

# Chombo Software Package for AMR Applications Design Document

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June 2, 2023

#### **Disclaimer**

This was produced by the Lawrence Berkeley National Laboratory, Berkeley, CA. Research at LBNL was supported financially by the Office of Advanced Scientific Computing Research of the US Department of Energy under contract number DE-AC02-05CH11231. This document was prepared as an account of work sponsored by the United States Government. While this document is believed to contain correct information, neither the United States Government nor any agency thereof, nor The Regents of the University of California, nor any of their employees, makes any warranty, express or implied, or assumes any legal responsibility for the accuracy, completeness, or usefulness of any information, apparatus, product, or process disclosed, or represents that its use would not infringe privately owned rights. Reference herein to any specific commercial product, process, or service by its trade name, trademark, manufacturer, or otherwise, does not necessarily constitute or imply its endorsement, recommendation, or favoring by the United States Government or any agency thereof, or The Regents of the University of California, The views and opinion authors expressed herein do not necessarily state or reflect those of the United States Government or any agency thereof, or The Regents of the University of California.

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# Chapter 1

# Introduction

In many problems in partial differential equations, one is confronted with problems having multiple length scales and strong spatial localizations. Examples include nonlinear systems of hyperbolic partial differential equations containing complex combinations of discontinuities and smooth flow. Also included are combustion problems in which, at any given instant, burning is taking place in a small subset of the problem domain and problems with complex geometries in which localized geometric features can generate strong, localized solution gradients. Finite difference calculation using block-structured adaptive mesh refinement (AMR) is a powerful tool for computing solutions to partial differential equations involving such multiple scales. In this approach, the underlying problem domain is discretized using a rectangular grid and a solution is computed on that grid. Regions requiring additional resolution are identified by computing some local measure of the original error and covered by a disjoint union of rectangles in the domain, which are then refined by some integer factor. The solution is then computed on the composite grid. This process may be applied recursively, and for time-dependent problems, the error estimation and regridding can be integrated with the time evolution and refinement applied in time as well as in space. Such an approach was first introduced by Berger and Oliger [6] for computing time-dependent solutions to hyperbolic partial differential equations in multiple space dimensions. Since that time, the approach has been extended to a variety of problems in applied partial differential equations [7] [29] [4] [2] [26] [17] [1] [19] [27] [18] [10] .

One of the principal disadvantages of block-structured AMR is its relative difficulty to implement compared to single-grid algorithms. The algorithms are more complex and the data structures are unfamiliar to traditional FORTRAN programmers.

To ameliorate these difficulties, we have developed Chombo, a set of C++ classes designed to support block-structured AMR applications. Chombo is based in part on the BoxLib toolkit and related work done by our colleagues at the Center for Computational Sciences and Engineering (CCSE) at LBNL [12] [28]. The Chombo package at the present time consists of the following components:

The BaseTools Library contains helper classes and functionality which is indepen-

dent of spatial dimension, such as Vector and List container classes, etc. The BaseTools library need not be specified in the Chombo makefiles, and is always included and linked to when compiling using the Chombo make system. In earlier versions of Chombo, the contents of the BaseTools library resided in the BoxTools library.

- The BoxTools Library includes the BoxLib rectangular array library. BoxTools also contains a full set calculus on  $\mathbb{Z}^n$ , and classes for defining data on unions of rectangles as well as mapping such data onto distributed memory systems.
- The AMRTools Library consists of classes which implement a number of operations that often appear in AMR algorithms: conservative interpolation, averaging between AMR levels, interpolation of boundary conditions at coarse-fine interfaces, and refluxing operations to maintain conservation at coarse-fine interfaces.
- The AMRTimeDependent library consists of classes which support the Berger-Oliger time stepping algorithm and examples of its use in solving systems of hyperbolic conservation laws..
- The AMRElliptic library consists of classes which support an AMR-multigrid algorithm for elliptic partial differential equations, and examples of its use in solving Poisson and Helmholtz equations.
- The MultiDim library includes functions which support Chombo in a multidimensional environment, as described in Chapter 11.
- The Sundials library includes interface classes which enable Chombo applications to link to the SUNDIALS library of linear and nonlinear integrators, as described in Chapter 9.
- Support for embedded boundary (EB) discretizations and algorithms is found in the EBAMRElliptic, EBAMRTimeDependent, EBAMRTools, EBTools, and Workshop libraries, which are described in a separate set of EB design documents. These are only built if EB=TRUE is specified during the build process.

In addition to these basic tools we have provided extensive documentation. There are some general comments regarding the use of the package, however, that are worth emphasizing here.

C++ rectangular array operations applied one point at a time in a for loop will
not produce high performance on bulk rectangular operations. For this reason,
there is an interface between the array classes that allows them to be passed to
FORTRAN routines. We have expanded this interface with a macro package. The
Chombo FORTRAN package additionally allows one to write dimension-independent
FORTRAN. Both the macro interface to Fortran and Chombo Fortran are described

in Chapter 10. On the other hand, sparse irregular calculations have generally been implemented in C++ directly using pointwise operations.

- We have attempted to leverage other related research activities. For example, HDF5 is an emerging standard for portable, self-describing, binary I/O. For this reason, we have based our I/O on HDF5. Some of our algorithms use FFTs, so we have also provided an interface which uses the FFTW package [14].
- We are trying to enable use of parts of Chombo which others might find useful by pursuing a component-based design approach. For example, it is possible to use our implementation of the Berger-Rigoutsos [8] grid generation algorithm or the parallel data distribution support without using the rectangular array library.

Finally, we want to emphasize that the developers of this package are themselves using Chombo to develop new algorithms and packages. This means that we will be actively adding capability that we expect to make available to other users of this package.

# 1.1 Requirements

Before discussing the installation procedure, we must discuss what other software needs to be installed on a system in order to build Chombo.

 To build Chombo, the GNU version of make (GNUmake) must be installed. The Chombo makefile system requires GNU make version 3.77 or later. GNU software can be downloaded from many places, including:

```
ftp://ftp.gnu.org/gnu/make
```

- A functioning MPI-1.2 (or higher) compliant C-binding is needed to build Chombo for parallel processing. This is only necessary if Chombo is compiled with the MPI=TRUE option. See section 1.2 for the various compilation options for Chombo.
- A C++ compiler is required. Chombo makes heavy use of the ISO/IEC 14882 C++
  Standard. Chombo now utilitzes features from the C++11 standard and users are
  now free to step up to this standard. Some compilers are not fully compliant with
  this specification, although most are. If hybrid parallelism is desired the compiler
  should also support OpenMP API. Chombo has been compiled and tested with
  - GNU g++ 4.48 or higher.
  - Intel icc v15 or higher on Linux.
  - IBM xIC version 8 or later.
  - LLVM clang++, pretty much every version seems to work. Apple's version of clang++ does \*not\* support OpenMP builds.

- Cray compilers now appear to be based on LLVM clang++ 9.0.
- At least a Fortran 77 compiler is required. Chombo has been compiled and tested with:
  - g95, gfortran, g77
  - Intel ifc on Linux
  - IBM xlf, xlf90 on AIX
  - Portland Group pgf77, pgf90
  - HP/Compaq/DEC f77,f90,f95
  - Sun f77,f90
- Perl version 5.0 or higher is required. The Chombo Fortran system uses perl to produce dimension-independent Fortran code. Perl 5 can be downloaded from:

```
http://www.perl.org/get.html
```

 HDF5 must be installed. This provides Chombo a mechanism for portable and parallel self-describing binary output. HDF5 can be downloaded from:

```
http://www.hdfgroup.org/downloads/
```

We recommend using HDF5 version 1.8.x or higher, although Chombo can be built using the HDF5 version 1.6 API if desired by including <code>-DH5\_USE\_16\_API</code> in the compilation flags during the Chombo build process as detailed in section 1.2.2. We suggest configuring HDF5 with the option "--enable-build-mode=production" for best performance.

If you are planning to build Chombo to use MPI, a parallel version of the HDF5 libraries must also be built. Configure HDF5 with "--enable-parallel" for a parallel version. Make sure to install it into a different directory from the serial version, since the libraries have the same names in both cases. (for example, you can configure hdf5 with --prefix=/usr/local/hdf5/serial for the serial build, and --prefix=/usr/local/hdf5/parallel for the parallel (MPI-enabled) one.)

# 1.2 Installation

# 1.2.1 Downloading Chombo

Previous versions of Chombo were distributed as compressed tar files. The Chombo team is now using a more continuous distribution process using branches and the Subversion revision control system. We also are maintaining a mirror of the Chombo release on github, although svn remains our primary distribution channel.

#### **SVN**

To check out Chombo you will use the svn Subversion client. Subversion is already installed on most unix-like operating systems. In order to access the Chombo svn repository, first go to https://anag-repo.lbl.gov and register to establish a username and password. You will then be redirected to a page with specific download instructions. There is also a link on https://anag-repo.lbl.gov which has the same information without registering. Future changes to the Chombo release may be obtained through svn updates.

#### Git

We are maintaining a mirror of the Chombo release on GitHub, which can be found at https://github.com/applied-numerical-algorithms-group-lbnl/Chombo\_3.2

# 1.2.2 Configuring Chombo for a Particular System

Before Chombo can be built, the makefile system must be configured for the computer it is to be built on. The major configuration parameters relate to the locations of the other software on which Chombo depends and the compilers.

All Chombo configuration is done by setting variables that the makefiles use. The makefile system sets as many of these variables as possible, but some are hard to determine and others are purely a matter of user preference. All customizations of the makefile variables are done in a single file:

Chombo/lib/mk/Make.defs.local

This file does not exist in the Chombo distribution tar file. There are several ways to create it:

- copy the file Chombo/lib/mk/Make.defs.local.template
- copy (or create a symbolic link to) the system-specific customization file from Chombo/lib/mk/local if you are using a computer we already use (Franklin, Hopper, Jaguar, etc.)

The first two option produces a customization file with no variables defined, but with the important variables documented in comments. The resulting Make.defs.local file must be edited to set the HDFINCFLAGS and HDFLIBFLAGS variables (see below) if either of these options is used. The system-specific files in the second option set the variables for particular supercomputers that many people use and should work on those systems without modification.

The makefile variables that are most commonly customized include:

DIM the number of spatial dimensions in the calculations (=1, 2, 3, 4, 5, or 6). The default is 2. There is limited functionality for high dimensions (DIM > 3).

- PRECISION determines the size of floating point variables; the acceptable values are FLOAT and DOUBLE. The default is DOUBLE.
  - DEBUG determines whether to compile with a symbol table (=TRUE) or not (=FALSE). The default is TRUE.
    - OPT determines whether to compile with optimization (=TRUE) or not (=FALSE). The default is FALSE. OPT can also be set to HIGH in which case asserts are removed from the code and FArrayBox memory is initialized to zero (instead of a large positive value) during memory allocation. It is recommended that OPT=HIGH not be used unless absolutely necessary.
  - PROFILE determines whether to compile for performance profiling (=TRUE) or not (=FALSE). The default is FALSE.
    - CXX the command to run the C++ compiler (include path and options, if necessary). The default is g++, except on systems with a usable vendor-provided compiler.
    - FC the command to run the Fortran compiler (ditto). The default is gfortran, except on systems with a usable vendor-provided compiler.
    - MPI determines whether to compile for parallel (=TRUE) or serial (=FALSE) execution. The default is FALSE.
    - OMP determines if hybrid parallelism is in use (=TRUE). The default is FALSE.
  - MPICXX when \$MPI is TRUE, this specifies the command name of the parallel C++ compiler. The default is mpiCC, except on systems with a usable vendor-provided compiler.
- OBJMODEL an optional flag value that specifies a special way to compile the Chombo code (e.g. for 64bits or dynamic libraries). The actual values are defined in the makefiles for the individual compilers. Most users will not need to set this. The default is blank.
- XTRACONFIG an additional identification string to be added to filenames generated by the makefiles. This allows the user to build separate libraries based on parameters other than those specified by the makefile system. This string is empty by default.
  - LD the command to run the linker, if different from CXX
- HDFINCFLAGS the C++ compiler options to compile with HDF (usually -I < hdf\_dir>/include, where < hdf\_dir> is the root directory of the HDF installation). The default is blank, but that usually will not work, so this variable must be set. For serial builds against a standard installation of HDF5. It is no longer necessary to use the -DH5\_USE\_16\_API flag.
- HDFLIBFLAGS the linker options to access the HDF libraries (usually -L<hdf\_dir>/lib -lhdf5 -lz)

HDFMPIINCFLAGS same as HDFINCFLAGS, except for the parallel version of HDF. The default is blank. (this should be blank if parallel HDF is not installed)

HDFMPILIBFLAGS same as HDFLIBFLAGS, except for the parallel version of HDF (this should also be blank if parallel HDF is not installed)

The first 10 variables in this list (from DIM to XTRACONFIG) are called the "configuration" variables. The Chombo makefiles allow for different configurations to exist simultaneously by using the configuration in the names of the library and executable files. The normal procedure is to define a default configuration in the Make.defs.local file (or use the standard configuration defined in the Chombo/lib/mk/Make.defs.defaults file) and build alternate configurations by specifying the configuration variables explicitly on the make command line. For example:

```
make DIM=3 DEBUG=FALSE OPT=TRUE all
```

will build Chombo in 3 dimensions with compiler optimization enabled.

If the compilers on the system are not already known to the makefiles, it also may be necessary to set variables that determine what options to use in the compile commands. Variables for known compilers are set in files in the directory Chombo/lib/mk/compiler. The files in this directory have names of the form Make.defs.compiler\_name. The compiler\_name is taken from the CXX and FC configuration variables.

The recommended approach to setting compiler variables for unknown compilers is to first try to build the Chombo libraries and programs with the default compiler options variables, and if that doesn't work, to customize the variables in the Make.defs.local file.

The compiler variables are:

cppdbgflags options for the preprocessor step of C++ and Fortran compiles when DEBUG=TRUE (default is blank)

cppoptflags options for the preprocessor step of C++ and Fortran compiles when OPT=TRUE (default is blank)

cxxdbgflags options for C++ compiler and linker when DEBUG=TRUE (default is -g)

cxxoptflags options for C++ compiler and linker when OPT=TRUE (default is -O)

fdbgflags options for Fortran compiler when DEBUG=TRUE (default is -g)

foptflags options for Fortran compiler when OPT=TRUE (default is -O)

1ddbgflags options for linker only when DEBUG=TRUE (default is blank)

ldoptflags options for linker only when OPT=TRUE (default is blank)

cxxprofflags options for C++ compiler and linker when PROFILE=TRUE (default is -pg)

fprofflags options for Fortran compiler when PROFILE=TRUE (default is -pg)

These variables can be overridden on the make command line by setting the variables:

CPPFLAGS CXXFLAGS FFLAGS LDFLAGS

# 1.2.3 Compiling Chombo's Libraries

Once the makefile variables are properly customized in the Make.defs.local file, the libraries can be built. The commands to do this are:

```
cd Chombo/lib
make lib
```

Add to the "make" command any non-default definitions of configuration variables you wish to use.

This will produce a lot of output, most of which is compile commands and messages from make. Depending on which compilers are used, there may be some compiler warnings about unused variables and invalid offsets, but these can be safely ignored. None of the compiles should produce error messages. If this occurs, you have a problem. Usually the solution is to fix the compiler options. Problems with the files in Chombo/lib/mk/compiler should be reported to <chombo@lbl.gov>.

# 1.2.4 Compiling and running Chombo's test programs

Once the libraries are successfully compiled, the test programs should be built and run. The commands to do this are:

```
cd Chombo/lib
make test
make run
```

Of course, any variables you defined on the command line when build the libraries also should be defined for these "make" commands.

The first "make" command compiles and links the test programs. Errors are usually in the link step, since the test code will usually compile if the library code compiles. Link errors are commonly due to bad or missing libraries, bad template instantiation by the compiler or problems with Fortran libraries. Make sure that the HDFLIBFLAGS variable has the correct value and that the HDF libraries were compiled with the same compiler that the Chombo build is using. Undefined references to routines named H5\* is a symptom of problems with the HDF library. Other problems should be referred a local guru or, failing that, to <chombo@lbl.gov>.

The "make run" command executes all the Chombo test programs, in a mode that produces minimal output. Successful execution of a test program is indicated by the message "... testFoo finished with status 0". If all you care about is whether the tests succeed or not, the command to use is:

```
make run | grep 'finished with'
```

If the status is anything other than 0, the test failed and you should rerun it by hand in verbose mode.

To do this, find the message that starts "make –no-print-directory –directory" and occurs before the output of the test that failed. The word after "–directory" is the directory containing the failed test. Change directory into "test" then change into that directory. Run the command "make run VERBOSE=-v" and save the output. Then do "cd ../.." and run the command "make vars". Email the output from both commands to <chombo@lbl.gov> and we'll try to suggest a solution.

# 1.2.5 Compiling and running Chombo's example applications

The test programs are all simple codes that exercise small pieces of the Chombo libraries, usually just single classes. They are not intended to show how the Chombo software should be used in a real application. For that, there are example applications.

Building and running the examples is the same as the test programs. The commands are:

```
cd Chombo/releasedExamples make all make run
```

As before, any variables defined on the command line when you built the libraries should be added to this "make" command too.

This "make all" step usually succeeds if the libraries and tests built successfully. Errors should be reported.<sup>1</sup>

The "make run" step produces a lot of output, and can take a long time to run, depending on the computer and configuration. As with the tests, the command:

```
make run | grep 'finished with'
```

will print out a minimum of output and still indicate whether all the example programs ran successfully. It is recommended to run the examples once and look at the output to ensure that the programs actually ran correctly.

Some of the example programs use a lot of memory and take a long time to run in 3d: (e.g. AMRNodeElliptic/execPolytropic). The command:

```
cd Chombo/releasedExamples make usage
```

will list all the example targets. An individual example can be run by running "make" with that target.

<sup>&</sup>lt;sup>1</sup>Exception: the IBM xIC compiler complains about multiple definitions. Our experience has been that it is safe to ignore these complaints as long as the executable files (\*.ex) are created and run successfully.

```
## path to the 'Chombo/lib' directory
CHOMBO_HOME := ../../../Chombo/lib

## defines the Chombo variables
include $(CHOMBO_HOME)/mk/Make.defs

## this is the name of the file with 'main()'
ebase := main  # change this !!

## Chombo libraries needed by this program
LibNames := AMRElliptic AMRTools BoxTools

## Other libraries needed by this program (using -L and -l options)
XTRALIBFLAGS :=

## 'all' is the default target and 'all-test' is defined in Make.rules
all: all-test

## defines the rules to build everything
include $(CHOMBO_HOME)/mk/Make.rules
```

Figure 1.1: Sample Chombo example makefile

# 1.2.6 Building an application using Chombo

There are two ways to build an application with the Chombo library: by using the Chombo makefiles and building the application as if it was a Chombo application, or by treating Chombo as just another library in an existing makefile.

## 1.2.6.1 Using the Chombo application makefiles

The Chombo makefiles support two different ways of building an application. One assumes all the source code for the application (aside from whatever libraries it uses) are stored in a single subdirectory. The other allows for source code in multiple directories. The former is simpler.

Figure 1.1 shows an example of a "GNUmakefile" that builds an application with all of its code contained in a single directory. It is based on the GNUmakefile in Chombo/releasedExamples/AMRPoisson/execCell, with unnecessary lines removed for simplicity.

The first line defines the CHOMBO\_HOME variable, which tells the rest of the makefiles where the Chombo/lib home directory is. The directory containing the application code can be anywhere relative to the Chombo directory, but the CHOMBO\_HOME variable must be defined appropriately. The next line uses that variable to define the variables the rest of the makefiles need. The next line is application-specific, defining the primary build target. For the example shown here, there should be a file named "main.cpp" in the directory containing this makefile. The Chombo makefile system will compile this file and all the other source files in the directory. Source files are identified for compilation by using a set of suffix rules; any files in the directory with common suffixes like "cpp", "F", etc. will be compiled.<sup>2</sup> The next line specifies which Chombo libraries are needed, which will vary depending on the application. Note that most linkers require that the libraries be listed in the order they will be searched. For Chombo, that generally means that the more basic libraries (like BoxTools) must come after libraries (like AMRTools) which reference them. The XTRALIBFLAGS variable should be defined to specify additional libraries if needed. The value should contain the options to be given to the linker to access the libraries (e.g "-Lsomedir -lsomelib"). The next line defines the "all" target, which will be the default target. The Chombo makefile system handles everything when the target is invoked. Actually, the "all-test" target does the work. The final line includes the makefile that defines this target, and all the rules that it needs.

For more complex applications, a slightly more involved approach allows the Chombo makefile system to build an application with source code in multiple directories. The file

Chombo/releasedExamples/AMRGodunov/execPolytropic/GNUmakefile

demonstrates how this is done. In general, the GNUmakefile looks similar to the description above. The major difference is the addition of a src\_dirs variable which lists the other directories containing source code, along with a base\_dir variable (usually ".") which defines the directory where the source file containing main is located. The final line includes another Chombo makefile (Chombo/lib/mk/Make.example) that defines all the targets and rules. The "make all" command would be used to build the application, just as with the Chombo tests and examples.

#### 1.2.6.2 Using an existing application makefile

To use an existing makefile, it is necessary to modify the rules for compiling C++ code that uses Chombo classes and to modify the link rule to use the Chombo libraries.<sup>3</sup>

The compile rules must be modified to add the "Chombo/lib/include" subdirectory to the search path for C++ header files (-I option for most compilers) and to define some C-preprocessor macro variables that the Chombo header files use. The compiler options for this will usually look something like:

```
-IChombo/lib/include -I<HDF_DIR>/include -DCH_SPACEDIM=<dim>
-DCH_USE_precision> -DCH_<system> -DCH_LANG_CC -DHDF5 -DMPI
```

<sup>&</sup>lt;sup>2</sup>The complete list of suffixes may be found in Chombo/lib/mk/Make.rules

<sup>&</sup>lt;sup>3</sup>This assumes that the Chombo libraries are compiled before trying to compile the application.

The link rule must be modified to add the Chombo "lib" directory to the search path for libraries and to specify the Chombo libraries to be searched. The linker options for this will usually look something like:

```
-LChombo/lib -lamrelliptic<config> -lamrtimedependent<config> -lamrtools<config> -lboxtools<config> -L<HDF_DIR>/lib -lhdf5 -lz
```

where <HDF\_DIR> is as above, "-lhdf5 -lz" are the HDF5 libraries and <config> is the Chombo configuration string. See "Chombo/lib/mk/Make.defs.config" to see how to compute the configuration string.

# 1.2.7 Building Chombo's Doxygen documentation

C++ header files in Chombo libraries are annotated with comments which are compatible with the Doxygen (http://www.doxygen.org) software documentation system. To create html documentation of the Chombo libraries,

```
cd Chombo/lib
make doxygen
```

which will install doxygen-generated html documentation in Chombo/lib/doc/doxygen.

To access it once built, point a web browser at Chombo/lib/doc/doxygen/html/index.html.

To include EB documentation, add EB=TRUE to make doxygen.

Doxygen documentation of the current release is also available online at http://chombo.lbl.gov.

# 1.2.8 Namespace

Chombo can be embedded in its own C++ namespace. While this was part of previous versions of the code, it is more relevant now that Chombo is part of larger development projects. This is controlled by the NamespaceHeader.H and NamespaceFooter.H files that are included in every header and source file in Chombo.

By default, Chombo builds with NAMESPACE=FALSE which turns these include files into empty files, and they have no effect. If you build Chombo with

```
>make NAMESPACE=TRUE
```

Then every class and function defined between NamespaceHeader.H and NamespaceFooter.H will be placed in the Chombo namespace. In practice, this means that one should place

```
#include "NamespaceHeader.H"
```

after all other include files, but before any C++ code in a file, and then

#include "NamespaceFooter.H"

after all C++ code in a file (but within any multiple-inclusion guards, if in a C++ header file). Alternatively,

#include "UsingNamespace.H"

will ensure that the C++ code in a given file is not embedded in a Chombo namespace, but will be able to use Chombo code which is in the Chombo namespace. This is most useful in an application code.

If instead a user uses

#### >make NAMESPACE=MULTIDIM

Then Chombo creates a dimension-dependent namespace Chombo::XD where X is the dimensionality in which Chombo is being built. This can be used to build Chombo in multiple dimensions and use them together in the same application. (see Chapter 11.)

# Chapter 2

# BaseTools and BoxTools

# 2.1 AMR Spatial Discretization

The underlying discretization of space is given as points  $(i_0,...,i_{\mathbf{D}-1})=i\in\mathbb{Z}^{\mathbf{D}}$ . The problem domain is discretized using a grid  $\Gamma\subset\mathbb{Z}^{\mathbf{D}}$  that is a bounded subset of the lattice.  $\Gamma$  is used to represent a cell-centered discretization of the continuous spatial domain into a collection of control volumes:  $i\in\Gamma$  represents a region of space  $[x_0+(i-\frac{1}{2}u)h,x_0+(i+\frac{1}{2}u)h]$ , where  $x_0\in\mathbb{R}^{\mathbf{D}}$  is some fixed origin of coordinates, h is the mesh spacing, and  $u\in\mathbb{Z}^{\mathbf{D}}$  is the vector whose components are all equal to one. We can also define various face-centered and node-centered discretizations of space based on those control volumes. For example, we denote by  $\Gamma^v$  the set of points in physical space of the form  $x_0+(i\pm\frac{1}{2}v)h, i\in\Gamma$ . Here v can be any vector whose entries are equal to either zero or one.

We will find it useful to define a number of operators on points and subsets of  $\mathbb{Z}^{\mathbf{D}}$ . We denote by  $|\boldsymbol{i}| = \max_{d=0\dots\mathbf{D}-1}(|i_d|)$ ,  $\Gamma+\boldsymbol{i}$  as the translation of a set by a point in  $\mathbb{Z}^{\mathbf{D}}$ , and  $\mathcal{G}(\Gamma,r)$  to the set of all points within a  $|\cdot|$ -distance r of  $\Gamma$ 

$$\mathcal{G}(\Gamma,r) = \underset{|m{i}| \leq r}{\cup} \Gamma + m{i}$$

We define a coarsening operator by  $\mathcal{C}_r: \mathbb{Z}^\mathbf{D} \to \mathbb{Z}^\mathbf{D}$ ,

$$\mathcal{C}_r(oldsymbol{i}) = (\lfloor rac{i_0}{r} 
floor, ..., \lfloor rac{i_{d-1}}{r} 
floor)$$

where r is a positive integer. The coarsening operator acting on subsets of  $\mathbb{Z}^{\mathbf{D}}$  can be extended in a natural way to the other grid centerings:  $\mathcal{C}_r(\Gamma^v) \equiv (\mathcal{C}_r(\Gamma))^v$ .

We extend this discretization of space to represent a nested hierarchy of grids that discretize the same continuous spatial domain. We assume that our problem domain can be discretized by a nested hierarchy of grids  $\Gamma^0...\Gamma^{lmax}$ , with  $\Gamma^{l+1}=\mathcal{C}^{-1}_{n^l_{ref}}(\Gamma^l)$ , and  $\boldsymbol{x}^l_0-\frac{1}{2}\boldsymbol{u}h^l$  independent of l. The integer  $n^l_{ref}$  is the refinement ratio between level l and

l+1. We also assume that the mesh spacings  $h^l$  associated with  $\Gamma^l$  satisfy  $\frac{h^l}{h^{l+1}}=n^l_{ref}$ . These conditions imply that the underlying continuous spatial domains defined by the control volumes are all identical.

Adaptive mesh refinement calculations are performed on a hierarchy of meshes  $\Omega^\ell \subset \Gamma^\ell$ , with  $\Omega^\ell \supset \mathcal{C}_{n^\ell_{ref}}(\Omega^{\ell+1})$ . Typically,  $\Omega^l$  is decomposed into a disjoint union of rectangles in order to perform calculations efficiently. We denote such a decomposition by  $\mathcal{R}(\Omega^l) = \{\Omega^l_k\}_{k=1}^{N^l_{grids}}$ , where the  $\Omega^l_k$ 's are rectangles and  $\Omega^l_k \cap \Omega^l_{k'} = \emptyset$  if  $k \neq k'$ . We say that  $\mathcal{R}(\Omega^l)$  is p-blocked, p>1, if  $\Omega^l_k = \mathcal{C}_p^{-1}(\mathcal{C}_p(\Omega^l_k))$  for all k. We will assume throughout that  $\Omega^l$  admits a decomposition  $\mathcal{R}(\Omega^l)$  that is  $n^{l-1}_{ref}$ -blocked for all l>0. In particular, the control volume corresponding to a cell in  $\Omega^{l-1}$  is either completely contained in the control volumes defined by  $\Omega^l$  or its intersection has zero volume. We will also assume that there is at least one level l cell separating level l+1 cells from level l-1 cells:  $\mathcal{G}(\mathcal{C}_{n^l_{nref}}(\Omega^{l+1}),1)\cap \Gamma^l\subseteq \Omega^l$ . We will refer to grid hierarchies that meet these two conditions as being p-roperly nested. We emphasize that this form of proper nesting is a minimum requirement for the AMR algorithms discussed in this document. For some applications, it may be necessary to impose more stringent conditions on the grid hierarchy.

A discretized dependent variable in AMR is a level array

$$\varphi^l:\Omega^l\to\mathbb{R}^m$$

We denote by  $\varphi_i \in \mathbb{R}^m$  the value of  $\varphi$  at cell  $i \in \Omega^l$ . We can also define level arrays on other grid centerings, i.e.,  $\psi: \Omega^{l,v} \to \mathbb{R}^m$ . In that case, we denote the indexing operation by  $\psi_{i+\frac{1}{3}v} \in \mathbb{R}^m$ . In particular, we can define vector fields at a level

$$ec{F}^l = (F_0^l, ..., F_{\mathbf{D}-1}^l)$$
 ,  $F_d^l: \Omega^{l, oldsymbol{e}^d} 
ightarrow \mathbb{R}^m$ 

We will be interested in operations on pairs of refined grids that are not necessarily contained in an AMR mesh hierarchy (e.g., during regridding). In those cases, we will denote by  $\Gamma^f$ ,  $\Gamma^c$  the fine and coarse problem domains,  $n_{ref}$  the refinement ratio between the two levels,  $\Omega^f$ ,  $\Omega^c$  the refined regions in the two domains, and  $\varphi^f$ ,  $\varphi^c$ , etc., level arrays defined on  $\Omega^f$ ,  $\Omega^c$ . We will always assume that the two levels are properly nested.

In the remainder of this section, we will describe *BoxTools*, a set of abstractions for defining points and regions in a multidimensional integer lattice index space, and representing aggregate data in such regions. The classes defined in the remainder of this chapter correspond to the mathematical objects described above in the following fashion.

