates. Although the Cartesian grids are poor matches to the spherical symmetry of the physical problem, they do have the virtue of eliminating the coordinate singularity at the origin created by spherical coordinates.

Figure 7. Shock tube test problem in which the local patch used cylindrical coordinates and the global patch used Cartesian coordinates, observed at $t = 2500$. The rest-mass density ($\rho$, top row), pressure ($p$, middle row), and $x$-component of the velocity ($v_x$, bottom row) are shown. Their full 2D contours (right column) are shown next to slices (left column) taken along $y = 8.23$, where departures from the exact solution are the largest. In the 2D plots, the local patch data (outlined in dark long dashes) are shown on top of the global patch data, and the location of the slices is displayed (light, short dashes). The line plots use the same conventions used in Figure 4.
A blast wave is formed when a large amount of energy \( E \) is deposited in a small region. If the ambient gas is motionless, a spherical shock wave travels rapidly outward. Once the mass swept up by the shock exceeds the mass originally located in the small energy-deposition region, the shock front’s radial position as a function of time is given by

\[
R_s = \left(\frac{\xi E}{\rho}\right)^{1/5} t^{2/5}
\]

until \( E/R_s^3 \) is small enough to be comparable with the ambient pressure. Here, \( \rho \) is the initial (uniform) density of external gas, and \( \xi \) is a dimensionless number \( \sim 1 \).

To simulate this, we follow Fryxell et al. (2000) and divide the initial state into two regions. As in the Sod shock tube problem, we choose the unit of velocity to be \( c \), but use arbitrary code-units for length and mass. In terms of these units, region 1 is a small sphere of radius \( \delta r = 25 \). Its initial pressure \( \rho_1 = (\gamma - 1)E/(4\pi\delta r^3) = 1 \) for adiabatic index \( \gamma \) (again \( = 1.4 \)), while its density \( \rho_1 = 10^{-3} \). Region 2 is everything outside \( r = \delta r \). Here, the initial pressure \( \rho_2 = 10^{-10} \) and initial density \( \rho_2 = \rho_1 \). The dimensionless coefficient of Equation (1) is a function of \( \gamma \); for \( \gamma = 1.4 \), it is 1.175 (Ostriker & McKee 1988).

For the monopatch, the computational domain is a cube of dimension 2500 having \( N_{\text{mono}} = 400 \) equal-volume cubical zones with side length 6.25. We perform two multipatch simulations in order to illustrate its dependence on grid-scale contrast. In both, the local patch is a cube of side length 600 centered on the origin with 1203 equal-volume cubical cells of side length 5, similar to the Cartesian cell size in the monopatch simulation. Likewise, in both simulations, the global patch is a sphere of radius 1000 described in spherical coordinates, but with two cutouts: a sphere of radius 290 surrounding the origin and a bi-cone of half-opening angle \( \pi/10 \) surrounding the polar axis. The two multipatch simulations differ in global patch resolution. In the “low-contrast” case, the angular grid is uniform, with 120 cells in polar angle \( \theta \) and 320 cells in azimuthal angle \( \phi \), but the radial grid has 80 logarithmically spaced cells. In this case, the radial cell size in the global patch at the patch boundary \( r \approx 300-400 \) is similar to the local patch cell size. In the “high-contrast” case, the cell counts in all three dimensions are a factor of 4 smaller, so that radial cells at the patch boundary are separated by \( \approx 33 \), roughly a factor of 4 larger than local patch cells.

We portray how well the multipatch simulations do, relative to both the analytic solution and the monopatch simulation, in Figure 8, which again shows the situation at three different times. At the earliest time, the shock front is entirely within the local patch, while it is a short distance outside the local patch in the middle time, and far outside the local patch at the last time. At the earliest time, the data for the Cartesian local patch and the Cartesian monopatch are, not surprisingly, nearly identical; the entire global patch remains in the initial state at this time. Interestingly, the shock at this time is at a slightly larger radius than predicted by the analytic solution in both the monopatch and the multipatch simulations.

At the middle time, the local patch and monopatch still closely agree, but the shock region is located in the global patch. As is clear from the curve showing the analytic solution, when the shock has left the local patch, a grid scale \( \lesssim 10 \) is a prerequisite for describing the density and pressure profiles. The low-contrast multipatch case therefore does reasonably well, slightly outperforming the monopatch; where the high-contrast multipatch case samples the profile, it is in general in good agreement with both the monopatch and low-contrast multipatch data, but its sampling is too sparse to resolve the actual profile. This pattern persists to late times: monopatch and low-contrast multipatch behave very similarly to one another; high-contrast multipatch points are placed too sparsely to resolve the profiles, and their error levels are a bit greater than for the other two simulations.