- ullet Points in the rectangular lattice  $i\in\mathbb{Z}^{\mathrm{D}}\Leftrightarrow$  the class IntVect.
- $\bullet$  Rectangular subsets  $\Gamma \subset \mathbb{Z}^{\mathbf{D}} \Leftrightarrow$  the class Box.
- $\bullet$  Arbitrary subsets  $\mathcal{I} \subset \mathbb{Z}^\mathbf{D} \Leftrightarrow$  the class <code>IntVectSet</code>.
- Rectangular arrays  $\varphi: \Gamma \to \mathbb{R}^m \Leftrightarrow$  the class FArrayBox.
- ullet Unions of rectangles at a fixed level of refinement  $\Omega, \mathcal{R}(\Omega)$ , and their distribution onto processors  $\Leftrightarrow$  the class DisjointBoxLayout.
- ullet Level arrays  $\varphi:\Omega o\mathbb{R}^m\Leftrightarrow$  the class LevelData<FArrayBox>.

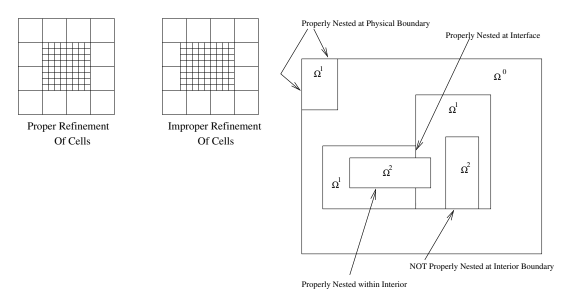


Figure 2.1: Examples illustrating proper nesting requirements for locally refined grids.

# 2.2 Points, Regions and Rectangular Arrays

BoxTools is a set of abstractions for defining points and regions in a multidimensional integer lattice index space, and representing aggregate data in such regions. The dimensionality  $\mathbf D$  of the index space is a compilation-time constant. It is accessed as a macro CH\_SPACEDIM, which is set in the Make process, and is propagated into Fortran .F or .ChF files in applying the C preprocessor. A second compile-time constant in BoxTools is that of the precision of floating-point variables. BoxTools provides a type Real, which is set using a typedef declaration to either float or double at compile time. The macro REAL\_T serves the same function in Fortran. This macro is defined in the file REAL\_H, which can be included as a header for both C++ and Fortran files.

## 2.2.1 Class IntVect

IntVects represent points in the rectangular lattice  $\mathbb{Z}^D$ .

**Operations on** IntVects. In the following definitions i, j are IntVects and s, d are integers,  $0 \le d < \mathbf{D}$ .

- <u>Constructors.</u> IntVect has the usual default and copy constructors, as well as constructors that take tuples of integers as arguments, e.g., IntVect( $i_0, i_1$ ) (two dimensions), IntVect( $i_0, i_1, i_2$ ) (three dimensions).
- Arithmetic operators.  $i \oplus j, i \oplus s, \oplus \in \{+, -, *, /\}$  produce IntVects by operating componentwise on the inputs. +=, -=, \*=, /=, perform the same operations in place. e.g., i+=j is the same as i=i+j. IntVect also provides component-wise  $\min(i,j), \max(i,j)$  operators.

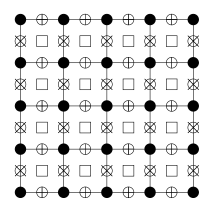


Figure 2.2: Vertex  $(\bullet)$ , cell  $(\Box)$  and face  $(\oplus$  and  $\otimes)$  locations on a grid.

- Logical operators.  $i_1 == i_2$ ,  $(i_1! = i_2)$  Test for mathematically equal (unequal) IntVects. Comparison operators are defined element-wise: >,>=,<,<=, i < j iff  $i_d < j_d$ . Lexicographic ordering operators i.lexlt(j), i.lexgt(j) are also provided.
- <u>Static members.</u> Unit is the IntVect consisting of all ones. Zero is the vector consisting of all zeros. BASISV(d), d=0,..., D-1 returns the unit IntVect in the d direction.
- Indexing operations. i[d] returns the component of i, and can be used to assign values to components: i[d] = q.

# **2.2.2 Class** Box

A Box represents a rectangular region in  $\mathbb{Z}^{\mathbf{D}}$ , defined by specifying the IntVects defining its low and high corners. For each coordinate direction, a Box can be *cell-centered* or *node-centered*. This allows one to represent the various face-, edge-, and node-centered rectangular grids (figure 2.2).

**Operations on** Boxes. In what follows, B,  $B_1$ ,  $B_2$  are Boxes, i,  $i_1$ ,  $i_2$ , v are IntVects, v having components equal to zero or one, and s, d are integers,  $0 \le d < D$ .

- Constructors.  $B(i_1, i_2, v = {\tt Zero})$  Constructs a Box with low and high corners  $i_1, i_2$ , and centering defined by v. If  $v_d = 0$ , then the Box is cell-centered in the d direction; if  $v_d = 1$ , then the Box is node-centered in the d direction. In particular, the default centering is cell-centered in all three directions. Box has a copy constructor and assignment operator. One can reset the low and high corners of the Box (setSmall(i), setBig(i)).
- Logical functions.  $B_1 == B_2$ ,  $B_1 != B_2$  test whether  $B_1$  and  $B_2$  are equal or unequal, including having the same centering.  $B.\mathtt{isEmpty}()$  tests whether B is empty.  $B.\mathtt{contains}(B_1)$ ,  $B.\mathtt{contains}(\boldsymbol{i})$ , tests whether B contains  $B_1, \boldsymbol{i}$ .  $B_1.\mathtt{intersects}(B_2)$  checks whether  $B_1 \cap B_2 \neq \emptyset$ .  $B_1.\mathtt{sameType}(B_2)$ ,  $B_1.\mathtt{sameSize}(B_2)$  check whether  $B_1$  and  $B_2$  have the same centering, or whether  $B_1 = B_2 + \boldsymbol{i}$  for some  $\boldsymbol{i}$ .  $B_1 < B_2$  if  $B_1.\mathtt{smallEnd}().\mathtt{LexLT}(B_2.\mathtt{smallEnd}())$ .

- Shifting and Centering.  $B.\mathtt{convert}(v)$  changes the centering of B to that specified by v, as in the constructor. One can also change the centering in one direction.  $B.\mathtt{surroundingNodes}()$  converts all the cell-centered directions to node-centered, and increments the high corner in those directions by 1.  $B.\mathtt{surroundingNodes}(d)$  performs the same operation in the d coordinate direction.  $B.\mathtt{enclosedCells}()$ ,  $B.\mathtt{enclosedCells}(d)$  perform the opposite operation, converting the centerings to be cell-centered, and decrementing by 1 the high corner of the box for those directions for which the centering changed. There are also corresponding friend functions, e.g.,  $\mathtt{surroundingNodes}(B)$  that return a new box with the appropriately modified centerings. The various grids depicted in figure 2.2 can be obtained from on another by application of the member functions  $\mathtt{surroundingNodes}$  and  $\mathtt{enclosedCells}$ .  $B.\mathtt{shift}(i)$ , B+=i perform the identical operation of replacing B with  $\mathtt{Box}(B.\mathtt{smallEnd}()+i, B.\mathtt{bigEnd}()+i)$ .  $B.\mathtt{shift}(d,s)$  is the same as B+=s\*BASISV(d). B-=i is the same as B+=(-i).
- Size functions. B.smallEnd(), B.bigEnd(), B.size(), return IntVects containing the low corner, high corner, and size in each direction. The same functions called with an integer argument (B.size(s)) returns the s-th component of those IntVects. B.numPts() returns the discrete volume of  $B.\ B.loVect(), B.highVect()$ , return pointers to the D-tuples of integers defining the low and high corners of B in order to pass them to Fortran.
- Set operations. Although it is not possible to define a complete set calculus on Boxes (the union of two rectangles is not always a rectangle), Box provides many of the set functions most commonly required.  $B_1\&=B_2$  sets  $B_1=B_1\cap B_2$ .  $B.\min Box(B_2)$  sets  $B_1$  to be the minimum sized Box containing  $B_1,B_2$ .  $B.\operatorname{grow}(s)$  grows B in all directions by a size s (s can be negative corresponding to shrinking).  $B.\operatorname{grow}(i)$  grows B by  $i_d$  in the dth direction, and  $B.\operatorname{grow}(d,s)=B.\operatorname{grow}(s*BASISV(d))$ .  $B.\operatorname{coarsen}(s)=\mathcal{C}_s(B)$ ,  $B.\operatorname{refine}(s)=\mathcal{C}_s^{-1}(B)$ . B can also be coarsened and refined by different amounts in the various coordinate directions using  $B.\operatorname{coarsen}(i)$ ,  $B.\operatorname{refine}(i)$ . Grow, minBox, coarsen, refine all have corresponding friend functions that return a new Box on which the operation has been performed, e.g.,  $\min Box(B_1,B_2)$ .  $\operatorname{adjCellLo}(B,d,s=1)$  returns the cell-centered box of width s direction adjacent to s0 on the low side in the s1 the s2 direction adjBdryLo, s3 direction on the high side of s3 in the s4 direction and s4 direction and s5 adjBdryHi return the corresponding node-centered Boxes.

# 2.2.3 Class IntVectSet

IntVectSet represents an irregular region in an integer lattice **D**-dimensional index space as an arbitrary collection of IntVects. A full set calculus is defined.

**Operations on** IntVectSets. In the following,  $\mathcal{I}, \mathcal{I}_1, \mathcal{I}_2$  are IntVectSets, B is a Box, and s is an integer.

Constructors. The default constructor constructs an empty IntVectSet. They can

also be initialized at construction with an IntVect, a Box or another IntVectSet. An existing IntVectSet can also be re-initialized with any of those three objects using the member functions define. IntVectSet has an assignment operator.

- Set Operations. IntVectSets can be updated in place by taking unions  $(\mathcal{I}_1|=\mathcal{I}_2, \mathcal{I}|=B, \mathcal{I}|=i)$  intersections  $(\mathcal{I}_1\&=\mathcal{I}_2, \mathcal{I}\&=B, \mathcal{I}\&=i)$  and set-theoretic differences  $(\mathcal{I}_1-=\mathcal{I}_2, \mathcal{I}_2-=B, \mathcal{I}_3-=i)$  with another IntVectSet, a Box or an IntVect.  $\mathcal{I}.\mathsf{coarsen}(s)$  sets  $\mathcal{I}$  to  $\mathcal{C}_s(\mathcal{I})$ ,  $\mathcal{I}.\mathsf{refine}(s)$  changes  $\mathcal{I}$  to  $\mathcal{C}_s^{-1}(\mathcal{I})$ .  $\mathcal{I}.\mathsf{grow}(s)$  changes  $\mathcal{I}$  to  $\bigcup_{i:|i|\leq s}(\mathcal{I}+i)$ . Union, intersection, difference, coarsen, and refine all have associated friend functions that return a new IntVectSet suitably modified. For example,  $\mathcal{I}_1|\mathcal{I}_2$  returns  $\mathcal{I}_1\bigcup\mathcal{I}_2$ , leaving  $\mathcal{I}_1$  and  $\mathcal{I}_2$  unchanged.  $\mathsf{shift}(\mathcal{I},i)$  returns  $\mathcal{I}_1$ .
- Other functions.  $\mathcal{I}.$ isEmpty() returns true if  $\mathcal{I} = \emptyset$ .  $\mathcal{I}.$ minBox() returns the minimum cell-centered Box containing  $\mathcal{I}$ .  $\mathcal{I}.$ contains(B),  $\mathcal{I}.$ contains(i) returns true if  $B \subset \mathcal{I}, i \in \mathcal{I}$ .

**Performance Issues**. IntVectSet uses two representations internally: a fast bitmap for small sets, and a slower tree representation for large sets. The heuristic employed between switching between the two representations is to use bitmaps for sets that are initialized to be rectangles, or that are obtained by applying intersection, set-theoretic difference, coarsen, refine or grow to IntVectSet(s) that are represented by bitmaps. However, the use of the union function causes the representation to be converted irreversibly to a tree representation, with a significant performance penalty. For that reason, the union operations should be used sparingly.

## 2.2.4 Box and IntVectSet Iterators

A BoxIterator or IVSIterator traverses a sequence of IntVects that comprise a given Box or IntVectSet. Each IntVect appears exactly once in the sequence. There is no guarantee that the IntVects will appear in any particular order.

**Operations on** BoxIterator, IVSIterator. In what follows, B is a Box,  $\mathcal{I}$  is an IntVectSet, and iter is either a BoxIterator or an IVSIterator.

- <u>Construction</u> The iterators can be constructed with object to be iterated over (iter(B), iter( $\mathcal{I}$ )), or null-constructed and defined later (iter.define(B), iter.define( $\mathcal{I}$ )).
- <u>Iteration</u>. iter.begin() sets iter to the beginning of the iteration sequence, ++iter advances iter to the next iterate, and iter.ok() checks to see if the current iterate is valid. A null-constructed iterator, or an iterator constructed with an empty Box or IntVectSet will always return false. iter() returns an IntVect containing the current value of the iterate.

# 2.2.5 Class Interval

An Interval consists of two ordered integers. An Interval can be created only by specifying its endpoints. The only operations that can be performed are to extract its

endpoints or determine its size, which is the number of integers it contains. If the endpoints are equal, the size is one. It is permitted to define an Interval with zero or negative size. It is entirely the responsibility of the user to determine whether this is valid. Interval interacts only weakly with the other abstractions and is exclusively used to specify data component ranges in Chombo (see sections 2.2.6 and 2.4.3).

# 2.2.6 Rectangular arrays

A BaseFab<T> is a templated container class for multidimensional array data. It consists of three major elements: a Box to define the range of spatial indices over which the array is defined; an integer specifying the number of components; and a T\* pointing to a contiguous block of array elements. The data is stored in Fortran order so that the pointer can be passed to a Fortran routine where it can be accessed as a multidimensional array.

A BaseFab is defined by specifying a domain in the form of a Box, which can have any centering, and the number of components, ncomps. This is intended to represent a  $\mathbf{D}+1$  dimensional array in Fortran. A component index is an integer in the range 0 to ncomps-1 which is used to specify or select a component of a BaseFab. A range of component indices is often represented by an Interval. An already defined BaseFab can be redefined with a new domain and number of components. The behavior of existing data is undefined during redefinition. BaseFabs are large aggregate objects containing pointer data, so that shallow copy can lead to subtle bugs, and deep copying is expensive. For that reason, the assignment operator and copy constructor have been rendered inaccessible to the user by making them private. In particular, it is necessary to pass BaseFabs as reference parameters in procedure calls.

**Operations on** BaseFabs. In the following  $\mathcal{A}, \mathcal{A}'$  are BaseFabs, B is a Box, i is an IntVect, and  $n_{comp}, d, s$  are integers, with  $0 \le d < \mathbf{D}, n_{comp} > 0$ .

- <u>Constructors</u>. BaseFab has a default constructor, as well as a constructor BaseFab(B) that completely defines the BaseFab.  $\mathcal{A}.\mathtt{resize}(B,n_{comp})$  resets  $\mathcal{A}$  to be defined over a Box B and with n components. Any data contained in  $\mathcal{A}$  previously is discarded, and the data  $\mathcal{A}$  is assumed to be uninitialized.
- Accessors.  $\mathcal{A}(i,s)$  is an indexing operator, returning a reference of type T& to the storage location for the value at point i and component s. For a BaseFab that is node-centered in one or more of the coordinate directions, the convention for indexing with an IntVect (which does not have centering) is that  $\mathcal{A}(i,s)$  returns the reference corresponding to  $i-\frac{1}{2}v$ , where v is the IntVect of zeros and ones defining the centering, i.e., the cell center and the node-centered points on the low side have the same index.  $\mathcal{A}.\text{box}()$  returns the Box over which  $\mathcal{A}$  is defined, and  $\mathcal{A}.\text{ncomp}()$  the number of components. BaseFab provides an interface to the Box member functions smallEnd(), bigEnd(), loVect(), hiVect():  $\mathcal{A}.\text{smallEnd}() = \mathcal{A}.\text{box}().\text{smallEnd}()$ , etc.  $\mathcal{A}.\text{dataPtr}(s)$  returns a pointer of type T\* to the data in  $\mathcal{A}$  beginning at the nth component; n defaults to

zero. A.nCompPtr() returns a pointer to an integer containing the number of components. loVect, hiVect, dataPtr, nCompPtr are to be used in calling Fortran.

- Data Modification Functions.  $\mathcal{A}.\mathtt{setVal}(t)$  sets all of the data values in  $\mathcal{A}$  to the single value t.  $\mathcal{A}.\mathtt{copy}(\mathcal{A}_2)$  copies all of the values in  $\mathcal{A}_2$  into the part of  $\mathcal{A}_1$  defined on  $\mathcal{A}.\mathtt{box}()\&\mathcal{A}_2.\mathtt{box}()$ .  $\mathcal{A}_1$  and  $\mathcal{A}_2$  must have the same number of components. Both setVal and copy have overloaded versions that permit the operations to be performed on a specified sub-rectangle and over subsets of the component ranges.
- <u>Domain Modification Functions</u>  $\mathcal{A}.\mathtt{shift}(i)$  changes the Box over which  $\mathcal{A}$  is defined to  $\mathcal{A}.\mathtt{box}()+i$ , leaving the data unmodified. Mathematically,  $\mathcal{A}$  becomes  $\mathcal{A}'$ , with  $\mathcal{A}'(j,s)==\mathcal{A}(j-i,s), \forall j\in\mathcal{A}'.\mathtt{box}()$ . The shift function is overloaded to shift  $\mathcal{A}$  by some distance in a single coordinate direction  $(\mathcal{A}.\mathtt{shift}(d,s))$ .  $\mathcal{A}.\mathtt{shiftHalf}(i)$  shifts the domain of  $\mathcal{A}$  by i "halves" in each direction, where a half-shift changes the centering to the adjacent nodes/ cells centered Box in that direction.

# **FArrayBox**

An FArrayBox is-a BaseFab<Real> which contains floating-point data. A number of aggregate floating-point arithmetic operations are provided. FArrayBox is implemented as a derived class from BaseFab<Real>. In addition to BaseFab operations, FArrayBox has a collection of operations that are specialized to real-valued arrays.

**Note:** All FArrayBox objects are initialized upon creation, using setVal with an argument of 1.23456789e10, whether the Chombo libraries are compiled with debugging on or off.

- Pointwise Arithmetic Operators  $\mathcal{A}_1 \oplus = \mathcal{A}_2$ ,  $\oplus \in \{+, -, *, /\}$ , updates in place the values of  $\mathcal{A}_1(i,s)$  with  $\mathcal{A}_1(i,s) \oplus \mathcal{A}_2(i,s)$ , for  $0 \le s < \mathcal{A}_1.n\text{Comps}() = \mathcal{A}_2.n\text{Comps}()$ , and  $i \in \mathcal{A}_1.\text{box}() \cap \mathcal{A}_2.\text{box}()$ . There are also a collection of member functions plus, minus, mult, divide, that perform these operations over subboxes and subranges of the components.  $\mathcal{A}.\text{abs}()$  updates in place the values of  $\mathcal{A}$  with their absolute values. abs is overloaded with versions specifying subbox and a single component. The unary operators negate and invert behave in a similar fashion to abs.
- Reduction Operators  $\mathcal{A}.\mathtt{sum}(s)$ ,  $\mathcal{A}.\mathtt{min}(s)$ ,  $\mathcal{A}.\mathtt{max}(s)$ , return real values containing the sum, minimum, and maximum of the values of the s-th component of  $\mathcal{A}$ .  $\mathcal{A}.\mathtt{minIndex}(s)$ ,  $\mathcal{A}.\mathtt{maxIndex}(s)$  return IntVects corresponding to one of the locations  $\boldsymbol{i}$  such that the minimum or maximum is attained.  $\mathcal{A}.\mathtt{norm}(p,s), p \geq 1$  returns the discrete p norm of the s-th component of  $\mathcal{A}$ .  $\mathcal{A}.\mathtt{norm}(p,s) = (\sum_{\boldsymbol{i} \in \mathcal{A}.\mathtt{box}()} |\mathcal{A}(\boldsymbol{i},s)|^p)^{1/p}$ . There are also overloaded versions of these functions that perform their operations over a subbox, or for a range of components.
- Mask Functions  $\mathcal{A}.\mathtt{maskLT}(M,a,s)$  sets the values of the input BaseFab<int> M to one or zero, depending on whether  $\mathcal{A}(\boldsymbol{i},s) < a$  or not. M is resized by the function so that  $M.\mathtt{box}() = \mathcal{A}.\mathtt{box}()$ . maskLT also returns the integer number of non-zero entries in M. maskLE, maskEQ, maskGT, maskGE are defined similarly.

The Fortran Interface. The collection of values taken on by a BaseFab  $\mathcal A$  is stored in a contiguous block of storage beginning at  $\mathcal A.\mathtt{dataPtr}()$ . The data is stored in Fortran ordering corresponding with the spatial indices first, followed by the component index. Specifically, if a Fortran routine is called from C++

```
extern "C" {foo_(real*, int*, int*, int*);}

FArrayBox A(B,nc);
foo_(A.dataPtr(),A.loVect(),A.hiVect(),A.nCompPtr())
```

The indexing of A in the Fortran routine is given by

```
subroutine foo(a,lo,hi,nc)
integer lo(0:CHF_SPACEDIM-1)
integer hi(0:CHF_SPACEDIM-1)
real_t a(lo(0):hi(0),lo(1):hi(1),0:nc-1)

do ic =0,nc-1
    do j = lo(1),hi(1)
    do i = lo(0),hi(0)
```

For further details on the Fortran interface, see Chapter 8.

## 2.2.6.1 **Aliasing**

Often, one wants to use a subset of the components of a larger BaseFab<T> in a particular part of an algorithm. Rather than defining a completely new object and then copying back and forth, BaseFab<T>, and by inheritance FArrayBox, can also be built as an *alias* of another BaseFab<T> (where T is the same for the two objects). For example:

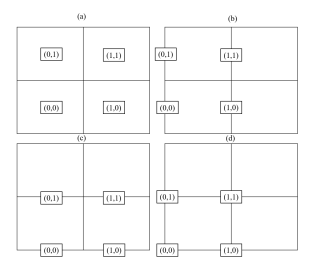


Figure 2.3: Four data centerings and corresponding Chombo index conventions associated with a  $2\times 2$  cell-centered 2D box from (0,0) to (1,1). (a) Cell-centered data, (b) x-face centered data, (c) y-face centered data, and (d) node-centered data.

This BaseFab does not allocate any memory, but sets its data pointer into the memory pointed to by the argument BaseFab. It is the users responsibility to ensure this aliased BaseFab is not used after the original BaseFab has deleted its data member (resize, define(..) called, or destruction, etc.).

This is similar to using an offset pointer into an array. The offset pointer is only valid as long as the original array is valid.

This aliased BaseFab will also generate side effects (modifying the values of data in one will modify the other's data). Deleting the alias will not affect the original.

This aliased BaseFab will have subcomps.size() components, starting at zero. The aliased BaseFab can only have the same Box domain as the original.

## 2.2.7 Dataholders for non-cell-centered data

While the primary centering used in Chombo is cell-centered data, support for data which uses the alternate centerings described in section 2.2.2 and shown in Figure 2.2 is also provided for convenience. This makes it convenient to define and use differently-centered data associated with the cell-centered Boxes in a BoxLayout or DisjointBoxLayout (described in section 2.4.2.1) in a way that maps naturally to the way algorithms are often defined. The various data centerings associated indexing conventions are shown in Figure 2.3.

#### 2.2.7.1 FluxBox

The FluxBox class encapsulates data on the faces of a cell-centered box. It is essentially made up of a Vector of SpaceDim FArrayBoxes, one for each set of faces generated by a surroundingNodes(dir) operation on the cell-centered Box used to define the FluxBox. The normal usage pattern for FluxBox is to extract the face-centered FArrayBox for a given face direction and then proceed in a similar way to the way one uses a cell-centered FArrayBox, with the convention that in the ith direction, the face center is centered on the low-side of the corresponding cell center. For example, in the x-direction (dir = 0), the cell-center of cell (i,j,k) in 3D is located at  $(x,y,z) = \Delta x(i+\frac{1}{2},j+\frac{1}{2},k+\frac{1}{2})$ , and the face-center is located at  $(x_{face},y_{face},z_{face}) = \Delta x(i,j+\frac{1}{2},k+\frac{1}{2})$ . Note that for face-centered data there is one node-centered coordinate and SpaceDim-1 cell-centered coordinates.

One accesses the dir-face FArrayBox in a FluxBox by using the [] operator:

```
// define cell-centered box.
Box b;
// define FluxBox on faces associated with b
FluxBox fluxes(b, 1);
// access the x-direction face-centered data in fluxes
FArrayBox& xFaceData = fluxes[0];
```

# 2.2.7.2 EdgeDataBox

The EdgeDataBox class encapsulates data on the edges of a cell-centered box. It is essentially made up of a Vector of SpaceDim FArrayBoxes, one for each set of edges generated from a cell-centered Box by a surroundingNodes() operation (in all directions) followed by an enclosedCells(dir) operation. As with FluxBox, the normal usage pattern for EdgeDataBox is to extract the edge-centered FArrayBox for a given edge direction and then proceed to use in a way similar to a cell-centered FArrayBox, with the convention that in the ith edge direction, the edge center is centered to the low-side of the corresponding cell center. For example, in the x-direction (dir = 0), the cell-center of cell (i,j,k) in 3D is located at  $(x,y,z) = \Delta x(i+\frac{1}{2},j+\frac{1}{2},k+\frac{1}{2})$ , and the edge-center is located at  $(x_{edge},y_{edge},z_{edge}) = \Delta x(i+\frac{1}{2},j,k)$ . Note that in this case there is one cell-centered coordinate and SpaceDim-1 node-centered coordinate.

One accesses the dir-face FArrayBox in a EdgeDataBox by using the [] operator:

```
// define cell-centered box.
Box b;
// define EdgeDataBox on edges associated with b
edgeDataBox edgeData(b, 1);
// access the x-direction edge-centered data in edgeData
FArrayBox& xEdgeData = edgeData[0];
```

# 2.2.7.3 NodeFArrayBox

The NodeFArrayBox encapsulates data on the nodes of a cell-centered box, constructed by calling surroundingNodes() on the cell-centered Box. Data is located at  $(x_{node}, y_{node}, z_{node}) = \Delta x(i, j, k)$ . Note that in this case all SpaceDim coordinates are node-centered. Accessing the node-centered FArrayBox is via the getFab() function:

```
// define cell-centered box.
Box b;
// define NodeFArrayBox on nodes associated with b
NodeFArrayBox nodeData(b, 1);
// access the node-centered data in in nodeData
FArrayBox& data = nodeData.getFab();
```

# 2.3 Class ProblemDomain

ProblemDomain is a class to handle interaction with boundary conditions at the edge of the computational domain, either physical boundary conditions or periodic ones. This class contains much of the functionality of the Box class, since logically the computational domain is generally a Box.

Intersection with a ProblemDomain object will result in only removing regions which are outside the physical domain in non-periodic directions. Regions outside the logical computational domain in periodic directions will be treated as ghost cells which can be filled with an exchange() function or through suitable interpolation from a coarser domain.

Since ProblemDomain will contain a Box, it is a dimension-dependent class, so SpaceDim must be defined appropriately when compiling.

Note that this implementation of ProblemDomain is inherently cell-centered.

The user interface for ProblemDomain is as follows:

• ProblemDomain()

Default constructor – the object is defined in an unusable state until the user calls the define function.

ProblemDomain(const Box& domBox, const bool\* isPeriodic)
 Full constructor. Places the ProblemDomain object in a usable state.

#### **Arguments:**

- domBox Computational domain.
- isPeriodic SpaceDim array of bools which defines whether BC's are physical or periodic in each coordinate direction.

• ProblemDomain(const Box& domBox)

Partial constructor, creates non-periodic (in any coordinate direction) ProblemDomain.

## **Arguments:**

- domBox Computational domain.
- ProblemDomain(const IntVect& small, const IntVect& big, const bool\* isPeriodic)

Full constructor, creates ProblemDomain.

## **Arguments:**

- small Location of lower-left corner of domain box
- big Location of upper-right corner of domain box
- isPeriodic SpaceDim array of bools which defines whether BC's are physical or periodic in each coordinate direction.
- ProblemDomain(const IntVect& small, const IntVect& big)
   Partial constructor, creates non-periodic (in any coordinate direction) ProblemDomain.

## **Arguments:**

- small Location of lower-left corner of domain box
- big Location of upper-right corner of domain box
- ProblemDomain(const IntVect& small, const int\* vec\_len, const bool\* isPeriodic)

Full constructor, creates ProblemDomain.

#### **Arguments:**

- small Location of lower-left corner of domain box
- vec\_len Size of domain in each direction.
- isPeriodic SpaceDim array of bools which defines whether BC's are physical or periodic in each coordinate direction.
- ProblemDomain(const IntVect& small, const int\* vec\_len)
   Partial constructor, creates ProblemDomain with non-periodic boundary conditions by default.

## **Arguments:**

- small Location of lower-left corner of domain box
- vec\_len Size of domain in each direction.
- ProblemDomain(const ProblemDomain& src)
   Copy constructor
- const Box& domainBox() const
   Returns logical computational domain.
- bool isPeriodic(int dir) const
   Returns true if boundary condition is periodic in direction dir.
- bool isPeriodic() const
   Returns true if boundary condition is periodic in any direction.
- ShiftIterator shiftIterator() const

Returns ShiftIterator for this problem domain. ShiftIterator is a utility class to aid with periodic boundary conditions whose use is mostly internal to BoxLayout, Copier, etc which allows looping over IntVects which used to shift the domain for enforcing periodic boundary conditions.

- bool isEmpty() const
   Returns true if this ProblemDomain has an empty domainBox.
- bool contains(const IntVect& p) const
   Returns true if argument is contained within this ProblemDomain. An empty ProblemDomain does not contain and is not contained by any ProblemDomain. In a periodic direction, all locations are contained, since a periodic domain is an infinite domain. If periodic in all directions, this will always return true.
- bool contains(const Box& b) const

Returns true if argument is contained within this ProblemDomain. An empty ProblemDomain does not contain and is not contained by any ProblemDomain. In a periodic direction, all locations are contained, since a periodic domain is an infinite domain. If periodic in all directions, this will always return true.

bool intersects( const Box& a\_box) const
 Returns true if this ProblemDomain and the argument have non-null intersections.
 It is an error if a\_box is not cell-centered. An empty ProblemDomain does not

intersect any Box. Boxes always intersect in periodic dimensions, since a periodic domain is an infinite domain. If periodic in all directions, this will always return true.

• bool intersectsNotEmpty (const Box& a\_box) const

Returns true if this ProblemDomain and the argument have non-null intersections. It is an error if  $a\_box$  is not cell-centered. This routine does not perform the check to see if \*this or b are empty boxes. It is the callers responsibility to ensure that this never happens. If you are unsure, the use the .intersects(..) routine. In periodic directions, will always return true.

- bool intersects(const Box& box1, const Box& box2) const Returns true of box1 and box2 and any of their periodic images intersect. (This is useful for checking disjointness).
- ProblemDomain& operator= (const ProblemDomain& b)
   Assignment operator.
- void setPeriodic(int a\_dir, bool a\_isPeriodic)
   Sets whether boundary condition is periodic in direction a\_dir (true is periodic).
- friend

Box bdryLo(const ProblemDomain& a\_pd, int a\_dir, int a\_len=1)

Returns the edge-centered box (in direction  $a_{-}$ dir) defining the low side of the domainBox in the argument ProblemDomain. The neighbor of an empty ProblemDomain is an empty Box of the appropriate type. If dir is a periodic direction, will return an empty box.