Our conclusion from this comparison is that the multipatch method performs very similarly to a monopatch method. The poorer performance of the high-contrast case is due entirely to its overly coarse grid, a failing that would have very much the same effect if this grid had been used in a conventional monopatch simulation.

Results from an absolute test may be seen in Figure 9. Here we show how well the high-contrast multipatch simulation is able to support the intrinsic spherical symmetry of the physical problem. When the shock still lies within the Cartesian local patch (left panel), its outline is very nearly circular, but there are small departures from perfect azimuthal symmetry due to the underlying Cartesian grid. When the shock lies partly in the spherical global patch and partly in the Cartesian local patch (middle panel), the shock is almost perfectly azimuthally symmetric in the global patch, but where it passes through the local patch, it retains the same level of small-scale noise as at the earlier time. At late times, when the shock is entirely within the global patch, it shows a very high degree of azimuthal symmetry. Thus, in these tests, the multipatch system induces no departures from the true geometric symmetry; such small errors as exist are due entirely to the symmetry of the grid.

### 3.3. Interpatch Conservation

As remarked earlier, our interpolation scheme is not strictly conservative, even though many hydrodynamics codes that can be used in concert with our multipatch method are. That contrast makes it worthwhile to examine how large an error may be induced by non-conservative interpolation, and how that error depends on the interaction between problem character and details of multipatch implementation.

In principle, this error could depend on many variables. To simplify the discussion and focus on what we believe is the principal issue, we study an idealized problem, one in which matter flows from a patch in which it has acquired an order-unity amplitude fine-scale structure into a more coarsely resolved patch. The parameter that appears to affect conservation errors the most is the ratio between the length scale of the structure and the resolution scale of the coarser patch.

To illustrate this dependence, we construct a 3D system in which the problem volume extends from \( x = -30 \) to \( x = +50 \) in Cartesian coordinates, but which in the transverse directions \( y \) and \( z \) spans only the range \([-20, +20]\). The local patch is stationary and occupies the region \( -30 \leq x \leq 0 \) in global coordinates. Both patches have uniform cubical grids that are parallel to each other, but the cell sizes of the global patch (5) are \( 10 \times \) that of the local patch (0.5).

At \( t = 0 \), all of the fluid is traveling at \( V_y = 0.1 \) in the \( x \)-direction, but its density and pressure differ sharply across the line \( x = +10 \), located a short distance into the global patch from the local patch boundary. To the left of that line, \( \rho_L = 1 + \sin^2(\omega_n y) \) and \( p_L = 10^{-12} \rho_L \) (as in the previous
tests, \(c = 1\) in our units), while on the right \(\rho_R = 10^{-16}\) and \(p_R = 10^{-28}\). The sharp pressure contrast induces a flow from left to right in the frame of the bulk flow. Because the sound speed on the left is so small (~\(10^{-6}\)), even in 1000 time units the high-pressure gas expands only a very slight distance to the right in the moving frame of the fluid. Thus, the density and pressure modulation across the patch boundary at \(x = 0\) is essentially constant throughout the run of the test.

The frequencies \(\omega_n = n\omega_p\) are chosen as multiples of the spatial Nyquist frequency of the global grid resolution, \(\omega_p = \pi/\Delta x\); this definition also ensures that the total mass in the entire computational domain is the same for each \(\omega_p\). The case illustrated in Figure 10 is for \(n = 0.7\); as can be seen, the coarse grid drastically smooths the modulation.

Outflow boundary conditions are enforced at \(x = 50\) while reflecting boundary conditions are applied at all other physical
boundaries. In the absence of numerical error, these boundary conditions (and the extremely low value of $\rho R$ in the initial state) guarantee that the total mass, energy, and momentum on the grid remain constant until the front arrives at the far-right edge at $t = 400$.

In Figure 11, we show the ratio $\Delta M(t)/M_{\text{flow}}(<t)$ between the change in total mass and the amount of mass that has flowed across the patch boundary up to that time for initial density modulations with frequencies $\omega_n = \{0, 0.01, 1.0, 2.0, 2.6\} \times \omega_\nu$. As is appropriate to the steady-state flow we are studying, the fractional error is independent of time for all cases. Also not surprisingly, when $n < 1$, so that the global grid resolves the modulation well, the error is small: $\lesssim 10^{-4}$ for $n = 0.01$, when there is almost no modulation, and $\approx 1.5 \times 10^{-3}$ for $n = 0.1$. Once $n \geq 1$, when the global grid can no longer support the modulation, the pattern changes. The error for $n = 2.6$ is larger than for $n < 1$, but still tolerable ($\approx 1.5 \times 10^{-2}$). However, the error for both $n = 1$ and $n = 2$ is uncomfortably large: $\approx 0.3$. These two values of $n$ are special cases: the modulation is resonant with the global grid pattern, so the error in the mass flow depends strongly on the phase of the modulation at cell centers, which is the same for all global cells. The value for $n = 2.6$ should therefore be more characteristic of generic modulations.

We have also examined the fractional conservation errors for $x$-momentum and internal energy, but we do not present them explicitly because the figures are virtually indistinguishable from those shown in Figure 11. That they should be so similar is to be expected because they are interpolated by identical procedures.

In addition, we have performed simulations of the same problem with a different global patch, one with polar coordinates. In this case, we are testing the robustness of conservation properties with respect to change of grid symmetry, rather than with respect to change of grid resolution. For this reason, the local patch and initial condition structure are identical to those of the previous conservation test, but the global patch is given a cylindrical grid (see Figure 12). The origin of the cylindrical coordinates is placed at $(x, y) = (20, 0)$; its radial cells have width 0.5 and its azimuthal cells have width 0.025 radians so that both cell dimensions near the patch boundary match those of the local patch. The global grid
frequency $\omega_2$ for this test is defined so that it is equivalent to the one used in the first set of tests, i.e., $\omega_2 = \pi/(10\Delta_{\text{local}})$. The polar grid near the patch boundary should therefore be able to support modulations with values of $n$ similar to those used before. On the other hand, even though the azimuthal cell size becomes even finer near the global patch origin, the mismatch between polar cell shapes and the rectangular modulation induces larger errors closer to the global patch origin (also shown in Figure 12).

The results (seen in Figure 13) are, nonetheless, comparable to those from the Cartesian–Cartesian tests. For all $n \lesssim 2$, the fractional error is $\approx 2 \times 10^{-3}$, again almost independent of time. Unlike the Cartesian–Cartesian case, however, the polar grid eliminates the possibility of resonant response for integer $n$. With a polar global grid, the error for $n = 2.6$ rises over time; we believe that this increase is due to the mismatch between the flow properties and the global grid geometry, a mismatch exacerbated by higher-modulation wavenumbers. If so, it is not a product of errors created as information is transferred across the patch boundary, but rather one intrinsic to the inappropriate symmetry of the polar grid. The evidence for this supposition is that if we define the total mass on the grid as

$$ M = \int_{-30}^{50} dx \int dy \int dz \rho, $$

we find that the fractional error grows as $x_\ast$ increases toward $x = 20$, the $x$-coordinate of the polar grid origin. The values shown in Figure 12 are for $x_\ast = 18$. Again, just as for the Cartesian–Cartesian case, the error numbers for momentum and energy are virtually identical to those for mass.

In more realistic problems, we have found that dynamical effects can enlarge these errors, and in some cases positive feedback loops can develop. However, the driving factor appears to be similar: the inability of a coarse grid to support variations occurring on length scales too small for it to resolve. Devices to curb this sort of interpolation error are problem dependent. For example, in our tidal disruption test run (mentioned at the end of Section 1.1), we found that smoothing the hydrodynamic variables over the three ghost cells at the interpatch boundary successfully damped a growing departure from mass conservation, holding the error in total mass to $<1\%$ over a time in which the star shed 95\% of its mass.

4. Computational Efficiency and Parallelization Scaling

Lastly, we present data on computational efficiency and parallelization scaling. When discussing these issues in the context of ordinary monopatch operation, the principal questions generally have to do with the fundamental efficiency of the computational algorithm and the ratio between time spent exchanging boundary condition information and computing updates. The former sets the basic scale in terms of zone cycles per processor per unit time; the latter is determined by the additional cost incurred by interprocessor communication. When a hydrodynamics code parallelizes well, the fraction of total processor-hours devoted to communication is nearly independent of the total number of processors. Thus, to gauge how much overhead is created by multipatch operation and how efficiently the multipatch system makes use of parallelization, we must contrast multipatch benchmarks with monopatch benchmarks treating the same physics problem and do so as a function of total number of processors. In addition, we will explore how much our heterogeneous time-step option improves efficiency by contrasting its performance with matched homogeneous time-step runs.