#### **Arguments:**

- a\_pd input ProblemDomain
- a\_dir normal direction of edge to return. Directions are zero-based and must be  $0 \le a_dir < SpaceDim$ .
- a\_len Width of returned box in normal direction a\_dir.

#### • friend

Box bdryHi(const ProblemDomain& a\_pd, int a\_dir, int a\_len=1)

Returns the edge-centered box (in direction a\_dir) defining the high side of the domainBox in the argument ProblemDomain. The neighbor of an empty ProblemDomain is an empty Box of the appropriate type. If dir is a periodic direction, will return an empty box.

#### **Arguments:**

- a\_pd input ProblemDomain
- a\_dir normal direction of edge to return. Directions are zero-based and must be  $0 \le a_dir < SpaceDim$ .
- a\_len Width of returned box in normal direction a\_dir.

#### • friend

Box adjCellLo(const ProblemDomain& a\_pd, int a\_dir, int a\_len=1)

Returns the cell-centered box (in direction a\_dir) adjacent to the low side of the domainBox in the argument ProblemDomain. The neighbor of an empty ProblemDomain is an empty Box of cell-centered type. If dir is a periodic direction, will return an empty box.

## **Arguments:**

- a\_pd input ProblemDomain
- a\_dir normal direction of side to return. Directions are zero-based and must be  $0 \le a_dir < SpaceDim$ .
- a\_len Width of returned box in normal direction a\_dir.

#### • friend

Box adjCellLo(const ProblemDomain& a\_pd, int a\_dir, int a\_len=1) Returns the cell-centered box (in direction a\_dir) adjacent to the low side of the domainBox in the argument ProblemDomain. The neighbor of an empty ProblemDomain is an empty Box of cell-centered type. If dir is a periodic direction, will return an empty box.

#### **Arguments:**

- a\_pd input ProblemDomain
- a\_dir normal direction of side to return. Directions are zero-based and must be  $0 \le a_dir < SpaceDim$ .
- a\_len Width of returned box in normal direction a\_dir.
- Box operator& (const Box& b) const

Returns the Box that is the intersection of the input box b and the ProblemDomain. The Box b must be cell-centered. The intersection of an empty ProblemDomain and any box is the Empty Box. This operator does nothing in periodic directions (since a periodic domain is an infinite domain).

• ProblemDomain& refine(int a\_refinement\_ratio)

Modifies this ProblemDomain by refining it by (the positive) a\_refinement\_ratio. The empty ProblemDomain is not modified by this function.

#### • friend

Returns a ProblemDomain that is the argument ProblemDomain refined by (the positive) a\_refinement\_ratio. If a\_probdomain is an Empty ProblemDomain, then an empty problem domain is produced.

ProblemDomain& refine(const IntVect& a\_refinement\_ratio)

Modifies this ProblemDomain by refining it by the given refinement ratio in each direction. The empty ProblemDomain is not modified by this function.

#### friend

ProblemDomain refine (const ProblemDomain& a\_probdomain, const IntVect& a\_refinement\_ratio)

Returns a ProblemDomain that is the argument ProblemDomain refined by (the positive) a\_refinement\_ratio. If a\_probdomain is an Empty ProblemDomain, then an empty problem domain is produced.

• ProblemDomain& coarsen(int a\_refinement\_ratio)

Modifies this ProblemDomain by coarsening it by (the positive) a\_refinement\_ratio. The empty ProblemDomain is not modified by this function.

#### • friend

Returns a ProblemDomain that is the argument ProblemDomain coarsened by (the positive) a\_refinement\_ratio. If a\_probdomain is an Empty ProblemDomain, then an empty problem domain is produced.

ProblemDomain& coarsen(const IntVect& a\_refinement\_ratio)

Modifies this ProblemDomain by coarsening it by the given refinement ratio in each direction. The empty ProblemDomain is not modified by this function.

#### friend

ProblemDomain refine (const ProblemDomain& a\_probdomain, const IntVect& a\_refinement\_ratio)

Returns a ProblemDomain that is the argument ProblemDomain coarsened by (the positive) a\_refinement\_ratio. If a\_probdomain is an Empty ProblemDomain, then an empty problem domain is produced.

```
friend
std::ostream& operator<< (std::ostream& os, const ProblemDomain& b)
Writes and ASCII representation to the ostream.

friend
std::istream& operator<< (std::istream& is, ProblemDomain& b)
read from istream.
```

# 2.4 Data on Unions of Rectangles

This section describes our tools for doing calculations over unions of rectangles. These tools may be used either in serial or in parallel though the design reflects our parallel programming model. In this section, we explain the tools we use to describe unions of rectangles and the data which lives over these regions.

# 2.4.1 Introduction

We wish to represent data defined on unions of rectangles. Such data can be mapped naturally onto distributed memory by assigning boxes to processors, with data defined on those boxes stored on the processor to which the box is assigned. This approach has been used quite successfully. Berger and Bokhari [5], Kohn and Baden [21], Rendleman, et. al. [28], and others have used this technique. Our API is derived from joint work with Baden to develop an abstract version of KeLP [13]. It is implemented using the following three sets of classes:

- BoxLayout, DisjointBoxLayout—classes that represent unions of rectangles and the mapping of those rectangles to processors.
- LayoutData, BoxLayoutData, LevelData— templated classes for distributing data over processors.
- LayoutIterator/LayoutIndex, DataIterator/DataIndex— classes for iterating over and indexing into the classes above.

# 2.4.2 Layouts

The classes BoxLayout and DisjointBoxLayout represent unions of rectangles and the mapping of the rectangles onto processors. BoxLayout represents an arbitrary union of valid boxes. DisjointBoxLayout is a BoxLayout and has the additional property that none of the boxes intersect. Both types of layout have two states: open and closed. During construction, a layout is open. In its open state, a user can add boxes and modify the mapping of boxes to processors. When a user is finished changing a BoxLayout to her satisfaction, she invokes the close() function. After closing, the BoxLayout cannot be accessed in a non-const manner. There is no way to reopen a closed BoxLayout. The closed property propagates through assignment and copy construction. Only closed layouts may be used to build the distributed data classes.

# 2.4.2.1 Class BoxLayout

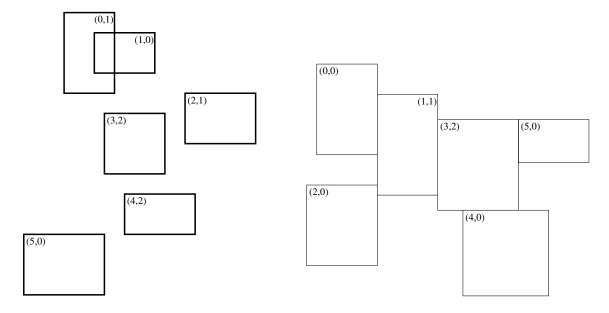


Figure 2.4: Left: Example of a BoxLayout. The first integer in the pair identifies the Box, and the second integer the processor ID. In this case we have the following assignments. Processor 0:  $B_1$ ,  $B_5$ . Processor 1:  $B_0$ ,  $B_2$ . Processor 2:  $B_3$ ,  $B_4$ . Note that  $B_0$  and  $B_1$  have a non-empty intersection. Right: Example of a disjoint BoxLayout. The first integer in the pair identifies the Box, and the second integer the processor ID. In this case we have the following assignments. Processor 0:  $B_0$ ,  $B_2$ ,  $B_4$ ,  $B_5$ . Processor 1:  $B_1$ . Processor 2:  $B_3$ . Note that a disjoint BoxLayout has empty intersections.

A BoxLayout is a collection of boxes. On parallel platforms, BoxLayout includes a mapping to processors. In both cases, the data holders LayoutData, BoxLayoutData, and LevelData define mappings from the Boxes in the BoxLayout to objects of the template type T. The important functions of BoxLayout are as follows:

### Construction

BoxLayout(const Vector<Box>& boxes, const Vector<int>& procIDs)
void define(const Vector<Box>& boxes, const Vector<int>& procIDs)
virtual void deepCopy(const BoxLayout& source)
DataIndex addBox(const Box& box, int iProc)
virtual void close()

The constructor and define functions construct a BoxLayout from a vector of Boxes and a vector of processor assignments. The input procIDs must all be in the range [0...numProcs()-1] where the function numProcs(), located in SPMD.H, returns the number of processors being used in the calculation. procIDs[i] is the processor number of the processor on which the data that maps to the box boxes[i] is stored. The processor assignment Vector must be the same length as the Vector<Box> argument. On exit, the BoxLayout will be closed. One can either null construct the BoxLayout and call the define function or construct and define at once. If the user is not using MPI, the procIDs argument is ignored. The new object created with deepCopy disassociates itself with original implementation safely. This object now is considered 'open' and can be non-const modified. There is no assurance that the order in which this BoxLayout is indexed corresponds to the indexing of source. addBox incrementally adds a box and its processor assignment to an open layout (if the layout has been closed, calling this function generates a run-time error) and returns a DataIndex object. The DataIndex object is valid both before and after close is called. It can be used later to access this box again, or access the data object (T) in a BoxLayoutData that is built from this BoxLayout object. close marks this BoxLayout as complete and unchangeable. It is here that the layout gets sorted. This must be called before any data containers are constructed using the layout.

### • Boolean functions.

bool operator==(const BoxLayout& rhs) const
bool check(const DataIndex& index) const
bool isClosed()

Equality for BoxLayout is a reference-counted pointer check. This returns true if these two objects share the same implementation. Important Warning: Two layouts can have the same boxes and same processor mapping and still return false if they were built separately. To force equality of two layouts, use the copy constructor. check returns true if the input DataIndex matches the layout. isClosed returns true if close() has been called.

### Accessors.

Box& operator[](const DataIndex& it)
DataIterator dataIterator() const
LayoutIterator layoutIterator() const

This allows access to an individual box through the iterator. One must be iterating through the correct layout (check must return true) in order for the accessor operator to work correctly. The member functions dataIterator, layoutIterator return the iterators associated with this layout.

# • Coarsening and Refinement Operations.

The functions coarsen, refine coarsens or refines each box in the layout by the input refinement ratio. Iterator objects that worked for the input will work for the output.

## 2.4.2.2 Class DisjointBoxLayout

DisjointBoxLayout is-a BoxLayout. The difference between them is that, for DisjointBoxLayout, closed also implies that the boxes in a DisjointBoxLayout have no non-trivial intersection with one another in index space. Any attempt to close a DisjointBoxLayout object with boxes which have non-trivial intersection will result in a run-time error. Coarsening may not preserve disjointedness, and applying the coarsen operator to a DisjointBoxLayout will generate also a run-time error if the new coarsened boxes aren't disjoint. Otherwise, all of the functions of BoxLayout carry over to DisjointBoxLayout.

If the problem domain is periodic, disjointness is tied to the periodicity of the domain – a box in the BoxLayout may intersect the periodic image of another box. To account for this, DisjointBoxLayout can also be defined with a ProblemDomain. In the default case, the domain is defined to be non-periodic in all directions. If the domain is periodic, then the periodicity of the domain is taken into account when checking for disjointness of the boxes in the BoxLayout.

The important extra functions of DisjointBoxLayout are as follows:

### Constructors

These functions are the same as the corresponding functions in BoxLayout, but with the addition of a ProblemDomain argument.

# Checking functions

bool checkPeriodic(const ProblemDomain& probDomain)

The checkPeriodic function returns true if the argument ProblemDomain is consistent with the ProblemDomain used to define the DisjointBoxLayout. Two ProblemDomains are consistent if they are periodic in the same directions, and they have same domain size in any periodic directions. In non-periodic directions, no consistency is required.

# 2.4.3 Templated Data Holders

LayoutData<T>, BoxLayoutData<T>, and LevelData<T> are templated data holders over a BoxLayout that hold one T at each box in the layout. Each class represents a different level of functionality. LayoutData<T> is a holder for creating local data corresponding to the part of the BoxLayout assigned to that processor. In particular, there is no support in Chombo for communicating LayoutData<T> information between processors. LevelData<T> implements an abstract form of a cell-centered level array, represented as a collection of rectangular "arrays" (i.e., objects of type T), each of which is defined on an element of  $\mathcal{R}(\Omega)$ . These arrays are distributed over processors using the rule encoded in the DisjointBoxLayout used to construct them. Finally, a BoxLayoutData<T> is a generalization of a LevelData<T>, in that the underlying BoxLayout is allowed to have overlapping Boxes. Thus one can copy from a LevelData<T> to a LevelData<T> or BoxLayoutData<T>, but not from a BoxLayoutData<T>, since the latter is not guaranteed to be single-valued on each cell.

# 2.4.3.1 Class LayoutData

LayoutData is a templated data holder for a collection of Box-oriented objects. A LayoutData can be built upon either a BoxLayout or a DisjointBoxLayout. The arrangement of the T objects is given by the underlying BoxLayout object. Each box in the BoxLayout will have a corresponding T object in the LayoutData object. The T objects contained within a LayoutData object should be accessed through a DataIterator. Non-local access to a LayoutData (access to a T that lives on another processor) is an error. Data in a LayoutData cannot be communicated to other processors. The class T must provide null construction.

The important parts of the LayoutData<T> API are as follows:

## • Construction.

```
LayoutData(const BoxLayout& dp);
void define(const BoxLayout& dp);
```

The constructor allocates a T object for every box in the BoxLayout dp using the T() (null) constructor. The function define performs the same task for a null-constructed LayoutData. The dp must be closed or a runtime error will occur.

### Accessors.

```
DataIterator dataIterator() const;
T& operator[](const DataIndex& index);
```

The input DataIndex for the indexing operator [] must match the BoxLayout which was used in construction of the LayoutData. It must also correspond to an element in the BoxLayout on myProc(). dataIterator returns an iterator which provides the DataIndex(es) which can be used to access the objects T which live at each box.

# 2.4.3.2 Class BoxLayoutData

**Requirements on the template class T**: BoxLayoutData<T> requires that T provides the following member functions, in addition to a null constructor for T:

# • Constructors.

```
T(const Box& box, int comps)
define(const Box& box, int comps)
```

Allocate all the memory for data given a region and the number of components. The data does not necessarily need to be initialized.

# • Copiers.

copy copies the data from source over the regionFrom to the regionTo in the destination. The two regions must be the same size, and must be valid in the source and destination, respectively. The component range specified by sourceInterval is copied to the component range specified by destInterval and the size of these two Intervals must be the same. linearIn/linearOut copy the object from/to the stream of bytes buf. This stream is assumed to be allocated by the calling program. size returns the size of the linearized object in bytes. preAllocateable returns:

- 1. if the size function is strictly a function of Box and Interval, and does not depend on the current state of the T object.
- 2. if size is symmetric, in that sender and receiver T object can size their message buffers, but a static object cannot.

3. if the object is truly dynamic. the message size is subject to unique object data.

A BoxLayoutData can be built upon either a BoxLayout or a DisjointBoxLayout. BoxLayoutData<T> is-a LayoutData<T> which means that it has all of the member functions of LayoutData<T>. The important extra functions of BoxLayoutData<T> are:

### • Constructors.

Defines the object from a layout and a number of components. Because of the semantics of inheritance, any DisjointBoxLayout can be used as an argument here instead of BoxLayout. The second define explicitly defines this BoxLayoutData from input. This includes copying the data values. The third define defines this BoxLayoutData to be on the same BoxLayout as the input da but only for the components defined by the Interval comps.

### • Accessors.

```
int nComp() const;
Interval interval() const;
```

nComp returns the number of components in the data holder. interval returns the component range of the data holder (0:nComp()-1).

#### 2.4.3.3 Class LevelData

A LevelData can be built only upon DisjointBoxLayouts. LevelData<T> has the same requirements on its T that BoxLayoutData<T> has. It also contains the important extra concepts of ghost values and data communication. Each box in the input layout is grown in each direction by the number of ghost cells in that direction. The data that lives on the input region (the part inside of the ghost cells) is considered "valid" data and the data on the ghost cells is considered "ghost" data. There are two data communication paradigms. One is the exchange function which copies data from the valid regions to the ghost regions where they intersect. The other function is copyTo which allows data communication between data holders. The source of a copyTo must be a LevelData<T>. The destination of copyTo may be either a LevelData<T> or a BoxLayoutData<T>. LevelData<T> is-a BoxLayoutData<T> (which is-a LayoutData<T>) which means that it has all of the member functions of BoxLayoutData<T> (and, by transitivity, all the member functions of LayoutData<T>). The important extra functions of LevelData<T> are as follows:

### • Constructors.

The construction functions work in the same way as the construction functions for BoxLayoutData. The main difference is that for each Box B in the BoxLayout, the object of type T is associated to the Box grown from B by ghost, i.e., T(grow(B,ghost),comps).

# • Copiers.

The first copyTo copies all of the data in the valid regions of this object to dest where the two BoxLayouts intersect. The length of the input and output intervals must be the same. The second version of copyTo copies to the LevelData dest filling the ghost cells of 'dest' with data from 'this' also (figure 2.5). The exchange function copies data from the valid regions to the ghost regions where they intersect (figure 2.5). If the DisjointBoxLayout used to define this LevelData is periodic in any direction, both copyTo and exchange will also fill cells from valid regions of the appropriate periodic images as necessary. ghostvect returns the IntVect defining the size of the ghost region.

# 2.4.3.4 Aliasing

For template classes that support an aliasing constructor, eg:

```
BaseFab<int> original(b, 4);
Interval subcomps(2, 3);
BaseFab<int> alias;
alias.define(subcomps, original);
```

then a user can alias an entire LevelData at once using the function

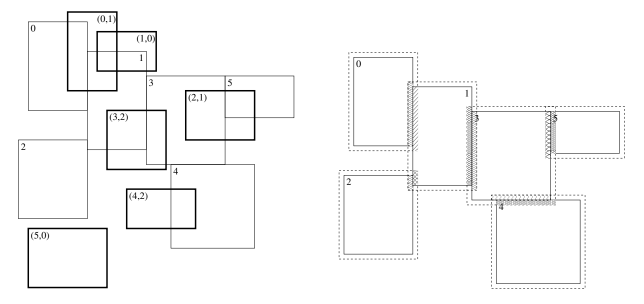


Figure 2.5: Left: copyTo example. This figure illustrates copying from a LevelData built on the DisjointBoxLayout in figure 2.4 to a BoxLayoutData built on top of the BoxLayout in figure 2.4. A single call to copy would perform the following data movements: Data from  $B_0$  copied to  $B_0'$ . Data from  $B_1$  copied to  $B_0'$ ,  $B_1'$ ,  $B_3'$ . Data from  $B_3$  copied to  $B_2'$ ,  $B_3'$ . Data from  $B_4$  copied to  $B_4'$ . Data from  $B_5$  copied to  $B_2'$ . No data is copied from  $B_2$  or to  $B_5'$ . Right: exchange example. This figure illustrates copying data from the valid regions of a LevelData built on top of the DisjointBoxLayout in figure 2.4 to ghost cell regions of the same LevelData. The dashed Boxes indicate which ghost cell regions will be filled by a single call to exchange.

See section 2.2.6.1 for semantics of aliasing.

# 2.4.4 Iterators

There are two iterators over multiple-box objects in Chombo, LayoutIterator and DataIterator. They each return objects (LayoutIndex, DataIndex) that can be used to index into layouts and data holders. A layout may be indexed into by either LayoutIndex or a DataIndex, while a data holder may only be indexed into using a DataIndex. In serial the iteration patterns of the two types of iterator are exactly the same. The iterators iterate through every box in the layout. In parallel, the LayoutIterator still iterates through every box in the layout but the DataIterator iterates through only boxes whose data resides upon the current processor.

**Principal Operations on** DataIterator, LayoutIterator. In the following, BL is a BoxLayout, DBL is a DisjointBoxLayout, and iter is either a DataIterator or a LayoutIterator.

- <u>Construction</u>. The iterators can be constructed with the object to be iterated over (iter(BL), iter(DBL)), or null-constructed and defined later (iter.define(BL), iter.define(DBL)).
- <a href="Iteration">Iter.begin()</a> sets iter to the beginning of the iteration sequence, ++iter advances iter to the next iterate, and iter.ok() checks to see if the current iterate is valid. iter() returns the current value of the iterate which is a DataIndex if iter() is a DataIterator or a LayoutIndex if iter() is a LayoutIterator.

We give examples of the use of LayoutIterator and DataIterator. In the first example, we iterate over all the Boxes in the layout to determine whether they cover the  ${\tt Box}\ B.$ 

```
Box B;
IntVectSet ivs(B);
BoxLayout bl;
LayoutIterator liter(bl);
for (liter.begin();liter.ok();++liter)
{
     ivs -= bl[liter];
}
if (ivs.isEmpty())
{
. . .
In the second example, we set the values of all the components in all the FArrayBoxes
in a BoxLayoutData<FArrayBox> to zero.
BoxLayout bl;
BoxLayoutData<FArrayBox> bld(bl,1);
DataIterator diter(bl);
for (diter.begin();diter.ok();++diter)
    bld[diter].setVal(0.0);
```

}

# Chapter 3

# **AMRTools**

# 3.1 Multilevel Operators.

In this section, we describe algorithmic and library support suitable for implementing extensions of second-order accurate discretizations of quasi-linear elliptic, parabolic, and hyperbolic PDE's in conservation form to AMR. Our approach will be to express the AMR discretizations in terms of the corresponding uniform grid discretizations at each level, using appropriate interpolation operators to provide ghost cell values for points in the stencil extending outside of the grids at that level. We will also define a conservative discretization of the divergence operator on multilevel data.

From a formal numerical analysis standpoint, a solution on an adaptive mesh hierarchy  $\{\Omega^l\}_{l=0}^{l_{max}}$  approximates the exact solution to the PDE only on those cells that are not covered by a grid at a finer level. We define the valid region of  $\Omega^l$ 

$$\Omega_{valid}^{l} = \Omega^{l} - \mathcal{C}_{n_{ref}^{l}}(\Omega^{l+1})$$

A composite array  $\varphi^{comp}$  is a collection of discrete values defined on the valid regions at each of the levels of refinement.

$$\varphi^{comp} = \{\varphi^{l,valid}\}_{l=0}^{l_{max}}$$
 ,  $\varphi^{l,valid}: \Omega^l_{valid} \to \mathbb{R}^m$ 

We can also define valid regions and composite arrays for other centerings.  $\Omega^{l,e^d}_{valid}=\Omega^{l,e^d}-\mathcal{C}_{n^l_{ref}}(\Omega^{l+1,e^d}).$  Thus  $\Omega^{l,e^d}_{valid}$  consists of d-faces that are not covered by the d-faces at the next finer level. A composite vector field  $\vec{F}^{comp}=\{\vec{F}^{l,valid}\}_{l=0}^{l_{max}}$  is defined as follows.

$$\vec{F}^{l,valid} = (F_0^{l,valid} \dots F_{\mathbf{D}-1}^{l,valid})$$
$$F_d^{l,valid} : \Omega_{valid}^{l,e^d} \to \mathbb{R}^m$$

Thus a composite vector field has values at level l on all of the faces that are not covered by faces at the next finer level.

We want to compute finite difference approximations to differential operators. For example, let L be a finite difference approximation to a linear differential operator  $\mathcal{L}$ . On a uniform grid, L typically takes the form

$$(L\varphi)_{i} = \sum_{|s| < S} c_{i,s} \varphi_{i+s} \tag{3.1}$$

Starting from this operator, we can extend L to be defined on an AMR grid hierarchy in the following fashion. For each  $\Omega_k^l \in \mathcal{R}(\Omega^l)$ 

$$\begin{array}{ll} \boldsymbol{\varphi}_{\pmb{i}}^{l,ext} & = & \boldsymbol{\varphi}_{\pmb{i}}^{l} \text{ on } \Omega_{valid}^{l} \\ & = & I(\boldsymbol{\varphi}^{comp}, \pmb{x}_0^l + \pmb{i}h^l) \text{ on } \mathcal{G}(\Omega_k^l, S) \cap \Gamma^l - \Omega_{valid}^l \end{array}$$

$$(L\varphi)_{\pmb{i}} = \sum_{|\pmb{s}| \leq S} c_{\pmb{i},\pmb{s}} \varphi_{\pmb{i}+\pmb{s}}^{l,ext} \text{ on } \Omega_k^l$$

Here  $I = I(\varphi^{comp}, x)$  is an interpolation operator that takes some combination of the valid composite data and constructs an interpolant at the point  $x \in \mathbb{R}^{\mathbf{D}}$ .

Let  $\psi$  be a smooth function on  $\mathbb{R}^{\mathbf{D}}$ , and define the level array

$$\psi^l = \psi({m x}_0^l + {m i} h^l)$$
 on  $\Gamma^l$ 

and composite array

$$\psi^{comp} = \{\psi^{l,valid}\}_{l=0}^{l_{max}}$$
 
$$\psi^{l,valid} = \psi^{l} \text{ on } \Omega^{l}_{valid}$$

Then the truncation error of the operator L can be computed as follows. For  ${\pmb i} \in \Omega^l_{valid}$ 

$$\begin{aligned} \tau_{i} &\equiv L^{comp}(\psi^{comp})_{i} - \mathcal{L}(\psi)(\boldsymbol{x}_{0}^{l} + \boldsymbol{i}h^{l}) \\ &= \sum_{|\boldsymbol{s}| \leq \mathcal{S}} c_{\boldsymbol{i},\boldsymbol{s}} \psi_{\boldsymbol{i}+\boldsymbol{s}} - \mathcal{L}(\psi)(\boldsymbol{x}_{0}^{l} + \boldsymbol{i}h^{l}) \\ &- \sum_{|\boldsymbol{s}| \leq \mathcal{S} \atop \boldsymbol{i}+\boldsymbol{s} \notin \Omega_{volid}^{l}} c_{\boldsymbol{i},\boldsymbol{s}}(\psi_{\boldsymbol{i}+\boldsymbol{s}} - I(\psi^{comp}, \boldsymbol{x}_{0}^{l} + (\boldsymbol{i}+\boldsymbol{s})h^{l})) \end{aligned}$$

The first sum is the truncation error on a uniform grid, while the second sum gives the effect of replacing the uniform grid values of the smooth function  $\psi$  by those obtained by interpolation.

Unfortunately, this process, when used by itself, becomes unwieldy for any but the simplest finite difference approximations. Typically, in order to obtain  $\tau = O(h^q)$  it is necessary to compute  $I(\varphi^{comp}, \boldsymbol{x}_0^l + \boldsymbol{i}h)$  to an accuracy of  $O(h^{p+q})$ , where p is order of the highest derivative of the operator, due to the contributions of the second summand

 $(\max_{|s| \leq S} (|c_{i,s}|) = O(h^{-p}))$ . To obtain interpolants of such accuracy, we must either use general polynomial interpolants using data located on multiple levels of refinement, or impose minimum distance requirements between grids at different levels of refinement. The alternative is to accept a larger truncation error near the boundary between levels of refinement. In the AMR algorithms that motivate the design of Chombo, we use a combination of all three techniques. This approach is motivated by the following mathematical and algorithmic considerations.

- Our target applications involve solving first and second order quasi-linear systems of PDE's of classical type, i.e., elliptic, parabolic, and hyperbolic.
- Our underlying uniform-grid discretizations are based on second-order accurate methods, mainly in discrete conservation form.

The latter property is one that we would like to preserve in the AMR versions of these algorithms. However, the requirement for discrete conservation form leads to a loss of accuracy at coarse-fine boundaries. Finite difference methods rely on the cancellation of truncation error terms in the differenced quantities in order to obtain a given accuracy on a uniform grid. This is the mechanism, for example, by which the second divided difference approximates the second derivative to  $O(h^2)$ , even though it is a divided difference of quantities that are themselves accurate only to  $O(h^2)$ . This mechanism fails at the interface between different levels of refinement. If one is to approximate the divergence operator with a divided difference of single-valued fluxes, it is not possible to compute the flux so that the truncation error cancels that of the fluxes on both the adjacent coarse and fine faces.

Fortunately, our choice of target applications makes this local loss of accuracy acceptable. For elliptic and parabolic problems, a truncation error of  $O(h^{p-1})$  on a set of codimension one induces a solution error of  $O(h^p)$ , due to a discrete form of elliptic regularity. In hyperbolic problems, a truncation error of  $O(h^{p-1})$  on a set of codimension one induces a total error of  $O(h^p)$  in the  $L_1$  norm (and in  $L_{\infty}$  as well if the boundary is non-characteristic).

In the following, we give the details of the algorithms for interpolating between levels that arise in this approach. They include averaging and interpolation methods for transferring information between levels; specialized operators for interpolating boundary conditions at boundaries between levels; and a conservative multilevel discretization of the divergence operator. For all of these cases, we will describe the algorithms for the case of data defined on two successive levels  $\Omega^f, \Omega^c_{valid}$ . The resulting operators can all be extended to the full AMR hierarchy by applying them to a pair of levels at a time, provided that appropriate nesting conditions are met. For the most part, only proper nesting is required. When that is not the case, we will explicitly state the nesting conditions required on grids at successive levels.

# 3.1.1 Interlevel Transfer Operators

# 3.1.1.1 Conservative Averaging.

This operator is used to average from finer levels on to coarser levels, or for constructing averaged residuals in multigrid iterations.

$$Average(\varphi, n_{ref})_{i^c} = \frac{1}{(n_{ref})^d} \sum_{i \in \mathcal{C}_{n_{ref}}^{-1}(i^c)} \varphi_i$$

This process produces values on the coarse grid that are an  $O(h^2)$  estimate of the solution on the fine grid.

## 3.1.1.2 Piecewise Constant Interpolation.

This operator is primarily used in multigrid iteration to interpolate the correction from the coarser level to the next finer level.

$$I_{pwc}(\varphi)_{i^f} = \varphi_i$$

where  $i = C_{n_{ref}}(i^f)$ . This method has an interpolation error of O(h).