However, all these tests can at best be indicative rather than definitive. Even in monopatch operation, hydrodynamic code speeds can be problem dependent, and there is every reason to expect that multipatch methods will, if anything, add new ways for code performance to be sensitive to the nature of the specific problem. For example, in multipatch problems the amount of computation required to perform a single zone cycle can depend on how much effort is necessary to compute the flow near the patch boundary successfully damped a growing departure
expect a significant contrast in overhead between cases in which the local patches move or are stationary, we will specifically examine that variety of problem dependence.

To finesse these complexities as best we can, we focus on a single simple test problem: evolving a hydrostatic gas in the absence of any external forces. The background spacetime is therefore Minkowski, and in the initial condition there is uniform density, pressure, and entropy. The fluid’s adiabatic index is $\gamma = 5/3$.

The problem volume is a 3D cube treated with two patches, a global patch and a local patch. Both use Cartesian coordinates with uniformly spaced grids. The local patch is a cube with side length $1/8$ the global patch’s side length, but has a grid scale that is also $1/8$ the global patch’s; the two patches therefore have the same number of cells. These choices produce a time-step ratio $\Delta t_g/\Delta t_l = 8$. These will be studied with two different numbers of zones per processor, $n = 20^3$ and $n = 40^3$. Each simulation is run for a fixed time duration, chosen to be just long enough that initialization time is negligible. Zero-gradient boundary conditions are used for the global patch; the local patch never encounters the problem boundary. All cases were performed on the same platform (Texas Advanced Computing Center, Sandy Bridge nodes on Stampede).

In our first set of benchmarks, we consider the case of a stationary local patch and a single time step for both patches. The results (obtained from STDLIB C “time(5)”) are shown in the two left-hand panels of Figure 14. In this set, we assign the same number of processors to each patch; that means both patches have the same number of cells per processor. When the number of cells per processor is relatively large, multipatch operations create a very modest overhead: the ratio of cycle-update speed for multipatch to monopatch with $40^3$ cells per processor is $\approx 0.75$. On the other hand, with fewer cells per processor ($20^3$), the ratio is closer to $\approx 0.4$.

Another view of the stationary patch computational efficiency may be seen in the lower-left panel of Figure 14, showing the ratio of time spent in communication relative to the total computational time. Here we define “communication” as any operations involving boundary condition exchange between processors. Examples of communication specific to multipatch operation include determination of client–server relations, transmission of ghost-cell coordinates, and interpolation of data to those coordinates. A fourth category of communication, data transmission from server to client, occurs in any sort of parallelized simulation. “Total time” is defined as all time spent on communication plus time spent on computing hydrodynamic updates; it does not include time spent on ancillary activities such as initialization or writing output. In terms of this measure, we find that for stationary patches the fraction of total time spent in communication is $\approx 2 \times 3 \times$ as great as for monopatch runs with the same number of cells per processor. This extra time can be largely attributed to the interpolation step because communication time in monopatch runs is due only to MPI data transmission, whereas in multipatch operations it also includes interpolation and client–router–server data exchange.

In the second set of comparisons (right panels of Figure 14), we examine what happens when the local patch moves. In this case, the contrast in zone-update rate is larger and is also a stronger function of the number of processors per patch. As before, larger numbers of cells per processor yield greater efficiency. With $40^3$ cells per processor, the update rate for the multipatch is $\approx 0.7 \times$ the monopatch rate for 512 processors per patch, but declines to $\approx 0.5 \times$ for 1728 processors per patch. The corresponding figures for $20^3$ cells per processor are $0.6 \times$ and $0.2 \times$ the monopatch rate. The fraction of time spent in communication is consistently $5 \times 6 \times$ larger in moving patches than for the corresponding monopatch case, nearly independent of the number of processors per patch.

The overhead and scaling behavior of moving patches differs from that of stationary patches because, in addition to data interpolation, it is also necessary to determine client–server relationships at each time step and to transmit fresh ghost-cell coordinate lists. These additional tasks both increase the total overhead and make parallelization scaling poorer. The underlying reason is that as the number of processors per patch grows, more processors must be queried to determine the correct set of client–server connections. The relative load this imposes is larger when the number of cells per processor is smaller because, just like all other boundary data considerations, it is sensitive to the processor domain’s surface/volume ratio.