# 3.1.1.3 Piecewise Linear Interpolation.

This method is primarily used to initialize fine grid data in newly-refined regions after regridding. Given a level array  $\varphi^c$  on  $\Omega^c$ , we want to compute  $I_{pwl}(\varphi)$  defined on an  $\Omega^f$  properly nested in  $\Omega^c$ .

$$I_{pwl}(\varphi)_{i^f} = \varphi_i + \sum_{d=0}^{\mathbf{D}-1} \left( \frac{(i_d^f + \frac{1}{2})}{n_{ref}} - (i_d + \frac{1}{2}) \right) (\delta^d \varphi)_i$$

$$(\delta^d arphi)_{m{i}} = egin{cases} \eta_{m{i}} (\delta^d_c arphi)_{m{i}} & ext{if both } m{i} \pm m{e}^d \in \Gamma^c \ arphi_{m{i} + m{e}^d} - arphi_{m{i}} & ext{if } m{i} - m{e}^d 
ot\in \Gamma^c \ arphi_{m{i} - m{\varphi}_{m{i} - m{e}^d} & ext{if } m{i} + m{e}^d 
ot\in \Gamma^c \end{cases}$$

$$\eta_{i} = \chi(min(\varphi_{i}^{max} - \varphi_{i}, \varphi_{i} - \varphi_{i}^{min}), \sum_{d=0}^{\mathbf{D}-1} |\delta_{c}^{d}\varphi|_{i})$$

$$(\delta_c^d \varphi)_{\boldsymbol{i}} = \begin{cases} \frac{1}{2} (\varphi_{\boldsymbol{i} + \boldsymbol{e}^d} - \varphi_{\boldsymbol{i} - \boldsymbol{e}^d}) & \text{if both } \boldsymbol{i} \pm \boldsymbol{e}^d \in \Gamma^c \\ \varphi_{\boldsymbol{i} + \boldsymbol{e}^d} - \varphi_{\boldsymbol{i}} & \text{if } \boldsymbol{i} - \boldsymbol{e}^d \not\in \Gamma^c \\ \varphi_{\boldsymbol{i}} - \varphi_{\boldsymbol{i} - \boldsymbol{e}^d} & \text{if } \boldsymbol{i} + \boldsymbol{e}^d \not\in \Gamma^c \end{cases}$$

$$\chi(a,b) = \begin{cases} \frac{a}{b} & \text{if } a < b \\ 1 & \text{otherwise} \end{cases}$$

$$\varphi_{i}^{max} = \max_{|s| \le 1} (\varphi_{i+s}) , \ \varphi_{i}^{min} = \min_{|s| \le 1} (\varphi_{i+s})$$

At cells adjacent to the boundary of the computational domain  $\Gamma^c$ , the maximum and minimum are taken over the points i+s that are contained in the computational domain. Also note, the arguments to  $\chi$  are always non-negative.

We use the limiter  $\eta$  to keep the interpolated values from exceeding local estimates of the maximum and minimum values of the solution on the coarser grid. As long as  $\eta=1$ , i.e., the limiter does not reduce the values of the slopes, the error in the interpolated values is  $O(h^2)$ .

# 3.1.2 Coarse-Fine Boundary Interpolation

When evaluating operators on refined levels, it's often convenient to employ a ring of p ghost cells around each grid box. The number of ghost cells p is normally chosen based on the specific operators in use and allows the fine-grid stencils to be completed near coarse-fine interfaces. We then fill the ghost cells with appropriately interpolated values, which may involve coarse data, fine data, or a combination of the two.

# 3.1.2.1 Piecewise Linear Interpolation

Assume there are two levels of grid  $\Omega^c$ ,  $\Omega^f$ , and data defined on the fine grid and on the valid region of the coarse grid:

$$\varphi^f: \Omega^f \to \mathbb{R}$$
$$\varphi^{c,valid}: \Omega^c_{valid} \to \mathbb{R}$$

We want to compute an extension  $\tilde{\varphi}^f$  of  $\varphi^f$  on  $\tilde{\Omega}^f=\mathcal{G}(\Omega^f,p)\cap\Gamma^f,p>0$ , which is accurate to  $O(h^2)$ , assuming only that  $\mathcal{C}_r(\tilde{\Omega}^f)\cap\Gamma^c\subset\Omega^c$ . There must be enough points on the coarse level to interpolate out to a distance of p fine cells from  $\Omega^f$ . One way to achieve this goal is by choosing an appropriate blocking factor, i.e., we assume that  $\Omega^f$  is  $n_{ref}(\lfloor \frac{p}{n_{ref}} \rfloor + min(1, p \ mod \ n_{ref}))$  - blocked. Combined with proper nesting, this ensures that there are sufficient cells in  $\Omega^c$  to perform the interpolation.

We perform this calculation in these steps

- (i) Extend  $\varphi^{c,valid}$  to  $\varphi^c$ , defined on all of  $\Omega^c: \varphi^c = Average(\varphi^f, n_{ref})$  on  $\mathcal{C}_{n_{ref}}(\Omega^f)$
- (ii) For each  $m{i}^f \in \tilde{\Omega}^f \Omega^f$ , compute a piecewise linear interpolant. For  $m{i} = \mathcal{C}_{n_{ref}}(m{i}^f)$ ,

$$\tilde{\varphi}_{i^f}^f = \varphi_i + \sum_{d=0}^{\mathbf{D}-1} \left( \frac{(i_d^f + \frac{1}{2})}{n_{ref}} - (i_d + \frac{1}{2}) \right) (\delta^d \varphi)_i \equiv I_{pwl}^B (\varphi^f, \varphi^{c,valid})_{i^f}$$

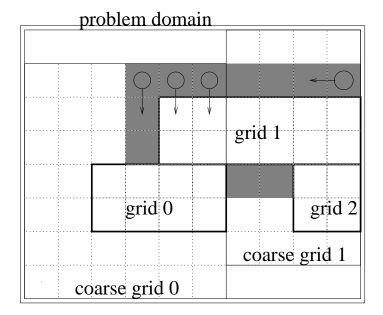


Figure 3.1: Interpolation on the coarse grid. One-sided differences are used at cells marked with circles.

Unlike in the interlevel transfer operator  $I_{pwl}$ , we use a minimal stencil for  $(\delta^d \varphi)_i$  (Figure 3.1).

$$(\delta^d arphi)_{m{i}} = egin{cases} \delta_{vL}(arphi_{m{i}-m{e}^d}, arphi_{m{i}}, arphi_{m{i}+m{e}^d}) & ext{if both } m{i} \pm m{e}^d \in \Omega^c \ arphi_{m{i}+m{e}^d} - arphi_{m{i}} & ext{if } m{i} - m{e}^d 
ot\in \Omega^c \ arphi_{m{i}} - arphi_{m{i}-m{e}^d} & ext{if } m{i} + m{e}^d 
ot\in \Omega^c \end{cases}$$

$$\delta_{vL}(a_l, a_c, a_r) = \begin{cases} min(\frac{1}{2}|a_l - a_r|, |a_l - a_c|, |a_r - a_c|) sign(a_r - a_l) & \text{if } (a_l - a_c)(a_c - a_r) > 0 \\ 0 & \text{otherwise} \end{cases}$$

(iii) 
$$\tilde{\varphi}^f = \varphi^f$$
 on  $\Omega^f$ 

The truncation error of this interpolation operator is  $O(h^2)$ , i.e., if  $\psi = \psi(\boldsymbol{x})$  is a smooth function, and

$$\begin{split} \psi_{\pmb{i}}^f &= \psi(\pmb{x}_0^f + \pmb{i}h^f) \text{ on } \Omega^f \\ \psi_{\pmb{i}}^{c,valid} &= \psi(\pmb{x}_0^c + \pmb{i}h^c) \text{ on } \Omega^{c,valid} \end{split}$$

then

$$\tilde{\psi}_{if}^f = \psi(\boldsymbol{x}_0^f + \boldsymbol{i}h^f) + O(h^2) \text{ for } \boldsymbol{i} \in \tilde{\Omega}^f - \Omega^f$$

where  $\tilde{\psi}^f$  is the extension of  $(\psi^f,\psi^c)$  computed using the process outlined above. The key point is that, as long as the extension of  $\psi^c$  to  $\mathcal{C}_{n_{ref}}(\Omega^f)$  is accurate to  $O(h^2)$ , the undivided difference formula approximates  $h^c \frac{\partial \psi}{\partial x^d}$  to  $O(h^2)$ , and differs from the Taylor expansion of  $\psi$  around  $(\boldsymbol{x}^c + \boldsymbol{i}h^c)$  by  $O(h^2)$ .

# 3.1.2.2 Quadratic Coarse-Fine Boundary Interpolation

This interpolation scheme is motivated by the requirements of constructing consistent discretizations of second-order operators. Given  $\varphi^f$ ,  $\varphi^{c,valid}$ , we want to compute a level vector field  $\vec{G}^f = (G_0^f, \ldots G_{\mathbf{D}-1}^f)$  that approximates the gradient to sufficient accuracy so that, when we take its divergence, we obtain at least an O(h) approximation to the Laplacian. For each  $\Omega_k^f \in \mathcal{R}(\Omega^f)$ , we construct an extension  $\tilde{\varphi}$  of  $\varphi^f$ .

$$\tilde{\varphi}: \tilde{\Omega}_k^f \to \mathbb{R}^m$$

$$\tilde{\Omega}_k^f = (\bigcup_{\substack{\pm = +, -\\ d = 0}}^{\mathbf{D}-1} \Omega_k^f \pm \mathbf{e}^d) \cap \Gamma^f$$

Then, for each  $i+\frac{1}{2}e^d$  such that both  $i,i+e^d\in\tilde{\Omega}_k^f$  we can compute a centered difference approximation to the gradient on a staggered grid

$$G_{d,oldsymbol{i}+rac{1}{2}oldsymbol{e}^d}^f = rac{1}{h^f} ( ilde{arphi}_{oldsymbol{i}+oldsymbol{e}^d} - ilde{arphi}_{oldsymbol{i}})$$

For this estimate of the gradient to be accurate to  $O(h^2)$ , it is necessary to compute an  $O(h^3)$  extension of  $\varphi^f$ . On  $\tilde{\Omega}_k^f \cap \Omega^f$ , the values for  $\tilde{\varphi}$  will be given by  $\tilde{\varphi}_i = \varphi_i^f$ . The values for the remaining points in  $\tilde{\Omega}_k^f - \Omega^f$  will be obtained by interpolating data from  $\varphi^f$  and  $\varphi^c$ .

To perform this interpolation, we first observe that, given  $i \in \tilde{\Omega}_k^f - \Omega^f$ , there is a unique choice of  $\pm$  and d, such that  $i \mp e^d \in \Omega_k^f$ . Having specified that choice, the interpolant is constructed in two steps (figure 3.2).

(i) Interpolation in the direction orthogonal to  $oldsymbol{e}^d$ . We compute

$$oldsymbol{x} = rac{oldsymbol{i} + rac{1}{2}oldsymbol{u}}{n_{ref}} - (oldsymbol{i}^c + rac{1}{2}oldsymbol{u})$$

where  $i^c = C_{n_{ref}}(i)$ . The real-valued vector x is the displacement of the cell center i on the fine grid from the cell center at  $i^c$  on the coarse grid, scaled by  $h^c$ .

$$\hat{\varphi}_{i} = \varphi_{i^{c}}^{c} + \sum_{d' \neq d} \left[ \left( x_{d'} (D^{1,d'} \varphi^{c})_{i^{c}} + \frac{1}{2} (x_{d'})^{2} (D^{2,d'} \varphi^{c})_{i^{c}} \right) + \sum_{d'' \neq d, d'' \neq d'} x_{d'} x_{d''} (D^{d'd''} \varphi^{c})_{i^{c}} \right]$$

The second sum has only one term if D = 3, and no terms if D = 2.

(ii) Interpolation in the normal direction.

$$\tilde{\varphi}_{i} = I_q^B(\varphi^f, \varphi^{c,valid}) \equiv 4a + 2b + c , \ \tilde{x_d} = x_d - \frac{1}{2}(n_{ref} + 3)$$

where a, b, c are computed to interpolate between the collinear data

$$((\mathbf{i} \pm \frac{1}{2}(n_{ref}^{l} - 1)\mathbf{e}^{d})h, \hat{\varphi}_{\mathbf{i}}),$$

$$((\mathbf{i} \mp \mathbf{e}^{d})h, \varphi_{\mathbf{i} \mp \mathbf{e}_{d}}^{l}),$$

$$((\mathbf{i} \mp 2\mathbf{e}^{d})h, \varphi_{\mathbf{i} \mp 2\mathbf{e}_{d}}^{l})$$

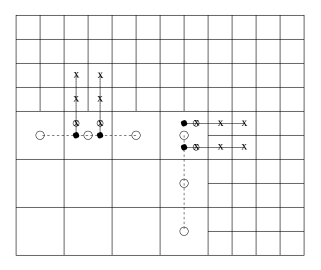


Figure 3.2: Interpolation at a coarse-fine interface. Left stencil is the usual stencil. Right stencil is the modified interpolation stencil; since the upper coarse cell is covered by a fine grid, use shifted coarse grid stencil (open circles) to get intermediate values (solid circles), then perform final interpolation as before to get "ghost cell" values (circled X's). Note that to perform interpolation for the horizontal coarse-fine interface, we need to shift the coarse stencil left.

In (i), the quantities  $D^{1,d'}\varphi^c, D^{2,d'}\varphi^c$  and  $D^{d'd''}\varphi^c$  are difference approximations to  $\frac{\partial}{\partial x_{d'}}, \frac{\partial^2}{\partial x_{d'}^2}$ , and  $\frac{\partial^2}{\partial x_{d'}\partial x_{d''}}$ , respectively.  $D^{1,d}\varphi$  must be accurate to  $O(h^2)$ , while the other two quantities need only be O(h). The strategy for computing these quantities is to use only values in  $\Omega^c_{valid}$  to compute these difference approximations. For the case of  $D^{1,d'}\varphi, D^{2,d'}\varphi$ , we use 3-point stencils, centered if possible, or shifted as required to consist of points on  $\Omega^c_{valid}$ .

$$(D^{1,d'}\varphi)_{\boldsymbol{i}} = \begin{cases} \frac{1}{2}(\varphi^c_{\boldsymbol{i}+\boldsymbol{e}^{d'}} - \varphi^c_{\boldsymbol{i}-\boldsymbol{e}^{d'}}) & \text{if both } \boldsymbol{i} \pm \boldsymbol{e}^{d'} \in \Omega^c_{valid} \\ \pm \frac{3}{2}(\varphi^c_{\boldsymbol{i}\pm\boldsymbol{e}^{d'}} - \varphi^c_{\boldsymbol{i}}) \mp \frac{1}{2}(\varphi^c_{\boldsymbol{i}\pm2\boldsymbol{e}^{d'}} - \varphi^c_{\boldsymbol{i}\pm\boldsymbol{e}^{d'}}) & \text{if } \boldsymbol{i} \pm \boldsymbol{e}^{d'} \in \Omega^c_{valid}, \ \boldsymbol{i} \mp \boldsymbol{e}^{d'} \not\in \Omega^c_{valid} \\ 0 & \text{otherwise} \end{cases}$$

$$(D^{2,d'}\varphi)_{\boldsymbol{i}} = \begin{cases} \varphi^c_{\boldsymbol{i}+\boldsymbol{e}^{d'}} - 2\varphi^c_{\boldsymbol{i}} + \varphi^c_{\boldsymbol{i}-\boldsymbol{e}^{d'}} & \text{if both } \boldsymbol{i} \pm \boldsymbol{e}^{d'} \in \Omega^c_{valid} \\ \varphi^c_{\boldsymbol{i}} - 2\varphi^c_{\boldsymbol{i}\pm\boldsymbol{e}^{d'}} + \varphi^c_{\boldsymbol{i}\pm 2\boldsymbol{e}^{d'}} & \text{if } \boldsymbol{i} \pm \boldsymbol{e}^{d'} \in \Omega^c_{valid}, \ \boldsymbol{i} \mp \boldsymbol{e}^{d'} \not\in \Omega^c_{valid} \\ 0 & \text{otherwise} \end{cases}$$

In the case of  $D^{d'd''}\varphi^c$ , we use an average of all of the four-point difference approximations  $\frac{\partial^2}{\partial x_{d'}\partial x_{d''}}$  centered at d',d'' corners adjacent to i such that all four points in the stencil are in  $\Omega^c_{valid}$  (Figure 3.3)

$$(D_{corner}^{d'd''}\varphi^c)_{\boldsymbol{i}+\frac{1}{2}\boldsymbol{e}^{d'}+\frac{1}{2}\boldsymbol{e}^{d''}} = \begin{cases} \frac{1}{h^2}(\varphi_{\boldsymbol{i}+\boldsymbol{e}^{d'}+\boldsymbol{e}^{d''}}+\varphi_{\boldsymbol{i}}-\varphi_{\boldsymbol{i}+\boldsymbol{e}^{d'}}-\varphi_{\boldsymbol{i}+\boldsymbol{e}^{d''}}) & \text{if } [\boldsymbol{i},\boldsymbol{i}+\boldsymbol{e}^{d'}+\boldsymbol{e}^{d''}] \subset \Omega_{valid}^c \\ 0 & \text{otherwise} \end{cases}$$

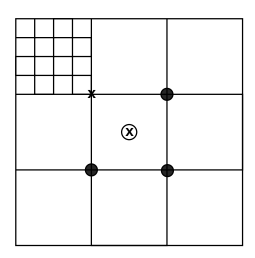


Figure 3.3: Mixed-derivative approximation illustration. The upper-left corner is covered by a finer level so the mixed derivative in the upper left (the uncircled x) has a stencil which extends into the finer level. We therefore average the mixed derivatives centered on the other corners (the filled circles) to approximate the mixed derivatives for coarse-fine interpolation in three dimensions.

$$(D^{2,d'd''}\varphi^c)_{\boldsymbol{i}} = \begin{cases} \frac{1}{N_{valid}} \sum_{s'=\pm 1} \sum_{s''=\pm 1} (D^{d'd''}\varphi^c)_{\boldsymbol{i}+\frac{1}{2}s'\boldsymbol{e}^{d'}+\frac{1}{2}s''\boldsymbol{e}^{d''}} & \text{if } N_{valid} > 0\\ 0 & \text{otherwise} \end{cases}$$

where  $N_{valid}$  is the number of nonzero summands. To compute (ii), we need to compute the interpolation coefficients a b, and c.

$$a = \frac{\hat{\varphi} - (n_{ref} \cdot |x_d| + 2)\varphi_{i \neq e^d} + (n_{ref} \cdot |x_d| + 1)\varphi_{i \neq 2e^d}}{(n_{ref} \cdot |x_d| + 2)(n_{ref} \cdot |x_d| + 1)}$$
$$b = \varphi_{i \neq e^d} - \varphi_{i \neq 2e^d} - a$$
$$c = \varphi_{i \neq 2e^d}$$

# 3.1.2.3 Level Divergence, Composite Divergence, and Refluxing

Let  $\vec{F}$  be a level vector field on  $\Omega$ . We define a discretized divergence operator as follows.

$$(D\vec{F})_{i} = \frac{1}{h} \sum_{d=0}^{D-1} (F_{d,i+\frac{1}{2}e^{d}} - F_{d,i-\frac{1}{2}e^{d}}), i \in \Omega$$
(3.2)

Let  $\vec{F}^{comp} = \{\vec{F}^f, \vec{F}^{c,valid}\}$  be a two-level composite vector field. We want to define a composite divergence  $D^{comp}(\vec{F}^f, \vec{F}^{c,valid})_{\pmb{i}}$  for  $\pmb{i} \in \Omega^c_{valid}$ . To do this, we construct an extension of  $\vec{F}^{c,valid}$  to the edges adjacent to  $\Omega^c_{valid}$  that are covered by fine level faces. On the valid coarse-level d-faces,  $\hat{F}_{d,\pmb{i}+\frac{1}{2}e^d} = F^{c,valid}_{d,\pmb{i}+\frac{1}{2}e^d}$ . On the faces adjacent to cells in  $\Omega^c_{valid}$ , but not in  $\Omega^{l,e^d}_{valid}$ , we set  $\hat{F}_d$  to be  $< F^f_d>$ , the average of  $F^f_d$  onto the next

coarser level.

$$< F_d^f>_{i+\frac{1}{2}e^d} = \frac{1}{(n_{ref})^{\mathbf{D}-1}} \sum_{i^f+\frac{1}{2}e^d \in \mathcal{F}^d} F_{d,i^f+\frac{1}{2}e^d}^f$$

$$m{i} + rac{1}{2}m{e}^d \in \zeta_{d,+}^f \cup \zeta_{d,-}^f$$

Here the sum is over the set of all fine level d-faces that are covered by  $[i+\frac{1}{2}e^d]$ , which is given as a rectangle in  $\Gamma^{f,e^d}$ .

$$\mathcal{F}^d = [m{i} \; n_{ref} + rac{1}{2}m{e}^d, (m{i} + (m{u} - m{e}^d))n_{ref} + rac{1}{2}m{e}^d]$$

 $\zeta_{d,\pm}^f$  consists of all the d-faces in  $\Omega^c$  on the boundary of  $\Omega^{l+1}$ , with valid cells on the low  $(\pm=-)$  or high  $(\pm=+)$  side.

$$\zeta_{d,\pm}^f = \{ m{i} \pm rac{1}{2} m{e}^d : m{i} \pm m{e}^d \in \Omega_{valid}^c, m{i} \in \mathcal{C}_{n_{ref}}(\Omega^f) \}$$

For both performance reasons and algorithmic reasons, it is useful to express  $D^{comp}$  as a succession of applications of the level divergence operator D applied to extensions of  $\vec{F}^{l,valid}$  to the entire level, followed by a step that corrects the cells in  $\Omega^c_{valid}$  that are adjacent to  $\Omega^f$ . We define a flux register  $\delta \vec{F}^f$  associated with the fine level

$$\delta \vec{F}^f = (\delta F_0^f, ..., \delta F_{\mathbf{D}-1}^f)$$
$$\delta F_d^f : \zeta_{d,+}^f \cup \zeta_{d,-}^f \to \mathbb{R}^m$$

Let  $\vec{F}^c$  be any coarse level vector field that extends  $\vec{F}^{c,valid}$ , i.e.

$$F_d^c = F_d^{c,valid}$$
 on  $\Omega_{valid}^{c,e^d}$ 

Then for  $i \in \Omega^c_{valid}$ ,

$$D^{comp}(\vec{F}^f, \vec{F}^{c,valid})_i = (D\vec{F}^c)_i + D_R(\delta \vec{F}^c)_i$$
(3.3)

Here  $\delta \vec{F}^c$  is a flux register, set to be

$$\delta F_d^f = \langle F_d^f \rangle - F_d^c \text{ on } \zeta_{d,+}^c \cup \zeta_{d,-}^c$$

 $D_R$  is the reflux divergence operator, given by the following for valid coarse level cells adjacent to  $\Omega^f$ .

$$D_R(\delta \vec{F}^f)_{\boldsymbol{i}} = \frac{1}{h^c} \sum_{d=0}^{\mathbf{D}-1} \sum_{\substack{\pm = +, -: \\ \boldsymbol{i} \pm \frac{1}{2} \boldsymbol{e}^d \in \zeta_{d, \mp}^f}} \pm \delta F_{d, \boldsymbol{i} \pm \frac{1}{2} \boldsymbol{e}^d}^f$$

For the remaining cells in  $\Omega^c_{valid}$ ,  $D_R(\delta \vec{F}^f)$  is defined to be identically zero.

Let  $\vec{H} = (H_0 \dots H_{\mathbf{D}-1})$ ,  $H_d : \mathbb{R}^{\mathbf{D}} \to \mathbb{R}^m$  be a smooth vector field, and define discrete level and composite vector fields by evaluating  $\vec{H}$  on the grid.

$$\vec{H}^f = (H_0^f \dots H_{\mathbf{D}-1}^f)$$
,  $H_{d, \mathbf{i} + \frac{1}{2} \mathbf{e}^d}^f = H_d(\mathbf{x}_0^f + (\mathbf{i} + \frac{1}{2} \mathbf{e}^d) h^f)$   
 $\vec{H}^c = (H_0^c \dots H_{\mathbf{D}-1}^c)$ ,  $H_{d, \mathbf{i} + \frac{1}{2} \mathbf{e}^d}^c = H_d(\mathbf{x}_0^c + (\mathbf{i} + \frac{1}{2} \mathbf{e}^d) h^c)$ 

We can then compute the truncation error of the composite divergence using (3.3).

$$D^{comp}(\vec{H}^f, \vec{H}^{c,valid}) = D(\vec{H}^c) + D_R(\delta \vec{H})$$
$$= \nabla \cdot \vec{H} + O(h^2) + D_R(\delta \vec{H})$$

Here  $H^{c,valid}_d$  is given by restricting  $H^c_d$  to  $\Omega^{c,e^d}_{valid}$ , and we make use of the observation that the centered difference approximation to the divergence given by (3.2) is second order accurate. Away from the cells adjacent to  $\Omega^f$ , the contribution from the reflux divergence is zero, and the truncation error is  $O(h^2)$ . To estimate the truncation error at cells adjacent to the coarse - fine interface, we note that

$$\delta H_d^f = \langle H_d^f \rangle - H_d^c = O((h^c)^2)$$

So that  $D_R(\delta \vec{H}) = O(h^c)$ , i.e., we lose one order of accuracy due to the correction to the divergence that maintains conservation form.

# Laplacian

Using the operators described above, we can now define a discretization of the Laplacian on an adaptive mesh hierarchy. Let  $\varphi^{comp}$  a composite array defined on an AMR grid hierarchy satisfying proper nesting. The Laplacian is defined as the divergence of the gradient:

$$(L^{comp}\varphi^{comp})_{i} \equiv D^{comp}(\vec{G}^{l+1,valid}, \vec{G}^{l,valid})_{i}$$
,  $i \in \Omega^{l}_{valid}$  (3.4)

where  $\vec{G}^{l,valid} = \vec{G}(\varphi^{l,valid}, \varphi^{l-1,valid})$  is computed using the algorithm in section 3.1.2.2. It is assumed here that the discrete gradients can be computed using the boundary conditions for the faces that lay on the boundary of the domain. It is not difficult to check that, if the grids are properly nested, that the stencil of  $(L^{comp}\varphi^{comp})_i$  is contained in the valid regions of the meshes at levels  $l,\ l\pm 1$ . Away from boundaries between levels, this discretization reduces to the standard  $2\mathbf{D}+1$  point discretization of the Laplacian. On grid interiors,  $L^{comp}$  has a truncation error of  $O(h^2)$  due to cancellation of error terms in the centered-difference stencil. At coarse-fine interfaces, this drops to O(h) due to division of the  $O(h^3)$  interpolant by  $h^2$  and the loss of centered-difference cancellations. However, if the discrete equation  $L^{comp}\varphi = \rho$  is solved using these operators, the resulting solution  $\varphi$  is second-order accurate, because this loss of accuracy occurs on a set of codimension one [20]. The dependencies of the Laplacian operators may again be expressed explicitly: if  $L^{comp,l}(\varphi^{comp})$  is  $L^{comp}(\varphi^{comp})$  restricted to  $\Omega^l_{valid}$ , then  $L^{comp,l}(\varphi^{comp}) = L^{comp,l}(\varphi^{l,valid}, \varphi^{l+1,valid}, \varphi^{l-1,valid})$ .

# 3.2 C++ Classes for Two-Level Operators

In this section, we document the user interfaces for a set of C++ classes that implement the operators described above. Typically, the interface has two parts. The constructor and define function construct the persistent data, such as interpolation coefficients and the IntVectSets defining the irregular regions where the operator must be applied. This can either be done by calling a defining constructor, or by calling a member function define with the same arguments on an object that has been constructed with a default constructor. Note that for classes where problem domain information is required for construction, there are generally two sets of constructors and define functions — one with a Box to represent the domain, the second with a ProblemDomain; if the functions with a Box are used, a non-periodic domain is assumed. The second part of the interface consists of the functions that actually apply the operator to the data. Once the operator has been defined, the user can apply it multiple times to different data sets. The operator must be redefined only when the grids change.

# **3.2.1** Class CoarseAverage

This class sets data on a level equal to an average of the data on a finer level of refinement wherever the finer level covers the coarse level, using the averaging operator in section 3.1.1. Averaging can be either arithmetic (the usual case) or harmonic.

#### • void

# **Arguments:**

- a\_fine\_domain (not modified): the fine-level grids (valid region).
- a\_crse\_domain (not modified): the coarse-level grids (valid region).
- a\_numcomps (not modified): the number of components of coarse and fine data sets.
- a\_ref\_ratio (not modified): the refinement ratio  $n_{ref}$ .
- a\_ghost\_vect (not modified): radius of ghost cells around coarse grids to be filled (normally none)

#### • void

Replaces coarse data with the arithmetic average of fine data, in the valid fine domain. **Arguments:** 

- a\_coarse\_data (modified): coarse data set, destination of averaging.
- a\_fine\_data (not modified): fine data set, source of averaging.
- void

Replaces coarse data with the harmonic average of fine data, in the valid fine domain. **Arguments:** 

- a\_coarse\_data (modified): coarse data set, destination of averaging.
- a\_fine\_data (not modified): fine data set, source of averaging.

# 3.2.2 Class CoarseAverageFace

Similar to CoarseAverage, but averaging face-centered data instead of cell-centered data. Data on coarse-level faces is computed as the average of the overlying fine-level faces. Both arithmetic and harmonic averaging are supported.

- void define(const DisjointBoxLayout& a\_fineGrids, int a\_nComp, int a\_nRef)
- void averageToCoarse(LevelData<FluxBox>& a\_coarse\_data, const LevelData<FluxBox>& a\_fine\_data) averages fine-level data to coarse level using arithmetic averaging
- void averageToCoarseHarmonic(LevelData<FluxBox>& a\_coarse\_data, const LevelData<FluxBox>& a\_fine\_data)
   averages fine-level data to coarse level using harmonic averaging

# 3.2.3 Class FineInterp

This class fills the valid region of a level of data by piecewise linear interpolation from data on a coarser level of refinement, using the piecewise linear interpolation operator described in section 3.1.1.

The piecewise-linear interpolation scheme used in the FineInterp class is limited to prevent the introduction of new maxima and minima. The user is able to set the limiting behavior near domain boundaries. Near non-periodic domain boundaries, there are four options when computing interpolated values, corresponding to the four possible values in BoundaryLimitType enumeration in FineInterp:

- limitSlopes normal piecewise-linear interpolation, with the standard vanLeer limiting of slopes to prevent new maxima. This requires that ghost-cell values be set on the coarse data at domain boundaries.
- noSlopeLimiting (default) piecewise-linear interpolation without limiting. This doesn't require coarse-level ghost cells be set, but may introduce new maxima/minima (or break positivity) for non-smooth functions.
- PCInterp piecewise-constant interpolation. Safest bet, since it's max/min-preserving without requiring that ghost cells be set but also least accurate.
- limitTangentialOnly limit slopes in the coordinate directions tangential to the boundary, while not limiting in the normal direction. This is useful when you don't have reasonable ghost-cell values but still need limiting.

The user can over-ride the default behavior in favor of what an application demands by resetting a static variable, FineInterp::s\_default\_boundary\_limit\_type. The default behavior can then be over-ridden for an individual instantiation of the FineInterp class by modifying the member variable m\_boundary\_limit\_type.

#### • void

- a\_fine\_domain (not modified): grids (valid region) on the fine level.
- a\_numcomps (not modified): number of components of the coarse and fine data.
- a\_ref\_ratio (not modified): the refinement ratio  $n_{ref} = \Delta x^c/\Delta x^f$ .
- a\_fine\_problem\_domain (not modified): the problem domain in the fine level index space. The version which uses a Box assumes a non-periodic domain.
- void interpToFine(LevelData<FArrayBox>& a\_fine\_data,

```
const LevelData<FArrayBox>& a_coarse_data,
bool a_averageFromDest=false););
```

Replaces fine data by interpolation from coarse data. Arguments:

- a\_fine\_data (modified): the fine data set, destination of interpolation.
- a\_coarse\_data (not modified): the coarse data set, source of interpolation.
- a\_averageFromDest: if true, first fill a projection of the fine grid with averaged values from a\_fine\_data before filling with coarse data and performing interpolation. This is often useful in operations like flattening an AMR hierarchy to a single resolution, in which the fine data may not be properly nested within the coarse data grids. Default is false.

#### • void

Replaces fine data by piecewise-constant interpolation from coarse data. Arguments:

- a\_fine\_data (modified): the fine data set, destination of interpolation.
- a\_coarse\_data (not modified): the coarse data set, source of interpolation.
- a\_averageFromDest: if true, first fill a projection of the fine grid with averaged values from a\_fine\_data before filling with coarse data and performing interpolation. This is often useful in operations like flattening an AMR hierarchy to a single resolution, in which the fine data may not be properly nested within the coarse data grids. Default is false.

# 3.2.4 Class FineInterpFace

This class fills face-centered data in the valid region of a level of data by piecewise linear interpolation from face-centered data on a coarser level of refinement. This interpolation is performed in two steps:

- 1. Data on fine-level faces which overlie coarse-level faces is interpolated using only the underlying co-planar coarse faces.
- Data on fine-level faces which do not overlie coarse-level faces is computed using linear interpolation in the normal direction between the two nearest fine-level faces which overlie coarse-level faces (and were filled in the previous step)
- void define(const DisjointBoxLayout& a\_fine\_domain, const int& a\_numcomps, const int& a\_ref\_ratio,

```
const ProblemDomain& a_fine_problem_domain)
```

# **Arguments:**

- a\_fine\_domain: the fine level domain
- a\_numcomps: the number of components
- a\_ref\_ratio: the refinement ratio
- a\_fine\_problem\_domain: problem domain on finest level
- void interpToFine(LevelData<FluxBox>& a\_fine\_data, const LevelData<FluxBox>& a\_coarse\_data, bool a\_averageFromDest=false)

Replaces fine data by piecewise-constant interpolation from coarse data. Arguments:

- a\_fine\_data (modified): the fine data set, destination of interpolation.
- a\_coarse\_data (not modified): the coarse data set, source of interpolation.
- a\_averageFromDest: if true, first fill a projection of the fine grid with averaged values from a\_fine\_data before filling with coarse data and performing interpolation. This is often useful in operations like flattening an AMR hierarchy to a single resolution, in which the fine data may not be properly nested within the coarse data grids. Default is false.

# 3.2.5 Class PiecewiseLinearFillPatch

This class fills some of the ghost cells of a level of data by piecewise linear interpolation from data on a coarser level of refinement. It is intended to be used in the context of a multilevel time-dependent adaptive mesh refinement (AMR) calculation. The algorithm used is that described in section 3.1.2.1. The interface described here is slightly more general, as it allows for the coarse grid data to be a linear combination of the form

$$\varphi^{c,valid} = \alpha \varphi^{c,old} + (1 - \alpha)\varphi^{c,new}$$

This can be useful, for example, when one has coarse-level data at two times ( $t^{c,old}$  and  $t^{c,new}$ ) and needs interpolated ghost cell data at an intermediate time  $t^{fine}=t^{c,old}+\alpha(t^{c,new}-t^{c,old})$ .

Ghost cells which lie inside the valid region of another fine grid are not filled. Also, note that cells outside the problem domain are never filled; it is the application developer's responsibility to fill them elsewhere according to the application-specific boundary conditions. Cells outside the computational domain in periodic direction, however, are considered to be inside the problem domain and are filled.

#### • void

Defines domains of the levels and other persistent data.

### **Arguments:**

- a\_fine\_domain (not modified): grids on the fine level.
- a\_coarse\_domain (not modified): grids on the coarse level.
- a\_num\_comps (not modified): number of components of state vector.
- a\_crse\_problem\_domain (not modified): problem domain on the coarse level.
- a\_ref\_ratio (not modified): refinement ratio.
- a\_interp\_radius (not modified): number of layers of fine ghost cells to fill by interpolation.
- a\_pwconst\_interp\_only (not modified) if true, only set up for piecewiseconstant interpolation in space.

#### • void

```
Real a_time_interp_coef,
int a_src_comp,
int a_dest_comp,
int a_num_comp);
```

Fills the ghost cells of the fine level data by interpolation.

## **Arguments:**

- a\_fine\_data (modified): fine data whose ghost cells are to be filled.
- a\_old\_coarse\_data (not modified): coarse level data at the old time.
- a\_new\_coarse\_data (not modified): coarse level data at the new time.
- a\_time\_interp\_coef (not modified): time interpolation coefficient,  $\alpha$ . It is required that  $0 \le \alpha \le 1$ .
- a\_src\_comp (not modified): starting coarse data component.
- a\_dest\_comp (not modified): starting fine data component.
- a\_num\_comp (not modified): number of data components to be interpolated.

#### • void

Fills the ghost cells of the fine level data by piecewise-constant interpolation.

## **Arguments:**

- a\_fine\_data (modified): fine data whose ghost cells are to be filled.
- a\_coarse\_data (not modified): coarse level data
- a\_src\_comp (not modified): starting coarse data component.
- a\_dest\_comp (not modified): starting fine data component.
- a\_num\_comp (not modified): number of data components to be interpolated.

### **3.2.6** Class PiecewiseLinearFillPatchFace

The PiecewiseLinearFillPatchFace class is similar to the PiecewiseLinearFillPatch class, but computes interpolated face-centered data to fill "ghost faces" around fine grids by interpolating from coarse-level face-centered data. This interpolation is performed in two steps:

1. Data on fine-level faces which overlie coarse-level faces is interpolated using only the underlying co-planar coarse faces.

- Data on fine-level faces which do not overlie coarse-level faces is computed using linear interpolation in the normal direction between the two nearest fine-level faces which overlie coarse-level faces (and were filled in the previous step)
- void

Defines domains of the coarse and fine levels and other persistent data.

# **Arguments:**

- a\_fine\_domain (not modified): grids on the fine level.
- a\_coarse\_domain (not modified): grids on the coarse level.
- a\_num\_comps (not modified): number of components of state vector.
- a\_crse\_problem\_domain (not modified): problem domain on the coarse level.
- a\_ref\_ratio (not modified): refinement ratio.
- a\_interp\_radius (not modified): number of layers of fine ghost faces to fill by interpolation.
- void

Fills fine-level ghost faces by linear interpolation.

- a\_fine\_data (modified): fine data whose ghost faces are to be filled.
- a\_old\_coarse\_data (not modified): coarse level data at the old time.
- a\_new\_coarse\_data (not modified): coarse level data at the new time.
- a\_time\_interp\_coef (not modified): time interpolation coefficient,  $\alpha$ . It is required that  $0 \le \alpha \le 1$ .
- a\_src\_comp (not modified): starting coarse data component.

- a\_dest\_comp (not modified): starting fine data component.
- a\_num\_comp (not modified): number of data components to be interpolated.

# 3.2.7 Class QuadCFInterp

The class QuadCFInterp interpolates data onto the ghost cells along the coarse-fine interface of a LevelData<FArrayBox>, using the algorithm described in section 3.1.2.2. It uses one-sided differencing in places where the stencil to do full centered differencing is partially covered by finer grids. The user interface of QuadCFInterp is given as follows.

 void define(const DisjointBoxLayout& a\_fineBoxes, const DisjointBoxLayout\* a\_coarBoxes, Real a\_dx, int a\_refRatio, int a\_nComp, const ProblemDomain& a\_domf);

Full define function. This makes all coarse-fine information and sets internal variables.

## **Arguments:**

- a\_fineBoxes (not modified): The grids at the current level.
- a\_coarBoxes (not modified): The grids for the next coarser level in the AMR hierarchy.
- a\_dx (not modified): The grid spacing at the current level.
- a\_refRatio (not modified): The refinement ratio between this level and the next coarser level in the AMR hierarchy.
- a\_nComp (not modified): The number of components in the data to be interpolated.
- a\_domf (not modified): The problem domain at the fine level.
- void coarseFineInterp(LevelData<FArrayBox>& a\_phif, const LevelData<FArrayBox>& a\_phic) const;

Coarse-fine interpolation operator. Fills all the ghost cells on all the faces of the LevelData<FArrayBox> a\_phif with values interpolated with a\_phic.

- a\_phif (modified): The solution at the current level.
- a\_phic (not modified): The solution at the next coarser level in the AMR hierarchy.

# 3.2.8 Class LevelFluxRegister

LevelFluxRegister manages the manipulations at coarse-fine boundaries associated with maintaining conservation form of cell-centered discretizations of the divergence operator, using the algorithm described in section 3.1.2.3. Unlike the previous operators, LevelFluxRegister holds data, corresponding to the flux register  $\delta \vec{F}^f$  defined in section 3.1.2.3. The class also manages the manipulation of that data.

The user interface for LevelFluxRegister is as follows.

Defines the internal state of the flux register, allocating space for the register itself, as well as the indexing information required to perform the other operations.

- a\_dbl (not modified): The grids at the current level.
- a\_dblCoarse (not modified): The grids at the next coarser level in the AMR hierarchy.
- a\_dProblem (not modified): The problem domain at the current level.
- a\_nRefine (not modified): The refinement ratio between this level and the next coarser level.
- a\_nComp (not modified): The number of variables used in the computation.
- a\_scaleFineFluxes (not modified): If true (default) then all fine fluxes are scaled by 1/a\_nRefine^(SpaceDim-1). Otherwise, no scaling is done.
- virtual void setToZero() Initializes the register to all zeros.

```
const Interval& a_dstInterval,
int a_dir);
```

Increments the register with data from a\_coarseFlux, multiplied by a\_scale (\$\alpha\$):  $\delta F_d^f := \delta F_d^f + \alpha F_d^c$ , for all of the d-faces where the input flux (defined on a single rectangle) coincides with the d-faces on which the flux register is defined. a\_coarseFlux contains fluxes in the a\_dir direction for the grid a\_dblCoarse[a\_coarsePatchIndex]. Only the registers corresponding to the low faces of a\_dblCoarse[a\_coarseDataIndex] in the a\_dir direction are incremented (this avoids double-counting at coarse-coarse interfaces. a\_srcInterval gives the Interval of components of a\_coarseFlux that correspond to a\_dstInterval of components of the flux register.

### **Arguments:**

- a\_coarseFlux (not modified): Flux to put into the flux register. This is not const because its box is shifted back and forth - no net change occurs.
- a\_scale (not modified): Factor by which to multiply a\_coarseFlux in flux register.
- a\_coarseDataIndex (not modified): DataIndex which corresponds to which box in the coarse-level DisjointBoxLayout (a\_dblCoarse in the define function) over which a\_coarseFlux was calculated.
- a\_srcInterval (not modified): The Interval of components to put into the flux register.
- a\_dstInterval (not modified): The Interval of components of the flux register which are incremented by the flux data. Should have the same size as a\_srcInterval.
- a\_dir (not modified): Direction of faces upon which fluxes live.
- virtual void incrementCoarse(FArrayBox& a\_coarseFlux,

Real a\_scale, const DataIndex& a\_coarseDataIndex, const Interval& a\_srcInterval, const Interval& a\_dstInterval, int a\_dir, Side:LoHiSide a\_sd);

Increments the register with data from a\_coarseFlux, multiplied by a\_scale (\$\alpha\$):  $\delta F_d^f := \delta F_d^f + \alpha F_d^c$ , for all of the d-faces where the input flux (defined on a single rectangle) coincides with the d-faces on which the flux register is defined. a\_coarseFlux contains fluxes in the a\_dir direction for the grid a\_dblCoarse[a\_coarsePatchIndex]. Only the registers corresponding to the low faces of a\_dblCoarse[a\_coarseDataIndex] in the a\_dir direction are incremented (this avoids double-counting at coarse-coarse interfaces.

a\_srcInterval gives the Interval of components of a\_coarseFlux that correspond to a\_dstInterval of components of the flux register. Only the register corresponding to the direction a\_dir and the side a\_sd is incremented.

## **Arguments:**

- a\_coarseFlux (not modified): Flux to put into the flux register. This is not const because its box is shifted back and forth - no net change occurs.
- a\_scale (not modified): Factor by which to multiply a\_coarseFlux in flux register.
- a\_coarseDataIndex (not modified): DataIndex which corresponds to which box in the coarse-level DisjointBoxLayout (a\_dblCoarse in the define function) over which a\_coarseFlux was calculated.
- a\_srcInterval (not modified): The Interval of components to put into the flux register.
- a\_dstInterval (not modified): The Interval of components of the flux register which are incremented by the flux data. Should have the same size as a\_srcInterval.
- a\_dir (not modified): Direction of faces upon which fluxes live.
- a\_sd (not modified) which side (low or high) to update
- virtual void incrementFine(FArrayBox& a\_fineFlux,

Real a\_scale,
const DataIndex& a\_finePatchIndex,
const Interval& a\_srcInterval,
const Interval& a\_dstInterval,
int a\_dir,
Side::LoHiSide a\_sd);

Increments the register with the average over each face of data from a\_fineFlux, scaled by a\_scale ( $\alpha$ ):  $\delta F_d^f = \delta F_d^f + \alpha < F_d^f >$ , for all of the d-faces where the input flux (defined on a single rectangle) covers the d-faces on which the flux register is defined. a\_fineFlux contains fluxes in the a\_dir direction for the grid a\_dbl[a\_finePatchIndex]. Only the register corresponding to the direction a\_dir and the side a\_sd is initialized. a\_srcInterval and a\_dstInterval are as above.

- a\_fineFlux (not modified): Flux to put into the flux register. This is not const because its box is shifted back and forth - no net change occurs.
- a\_scale (not modified): Factor by which to multiply a\_fineFlux in flux register.

- a\_finePatchIndex (not modified): Index which corresponds to which box in the fine-level DisjointBoxLayout (a\_dbl in the define function) over which a\_fineFlux was calculated.
- a\_srcInterval (not modified): The Interval of components to put into the flux register.
- a\_dstInterval (not modified): The Interval of components of the flux register which are incremented by the flux data.
- a\_dir (not modified): Direction of faces upon which fluxes live.
- a\_sd (not modified): Side of the fine face where coarse-fine interface lies.
- virtual void reflux(LevelData<FArrayBox>& a\_uCoarse, const Interval& a\_coarse\_interval, const Interval& a\_flux\_interval, Real a\_scale);

Increments a\_uCoarse with the reflux divergence of the contents of the flux register, scaled by a\_scale ( $\alpha$ ):  $U^c := U^c + \alpha D_R(\delta \vec{F})$ . a\_flux\_interval gives the Interval of components of the flux register that correspond to a\_coarse\_interval of components of a\_uCoarse.

## **Arguments:**

- a\_uCoarse (modified): LevelData<FArrayBox> which is modified by the refluxing operation.
- a\_coarse\_interval (not modified): The Interval of components to put into a\_uCoarse.
- a\_flux\_interval (not modified): The Interval of components to use from the flux register.
- a\_scale (not modified): Factor by which to scale the flux register.

# 3.2.9 Class LevelFluxRegisterEdge

In Magnetohydrodynamics (among other fields), there is a class of numerical schemes commonly known as "constrained transport" schemes in which the solenoidal property of a field is maintained by making use of the identity div(curl(u)) == 0. In MHD, for example, one can write the evolution of the magnetic field  $\vec{B}$  in terms of the curl of the electric field  $\vec{E}$ , which ensures that the  $div(\vec{B}) = 0$  down to roundoff. This is generally accomplished on a staggered mesh, in which  $\vec{B}$  is face-centered and  $\vec{E}$  is edge-centered.

At coarse-fine interfaces, one must perform a correction analogous to the reflux-divergence operation in order to maintain a divergence-free magnetic field. This correction is described, for example, in [3]. The LevelFluxRegisterEdge is a class designed to manage and apply this correction, and is analogous to the LevelFluxRegister. While

the LevelFluxRegister corrects a cell-centered field with a "reflux-divergence" of face-centered fluxes, the LevelFluxRegisterEdge corrects a face-centered field with a "reflux curl" of edge-centered "fluxes".

 virtual void define(const DisjointBoxLayout& a\_dbl, const DisjointBoxLayout& a\_dblCoarse, const ProblemDomain& a\_dProblem, int a\_nRefine, int a\_nComp)

Defines this object

- virtual void setToZero()
   Sets all registers to zero.
- virtual void incrementCoarse(FArrayBox& a\_coarseFlux,

Real a\_scale,
const DataIndex& a\_coarseDataIndex,
const Interval& a\_srcInterval,
const Interval& a\_dstInterval)

increments the register with data from coarseFlux, multiplied by a\_scale. a\_coarseFlux must contain the edge-centered (in 3d, node centered in 2d) coarse fluxes in the dir direction for the grid m\_coarseLayout[coarseDataIndex]. By convention, only the low side flux is used to avoid double-counting at coarse-fine interfaces.

• virtual void incrementFine(FArrayBox& a\_fineFlux,

Real a\_scale,
const DataIndex& a\_fineDataIndex,
const Interval& a\_srcInterval,
const Interval& a\_dstInterval)

increments the register with data from fineFlux (which is edge-centered in 3d, node-centered in 2d), multiplied by a\_scale, for all coarse-fine face directions associated with the grid box  $m_fineLayout[fineDataIndex]$ 

virtual void incrementFine(FArrayBox& a\_fineFlux,

Real a\_scale, const DataIndex& a\_fineDataIndex, const Interval& a\_srcInterval, const Interval& a\_dstInterval, int a\_dir, Side::LoHiSide a\_sd)

increments the register with data from fineFlux (which is edge-centered in 3d, node-centered in 2d), multiplied by a\_scale. a\_dir is the normal of the coarse-fine

interface, and a\_sd determines whether we're looking at the high-side or the low-side for the grid box m\_fineLayout[fineDataIndex]

 virtual void refluxCurl(LevelData<FluxBox>& a\_uCoarse, Real a\_scale)

increments uCoarse with the reflux "CURL" of the contents of the flux register. This is done for all components so a\_uCoarse has to have the same number of components as input a\_nComp. This operation is global and blocking.

## 3.3 Class BRMeshRefine

Derived from the generic MeshRefine parent class, BRMeshRefine is an object which produces a hierarchy of block-structured grids which obey proper-nesting requirements. See Berger and Colella [7] for an explanation of proper nesting. BRMeshRefine follows the algorithm of Berger and Rigoutsos [8] to generate the grids from tagged points in discrete index space. There are two interfaces for BRMeshRefine grid generation: one takes tags at all levels in the hierarchy and one takes tags only at the coarsest level. If the BRMeshRefine object is defined with a ProblemDomain which is periodic in one or more directions, grids generated will be properly nested in the periodic directions.

The user interface for BRMeshRefine is as follows (many functions actually live in the parent MeshRefine class:

BRMeshRefine();

Default constructor – the object is defined in an unusable state until the user calls the define function.

• BRMeshRefine(

```
const ProblemDomain& a_baseDomain,
const Vector<int>& a_refRatios,
const Real a_fillRatio,
const int a_blockFactor,
const int a_bufferSize,
const int a_maxSize);
```

#### BRMeshRefine(

```
const Box& a_baseDomain,
const Vector<int>& a_refRatios,
const Real a_fillRatio,
const int a_blockFactor,
const int a_bufferSize,
const int a_maxSize);
```

Full constructor. Places the BRMeshRefine object in a usable state by calling the corresponding define function.

#### **Arguments:**

- a\_baseDomain Problem domain at the coarsest (level 0) level. Output grids will be constrained to be within the computational domain on each level.
- a\_refRatios Refinement ratios between the levels. a\_refRatio[i] represents the refinement ratio between levels i and i+1. The vector indices must correspond to level number.
- a\_fillRatio Overall grid efficiency to be generated. If this number is set low, the grids will tend to be larger and less filled with tags. If this number is set high, the grids will tend to be smaller and more filled with tags. This controls the aggressiveness of agglomeration by box merging.
- a\_blockFactor Blocking factor. For each box B in the grids, this is the number Nref for which it is guaranteed to be true that refine(coarsen(B,Nref),Nref)==B. Default = 1. Note that this will also be the minimum possible box size.
- a\_bufferSize Proper nesting buffer size. This will be the minimum number of level  $\ell$  cells between any level  $\ell+1$  cell and a level  $\ell-1$  cell. Default = 1.
- a\_maxSize Maximum length of a grid in any dimension. An input value of 0 means the maximum value will be  $\infty$  (no limit).

## void define(

```
const ProblemDomain&
                                 a_baseDomain,
       const Vector<int>&
                                 a_refRatios,
       const Real
                                 a_fillRatio,
       const int
                                 a_blockFactor,
       const int
                                 a_bufferSize,
                                 a_maxSize );
       const int
void
define(
       const Box&
                                 a_baseDomain,
       const Vector<int>&
                                 a_refRatios,
       const Real
                                 a_fillRatio,
       const int
                                 a_blockFactor,
                                 a_bufferSize,
       const int
                                 a_maxSize );
       const int
```

Defines (or redefines) a BRMeshRefine object and places it in a usable state.

- a\_baseDomain Problem domain at the coarsest (level 0) level. Output grids will be constrained to be within the computational domain on each level.
- a\_refRatios Refinement ratios between the levels. RefRatio[i] represents the refinement ratio between levels i and i+1. The vector indices must correspond to level number.
- a\_fillRatio Overall grid efficiency to be generated. If this number is set low, the grids will tend to be larger and less filled with tags. If this number is set high, the grids will tend to be smaller and more filled with tags. This controls the aggressiveness of agglomeration by box merging.
- a\_blockFactor Blocking factor. For each box B in the grids, this is the number Nref for which it is guaranteed to be true that refine(coarsen(B,Nref),Nref)==B. Default = 1. Note that this will also be the minimum possible box size.
- a\_bufferSize Proper nesting buffer size. This will be the minimum number of level  $\ell$  cells between any level  $\ell+1$  cell and a level  $\ell-1$  cell. Default = 1.
- a\_maxSize Maximum length of a grid in any dimension. An input value of 0 means the maximum value will be  $\infty$  (no limit).

Inherited from MeshRefine (but which uses the Berger-Rigoutsos algorithm implemented in BRMeshRefine), the interface which takes tags at all levels and generates a new multilevel hierarchy of grids which covers the tags at each level while satisfying the proper nesting requirements. Note that the proper nesting requirement is an overriding constraint – if a tagged cell cannot be refined while satisfying proper nesting, it is not refined. (This is only an issue if a\_baseLevel > 0.). The grids produced by this function will also satisfy the constraints placed by the BlockFactor, FillRatio, and MaxSize. Returns the finest level on which grids are defined.

- a\_newmeshes The set of grids at every level. This is resized and filled in this function
- a\_tags Tagged cells on every level from a\_baseLevel to a\_topLevel-1.
   The vector indices must correspond to level number.

- a\_baseLevel Index of base mesh level. This is the finest level which does \*not\* change. For example, if all grids except level 0 are going to be changed by BRMeshRefine, a\_baseLevel = 0.
- a\_topLevel Index of top level of relevant tags which is the same as one level \*below\* the highest level of grids that will be produced. So if the AMR hierarchy has 9 levels and one wants all of them to change except level 0, then a\_baseLevel = 0 and a\_topLevel = 7 (highest level number is 8).
- a\_oldMeshes Grids before BRMeshRefine is called. If there are no previous grids, set a\_oldMeshes to be the problem domains. See the example shown in figure 3.4. The vector indices must correspond to level number.
- Returns the finest level on which grids are defined in a\_newmeshes.

## • int

```
regrid(
```

```
Vector<Vector<Box> >& a_newmeshes,
IntVectSet& a_tags,
const int a_baseLevel,
const int a_topLevel,
const Vector<Vector<Box> >& a_oldMeshes) const;
```

The interface for BRMeshRefine which takes only a single level of tags and generates a multilevel hierarchy of grids which covers those tags while satisfying the proper nesting requirements. Note that the proper nesting requirement is an overriding constraint – if a tagged cell cannot be refined while satisfying proper nesting, it is not refined. (This is only an issue if a\_baseLevel > 0.). The grids produced by this function will also satisfy the constraints placed by the BlockFactor, FillRatio, and MaxSize. Returns the finest level on which grids are defined (for this function, this will normally be TopLevel+1)

- a\_newmeshes The new set of grids at every level. This is resized and filled in the function.
- a\_tags Tagged cells on a\_baseLevel.
- a\_baseLevel Index of base mesh level. This is the finest level which does \*not\* change. For example, if all grids except level 0 are going to be changed by BRMeshRefine, a\_baseLevel = 0.
- a\_topLevel Index of top level of relevant tags which is the same as one level \*below\* the highest level of grids that will be produced. So if the AMR hierarchy has 9 levels and one wants all of them to change except level 0, then a\_baseLevel = 0 and a\_topLevel = 7 (highest level number is 8).

- a\_oldMeshes Grids before BRMeshRefine is called. If there are no previous grids, set a\_oldMeshes to be the problem domains. See the example shown in figure 3.4 The vector indices must correspond to level number.
- Returns the finest level on which grids are defined in newmeshes.

Figure 3.4 is a sample code to show the use of BRMeshRefine to create lists of grids from tags. For an explanation of how to use LoadBalance to transform these into DisjointBoxLayouts see section 9.4.

 const Vector<int>& refRatios() const;
 Returns the vector of refinement ratios

 const Real fillRatio() const;
 Returns the FillRatio.

 const int blockFactor() const;
 Returns the blocking factor.

const int bufferSize() const;Returns the proper nesting buffer size.

• const int
maxSize() const;

returns the maximum box length. A value of 0 means the maximum value is  $\infty$  (no limit).

 void refRatios(const Vector<int>& a\_nRefVect);
 Sets the vector of refinement ratios

 void fillRatio(const Real a\_fillRat);
 Sets the FillRatio.

 void blockFactor(const int a\_blockFactor);
 Sets the blocking factor.

```
int setGrids(
             Vector<Vector<Box> >&
                                           a_vectGrids,
             const Vector<ProblemDomain>& a_vectDomain,
             Vector<int>&
                                           a_vectRefRatio,
             int&
                                           a_numlevels,
             Real
                                           a_fillRat,
             int
                                           a_maxboxsize)
{
 Box btag = a_vectDomain[0].domainBox();
  int ishrink = btag.size(0)/4;
  btag.grow(-ishrink);
  IntVectSet tags(btag);
  Vector<Vector<Box> > VVBoxNew(a_numlevels);
  Vector<Vector<Box> > VVBoxOld(a_numlevels);
  for(int ilev = 0; ilev <a_numlevels; ilev++)</pre>
   VVBoxOld[ilev].push_back(a_vectDomain[ilev].domainBox());
  }
  int baseLevel = 0;
  int topLevel = a_numlevels - 2;
  int blockFactor = 2;
  int buffersize = 1;
  if(topLevel >= 0)
  {
   BRMeshRefine meshRefine(a_vectDomain[0], a_vectRefRatio,
                    a_fillRat, blockFactor, buffersize,
                             a_maxboxsize)
   meshRefine.regrid(VVBoxNew, tags, baseLevel, topLevel,
                      VVBoxOld):
 }
 else
    VVBoxNew = VVBoxOld;
  }
  a_vectGrids = VVBoxNew;
 return 0;
}
```

Figure 3.4: Sample code to show the use of BRMeshRefine to create lists of grids from tags which have been defined on the base level.

- void bufferSize(const int a\_buffSize);
   Sets the proper nesting buffer size.
- void
   maxSize(const int a\_maxSize);
   Sets the maximum box length. An input value of 0 means the maximum value will be ∞ (no limit).

## 3.3.1 domainSplit

There are many times when the physical domain on the coarsest AMR level (level 0) is larger than the maximum desired block size. In this case, the solution is to split the domain into more than one piece. This is especially useful for parallel computations. To simplify this process, the stand-alone function DomainSplit is provided (in BRMeshRefine.H):

```
void
domainSplit(const ProblemDomain& a_domain,
            Vector<Box>&
                                  a_vbox,
            int
                           a_maxsize,
                            a_blockfactor=1,
            int
            IntVect a_refineDirs=IntVect::Unit);
void
domainSplit(const Box&
                         a_domain,
            Vector<Box>& a_vbox,
            int
                   a_maxsize,
                   a_blockfactor=1,
            IntVect a_refineDirs=IntVect::Unit);
```

- a\_domain Physical domain
- a\_vbox Vector of boxes which satisfy the blocking factor and maxsize requirements which make up the decomposed domain.
- a\_maxsize Maximum allowable box size (0 means no limit).
- a\_blockfactor Blocking factor; has the same definition as in BRMeshRefine.
- a\_refineDirs Allows for splitting only in specific directions 1 if splitting the domain in dimension d, and 0 if not

## 3.4 Multilevel Utilities

In addition to the two-level AMRTools operator objects, there are a set of multilevel tools which have proved useful in various multilevel AMR codes.

## **3.4.1 Function** computeSum

This function computes the volume-weighted sum of  $\phi$  over an AMR hierarchy by including only valid-region data in the sum.

$$sum = \sum_{\ell=\ell_{base}}^{\ell_{max}} \sum_{\Omega_{valid}^{\ell}} (h^{\ell})^{D} \phi_{\mathbf{i}}^{\ell}$$
(3.5)

• void

function which returns the volume-weighted sum of a\_phi (essentially the integral of a\_phi) over the valid regions of all levels  $\ell \geq$  a\_lBase.

## **Arguments:**

- a\_phi (not modified): data to be summed.
- a\_nRefFine (not modified): Vector of refinement ratios, where a\_refFine[i] is the refinement ratios between levels i and i + 1.
- a\_dxCrse (not modified): cell spacing on level a\_1Base.
- a\_comps (not modified): components of a\_phi to be summed.
- a\_1Base (not modified): Coarsest level to be included in sum; sum will include all levels  $\ell \geq$  a\_1Base.

## 3.4.2 Function computeNorm

This function computes the norm of  $\phi$  over an AMR hierarchy by including only valid-region data in the computation.

• void

const int 
$$a_p = 2$$
, const int&  $a_1Base = 0$ );

function which returns the  $p{\rm -norm}$  of a\_phi over the valid regions of all levels  $\ell \geq {\rm a\_lBase}.$ 

- a\_phi (not modified): data to be summed.
- a\_nRefFine (not modified): Vector of refinement ratios, where a\_refFine[i] is the refinement ratios between levels i and i + 1.
- a\_dxCrse (not modified): cell spacing on level a\_1Base.
- a\_comps (not modified): components of a\_phi to be summed.
- a\_p (not modified): type of norm to be computed. a\_p=0 is the max (infinity) norm.
- a\_1Base (not modified): Coarsest level to be included in sum; sum will include all levels  $\ell \geq$  a\_1Base.

## Chapter 4

# AMREIliptic Algorithm and Implementation

## 4.1 Multigrid Algorithm

We want to solve the equation

$$L^{comp}\varphi^{comp} = \rho^{comp}$$

on an AMR hierarchy  $\{\Omega^l\}_{l=0}^{l_{max}}$  satisfying the nesting conditions described in [7]. The algorithm we use here is a natural extension of multigrid iteration. The particular version we describe here [23, 24] is a linear version of the algorithm used in [29] to compute steady incompressible flow, and has been used in a variety of settings [2, 1, 9, 10].