We close this part of the discussion by making an important remark regarding the interpretation of these benchmarking results. Although we expect the qualitative trends to be robust, their quantitative character is sensitive both to the specific problem and to the specific architecture of the computing system used. Different problems (and different algorithms applied to the same problem) can have different numbers of arithmetic operations per cell update, while different cluster architectures can give different rates of interprocessor data transmission. Because part of the multipatch overhead depends on additional data transmission, the relative speeds for these two sorts of processes can alter the multipatch/monopatch comparison at a quantitative level. As we discuss below (Section 5), technical improvements in the implementation of the multipatch method can also lead to quantitative changes in efficiency.

Next, we compare the computational expense for a multipatch program running with and without the heterogeneous time-step algorithm at a fixed total number of zones in each patch $N$. The time-step ratio between the two patches is $\Delta t_g/\Delta t_l \approx 8$. For ideal load balancing with a heterogeneous time step, the global patch should therefore have eight times as many zones per processor as the local patch. In Figure 15, we show how the number of processor-hours required to reach a fixed physical time depends on the number of zones per patch for both a homogeneous and a heterogeneous time step. As one might expect, both scale very closely to linearly with the number of zones per patch, but the heterogeneous time step requires a number of processor-hours $2 \times 3 \times$ smaller than the homogeneous time-step operation.

Note that the gains achieved by using heterogeneous time steps appear in a way quite distinct from typical gains in efficiency. They do not lead to any increase in zone cycles per processor per second; instead, they lead to a decrease in the total number of zone cycles required to accomplish the simulation. In this way, their effect resembles the use of non-uniform spatial grid cells, a device leading to economies in the total number of zones computed by concentrating them where they are most needed, rather than a conventional increase in computing efficiency.
5. Conclusions

We have presented the essential methods underlying our implementation of a new multipatch infrastructure, PATCHWORK, designed to support multiscale, multiphysics, and multireference frame fluid simulations. This method offers a number of advantages for the numerical study of complex fluid problems involving subregions with contrasting properties. Each patch can have its own coordinate system and spatial grid, differing in geometry and resolution from all of the other patches (one of the many ways this can be useful is that if the coordinate system preferable for a part of the problem contains coordinate singularities, they can be covered with a new patch). If different regions demand contrasting time steps, the independence of the processes evolving the patches permits them to have separately determined time steps, potentially saving significant amounts of computing. Although the method assumes that a fluid exists throughout the problem volume, if different auxiliary processes are important in different regions (e.g., chemical reaction networks or self-gravity), their patches can treat those processes without burdening the other regions. Lastly, but possibly most importantly, substructures within the problem may have differing preferred reference frames; these, too, can be accommodated easily.

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Figure 14. Computational efficiency as a function of numbers of processors per patch and for different numbers of cells per processor. Monopatch method data are plotted with open symbols, multipatch with filled symbols. Runs with 20³ cells per processor are shown with black circles, runs with 40³ cells per processor with red squares. Left (right) panels show multipatch simulations with a stationary (moving) patch. Top panels: processing speed in zone cycles per processor per second. Bottom panels: fraction of total wall-clock time spent on communication.

Figure 15. Computational expense of a multipatch simulation with a homogeneous (blue circles) and heterogeneous (orange ×’s) time step for a given total number of zones per patch $N$. 

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The patches are linked to one another solely through boundary condition exchange. Contrasting grid systems are reconciled through interpolation; contrasting grid geometries and reference frames are reconciled through coordinate transformations and ensuring that all transformed physical quantities are well-defined scalars, vectors, or tensors.

Parallelization is essential to modern large-scale computing. Arranging the exchange of boundary condition information between the correct processors can be a complex problem in a multipatch system when the patches move relative to one another. We have constructed a solution to this problem—a client–router–server framework—that updates these connections efficiently. When the patches are stationary relative to one another, the connections need to be identified only once, so the overhead due to multipatch operations is fairly small, especially for larger numbers of cells per processor. When they move, the overhead is more significant and scales with the number of processors per patch, producing a reduction in the cell-update rate of about a factor of \( \sim 1.4 \) for 512 processors per patch or a factor of \( \sim 2 \) for 1728 processors per patch when using \( 40^3 \) cells per processor. We note, however, that these comparisons assume that monopatch and multipatch approaches use the same total number of cells; because a multipatch operation permits tuning the grid to match local requirements, in practice multipatch simulations may use a much smaller total number of cells than would be required for a monopatch simulation of the same problem—if a monopatch simulation could deal with the problem at all.