A pseudo-code description of the algorithm is given in figure (4.3). The operators Average and  $I_{pwe}$  are described in section 3.1.1, and the operator  $L^{nf}$  is a two-level discretization of the Laplacian:

$$L^{nf}(\psi^f, \psi^{c,valid}) = D(\vec{G}^f(\psi^f, \psi^{c,valid})).$$

It computes a uniform grid  $2\mathbf{D}+1$  point discretization of the Laplacian applied to an extension of  $\psi^f$  obtained using the quadratic interpolation procedure in section 3.1.2.2. The smoothing operator  $\operatorname{mgRelax}(\varphi^f,R^f,r)$  performs a multigrid V-cycle iteration on  $\varphi^f$  for the operator  $L^{nf}$ , assuming the coarse-grid values required for the boundary conditions are identically zero.

## 4.2 The AMR Elliptic User Interface

The implementation of the AMRElliptic package follows the algorithm specification in section 4.1.

Figure 4.1: Recursive relaxation procedure.

```
procedure LevelGSRB(\varphi^f, R^f) {  \varphi^f := \varphi^f + \lambda(L^{nf}(\varphi^f, \varphi^c \equiv 0) - R^f) \text{ on } \Omega^{BLACK}   \varphi^f := \varphi^f + \lambda(L^{nf}(\varphi^f, \varphi^c \equiv 0) - R^f) \text{ on } \Omega^{RED}  }
```

Figure 4.2: Gauss-Seidel relaxation with red-black ordering. Here  $\lambda$  is the relaxation parameter.

```
R := \rho - L(\varphi)
while (||R|| > \epsilon ||\rho||)
         \mathsf{AMRVCycleMG}(l^{max})
         R := \rho - L(\varphi)
end while
Procedure AMRVCycleMG(level l):
if (l=l^{max}) then \overset{.}{R^{l}}:=\rho^{l}-L^{nf}(\varphi^{l},\varphi^{l-1})
if (l > 0) then
         \varphi^{l,save} := \varphi^l \text{ on } \Omega^l
         e^l := 0 on \Omega^l
        \begin{aligned} & \operatorname{mgRelax}(e^l, R^l, n_{ref}^{l-1}) \\ & \varphi^l := \varphi^l + e^l \end{aligned}
         \stackrel{\cdot}{e}^{l-1} := 0 \text{ on } \Omega^{l-1}
         R^{l-1} := Average(R^l - L^{nf}(e^l, e^{l-1})) \text{ on } \mathcal{C}_{n_{ref}^{l-1}}(\Omega^l)
         R^{l-1}:=\rho^{l-1}-L^{comp,l-1}(\varphi) \text{ on } \Omega^{l-1}-\mathcal{C}_{n_{ref}^{l-1}}(\Omega^l)
         \mathsf{AMRVCycleMG}(l-1)
         e^{l} := e^{l} + I_{pwc}(e^{l-1})

R^{l} := R^{l} - L^{nf,l}(e^{l}, e^{l-1})
         \delta e^l := 0 \text{ on } \Omega^l
         \begin{aligned} & \mathsf{mgRelax}(\delta e^l, R^l, n_{ref}^{l-1}) \\ & e^l := e^l + \delta e^l \end{aligned}
         \varphi^l := \varphi^{l,save} + e^l
else
        solve L^{nf}(e^0)=R^0 on \Omega^0. \varphi^0:=\varphi^0+e^0
end if
```

Figure 4.3: Pseudo-code description of the AMR multigrid algorithm.

#### 4.2.1 Overview

Code reuse is facilitated by using a *Template Method* design pattern. The purpose of the *Template Method* design pattern is to define an algorithm as a fixed sequence of steps but have one or more of the steps be variable. In our case, we have a hierarchy of algorithms that we wish to re-use across a family of applications. The hierarchy of algorithms is defined by the specific *variable steps* that an application must provide to complete the algorithm.

Our variable steps are supplied by a hierarchy of *Operator Interfaces*. In C++, a variable step is represented as a virtual function. Our algorithms are in the form of *Solver Templates*. Each Solver Template requires virtual functions provided by its corresponding Operator Interface. The data type used in these algorithms is supplied by a template parameter.

Various specific solvers derive from the appropriate interface class to utilize the desired solver algorithm.

An overview of our class structure for this design is presented here. Indentation implies inheritance

LinearOp<T>: User can utilize solvers that implement the LinearSolver<T> interface

MGLevelOp<T>: Users can utilize solvers that implement LinearSolver<T> and MultiGrid<T> interfaces.

AMRLevelOp<T> Users can utilize solvers that implement LinearSolver<T>, MultiGrid<T> and AMRMultiGrid<T> interfaces. Examples of instantiations of these interfaces are:

- \* PoissonOp (template data type FArrayBox). A single-level solver
- \* AMRPoissonOp (template data type LevelData<FArrayBox>) cell-centered AMR Poisson solver
- \* VCAMRPoissonOp (template data type LevelData<FArrayBox>) cell-centered variable-coefficient AMR Poisson and Helmholtz solver.
- \* EBPoissonOp (template data type LevelData<EBCellFAB>)
- \* EBAMRPoissonOp (template data type LevelData<EBCellFAB>)
- \* AMRNodeOp (template data type LevelData<NodeFArrayBox>) Nodecentered AMRPoisson and Helmholtz solver.
- \* ResistivityOp (template data type LevelData<FArrayBox>) cell-centered variable-coefficient operator to solve variable coefficient resistivity operator.
- \* ViscousTensorOp (template data type LevelData<FArrayBox>) cell-centered variable-coefficient operator to solve variable coefficient viscous tensor operator.

LinearSolver<T>

- uses LinearOp<T> interface functions to implement the algorithm's variable steps.
- some example implementations of this algorithm interface are:
  - BiCGStabSolver<T>
  - RelaxSolver<T>

#### MultiGrid<T>

- calls the MGLevelOp<T> interface
- uses a LinearSolver<T> as bottom solver.

#### AMRMultiGrid<T>

- uses AMRLevelOp<T> and MGLevelOp interfaces
- combines AMR coarse-fine operations with MultiGrid<T> and LinearSolver<T> operations.

## 4.3 Operator Interfaces

The variable steps of our *Template Method* are supplied through classes that implement the Operator Interfaces.

## 4.3.1 Class LinearOp

LinearOp is an operator class for representing L when solving  $L(\phi) = \rho$  This interface class serves two main purposes. First, It acts as a factory class for the template data type. Second, it provides the variable steps necessary for the family of LinearSolver<T> classes in Chombo.

- virtual void residual( T& lhs, const T& phi, const T& rhs, bool homogeneous = false) = 0;
   Compute the residual. For example, if solving L(phi) = rhs, then set lhs = L(phi) rhs. If homogeneous is true, evaluate the operator using homogeneous boundary conditions.
- virtual void preCond( T& cor, const T& residual) = 0;
   Given the current state of the residual and correction, apply your preconditioner to cor.
- virtual void applyOp( T& lhs, const T& phi, bool homogeneous = false)
   = 0;
   In the context of solving L(phi) = rhs, set lhs = L(phi). If homogeneous is true, evaluate the operator using homogeneous boundary conditions.

- virtual void create( T& lhs, const T& rhs) = 0; Create data holder lhs that mirrors rhs. You do not need to copy the data of rhs, just make a holder the same size.
- virtual void clear( T& 1hs);
   Perform any operations required before 1hs is destructed. In general, this function only needs to be defined if the create function called new. There is a default implementation of this function, which does nothing (which means that in most cases, classes derived from LinearOp will not need to define this function).
- virtual void assign( T& lhs, const T& rhs) = 0; Set lhs equal to rhs.
- virtual Real dotProduct(const T& a1, const T& a2) = 0; Compute and return the dot product of a1 and a2. In most contexts, this will return the sum over all data points of a1\*a2.
- virtual void incr ( T& lhs, const T& x, Real scale) = 0; Increment by scaled amount (lhs += scale\*x).
- virtual void axby( T& lhs, const T& x, const T& y, Real a, Real b)
   = 0;
   Compute a scaled sum (lhs = a\*x + b\*y).
- virtual void scale( T& lhs, const Real& scale) = 0;
   Multiply the input by a given scale lhs \*= scale).
- virtual Real norm(const T& rhs, int ord) = 0; Return the norm of rhs. If ord == 0, compute max norm, If ord == 1, compute  $L_1$  norm: sum(abs(rhs)). Otherwise, compute  $L_{ord}$  norm.
- virtual void setToZero(T& lhs) = 0; Set lhs to zero.

## 4.3.2 Class MGLevelOp

Class MGLevelOp handles the additional tasks of coordinating operations between this level and the next coarser 'level'. MGLevelOp provides the coarsening and interlevel operations needed for algorithms that implement the MultiGrid<T> solver class.

• virtual void createCoarser(T& coarse, const T& fine, bool ghosted) = 0; Create a coarsened (by two) version of the input data container. This does not include averaging the data. So, if fine is over a Box of (0,0,0) (63,63,63), coarse should be over a Box of (0,0,0) (31,31,31).

- virtual void relax(T& correction, const T& residual, int iterations) = 0; Apply relaxation operator to remove the high frequency wave numbers from the correction. A point relaxation scheme, for example, takes the form correction -= lambda\*(L(correction) residual).
- virtual void restrictResidual(T& resCoarse, T& phiFine, const T& rhsFine)
   = 0; calculate restricted residual
   resCoarse[2h] = I[h->2h] (rhsFine[h] L[h](phiFine[h])
- virtual void prolongIncrement(T& phiThisLevel, const T& correctCoarse)
   = 0; correct the fine solution based on coarse correction
   phiThisLevel += I[2h->h](correctCoarse)

## 4.3.3 Class MGLevelOpFactory

Factory class for generating MGLevelOps.

virtual MGLevelOp<T>\* MGnewOp(const ProblemDomain& FineindexSpace, int depth, bool homoOnly = true) = 0;
 Create an operator at an index space = coarsen(fineIndexSpace, 2depth).
 Return NULL if no such Multigrid level can be created at this depth. If homoOnly = true, then only homogeneous boundary conditions will be needed.

## 4.3.4 Class AMRLevelOp

The AMRLevelOp interface adds variable steps required by the AMRMultiGrid class of solvers. These pertain to operations between multigrid levels that do not form complete covering sets, and therefore require information from multiple levels simultaneously for coarse-fine boundary conditions.

- virtual int refToCoarser() = 0;
   Return the refinement ratio to next coarser level.
   Return 1 when there are no coarser AMR levels.

Compute the residual: residual = rhs - L(phiFine, phi, phiCoarse).

```
virtual void AMRResidualNF(T& residual,
                              const T& phi,
                              const T& phiCoarse,
                              const T& rhs,
                              bool homogeneousBC) = 0;
Compute residual = rhs - L(phi, phiCoarse) assuming no finer level.
virtual void AMRResidualNC(T& residual,
                              const T& phiFine,
                              const T& phi,
                              const T& rhs,
                              bool homogeneousBC,
                              AMRLevelOp<T>* finerOp) = 0;
Compute residual = rhs - L(phiFine, phi) assuming no coarser AMR level.
virtual void AMRRestrict(T& resCoarse,
                            const T& residual,
                            const T& correction,
                            const T& coarseCorrection) = 0;
Set resCoarse = I[h-2h]( residual - L(correction, coarseCorrection))
virtual void AMRProlong(T& correction,
                          const T& coarseCorrection) = 0;
Set correction += I[2h->h](coarseCorrection)
virtual void AMRUpdateResidual(T& residual,
                                  const T& correction,
                                  const T& coarseCorrection) = 0;
Set residual = residual - L(correction, coarseCorrection) .
virtual Real AMRNorm(const T& coarseResid,
                       const T& fineResid,
                       const int& refRat,
                       const int& ord) = 0;
Compute norm over all cells on coarse not covered by finer AMR levels.
```

Set the output to a coarsened (by the input refinement ratio) version of the finer.

## 4.3.5 Class AMRLevelOpFactory

Factory interface for AMRLevelOp generation.

virtual AMRLevelOp<T>\* AMRnewOp(const ProblemDomain& indexSpace)=0;
 Return a new operator object. This is done with a call to new; caller is responsible for deletion.

## 4.4 Solver Templates

Solver Template requires virtual functions provided by its corresponding Operator Interface. The data type used in these algorithms is supplied by a template parameter.

Various specific solvers derive from the appropriate interface class to utilize the desired solver algorithm.

#### **4.4.1 Class** LinearSolver

LinearSolver represents both a *Solver Algorithm* and an interface class. It is a Solver Algorithm with respect to the variable steps provided by the *Operator Interfaces*. Given an instantiation of a LinearSolver, any operator that implements the LinearOp interface can make invocations to its define and solve functions. It is also an interface, in that LinearSolver does not provide a default implementation, but instead is an interface to a variety of linear solver algorithms. MultiGrid and AMRMultiGrid provide a default implementation of geometric multigrid. Generic linear solver templatesBiCGStab and others are built on top of this.

virtual void define(LinearOp<T>\* operator,
bool homogeneous = false) = 0;

Define the operator and whether it is a homogeneous solver or not. The LinearSolver does not take over ownership of this operator object. It does not call delete on it when the LinearSolver is deleted. It is meant to be like a late-binding reference. If you created operator with new, you should call delete on it after LinearSolver is deleted if you want to avoid memory leaks.

- virtual void solve(T& phi, const T& rhs) = 0; Solve L(phi) = rhs (phi = L-1 (rhs)).
- virtual void setConvergenceMetrics(Real metric, Real tolerance) = 0; If appropriate, sets a metric to judge convergence, along with the solver tolerance. If not set, use default convergence metrics. This can be useful when using a LinearSolver as a bottom solver, since one may want to propagate the convergence metric and solver tolerance from the outer solver in to the bottom solver.

### 4.4.2 Class BiCGStabSolver

Elliptic solver using the BiCGStab algorithm.

- virtual void define(LinearOp<T>\* op, bool homogeneous);
   Define the solver.
  - op is the linear operator.
  - homogeneous is whether the solver uses homogeneous boundary conditions.
- virtual void solve(T& phi, const T& rhs); Solve the equation.
- bool m\_homogeneous public member data: whether the solver is restricted to homogeneous boundary conditions.
- LinearOp<T>\* m\_op: public member data: operator to solve.
- int m\_imax;
   public member data: maximum number of iterations.
- int m\_verbosity;
   public member data: how much screen output the user wants. set = 0 for no output.
- Real m\_eps;
   public member data: solver tolerance
- Real m\_hang; public member data: minimum rate that norm of solution should charge per iteration
- Real m\_small;
   public member data: what the algorithm should consider "close to zero"
- int m\_numRestarts; public member data: number of times the algorithm can restart
- int m\_normType;
   public member data: norm to be used when evaluating convergence.
   0 is max norm, 1 is L<sub>1</sub>, 2 is L<sub>2</sub>.

#### 4.4.3 Class GMRESSolver

Elliptic solver using the GMRES algorithm. Important parameters are:  $m_reps$ , the relative tolerance (default = 1.0e-12) and  $m_restrtLen$ : the restart length (default = 30).

- m\_reps: relative tolerance (default = 1.0e-12)
- m\_restrtLen: restart length (default = 30)
- virtual void define(LinearOp<T>\* op, bool homogeneous);
   Define the solver.
  - op is the linear operator.
  - homogeneous is whether the solver uses homogeneous boundary conditions.
- virtual void solve(T& phi, const T& rhs); Solve the equation.
- bool m\_homogeneous public member data: whether the solver is restricted to homogeneous boundary conditions
- LinearOp<T>\* m\_op: public member data: operator to solve.
- int m\_imax;
   public member data: maximum number of iterations.
- int m\_verbosity;
   public member data: how much screen output the user wants. set = 0 for no output.
- Real m\_eps;
   public member data: absolute solver tolerance
- Real m\_small;
   public member data: what the algorithm should consider "close to zero"
- int m\_normType;
   public member data: norm to be used when evaluating convergence.
   0 is max norm, 1 is L<sub>1</sub>, 2 is L<sub>2</sub>.
- Real m\_residual;
   public member data: current residual
- Real m\_initial\_residual;
   public member data: initial residual

Real m\_iter;
 public member data: current iteration number

#### 4.4.4 Class MultiGrid

MultiGrid is a class which executes a v-cycle of geometric multigrid. This class is not meant to be particularly user-friendly, and a good option for people who want something a tad less raw is to use AMRMultigrid instead.

Function to define a MultiGrid object:

- factory is the factory for generating operators.
- bottomSolver is called at the bottom of v-cycle.
- domain is the problem domain at the top of the v-cycle.
- maxDepth defines the location of the bottom of the v-cycle.

The v-cycle will terminate (hit bottom) when the factory returns NULL for a particular depth if maxdepth = -1. Otherwise the v-cycle terminates at maxdepth.

- virtual void solve(T& e, const T& res);
   Execute ONE v-cycle of multigrid. If you want the solution to converge, you will probably need to iterate this. See AMRMultiGrid for a more automatic solve() function. This operates residual-correction form of equation so all boundary conditions are assumed to be homogeneous. L(e) = res
- int m\_depth, m\_pre, m\_post, m\_cycle,m\_numMG; Public solver parameters. m\_pre and m\_post are the ones that usually get set and are the number of relaxations performed before and after multigrid recursion. See AMRMultiGrid for a more user-friendly interface.
- Vector< MGLevelOp<T>\* > getAllOperators();
   For changing coefficients not for the faint of heart.

#### 4.4.5 Class AMRMultiGrid

Class to execute geometric multigrid over an AMR hierarchy a-la Martin and Cartwright. [23]

 virtual void define(const ProblemDomain& coarseDomain, AMRLevelOpFactory<T>& factory, LinearSolver<T>\* bottomSolver, int maxAMRLevels);

Define the solver.

- coarseDomain is the index space on the coarsest AMR level.
- factory is the operator factory through which all special information is conveyed.
- bottomSolver is the solver to be used at the termination of multigrid coarsening.
- numLevels is the number of AMR levels.

Solve  $L(\phi) = \rho$  from 1\_base to 1\_max. To solve over all levels, 1\_base = 0, 1\_max = max\_level = numLevels-1.

Set parameters of the solve.

- pre is the number of smoothings before averaging.
- post is the number of smoothings after averaging.
- bottom is the number of smoothings at the bottom level.
- numMG = 1 for v-cycle, 2 for w-cycle and so on (in most cases, use 1).
- itermax is the max number of v cycles.
- hang is the minimum amount of change per vcycle.
- eps is the solution tolerance.
- normThresh is how close to zero eps\*resid is allowed to get.

## **4.5** The MultilevelLinearOp<T> class

The MultilevelLinearOp<T> class is a LinearOperator<Vector<LevelData<T>\* > > designed to support multilevel composite operators. This is useful when using a LinearSolver such as BiCGStabSolver to solve elliptic equations over a hierarchy of AMR levels. The MultilevelLinearOp is derived from LinearOp<Vector<LevelData<T>\* > >. Defining a MultilevelLinearOp requires an AMRLevelOpFactory<LevelData<T> > > which is used to define AMRLevelOps for each AMR level, which are then used to evaluate the multilevel operator, etc. An example which uses the MultilevelLinearOp<FArrayBox> in conjunction with the VCAMRPoissonOp to solve the variable-coefficient Helmholtz equation using multigrid-preconditioned BiCGStab is in

 ${\tt Chombo/example/AMRPoisson/variableCoefficientExec/VCPoissonSolve.cpp}$ 

#### Public member functions and member data:

- MultilevelLinearOp()
  - default constructor leaves object in undefined state
- void define(const Vector<DisjointBoxLayout>& vectGrids,
   const Vector<int>& refRatios,
   const Vector<ProblemDomain>& domains,
   const Vector<RealVect>& vectDx,
   RefCountedPtr<AMRLevelOpFactory<LevelData<T> > >& opFactory,
   int lBase)
  - full define function
    - vectGrids AMR hierarchy of grids
    - refRatios refinement ratios; refRatios[0] is refinement ratio between levels
       0 and 1.
    - domains problem domains for each AMR level.
    - vectDx cell-spacing on each AMR level
    - opFactory AMRLevelOpFactory used to define operators for performing multilevel linear solves.
    - 1Base base level
- - compute residual = L(phi) rhs
    - 1hs residual

- phi current approximation to the solution
- rhs rhs when solving L(phi) = rhs
- homogeneous if true, evaluate using homogeneous form of physical domain boundary conditions
- - Apply preconditioner to problem which is already in residual-correction form. If
    m\_use\_multigrid\_preconditioner is true (default case), then the preconditioner is m\_num\_mg\_iterations AMR V-cycles using an AMRMultiGrid solver.
    Otherwise, use whatever preconditioner is provided by the AMRLevelOp<LevelData<T> >.
    - cor correction (modified by the preconditioner)
    - residual residual (= L(phi) rhs)
- - Evaluate the operator, setting lhs = L(phi) using the AMRLevelOp<T>::AMROperator functions. If homogeneous is true, evaluate the operator using homogeneous physical domain boundary conditions.
- - Create data holder 1hs that mirrors rhs, using the appropriate AMRLevelOp::create functionality. Does not copy the data of rhs, just makes a holder the same size.
- virtual void clear(Vector<LevelData<T>\* >& lhs) const
  - Clear memory in data holder lhs. Note that the MultilevelLinearOp class requires this function be implemented because the create function calls new when allocating the Vector<LevelData<T>\* >.
- - Set 1hs equal to rhs.
- - Compute and return the volume-weighted AMR dot product of a\_1 and a\_2. Does this by calling the AMRLevelOp dotProduct functions for each AMR level and then scaling appropriately.

- virtual void incr(Vector<LevelData<T>\* >& lhs, const Vector<LevelData<T>\* >& x, Real scale)
  - Increment by scaled amount (lhs += scale\*x)
- - Set input lhs to a scaled sum (lhs = a\*x + b\*y).
- virtual void scale(Vector<LevelData<T>\* >& lhs, const Real& scale)
  - Multiply the input by a given scale (lhs \*= scale).
- virtual Real norm(const Vector<<LevelData<T>\* >& rhs, int ord)
  - Return the AMR norm of rhs (only counts valid regions for each level).
    - ord norm type: 0 is max norm, 1 is  $L_1$  norm sum(abs(rhs)), otherwise,  $L_{ord}$  norm.
- virtual void setToZero(Vector<LevelData<T>\* >& lhs)
  - Set Ihs to zero.
- bool m\_use\_multigrid\_preconditioner
  - if true (default value), use AMRMultiGrid multigrid V-cycles for preconditioner
- int m\_num\_mg\_iterations
  - number of multigrid v-cycles to do in preconditioner (only relevant if m\_use\_multigrid\_preconditioner is true).
- int m\_num\_mg\_smooth
  - parameter for AMRMultiGrid number of smoothing passes for each multigrid relaxation step. (only relevant if m\_use\_multigrid\_preconditioner is true).

## 4.6 Elliptic Examples

We provide several examples of elliptic operators that conform to the AMRLevelOp interface

- AMRPoissonOp is used to solve Poisson's equation with constant coefficients.
- ResistivityOp is used to solve the variable-coefficient resistivity equations that arise from MHD.
- ViscousTensorOp is used to solve the variable-coefficient elliptic equations that arise when solving the compressible Navier-Stokes equations with variable viscosity.

All of these examples use the boundary condition interface described in section 4.6.4.

#### 4.6.1 AMRPoisson

We provide an AMRPoisson, an example of an AMRLevelOp class. This class is designed to solve

$$(\alpha I + \beta \Delta)\phi = \rho \tag{4.1}$$

where  $\alpha$  and  $\beta$  are constants. The discretization of the Laplacian is the standard centered-difference approximation

$$(\Delta^h \phi)_{\boldsymbol{i}} = \frac{1}{h^2} \sum_{d=0}^{\mathbf{D}-1} (\phi_{\boldsymbol{i}+\boldsymbol{e}^d} + \phi_{\boldsymbol{i}-\boldsymbol{e}^d} - 2\phi_{\boldsymbol{i}})$$

Domain boundary conditions are enforced by setting ghost cell values as described in section 4.6.4. Fluxes through coarse-fine interfaces are needed for the Martin-Cartwright algorithm. The flux F through a face at  $i + \frac{1}{2}e^d$  is given by

$$F_{\boldsymbol{i}+\frac{1}{2}\boldsymbol{e}^d} = \frac{\beta}{h}(\phi_{\boldsymbol{i}+\boldsymbol{e}^d} - \phi_{\boldsymbol{i}})$$

## 4.6.1.1 AMRPoisson Factory Interface

An AMRPoissonOpFactory needs to be defined using the following function.

- coarseDomain is the domain at the coarsest level.
- grids is the AMR hierarchy.

- refRatios are the refinement ratios between levels. The ratio lives with the coarser level so refRatios [4] is the ratio between AMR levels 4 and 5.
- coarseDx is the grid spacing at the coarsest level.
- BCFunc holds the boundary conditions.
- alpha is the coefficient of the identity.
- beta is the coefficient of the Laplacian.

## 4.6.1.2 Code fragment

For those of us who find code easier to read than documents, we provide a simplified example of how to use AMRMultiGrid and AMRPoissonOp. In the example below, we start with a known AMR hierarchy and a right-hand side and we solve (4.1). For a description of the boundary condition routine along with its code fragment, see subsection 4.6.4. Complete examples (along with convergence tests) can be found in Chombo/example/AMRPoisson.

```
solveElliptic solves (alpha I + beta Laplacian) phi = rhs
  using AMRMultiGrid and AMRPoissonOp
  Inputs:
  rhs: Right-hand side of the solve over the level.
  grids: AMRHierarchy of grids
  refRatio: refinement ratios
  levelODomain: domain at the coarsest AMR level
  coarsestDx: grid spacing at the coarsest level
  alpha: identity coefficient
  beta: Laplacian coefficient
  phi = (alpha I + beta Lapl)^{-1}(rhs)
void solveElliptic( Vector<LevelData<FArrayBox>* >&
                   const Vector<LevelData<FArrayBox>* > rhs,
                   const Vector<DisjointBoxLayout>&
                   const Vector<int>&
                                                        refRatios.
                   const ProblemDomain&
                                                        levelODomain,
                   Real alpha, Real beta, Real coarsestDx)
 int numlevels = rhs.size();
 //define the operator factory
 AMRPoissonOpFactory opFactory;
 opFactory.define(levelODomain,
                  grids, refRatios, coarsestDx,
                  &ParseBC, alpha, beta);
 //this is the solver we shall use
 AMRMultiGrid<LevelData<FArrayBox> > solver;
 //this is the solver for the bottom of the muligrid v-cycle
 BiCGStabSolver<LevelData<FArrayBox> > bottomSolver;
 //bicgstab can be noisy
```

```
bottomSolver.m_verbosity = 0;

//define the solver
solver.define(levelODomain, opFactory, &bottomSolver, numlevels);

//we want to solve over the whole hierarchy
int lbase = 0;
//so solve already.
solver.solve(phi, rhs, numlevels-1, lbase);
```

## 4.6.2 ResistivityOp

ResistivityOp is the AMRLevelOp-derived class which solves the variable-coefficient equation For notation's sake, what we are solving is

$$L\vec{B} = \alpha \vec{B} + \beta \nabla \cdot F = \rho.$$

 $\alpha$  and  $\beta$  are constants and F is given by

$$F = \eta(\nabla \vec{B} - \nabla \vec{B}^T + I\nabla \cdot \vec{B})$$

where I is the identity matrix and  $\eta = \eta(\vec{x}) \ge 0$ . The discretization of the flux divergence is as follows.

$$(\nabla \cdot F)_{\boldsymbol{i}}^{h} = \frac{1}{h} \sum_{d=1}^{D} (F_{\boldsymbol{i} + \frac{1}{2} \boldsymbol{e}^{d}} F \phi_{\boldsymbol{i} - \frac{1}{2} \boldsymbol{e}^{d}})$$

We discretize normal components of the face-centered gradient using an average of cell-centered gradients for tangential components and a centered-difference approximation to the normal gradient.

$$(\nabla \vec{B})_{\mathbf{i}+\frac{1}{2}\mathbf{e}^d}^{d'} = \begin{pmatrix} \frac{1}{h}(\vec{B}_{\mathbf{i}+\mathbf{e}^d} - \vec{B}_{\mathbf{i}}) & \text{if } d = d' \\ \frac{1}{2}((\nabla \vec{B})_{\mathbf{i}+\mathbf{e}^d}^{d'} + (\nabla \vec{B})_{\mathbf{i}}^{d'}) & \text{if } d \neq d' \end{pmatrix}$$

where

$$(\nabla \vec{B})_{i}^{d} = \frac{1}{2h} (\vec{B}_{i+e^d} - \vec{B}_{i-e^d}).$$

## 4.6.2.1 ResistivityOp Factory Interface

An ResistivityOpFactory needs to be defined using the following constructor.

```
ResistivityOpFactory(const Vector<DisjointBoxLayout>&
                                                                             grids,
                      const Vector<RefCountedPtr<LevelData<FluxBox> > &
                                                                             eta,
                     Real
                                                                             alpha,
                     Real
                                                                             beta,
                      const Vector<int>&
                                                                             refRatios
                      const ProblemDomain&
                                                                             domainCoa
                      const Real&
                                                                             dxCoar,
                     BCFunc
                                                                             bc);
```

- domainCoar is the domain at the coarsest level.
- grids is the AMR hierarchy.
- refRatios are the refinement ratios between levels. The ratio lives with the coarser level so refRatios [4] is the ratio between AMR levels 4 and 5.
- dxCoar is the grid spacing at the coarsest level.
- bc holds the boundary conditions.
- alpha is the coefficient of the identity.
- beta is the coefficient of the divergence of the flux.
- eta is the variable coeff function. This ought to to be positive if you expect multigrid to converge.

## 4.6.3 ViscousTensorOp

ViscousTensorOp is the AMRLevelOp-derived class which solves the variable-coefficient equation For notation's sake, what we are solving is

$$L\vec{B} = \alpha \vec{B} + \beta \nabla \cdot F = \rho.$$

 $\alpha = \alpha(\vec{x})$  and  $\beta = \beta(\vec{x})$ . F is given by

$$F = \eta(\nabla \vec{B} + \nabla \vec{B}^T) + \lambda(I\nabla \cdot \vec{B})$$

where I is the identity matrix  $\eta = \eta(\vec{x})$ , and  $\lambda = \lambda(\vec{x})$ . The discretization of the flux divergence is as follows.

$$(\nabla \cdot F)_{i}^{h} = \frac{1}{h} \sum_{d=1}^{D} (F_{i+\frac{1}{2}e^{d}} - F_{i-\frac{1}{2}e^{d}})$$

We discretize normal components of the face-centered gradient using an average of cell-centered gradients for tangential components and a centered-difference approximation to the normal gradient.

$$(\nabla \vec{B})_{i+\frac{1}{2}e^d}^{d'} = \begin{pmatrix} \frac{1}{h}(\vec{B}_{i+e^d} - \vec{B}_i) \text{ if } d = d' \\ \frac{1}{2}((\nabla \vec{B})_{i+e^d}^{d'} + (\nabla \vec{B})_i^{d'}) \text{ if } d \neq d' \end{pmatrix}$$

where

$$(\nabla \vec{B})_{i}^{d} = \frac{1}{2h} (\vec{B}_{i+e^d} - \vec{B}_{i-e^d}).$$

## 4.6.3.1 ViscousTensorOp Factory Interface

An ViscousTensorOpFactory needs to be defined using the following constructor.

- domainCoar is the domain at the coarsest level.
- grids is the AMR hierarchy.
- refRatios are the refinement ratios between levels. The ratio lives with the coarser level so refRatios [4] is the ratio between AMR levels 4 and 5.
- dxCoar is the grid spacing at the coarsest level.
- bc holds the boundary conditions.
- alpha is the coefficient of the identity.
- beta is the coefficient of the divergence of the flux.
- eta is the variable coeff function that multiplies the gradient.
- lambda is the variable coeff function that multiplies the divergence.

## 4.6.4 Boundary Condition Interface

The value of the boundary condition is described by the BCValueFunc function interface.

The boundary condition function must conform to the BCFunc specification. We provide both Dirichlet and Neumann boundary condition examples.

A simple example of a custom boundary condition is given below.

```
/* this is the bc value func */
void ParseValue(Real* pos,
                 int* dir.
                 Side::LoHiSide* side,
                 Real* values)
  ParmParse pp;
  Real bcVal;
  pp.get("bc_value",bcVal);
  values[0]=bcVal;
/*Use ParmParse to select boundary condtions */
/* this is the bcfunc */
void ParseBC(FArrayBox& state,
              const Box& valid,
              const ProblemDomain& domain,
              Real dx,
              bool homogeneous)
  if(!domain.domainBox().contains(state.box()))
      std::vector<int> bcLo, bcHi;
      ParmParse pp;
      pp.getarr("bc_lo", bcLo, 0, SpaceDim);
      pp.getarr("bc_hi", bcHi, 0, SpaceDim);
      Box valid = valid;
      for(int i=0; i<CH_SPACEDIM; ++i)</pre>
          Box ghostBoxLo = adjCellBox(valid, i, Side::Lo, 1);
Box ghostBoxHi = adjCellBox(valid, i, Side::Hi, 1);
          if(!domain.domainBox().contains(ghostBoxLo))
             {
               if(bcLo[i] == 1)
                   pout() << "const neum bcs lo for direction " << i << endl;</pre>
                   NeumBC(state,
                           valid,
                           dx,
                           homogeneous,
                           ParseValue,
                           Side::Lo);
                 }
               else if(bcLo[i] == 0)
                   pout() << "const diri bcs lo for direction " << i << endl;</pre>
                   DiriBC(state,
                           valid.
                           dx,
```

```
homogeneous,
                      ParseValue,
                      Side::Lo);
             }
          else
               MayDay::Error("bogus bc flag lo");
        }
      if(!domain.domainBox().contains(ghostBoxHi))
          if(bcHi[i] == 1)
               pout() << "const neum bcs hi for direction " << i << endl;</pre>
               NeumBC(state,
                      valid.
                      dx.
                      homogeneous
                      ParseValue,
                      Side::Hi);
          else if(bcHi[i] == 0)
               pout() << "const diri bcs hi for direction " << i << endl;</pre>
               DiriBC(state,
                      valid,
                      dx,
                      homogeneous,
                      ParseValue,
                      Side::Hi);
             }
          else
               MayDay::Error("bogus bc flag hi");
        }
   }
}
```

## 4.7 Parabolic Equations – the TGA scheme

In this section, we describe our approach to solving parabolic equations of the form:

$$\frac{\partial \phi}{\partial t} = L(\phi) + S,\tag{4.2}$$

where L is a second-order linear elliptic operator, and S is a source term.

To evolve  $\phi$  in time from time  $t^n$  to time  $t^{n+1}=t^n+\Delta t$ , we use a variant of the  $L_0$ -stable Runge-Kutta scheme presented by Twizell, Gumel, and Arigu (TGA) [30], which is second-order in time. Our variation is based on time-centering the source term S at time  $t^{n+\frac{1}{2}}=t^n+\frac{\Delta t}{2}$ , and is also described in [22].

Following [30], we discretize (4.2) in time:

$$\phi^{n+1} = (I - \mu_1 L)^{-1} (I - \mu_2 L)^{-1} \left[ (I + \mu_3 L) \phi^n + \Delta t (I + \mu_4 L) S^{n + \frac{1}{2}} \right], \tag{4.3}$$

where  $\phi^n = \phi(n\Delta t)$ ,  $S^{n+\frac{1}{2}} = S((n+\frac{1}{2})\Delta t)$ , and the coefficients  $\mu_1, \mu_2, \mu_3, \mu_4$  are the values suggested in [30]:

$$\mu_1 = \frac{2a - 1}{a + discr} \Delta t,$$

$$\mu_2 = \frac{2a - 1}{a - discr} \Delta t,$$

$$\mu_3 = (1 - a) \Delta t,$$

$$\mu_4 = (\frac{1}{2} - a) \Delta t$$

$$a = 2 - \sqrt{2} - \epsilon,$$

$$discr = \sqrt{a^2 - 4a + 2},$$

where  $\epsilon$  is a small quantity (we use  $10^{-8}$ ). We use this to define the operator  $L^{TGA}(\phi^n,S^{n+\frac{1}{2}})$  as follows:

$$L^{TGA}(\phi^n, S^{n+\frac{1}{2}}) \equiv \frac{\phi^{n+1} - \phi^n}{\Delta t} - S^{n+\frac{1}{2}}$$

$$\approx (L\phi)\left((n + \frac{1}{2})\Delta t\right) + O(\Delta t^2).$$
(4.4)

where  $\phi^{n+1} = \phi^{n+1}(\phi^n, S^{n+\frac{1}{2}})$  is defined to be the expression (4.3).

To simplify the use of the TGA scheme, Chombo includes two classes which implement  $L^{TGA}$  — one for a single AMR level, and one for an entire AMR hierarchy. Note that computing  $L^{TGA}$  requires solving the heat equation and applying the operator L, these classes require that there be suitable AMRLevelOp-derived operator classes which discretize the appropriate Helmholtz operator.

## 4.7.1 The TGAHelmOp and LevelTGAHelmOp classes

Since implementing the TGA algorithm requires some additional functionality in addition to that specified in the AMRLevelOp<T> class, we have the TGAHelmOp<T> and LevelTGAHelmOp<T, TFlux> classes, which are general Helmholtz-type operators like  $(\alpha + \nabla \cdot \beta \nabla)$  publicly derived from AMRLevelOp<T>. The multilevel TGA implementation uses TGAHelmOp. The single-level TGA implementation requires two functions not needed by the multilevel solve, so it uses LevelTGAHelmOp<T,TFlux>, which is publicly derived from TGAHelmOp<T> and includes the extra required functions. The AMRPoissonOp, ResistivityOp, and ViscousTensorOp classes are derived from LevelTGAHelmOp, and so may be used by both of the TGA solvers.

Additional functions required by TGAHelmOp<T> and LevelTGAHelmOp<T,TFlux>:

- virtual void setAlphaAndBeta(const Real& a\_alpha, const Real& a\_beta) = 0
  - Set the Helmholtz equation constants in the operator.
- virtual void diagonalScale(T& a\_rhs) = 0
   virtual void diagonalScale(T& a\_rhs, bool a\_kappaWeighted) = 0
  - Set the diagonal scaling of the operator. For example, if solving  $\rho(x)\frac{\partial \phi}{\partial t} = L(\phi)$ , the diagonal Scale would be  $\rho$ . In EB applications, even for constant coefficients, it means multiplication by  $\kappa$ .
- virtual void applyOpNoBoundary(T& a\_ans, const T& a\_phi) = 0
  - Apply operator without any boundary or coarse-fine boundary conditions and no finer level. This implies that we've set ghost-cell values outside the operator and want to use them in the operator evaluation.

#### Additional functions required by LevelTGAHelmOp<T,TFlux>:

- virtual void fillGrad(const T& a\_phi)
  - This a function used in operators of fairly complex flux functions in which the gradient of the solution is computed and stored separately. This function is to signal the operator to do said computation with the input data. The default implementation does nothing.

- compute face-centered flux.
  - flux face-centered dataholder into which to place flux
  - data cell-centered data used to compute the flux
  - grid cell-centered box over which to compute face-centered flux.
  - dit DataIndex of grid box we're working with
  - scale scaling factor to multiply flux.

## 4.7.2 The AMRTGA class

The AMRTGA<T> class is a templated implementation of the TGA algorithm designed to advance an entire multilevel hierarchy of AMR grids by one (non-subcycled) timestep where T is the data holder for a single AMR level (like a LevelData<FArrayBox>.

#### Public member functions:

#### - constructor

- solver AMRMultiGrid solver used to do the elliptic solves. This should be predefined with the appropriate TGAHelmOp operators.
- factory Factory which can be used to create TGAHelmOps for this problem.
- levelODomain problem domain on the coarsest level
- refRat vector of refinement ratios
- numLevels number of AMR levels
- verbosity how much output is written out. Higher number is more verbose (default is 0).

```
    void oneStep(Vector<T*>& a_phiNew,
    Vector<T*>& a_phiOld,
    Vector<T*>& a_source,
    const Real& a_dt,
    int a_lbase,
    int a_lmax)
```

- advances a parabolic PDE one timestep using TGA on a non-moving domain.
  - phiNew new-time solution computed using TGA.
  - phi01d old-time solution
  - source source term at the half-time
  - dt timestep
  - 1base coarsest level to be advanced
  - lmax finest level to be advanced

#### 4.7.3 The BaseLevelTGA and LevelTGA classes

The BaseLevelTGA<T, TFlux, TFR> class is a pure-virtual base class which implements the basic TGA scheme on a DisjointBoxLayout, which corresponds to solving on a single AMR level. The BaseLevelTGA class is templated on T, a (cell-centered) data holder over the entire DisjointBoxLayout (which is the same template type T used by the AMRLevelOp-derived operator, and which is the datatype of the solution on a single

level), TF1ux, a face-centered data holder on a single patch, and TFR, the appropriate FluxRegister type of object to store diffusive fluxes along any coarse-fine interfaces.

The simplest example of a BaseLevelTGA-derived class is the LevelTGA class, which is a publicly-derived BaseLevelTGA<LevelData<FArrayBox>, FluxBox, LevelFluxRegister>. Note that the BaseLevelTGA objects are defined over the entire AMR hierarchy at once, and then advance a single-level as specified.

#### Public member functions:

#### - constructor

- grids grids over the entire AMR hierarchy
- refRat refinement ratios
- level0domain problem domain on coarsest level
- opFact factory used to create LevelTGAHelmOps for this problem.
- solver AMR Multigrid solver over entire AMR hierarchy, pre-defined with the appropriate LevelTGAHelmOp-derived operators.

```
• void updateSoln(T&
                                       a_phiNew,
                   T&
                                       a_phi0ld,
                   T&
                                       a_src,
                  LevelData<TFlux>& a_flux,
                  TFR*
                                       a_FineFluxRegPtr,
                   TFR*
                                       a_CrseFluxRegPtr,
                   const T*
                                       a_crsePhiOldPtr,
                   const T*
                                       a_crsePhiNewPtr,
                   Real
                                       oldTime,
                   Real
                                       crseOldTime,
                   Real
                                       crseNewTime,
                   Real
                                       dt,
                   int
                                       a_level,
                   bool
                                       a_zeroPhi = true,
                                       a_fluxStartComponent = 0);
                   int
```

- update ALL components of phi
  - phiNew updated solution at oldTime + dt
  - phi01d solution at oldTime

- src source term at (oldTime + 0.5\*dt)
- flux -
- FineFluxRegPtr flux register for diffusive fluxes along coarse-fine interface between this level (level) and the next-finer level (level+1).
- CrseFluxRegPtr flux register for diffusive fluxes along coarse-fine interface between this level (level) and the next-coarser level (level-1).
- crsePhiOldPtr pointer to old-time coarse data (for coarse-fine boundary conditions)
- crsePhiNewPtr pointer to new-time coarse data (for coarse-fine boundary conditions)
- oldTime time centering of phiOld
- crseOldTime time centering of crsePhiOldPtr
- crseNewTime time centering of crsePhiNewPtr
- dt timestep
- level AMR level which we're advancing
- zeroPhi -
- fluxStartComponent component at which to place first component of flux

```
• void computeDiffusion(T&
                                            a_DiffusiveTerm,
                                            a_phi0ld,
                         T&
                                            a_src,
                         LevelData<TFlux>& a_flux,
                         TFR*
                                            a_FineFluxRegPtr,
                         TFR*
                                            a_crseFluxRegPtr,
                         const T*
                                            a_crsePhiOldPtr,
                         const T*
                                            a_crsePhiNewPtr,
                         Real
                                            a_oldTime,
                         Real
                                            a_crseOldTime,
                         Real
                                            a_crseNewTime,
                         Real
                                            a_dt,
                                            a_level);
                         int
```

- compute time-centered  $L^{TGA}(\phi,S)$  for use in subsequent update operations. In this case, we do solve for phiNew, then subtract source and phiOld back out to get L(phi). Function arguments have the same meanings as for the updateSoln function.
  - DiffusiveTerm  ${\cal L}^{TGA}(\phi,S)$  as computed by this function.

# 4.8 TGA addendum for EB and non-unity identity coefficients

Consider the partial differential equation

$$a\frac{\partial \phi}{\partial t} = L\phi + S$$

$$a = a(x), \quad a > 0.$$
(4.5)

We can extend the TGA algorithm by defining

$$L_a \phi \equiv \frac{L\phi}{a}$$

$$S_a \phi \equiv \frac{S}{a}$$

The solution is given by

$$(I - \mu_1 L_a)(I - \mu_2 L_a)\phi^{n+1} = (I + \mu_3 L_a)\phi^n + \Delta t(I + \mu_4 L_a)S_a$$

Because a (though non-zero), can be quite small, we are reluctant to have it in the denominator. We use the handy identity  $(AB)^{-1}=B^{-1}A^{-1}$  to get

$$(I-L)^{-1} = (aI - aL)^{-1}(a)$$
(4.6)

to derive

$$\phi^{n+1} = (aI - \mu_1 L)^{-1} (a(aI - \mu_2 L)^{-1} [(aI + \mu_3 L)\phi^n + \Delta t(aI + \mu_4 L)\frac{S}{a}]).$$
 (4.7)

Note that we still have to divide the source term by the coefficient but, since this is typically density, it should be floored reasonably above zero.

Finally, with embedded boundaries, we must also multiply out factors of the volume fraction  $\kappa$ . L is divided by  $\kappa$  and  $\kappa$  can be arbitrarily small. We can reuse the above identity to obtain the following.

$$\phi^{n+1} = (\kappa aI - \kappa \mu_1 L)^{-1} \kappa a \left[ (\kappa aI - \kappa \mu_2 L)^{-1} \left[ (\kappa aI + \kappa \mu_3 L) \phi^n + \Delta t (\kappa aI + \kappa \mu_4 L) \frac{S}{a} \right] \right].$$

Mercifully, we never have to divide by the volume fraction.

To accommodate these variable coefficient issues LevelTGAHelmOp needs some extra functions.

virtual void diagonalScale(T& a\_rhs, bool a\_kappaWeighted)
 This function multiplies in place the input by the identity coefficient. In the case of EB, setting a\_kappaWeighting to true means the input gets multiplied by κa.

virtual void diagonalScale(T& a\_rhs)

This version of diagonal Scale assumes that  $\kappa$  weighting is desired in EB applications.

virtual void divideByIdentityCoef(T& a\_S)
 This function divides the input in place by the a coefficient.

# 4.9 TGA with time-dependent operator coefficients

Consider a parabolic partial differential equation in which a quantity  $a\phi$  is conserved and in both a and  $\phi$  are time dependent:

$$\frac{\partial(a\phi)}{\partial t} = L\phi + S$$

$$a = a(\vec{x}, t), \quad a > 0.$$
(4.8)

For the purposes of this discussion, we assume that a can be updated explicitly, and that  $a^n$  and  $a^{n+1}$  are known and may be used in the solution of (4.8) to obtain  $\phi^{n+1}$  from  $\phi^n$ .

Equations of this form arise in the diffusion of energy and momentum in compressible hydrodynamic flows. In fluid flows with heat conduction, for example, the energy per unit volume E is related to both the mass density  $\rho$  and the temperature T by the expression

$$E = c_n \rho T \tag{4.9}$$

where  $c_p$  is the specific heat as measured under constant pressure. If we assume that  $c_p$  is constant within a material, the transfer of energy by heat conduction throughout a body with thermal conductivity K is described by the conservation equation

$$c_p \frac{\partial(\rho T)}{\partial t} = \nabla \cdot (K \nabla T) + S \tag{4.10}$$

where S is an external source of energy. The total energy is conserved in this process.

# 4.9.1 Implicit TGA update for $\phi$

We begin by expressing (4.8) in a form that more closely resembles (4.5). The time change in  $a\phi$  is

$$\frac{\partial(a\phi)}{\partial t} = \left[ \left( \frac{\partial a}{\partial t} \right) \phi + a \left( \frac{\partial \phi}{\partial t} \right) \right]. \tag{4.11}$$

Thus, we can express the evolution equation for  $\phi$  as

$$a\frac{\partial\phi}{\partial t} = L(\phi) + S - \left(\frac{\partial a}{\partial t}\right)\phi \tag{4.12}$$

Since we have explicitly updated a and have  $a^{n+1}$  as well as  $a^n$ , we may explicitly estimate the second term on the right hand side of (4.12) as a source using the first-order finite difference

$$\left(\frac{\partial a}{\partial t}\right)\phi \approx \frac{(a^{n+1} - a^n)}{\Delta t}\phi^n. \tag{4.13}$$

Because this expression appears on the right hand side of (4.12), it is then multiplied by a factor of  $\Delta t$  and rendered second-order accurate.

In order to conserve  $a\phi$ , it is crucial that we use values of a at the correct time centerings in the TGA update for  $\phi$ . We define

$$L_a(t) \equiv \frac{L(\phi, t)}{a(t)} \tag{4.14}$$

$$S_a(t) \equiv \frac{S(t) - \phi^n (a^{n+1} - a^n)/\Delta t}{a(t)}$$
 (4.15)

where we have added t as a parameter to all time-dependent quantities to emphasize their time dependence. The TGA update for  $\phi$  is then written

$$\phi^{n+1} = [I - \mu_1 L_a(t^{n+1})]^{-1} [I - \mu_2 L_a(t^*)]^{-1} \times \left( [I + \mu_3 L_a(t^n)] \phi^n + \Delta t [I + \mu_4 L_a(t^{n+\frac{1}{2}})] S_a(t^{n+\frac{1}{2}}) \right).$$
(4.16)

where  $t^*$  is the "intermediate" time used in the TGA update. Hereafter, we use abbreviated superscripts on a, L, and S to refer to their time centerings.

Using the identity (4.6) and multiplying the last two terms by  $a^n/a^n$  and  $a^{n+\frac{1}{2}}/a^{n+\frac{1}{2}}$  respectively, we obtain the expression

$$\phi^{n+1} = \left[ a^{n+1}I - \mu_1 L^{n+1} \right]^{-1} a^{n+1} \left[ a^*I - \mu_2 L^* \right]^{-1} a^* \times \left( \frac{1}{a^n} \left[ a^n I + \mu_3 L^n \right] \phi^n + \frac{\Delta t}{a^{n+\frac{1}{2}}} \left[ a^{n+\frac{1}{2}} I + \mu_4 L^{n+\frac{1}{2}} \right] \left[ \frac{S^{n+\frac{1}{2}} - \phi^n (a^{n+1} - a^n)/\Delta t}{a^{n+\frac{1}{2}}} \right] \right).$$
(4.17)

This expression is more complicated than (4.7) because the various factors of a are evaluated at different times and thus do not cancel.

#### 4.9.1.1 Embedded boundary considerations

The same analysis can be carried out in the presence of irregular cells near embedded boundaries, but since the volume fraction  $\kappa$  is constant in time, the same factors appear as in (4.7). In the Chombo implementation of this algorithm, factors of a and L in the numerator each include a factor of  $\kappa$ , but factors of a in the denominator do not, so we must take special care in computing  $\phi^{n+1}$ .

To illustrate how these factors must be treated in Chombo, we include all factors of  $\kappa$  instead of accounting for cancellations. In this case the expression for  $\phi^{n+1}$  within irregular cells with the appropriate factors of  $\kappa$  is

$$\phi^{n+1} = \left[ \kappa a^{n+1} I - \mu_1 \kappa L^{n+1} \right]^{-1} \kappa a^{n+1} \left[ \kappa a^* I - \mu_2 \kappa L^* \right]^{-1} \kappa a^* \times \left( 4.18 \right) \left( \frac{1}{\kappa a^n} \left[ \kappa a^n I + \mu_3 \kappa L^n \right] \phi^n + \frac{\Delta t}{\kappa a^{n+\frac{1}{2}}} \left[ \kappa a^{n+\frac{1}{2}} I + \mu_4 \kappa L^{n+\frac{1}{2}} \right] \left[ \frac{S^{n+\frac{1}{2}} - \phi^n (a^{n+1} - a^n)/\Delta t}{a^{n+\frac{1}{2}}} \right] \right).$$

The factors of  $\kappa$  in the denominator cancel with the factor, say, in front of  $a^*$ . We can implement this cancellation by calling diagonalScale with a a\_kappaWeighted=false to indicate that  $a^*$  should *not* be multiplied by  $\kappa$ . Our final expression for  $\phi^{n+1}$  for operators with time-dependent coefficients in irregular cells is

$$\phi^{n+1} = \left[ \kappa a^{n+1} I - \mu_1 \kappa L^{n+1} \right]^{-1} \kappa a^{n+1} \left[ \kappa a^* I - \mu_2 \kappa L^* \right]^{-1} a^* \times \left( 4.19 \right)$$

$$\left( \frac{1}{a^n} \left[ \kappa a^n I + \mu_3 \kappa L^n \right] \phi^n + \frac{\Delta t}{a^{n+\frac{1}{2}}} \left[ \kappa a^{n+\frac{1}{2}} I + \mu_4 \kappa L^{n+\frac{1}{2}} \right] \left[ \frac{S^{n+\frac{1}{2}} - \phi^n (a^{n+1} - a^n)/\Delta t}{a^{n+\frac{1}{2}}} \right] \right).$$

# **4.9.2** Conservative calculation of $L(\phi)$

After  $\phi^{n+1}$  is computed, we obtain a conservative value for  $L(\phi)$  by calculating

$$L(\phi) = \frac{(a^{n+1}\phi^{n+1} - a^n\phi^n)}{\Delta t} - S.$$
 (4.20)

## 4.9.3 Setting the time centering of a LevelTGAHelmOp

(4.17) is correct for any time-dependent entities a(t) and L(t), but we need a way to obtain these properly-centered values in practice. Since our algorithm is second-order accurate, we can compute a time-centered value for  $a^{n+\mu}$  or  $L^{n+\mu}$  by interpolating linearly between the beginning-of-step values  $a^n$ ,  $L^n$  and their end-of-step values  $a^{n+1}$ ,  $L^{n+1}$  if we know their values at  $t^{n+1}$  beforehand. The TGA algorithm sets the time centering of a LevelTGAHelmOp by calling the setTime method:

virtual void setTime(Real a\_oldTime, Real a\_mu, Real a\_dt)

Here, a\_oldTime is the beginning-of-step time  $t^n$ , a\_mu is the fraction of the time step that has elapsed, and a\_dt is the step size. With these three parameters, one can compute  $t^{n+\mu}$  and can interpolate a or L between  $t^n$  and  $t^{n+1}$  as needed. For example, the time-interpolated value for a at a fraction  $\mu$  through the step is

$$a^{n+\mu} = a^n + \mu \Delta t \left( \frac{a^{n+1} - a^n}{\Delta t} \right)$$
  
=  $(1 - \mu)a^n + \mu a^{n+1}$  (4.21)

#### Implementation details

Currently, the TGA solver assumes that a LevelTGAHelmOp is time independent unless otherwise specified. If you wish to create a time-dependent subclass of LevelTGAHelmOp, that subclass must call the single-argument LevelTGAHelmOp constructor that specifies that it is time-dependent:

LevelTGAHelmOp(bool a\_isTimeDependent)

This allows the TGA solver to query the operator via its isTimeDependent method:

bool isTimeDependent() const

Note that LevelTGAHelmOp::isTimeDependent is *not* a virtual method—it simply returns the flag passed to the above constructor. The time dependence of an operator object must be established at the time of its construction.

Additionally, if the operator needs to interpolate any of its coefficients over the time step, it should store the beginning-of-step and end-of-step values for such coefficients and ensure that they are valid before the TGA solve.

# Chapter 5

# **AMRTimeDependent**

# 5.1 Hyperbolic Systems of Conservation Laws

In this section, we will describe a general framework for solving time-dependent problems using AMR, including refinement in time. In order to motivate that framework, we first describe in detail the AMR algorithm in [7] for solving systems of hyperbolic conservation laws.

We want to solve a system of equations of the form

$$\frac{\partial V}{\partial t} + \nabla \cdot \vec{\mathcal{F}} = 0$$

$$V = V(\boldsymbol{x}, t) \in \mathbb{R}^{m}$$

$$\vec{\mathcal{F}} = (\mathcal{F}^{0}(V), ..., \mathcal{F}^{\mathbf{D}-1}(V))$$

$$\mathcal{F}^{d} : \mathbb{R}^{m} \to \mathbb{R}^{m}.$$

We assume that the system is hyperbolic, i.e., that the matrices  $\sum_{d=0}^{\mathbf{D}-1} \xi_d \nabla_v \mathcal{F}^d$  have real eigenvalues and a complete set of eigenvectors for all  $\xi \in \mathbb{R}^{\mathbf{D}}$ . If the system is hyperbolic, we expect that specifying initial conditions of the form

$$V(\boldsymbol{x},0) = \Psi(\boldsymbol{x})$$

leads to a well-posed problem.

A variety of multiple-scale phenomena arise in solutions to hyperbolic systems of conservation laws arising from continuum mechanics problems that make AMR an attractive option. These include dynamics of shocks and interfaces, shock-shock intersections, and nonlinear wave focusing.

We assume that the underlying discretization of the above hyperbolic system of equations on a uniform grid is an explicit finite-difference method in discrete conservation form:

$$U^{new} = U^{old} - \Delta t \ D\vec{F} \ \ {\rm on} \ \Gamma$$

where  $\vec{F}$  is a staggered-grid vector field on  $\Gamma$ , and D is the discrete divergence operator defined in section 3.1.2.3.  $\vec{F}$  is a function of  $U^{old}$ , with a finite domain of dependence:  $F_d(U^{old})_{i+\frac{1}{2}e^d}$  depends only on  $\{U^{old}_{i+s}\}_{|s-\frac{1}{2}e^d|\leq p}$  where p is independent of the mesh spacing.

We extend this method to an adaptive mesh hierarchy using the Berger–Oliger algorithm [6]. We define

$$\{U^l\}_{l=0}^{l_{max}}, \quad U^l: \Omega^l \to \mathbb{R}^m$$

where  $U^l=U^l(t^l)$ . Here  $\{t^l\}$  are a collection of discrete times that satisfy the temporal analogue of proper nesting;  $\{t^l\}=\{t^{l-1}+k\Delta t^l:0\leq k< n^l_{ref}\}$ . The algorithm in [7] for advancing the solution in time is given in pseudo-code in figure 5.1. The discrete fluxes  $\vec{F}$  are computed by using the piecewise linear interpolation function in section 3.1.1.3 to define an extended solution on

$$\tilde{\Omega} = \mathcal{G}(\Omega^l, p) \cap \Gamma^l,$$

namely  $\tilde{U}: \tilde{\Omega} \to \mathbb{R}^m$ , where

$$\begin{split} \tilde{U}_{\pmb{i}} &= \begin{cases} U_{\pmb{i}}^l(t^l) & \text{for } \pmb{i} \in \Omega^l; \\ I_{pwl}((1-\alpha)U^{l-1}(t^{l-1}) + \alpha \ U^{l-1}(t^{l-1} + \Delta t^{l-1}))_{\pmb{i}} & \text{otherwise} \end{cases} \end{split}$$
 where 
$$\alpha = \frac{t^l - t^{l-1}}{\Delta t^{l-1}}.$$

Regridding is performed in the following steps.

- 1. Construct  $\mathcal{I}^l \subset \Omega^l, l = l_{base}, ..., l_{max} 1$  corresponding to those cells for which a user-specified measure of the error exceeds a specified tolerance.
- 2. Generate new grids  $\Omega^{l,new}, l=l_{base}+1,...,l_{max}$  on which the new solution is to be defined. These new grids should satisfy  $\mathcal{C}_{n_{ref}^l}(\Omega^{l+1,new})\supset \mathcal{I}^l$ , and should be properly nested, as well as satisfying any other required nesting conditions. If  $l_{base}>0$ , these conditions impose some constraints on  $\mathcal{I}^l$ , which are met typically by reducing the size of the  $\mathcal{I}^l$ 's prior to the grid generation step.
- 3. Initialize the new data  $U^{l,new}(t^l)$ . For  $l=l_{base}, U^l=U^{l,new}$ . For  $l=l_{base}+1,...,l_{max}$ ,

$$U^{l,new}(t_{regrid})_{\boldsymbol{i}} = \begin{cases} U^{l,new}(t_{regrid})_{\boldsymbol{i}} = U^l(t_{regrid})_{\boldsymbol{i}} & \boldsymbol{i} \in \Omega^l \cap \Omega^{l,new}; \\ I_{pwl}(U^{l-1,new}(t_{regrid}))_{\boldsymbol{i}} & \text{otherwise}. \end{cases}$$

```
procedure advance (l) U^l(t^l+\Delta t^l)=U^l(t^l)-\Delta tD\vec{F}^l \text{ on } \Omega^l \text{ if } l < l_{max} \qquad \delta F_d^{l+1}=-F_d^l \text{ on } \zeta_{+,d}^{l+1}\cup \zeta_{-,d}^{l+1}, d=0,...,\mathbf{D}-1 \text{ end if } \\ \text{if } l>0 \qquad \delta F_d^l:=\delta F_d^l+\frac{1}{n_{ref}^{l-1}}\langle F_d^l\rangle \text{ on } \zeta_{+,d}^l\cup \zeta_{-,d}^l, d=0,...,\mathbf{D}-1 \text{ end if } \\ \text{for } q=0,...,n_{ref}^l-1 \qquad \text{advance}(l+1) \\ \text{end for } \\ U^l(t^l+\Delta t^l)=Average(U^{l+1}(t^l+\Delta t^l),n_{ref}^l) \text{ on } \mathcal{C}_{n_{ref}^l}(\Omega^{l+1}) \\ U^l(t^l+\Delta t^l):=U^l(t^l+\Delta t^l)-\Delta t^lD_R(\delta F^{l+1}) \\ t^l:=t^l+\Delta t^l \\ n_{step}^l:=n_{step}^l+1 \\ \text{if } (n_{step}^l=0 \text{ mod } n_{regrid}) \text{ and } (n_{step}^{l-1}\neq 0 \text{ mod } n_{regrid}) \\ \text{ regrid}(l) \\ \text{end if} \\ \end{cases}
```

Figure 5.1: Pseudo-code description of the Berger–Colella AMR algorithm for hyperbolic conservation laws.