Many extant fluid codes are automatically consistent with this infrastructure. Its sole substantive stipulation is that the dependent variables involved in boundary data exchange should be consistent in all patches. Although we were motivated to build this system by relativistic problems and our transformation methods are familiar because of their frequent application to relativistic dynamics, in fact they really stem from more general considerations of differential geometry; they therefore apply to any context in which scalars, vectors, and tensors can be defined.

PATCHWORK may be refined and extended, both in terms of its computational efficiency and the span of physical problems on which it can be used. Communication between local patches (as opposed to only local–global communication) can substantially extend the dynamic range of length scales treated. The amount of time spent on interpolation and interpatch data transmission can be reduced by minimizing the number of arrays transferred or by eliminating unnecessary steps in the coordinate transformations. Moving from an MPMD environment to one in which a single program employs task-based parallelization will permit dynamical processor reassignment, amplifying the economies in the total zone cycles that accrue from the use of heterogeneous time steps. Task-based parallelization may also improve interpolation efficiency because, for any single time step, only a fraction of the processors assigned to an individual patch are involved in interpatch data exchange. Another improvement will be to add interpolation options that, over a broader range of circumstances, more nearly conserve quantities that should be conserved, such as mass and momentum. Given suitable patch resolution, our current default method does not create significant errors, but it would be valuable to create new schemes, more nearly conservative, that would permit greater freedom in resolution choices. Similarly, some special devices will be necessary to extend our multipatch method to MHD problems in a way that preserves a divergence-free magnetic field. We are currently developing algorithms to achieve this and hope to report on them in the not-too-distant future.

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Appendix

Implementation of the PATCHWORK system is accomplished through a set of functions enabling an existing grid-based hydrodynamics code to run multipatch simulations. The main tasks of these additional functions may be organized according to three categories: setting up data infrastructure, performing boundary condition exchange between patches, and a number of other utilities specific to multipatch operation (e.g., managing heterogeneous time steps).

In addition to grouping the multipatch routines in terms of function, it is also convenient to group the changes necessary to convert a conventional hydrodynamics code in terms of user control and responsibility. Here, too, there are three categories: “permanent” routines, user-specified routines, and selected additional function calls. The routines we designate as “permanent” are those that remain fixed in character, independent of the application. These routines

1. initialize PATCHWORK’s data infrastructure,
2. carry out interpatch boundary data exchanges including interpolation and vector transformation,
3. restart the simulation when a patch has been added or removed (Section 2.4),
4. move patches, and
5. coordinate time steps, making allowances when necessary for heterogeneous time steps (Section 2.6).

Several routines must be supplied by users because they are specific to the problem. Their calling sequences must have a specified form, but their contents are up to the user. They

1. specify the geometric character of the patch and the transformation linking its coordinate system to the background coordinates,
2. create initial condition data (note that it is the user’s responsibility to ensure that initial data near patch boundaries are consistent with initial data in adjacent patches),

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3. implement additional physics (when needed), and
4. initialize the configuration and motion of the patch.

Lastly, we made our best effort to minimize modifications to the underlying fluid code by implementing the PATCHWORK system as a wrapper. For example, the hydrodynamics part of the code is almost never touched. However, a few modifications must be made.

The great majority of these have to do with introducing MPMD features in MPI communication-related routines. In particular, it is necessary to redefine the term “global.” In conventional parallelized codes, “global” denotes the entire problem volume and includes all processors associated with the run. However, in multipatch operation, it is necessary to distinguish the entire volume of a patch from the entire volume of the problem. Making this distinction means that the global MPI communicator in the underlying fluid code must be redesignated “local” so that it refers only to the processors associated with an individual patch. In addition, there are often a number of variables and functions with both “local” and “global” versions; all of these need to be renamed (in a way chosen by the user) to distinguish truly local (in the domain of an individual processor) from patch-global (throughout a single patch’s volume) to problem-global (covering the entire problem volume).

There are also a number of places where the fluid code must call one of the PATCHWORK routines, sometimes one of the permanent routines, sometimes one of the user-supplied routines. Their purpose is to

1. initialize PATCHWORK infrastructure,
2. move the patch to its next location (when necessary, this is done at the end of each time step), and
3. construct patch boundary data (this happens at the initialization, restart, and end of each time step).

In addition, if optional features are used (e.g., adding/removing patches or heterogeneous time steps), the user must likewise insert calls for them at appropriate places.

To close, we note that the output from each patch can be handled by whatever means the fluid code for that patch uses. If, however, the user wishes to merge the data sets, it is up to the user to write the software to accomplish it; the way such merges are done is very problem specific.

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