#### Refinement Criteria

Generally speaking, there are two approaches to determining which cells are to be tagged for refinement. One is to tag points at which some local function of the dependent variables or their derivatives exceeds some threshold. The second approach is to compute a local estimate of the truncation error, and tag points at which the magnitude of that error exceeds a given threshold. The first approach can be used very successfully in cases where the user can exploit application-specific information. The second approach is more general and more difficult to implement correctly. For that reason, we will discuss it in some detail here.

Let  $L^h(\varphi^h):\Gamma\to\mathbb{R}$  be a finite-difference approximation on a uniform grid to a differential operator  $\mathcal L$  defined for any  $\varphi^h:\Gamma\to\mathbb{R}^m$ . We define the truncation error for L to be

$$au_{m{i}}^h = L^h(\psi^h)_{m{i}} - \mathcal{L}(\psi)(m{x}_0 + m{i}h)$$

where  $\psi^h = \psi(\boldsymbol{x}_0 + \boldsymbol{i}h)$ , and  $\psi$  is a smooth function  $\psi: \mathbb{R}^{\mathbf{D}} \to \mathbb{R}^m$ . To compute the truncation error in a numerical solution to a system of PDE's,  $\psi$  would be the particular solution being computed. The difficulty is that we don't know the exact solution to the PDE. Instead, we can compute the Richardson estimate to the truncation error,

$$\tau_{\pmb{i}}^{R,2h} = (L^{2h}(Average(\varphi^h,2))_{\pmb{i}} - Average(L^h(\varphi^h),2)_{\pmb{i}})$$

for  $\pmb{i} \in \mathcal{C}_2(\Gamma)$ . Here  $\tau_{\pmb{i}}^{R,2h} = C \tau_{\pmb{i}}^h + O(h^{p+1})$  where p is the order of accuracy of the operator  $L: \tau_{\pmb{i}} = O(h^p)$ . It is straightforward to extend the definition of  $\tau^R$  to an AMR grid hierarchy. In that case, one can tag points to be refined based on whether  $|\tau_{\pmb{i}}^R|$  exceeds some threshold.

We can see two difficulties with this approach. The first is that finite-difference operators often have a lower-order accurate truncation error at the problem domain boundaries. In addition, the discussion in section 3.1 indicates that the truncation error is of lower-order accuracy at coarse-fine boundaries as well. However, as indicated previously, the effect of the truncation error at boundaries on the solution error is typically much smaller than the magnitude of the former would indicate. To compute an error estimator that appropriately reflects this fact, we rescale the tagging criterion at cells adjacent to the boundaries.

## **5.2 Classes** AMR and AMRLevel

The class AMR implements a framework for the Berger-Oliger adaptive mesh refinement (AMR) algorithm for time-dependent simulations [6]. The data is organized on a hierarchy of levels of refinement, stored as a collection of AMRLevels. The class AMRLevel is an abstract base class from which must be derived a concrete class which defines and contains the data representation for one level and implements the algorithms for advancing one

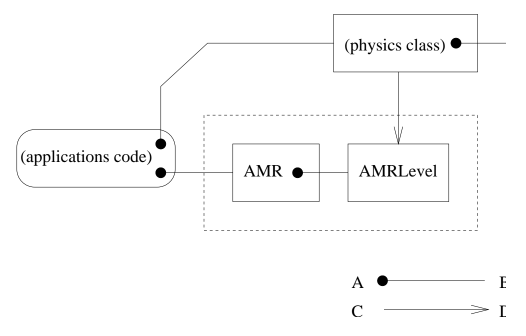


Figure 5.2: Class structure. AMR contains some AMRLevels. Physics class is derived from AMRLevel and contains the data representation.

level in time. The class AMR implements refinement of the time step  $\Delta t$  based on the refinement of the grid.

#### 5.2.1 Class structure

The class AMR manages the entire hierarchy of levels. The levels are represented in the class AMR as a collection of AMRLevels. The class AMRLevel is an abstract base class from which the applications implementer must derive a concrete *physics class* which defines the form of the solution data and, for a single level, implements algorithms for advancement by  $\Delta t$ , calculation of a stable  $\Delta t$ , initialization, input and output, etc. The class AMR has no knowledge of the form of the solution data. The applications implementer must instantiate an object of type AMR and an object of the physics class type. The physics class object is required input to the definition of an AMR. See figure 5.2.

Applications code constructs and invokes public member functions of the class AMR. The class AMR controls the entire hierarchy of levels. It constructs and invokes member functions of the AMRLevels. It requires a physics class derived from AMRLevelFactory as input.

#### **5.2.2 Class** AMR.

The AMR class is a framework for Berger-Oliger timestepping for adaptive mesh refinement of time-dependent problems. It is applicable to both hyperbolic and parabolic problems. It represents a hierarchy of levels of refinement as a collection of AMRLevels. The usage pattern of this class is as follows:

- Call define to define the parameters that do not change throughout the run (max level, refinement ratios, domain, and operator).
- Modify any parameters you like (blocking factor and so forth) using access functions.
- Call any one of the three setup functions so AMR can set up all its internal data structures.
- Call run to run the calculation.
- Call conclude to produce statistical output, e.g., how many cells were updated.

The important functions of the public interface for the AMR class are:

Defines this object. User must call a setup function before running.

#### **Arguments:**

- a\_max\_level (not modified): The maximum level number allowed, where the base level is zero. There may be a total of a\_max\_level+1 levels, since level zero and level a\_max\_level can both exist. Note that it while this is the maximum possible level, it is possible that fewer levels are actually defined, depending on the problem and the method and tolerances used to tag cells for refinement.
- ref\_ratios (not modified): Refinement ratios. There must be at least a\_max\_level+1 elements or an error will result. Element zero is the base level.
- a\_prob\_domain (not modified): Problem domain on the base level.

- a\_amrLevelFact (not modified): Pointer to a physics class factory object.
   The object it points to is used to construct the collection of AMRLevels in this AMR as objects of the physics class type.
- void setupForRestart(HDF5Handle& a\_handle);
   Sets up this object from checkpointed data. User must have previously called define. A user needs to call either this function or setupForNewAMRRun or setupforFixedHierarchyRun before she calls run.
- void setupForNewAMRRun();
   Sets up this object for cold start. User must have previously called define. Need to call this function or setupForRestart or setupforFixedHierarchyRun before you run.

This sets the grid hierarchy and sets regrid\_intervals to -1 (turns off regridding). If you want to keep regridding on, reset regridIntervals after this call.

void run(Real a\_max\_time, int a\_max\_step);
 Run the calculation. User must have previously called both the define function and a setup function in order to call this.

#### **Arguments:**

- a\_max\_time : Time to stop the calculation.
- a\_max\_step: Maximum number of iterations.
- void conclude() const: The user should call this last. It writes the last checkpoint file and performs other housekeeping functions.

There are also functions in the AMR class which allow the user to reset various parameters of the run (blocking factor, regridding intervals, checkpointing intervals, etc. See the reference manual for details. Examples of applications that use the AMR class to implement an adaptive Godunov method are given in Chombo/example/AMRGodunov.

#### **5.2.3 Class AMRLevel**

AMRLevel is an abstract base class for data at the same level of refinement within a hierarchy of levels. The concrete class derived from AMRLevel is called a *physics class*. The domain of a level is a disjoint union of rectangles in a logically rectangular index space. Data is defined within this domain. There is also a problem domain, which may be larger, within which data can, in theory, be interpolated from some coarser level.

AMRLevel is the interface that the class AMR uses to call the physics class. The important parts of the public interface to AMRLevel are:

Defines this AMRLevel.

#### **Arguments:**

- coarser\_level\_ptr (not modified): Pointer to next coarser level object.
- problem\_domain (not modified): Problem domain of this level.
- level (not modified): Index of this level. The base level is zero.
- ref\_ratio (not modified): The refinement ratio between this level and the next finer level.
- virtual Real advance() = 0;
   Advance this level by one time step. Return an estimate of the new time step at this level.
- virtual void postTimeStep() = 0;
   Do all operations that are required after a timestep is completed. Refluxing happens here.
- virtual void tagCells(IntVectSet& a\_tags) const = 0;
   Create tagged cells for dynamic mesh refinement.
- virtual void tagCellsInit(IntVectSet& a\_tags) const = 0;
   Create tagged cells for mesh refinement at initialization.

Perform any pre-regridding operations which are necessary. This is not a pure virtual function, to preserve compatibility with earlier versions of AMRLevel. The AMRLevel::preRegrid() instantiation does nothing.

virtual void regrid(const Vector<Box>& a\_new\_grids) = 0;
 Redefines this level to have the specified grids as its defined union of rectangles.

- virtual void postRegrid(int a\_base\_level);
   Perform any post-regridding operations which are necessary. This is not a pure virtual function to preserve compatibility with earlier versions of AMRLevel. The AMRLevel::postRegrid() instantiation does nothing.
- virtual void initialGrid(const Vector<Box>& a\_new\_grids) = 0;
   Initialize this level to have the specified domain a\_new\_grids.
- virtual void initialData() = 0;
   Initialize the data
- virtual void postInitialize() = 0;
   Do any operations that are required just after initialization.
- virtual Real computeDt() = 0;
   Returns maximum stable time step for this level.
- virtual Real computeInitialDt() = 0;
   Returns maximum stable time step for this level with initial data.
- virtual void writeCheckpointHeader(HDF5Handle& a\_handle) const = 0; Write the header to the checkpoint file handle.
- virtual void writeCheckpointLevel(HDF5Handle& a\_handle) const = 0;
   Write checkpoint data for this level.
- virtual void readCheckpointHeader(HDF5Handle& a\_handle) = 0;
   Read checkpoint header.
- virtual void readCheckpointLevel(HDF5Handle& a\_handle) = 0;
   Read checkpoint data for this level.
- virtual void writePlotHeader(HDF5Handle& a\_handle) const = 0;
   Write plot file header for this level.
- virtual void writePlotLevel(HDF5Handle& a\_handle) const = 0;
   Write plot file data for this level.

These are all the pure virtual functions of the AMRLevel interface and therefore all the functions that the user **must** define for her application. There are other ancillary functions in the interface that have reasonable defaults. Most of these involve data member access and modification.

### **5.2.4 Class** AMRLevelFactory

The class AMRLevelFactory is a pure virtual base class, with only one member function:

• virtual AMRLevel\* new\_amrlevel() const = 0;

This is the only member function of AMRLevelFactory, and it must be defined by the user in a derived class. The derived function will return a pointer to a physics-specific class derived from AMRLevel.

A pointer to an object of this class is passed to the define function of an AMR object, which uses it to construct the various AMRLevel objects that it requires.

# 5.3 AMRTimeDependent Example: Advection-Diffusion

We provide an example which uses the AMRTimeDependent infrastructure to solve the advection-diffusion equation. Given an analytic velocity field  $\vec{v}$  and a diffusion coefficient  $\nu$ , we evolve an advected and diffused scalar  $\phi$  using the advection-diffusion equation:

$$\frac{\partial \phi}{\partial t} + \nabla \cdot (\vec{v}\phi) = \nu \Delta \phi.$$

The algorithm is mostly as described in previous sections. The solution to the Riemann problem is given by simple upwinding.  $F_d$ , the advective flux in the d direction, is given by  $F_d = v^d \phi$ . The characteristic values are the velocity components.

The main difference between the algorithm used in this example and that in Figure 5.1 is in the approach taken to flux correction during the synchronization step. The total flux is the advective flux minus the diffusive flux:  $F_d^{tot} = v^d \phi - \nu \nabla \phi$ . Because the fluxes used to update the solution include both advective and diffusive fluxes, simply performing the explicit refluxing correction step:

$$\phi^l(t^l + \Delta t^l) := \phi^l(t^l + \Delta t^l) - \Delta t^l D_R(\delta F^{l+1})$$

will be unstable because it represents a forward-Euler update for the heat equation (which requires that  $\Delta t \approx O(h^2)$  for stability). To preserve stability, we instead apply the flux correction implicitly, as described in [22]:

We first solve a Helmholtz equation for a correction:

$$(I - \Delta t^{\ell_{base}} \nu \Delta^{comp}) \delta \phi = \Delta t^{\ell} D_R(\delta \vec{F}^{tot,\ell+1}) \quad \text{for } \ell \ge \ell_{base}$$
 (5.1)

Then, the correction is added to the solution:

$$\phi^{\ell} := \phi^{\ell} + \delta \phi^{\ell} \quad \text{for} \quad \ell \ge \ell_{base}. \tag{5.2}$$

### 5.3.1 Advection Diffusion-Specific Classes

The AMR advection-diffusion application in Chombo/example/AMRGodunov uses the AMRTimeDependent infrastructure extensively. These are the classes that are specific to this application.

- AdvectPhysics is the GodunovPhysics-derived class. It provides PatchGodunov the physics-specific information needed to advance the solution.
- LevelAdvect is used to advance an advection-diffusion solution one step:

$$\phi^{new} = \phi^{old} - \Delta t (\nabla \cdot (\vec{v}\phi))^{n + \frac{1}{2}}.$$

In the case of non-zero diffusion, we use this advance as an input to a semi-implicit advance using the LevelTGA class described in section 4.7. In the pure advection case, we can just use this update.

- AMRLevelAdvectDiffuse is the AMRLevel-derived class that drives the calculation.
- AMRLevelAdvectDiffuseFactory is the factory class that the user sends to AMR. This class generates AMRLevelAdvectDiffuse objects for AMR.

## 5.3.2 Usage Pattern

In broad strokes, the driver for the AMRLevelAdvectDiffuse example looks something like this:

```
// read inputs
ParmParse ppgodunov;

//define the domain of computation
ProblemDomain prob_domain;
getProblemDomain(prob_domain);

//define the initial and boundary conditions
RefCountedPtr<AdvectTestIBC> ibc;
getAdvectTestIBC(ibc);

//define the AdvectPhysics object
AdvectPhysics advPhys;
advPhys.setPhysIBC(&(*ibc));

//pick an advection velocity function
AdvectionVelocityFunction velFunc;
getAdvectionVelocityFunction(velFunc);
```

```
//define the factory object
  RefCountedPtr<AMRLevelAdvectDiffuseFactory> amrg_fact;
  getAMRLADFactory(amrg_fact, velFunc, advPhys);
  //define the AMR object
  AMR amr;
  defineAMR(amr, amrg_fact, prob_domain, a_refRat);
  //sets all the amr parameters (as opposed to a fixed grid run)
  setupAMRForAMRRun(amr);
  // run
  amr.run(a_stopTime,a_nstop);
  // output last pltfile and statistics
  amr.conclude();
To use these functions directly (all of these functions exist in AdvectDiffuseUtils),
one must have an input file that looks something like this:
##cfl number and its initial time counterpart
cfl = 0.9
initial_cfl = 0.1
##nu
diffusion_coef = 0.001
##how much output one wants from the application
verbosity = 1
##max number of time steps
max\_step = 100
##final solution time
max time = 100.0
##number of buffer cells
tag_buffer_size = 3
##tagging threshold for undivided difference of solution
refine_thresh = 0.05
##how often we regrid
regrid_interval = 2 2 2 2 2 2
##how fast time step can grow and how far out of bounds it can get
max_dt_growth = 1.1
dt_tolerance_factor = 1.1
## size of the domain
domain_length = 1.0
## distance (in level 1 cells) between levels 1+1 and 1-1
grid_buffer_size = 1
##do periodic boundary conditions or not
periodic_bc = 1 1 1
```

```
##fixed time step size. negative if not fixing time step
fixed_dt
          = -1.0
##maximum AMR level number
max_level = 2
##coarsest AMRLevel grid size
n_{cell} = 64 64 64
##refinement ratios between levels
ref_ratio = 2 2 2 2
##blocking factor
block_factor = 4
##maximum size of grids
max\_grid\_size = 64
##fill ratio for Berger-Rigoutsos
fill_ratio = 0.75
## how often to checkpoint. negative if no checkpoints
checkpoint_interval = -1
## how often to dump plotfile. negative if no plotfiles.
plot_interval = 10
##prefix of plotfiles
plot_prefix = plt
##prefix of checkpoint files
chk_prefix = chk
blob_center = 0.75 \ 0.5 \ 0.5
blob_radius = 0.1
#0 = constant
#1 = rotating
advection_vel_type = 0
use_limiting = true
amrmultigrid.num_smooth = 8
amrmultigrid.tolerance = 1.0e-10
amrmultigrid.num_mg
                       = 1
amrmultigrid.norm_thresh = 1.0e-10
amrmultigrid.hang_eps
                      = 1.0e-10
amrmultigrid.max_iter
                         = 100
amrmultigrid.verbosity = 1
##1 = neumann, 0 = dirichlet
bc_lo
      = 1 1 1
bc_hi
      = 1 1 1
bc_value = 0.
```

# **5.3.3** AMRLevelAdvectDiffuseFactory Interface

We use the following interface to create an AMRLevelAdvectDiffuseFactory.

AMRLevelAdvectDiffuseFactory(const AdvectPhysics&

gphys,

AdvectionVelocityFunction advFunc, BCHolder bcFunc, const Real& cfl, const Real& domainLength, refineThresh, const Real& const int& tagBufferSize, const Real& initialDtMultiplier, const bool& useLimiting, const Real& nu);

- gphys The AdvectPhysics used to create the advection term.
- advFunc The function that provides the advection velocity. This function must be of the form

- bcFunc The boundary condition class sent to solve the diffusion equation. See section 4.6.4 for details.
- cfl The CFL number.
- domainLength The physical length of the domain.
- refineThresh Undivided gradient size over which a cell will be tagged for refinement.
- tagBufferSize Number of buffer cells around each tagged point that will also be tagged.
- initialDtMultiplier CFL number at the beginning of the calculation.
- useLimiting Whether to use vanLeer limiting. Turn this to true unless you are doing a convergence test.
- nu Diffusion coefficient.

#### **5.3.4** LevelAdvect Interface

LevelAdvect is used to advance an advection-diffusion solution one step. In the case of non-zero diffusion, we use this advance as an input to a semi-implicit advance using LevelTGA. In the pure advection case, we can just use this update. If a user is simply using the AMRLevelAdvectDiffuse application, this class is entirely internal to AMRLevelAdvectDiffuse. This discussion is provided for the benefit for those who want to use this object for other applications.

We define a LevelAdvect with the following interface

```
void define(const AdvectPhysics&
                                         gphys,
            const DisjointBoxLayout&
                                         thisDisjointBoxLayout,
            const DisjointBoxLayout&
                                         coarserDisjointBoxLayout,
            const ProblemDomain&
                                         domain,
            const int&
                                         refineCoarse,
            const bool&
                                         useLimiting,
            const Real&
                                         dx,
            const bool&
                                         hasCoarser,
            const bool&
                                         hasFiner);
```

- gphys The AdvectPhysics used to create the advection term.
- domain The domain of the computation.
- refineCoarse The refinement ratio to the next coarser level.
- useLimiting Whether to use vanLeer limiting. Turn this to true unless you are doing a convergence test.
- dx The grid spacing.
- hasCoarser Whether there is a coarser AMR level.
- hasFiner Whether there is a finer AMR level.

We present the interface function that advances the solution.

$$\phi^{new} = \phi^{old} - \Delta t (\nabla \cdot (\vec{v}\phi))^{n + \frac{1}{2}}$$

```
Real step(LevelData<FArrayBox>&
                                         U,
          LevelFluxRegister&
                                         finerFluxRegister,
          LevelFluxRegister&
                                         coarserFluxRegister,
          LevelData<FluxBox>&
                                         advectionVelocity,
          const LevelData<FArrayBox>&
                                         S,
          const LevelData<FArrayBox>&
                                         UCoarseOld,
                                         TCoarseOld,
          const Real&
          const LevelData<FArrayBox>&
                                         UCoarseNew,
                                         TCoarseNew,
          const Real&
          const Real&
                                         time,
          const Real&
                                         dt);
```

- U The solution. Input is the old solution  $\phi^{old}$ . Input is the new solution  $\phi^{new}$ .
- finerFluxRegister The flux register with the next finer level.
- coarserFluxRegister The flux register with the next coarser level.

- advectionVelocity The holder for the advection velocity. This is held in a LevelData to make this object of somewhat more general utility.
- $\bullet$  S Source term. In the case of advection-diffusion,  $S=\nu\Delta\phi.$
- UCoarseOld Solution at next coarser level at the old time.
- TCoarseOld Time at which UCoarseOld exists.
- UCoarseNew Solution at next coarser level at the new time.
- TCoarseNew Time at which UCoarseNew exists.
- time Time at which U exists before advance happens  $(t^{old})$ .
- dt Time step.

# Chapter 6

# **HDF5 I/O with Chombo**

# 6.1 HDF5 I/O

We have developed a user interface for file I/O based on version 5 of the Hierarchical Data Format library (HDF5) developed at The National Center for Supercomputing Applications (NCSA). HDF5 provides efficient and flexible mechanisms for handling I/O of large scientific datasets, and is becoming a standard in the scientific community for binary portable data files. We exploit a number of features provided by HDF5, including the portability of data across platforms and the ability to read and write files on distributed memory parallel systems. HDF5 also has a number of useful utilities, such as h5dump, which produces a human-readable formatted ASCII output of an HDF5 file.

HDF5 has three main user abstractions: *group*, *dataset*, *attribute*. Group abstracts the notion of the location in a file, while dataset and attribute are different types of data that can be stored in an HDF5 file. We provide an API for creating HDF5 files and for reading from and writing into such files. These are implemented using two classes, plus a collection of stand-alone functions.

#### **6.1.1** Class HDF5Handle

HDF5Handle is a class that manages the accessing of and navigation within an HDF5 file.

#### • Constructors.

```
HDF5Handle();
HDF5Handle(const std:string& a_filename, mode);
int open(const std:string& a_filename, mode);
bool isOpen();
void close();
enum mode {CREATE,OPEN_RDONLY,OPEN_RDWR}
```

A HDF5Handle requires a a\_filename supplied either at construction using the second constructor, or by a call to open. filename follows the semantics of fopen from the

<stdio.h> of libc. It is an error if a file has already been opened by the HDF5Handle. It is also illegal to open a single file using two different HDF5Handles. The enumeration class mode specifies the access permissions. If mode = CREATE, the a file is created, deleting the previously existing copy of that file if necessary. If mode = OPEN\_RDONLY, an existing file is opened with read-only access. If mode = OPEN\_RDWR an existing file is opened with read-write access. In the latter two cases, if the file doesn't exist, then the open operation fails: there is no file bound to the HDF5Handle, and a call to isOpen would return false. HDF5Handle objects must be explicitly closed by the user, just like file pointers in standard C. This is done with the close function. You can inquire whether a handle is open or closed with the isOpen() function. Once close has been called, it is possible open a new file with the same HDF5Handle using open.

#### File Navigation.

```
int HDF5Handle::setGroup(const std::string& a_groupAbsPath);
const std::string& HDF5Handle::getGroup() const;
```

The function setGroup sets the group (i.e., the location in the file) to be that labeled with the string a\_groupAbsPath. If such a group does not yet exist, setGroup creates such a group. The function getGroup returns the string corresponding to the group to which the HDF5Handle is currently set. The input and output strings to which the groups are set are assumed to be of the form of an Unix absolute directory path, e.g., "/foo", "/level\_1/info", etc. There is a distinguished root group "/" to which the HDF5Handle is initialized when a file is opened. setGroup can be thought of as analogous to a Unix "cd" command. getGroup can be thought of as analogous to the Unix "pwd" command. We should emphasize that the setting of the group in an HDF5Handle is usually unrelated to the actual physical file layout. It just represents an evocative and convenient notation for navigating within an HDF5 file. setGroup returns 0 on success, a negative number if HDF5 had an error. If the group doesn't already exist, then it is created if the file is write-enabled (CREATE or OPEN\_RDWR). In the event of error, file remains open and setGroup can be called again, but HDF5Handle object is not capable of processing reads or writes until a successful setGroup has been performed. (except immediately after file opening when the root group is valid for writing).

#### 6.1.2 Class HDF5HeaderData

The class HDF5HeaderData provides an interface for writing collections of reals, integers, strings, Boxes and IntVectSets. In this interface, one must associate a name (in the form of a character string) for each object. The internal treatment of this data assumes that these are small collections of "metadata", where the efficiency of storage is not a serious concern.

```
class HDF5HeaderData
{
```

```
public:
  int writeToFile(HDF5Handle& a_handle) const;
  int readFromFile(HDF5Handle& a_handle);
  void clear();
  map<std::string, Real>
                                 m_real;
  map<std::string, int>
                                 m_int;
  map<std::string, std::string> m_string;
  map<std::string, IntVect>
                                 m_intvect;
  map<std::string, RealVect>
                                 m_realvect;
  map<std::string, Box>
                                 m_box;
};
```

Once an HDF5HeaderData object is created, the user adds objects to to be stored by adding values to the STL maps that are contained as member data. For example,

```
HDF5HeaderData metaData;
metaData.m_real["mesh spacing"] = dx;
```

If there is already a value in the map corresponding to the string "mesh spacing", the value is overwritten. One queries to see if an if an attribute is entered in one of the maps as follows.

```
bool ghost_exists =
  (metaData.m_intvect.find("ghost") != metaData.m_intvect.end());
```

Finally, one deletes an attribute from a map as follows:

```
metaData.m_real.erase("mesh spacing");
```

Once the user finishes filling in an HDF5HeaderData object, the member functions are writeToFile and readFromFile write and read group attributes from the group currently pointed to by a\_handle.

## **6.1.3 HDF5 I/O for** BoxLayoutData

We provide a set of function interfaces for writing out data defined on unions of rectangles. There are two sets of functions: one for reading and writing the unions of rectangles, the second for reading and writing BoxLayoutData objects.

### • BoxLayout I/O.

```
int write(HDF5Handle& a_handle, const BoxLayout& a_layout);
int read(HDF5Handle& a_handle, Vector<Box>& boxes);
```

The write function writes out the union of boxes corresponding to all of the BoxLayoutData objects to be written to that group. Consequently, one can only write one BoxLayout object for that group. The read function is not symmetric with the write function. The reason for this is that processor assignment is not written out to the file with the BoxLayout. The file is considered *parallel neutral*. Since a BoxLayout is a combination of Boxes and processor assignment, the read function does not have enough information to build a BoxLayout. It is the users responsibility to invoke the appropriate load balancing function after the boxes have been read in, and build a BoxLayout object

#### BoxLayoutData I/O.

write writes the collection of T objects in a\_data into an HDF5 dataset, linearizing each object into a into a set of 1D Arrays. The default implementation is to use the linearization function T::linearOut that is required to define the LevelData<T> class, but that will output data in terms of bytes, and in general will not be portable across platforms. The user may provide a more detailed linearization interface, in which case the HDF5 files can be made portable across platforms. Such an interface has been provided in Chombo for the case of T = FArrayBox. read reads a LevelData<T> object that had been previously written by the write function. On input, a\_layout is a null-constructed LevelData, which is then defined inside read. If redfineData == false, then the user takes responsibility for calling define is the correct manner for the LevelData<T> a\_data argument. The a\_layout argument must consist of the same collection of Boxes as that used to define a\_data, but may have a different mapping of boxes to processors than that of the data as it was written.

There are also versions of these functions for the case of BoxLayoutData<T>.

#### 6.1.4 HDF5 Out-Of-Core readers

Frequently, a user will generate a data file from a simulation run (particularly in parallel) that will exceed a single computers available RAM. This can make auxiliary post-processing programs impossible to run unless they too are programmed in parallel. The nature of

many post-processing operation, however, are more like data-mining than a full simulations. For those cases a simpler approach can be used where only a subset of the data file is read in and operated on.

```
template <class T>
int readLevel(HDF5Handle& a_handle,
              const int& a_level,
              LevelData<T>& a_data,
              Real& a_dx,
              Real& a_dt,
              Real& a_time,
              Box& a_domain,
              int& a_refRatio,
              const Interval& a_comps = Interval(),
              const IntVect& ghost = IntVect::Zero,
              bool setGhost = false);
int readBoxes(
              HDF5Handle& a_handle,
              Vector<Vector<Box> >& boxes);
int readFArrayBox(HDF5Handle& a_handle,
                  FArrayBox& a_fab,
                  int a_level,
                  int a_boxNumber,
                  const Interval& a_components,
                  const std::string& a_dataName = "data" );
```

The first function defaults to reading the entire range of components for a given level of data. If the user specifies Interval a\_comps (currently defaulting to the entire data range) then a user can select out just a particular range of components.

readFArrayBox is even more specific, in that it reads individual FArrayBox data's from the file by

level, index

reference.

# 6.2 AMR I/O routines

WriteAMRHierarchyHDF5 and ReadAMRHierarchyHDF5 are convenient global functions used in the AMR codes developed by ANAG. There are three main reasons for their use:

- 1. It relieves the user from having to learn about the HDF5 interface code.
- 2. It places the data into a format that can subsequently be read successfully by the Vislt post-processing and visualization tools.
- 3. They are symmetric and can be used for efficient checkpoint file generation.

### 6.2.1 Function WriteAMRHierarchyHDF5

#### void

#### Arguments:

No arguments are modified.

- filename: file to output to.
- a\_vectGrids : grids at each level.
- a\_vectData : data at each level.
- a\_vectNames: names of variables.
- a\_domain : domain at coarsest level.
- a\_dx : grid spacing at coarsest level.
- a\_dt : time step at coarsest level.
- a\_time: time.
- a\_vectRatio : refinement ratio at all levels (ith entry is refinement ratio between levels i and i + 1).
- a\_numLevels : number of levels to output.

filename is created if it doesn't already exist, and overwritten if it does exist. a\_vectGrids must match the grids that a\_vectData is defined over. a\_vectNames are the names you wish to be associated with the components of the a\_vectData. a\_domain is the covering domain box at the coarsest level. a\_numLevels is the number of levels, starting at level 0 that the user wishes to be output.

### 6.2.2 Function ReadAMRHierarchyHDF5

int

ReadAMRHierarchyHDF5(const string& filename,

Vector<DisjointBoxLayout>& a\_vectGrids,
Vector<LevelData<FArrayBox>\* > & a\_vectData,
Vector<string>& a\_vectNames,
Box& a\_domain,
Real& a\_dx,
Real& a\_dt,
Real& a\_time,
Vector<int>& a\_refRatio,
int& a\_numLevels,
const IntVect& ghostVector)

#### Arguments:

- filename: file to input from.
- a\_vectGrids : grids at each level.
- a\_vectData : data at each level.
- a\_vectNames: names of variables.
- a\_domain : domain at coarsest level.
- a\_dx : grid spacing at coarsest level.
- a\_dt : time step at coarsest level.
- a\_time: time.
- a\_vectRatio : refinement ratio at all levels.
- a\_numLevels : number of levels to read.
- a\_ghostVector : IntVect used to define a\_vectData

return codes:

```
0: success
-1: number of levels <= 0
-2: number of components <= 0
-3: error in readlevel function
-4: file open failed</pre>
```

the argument notes are the same as for WriteAMRHierarchyHDF5, with the addition of ghostVector ghostVector is passed in the argument list and is used in the definition of the LevelData<FArrayBox> definition of a\_vectData. This was most useful for data moving between a simulation code and a post-processing code where the ghost cell requirements can be different.

# 6.3 Other HDF5 I/O functions

To aid in debugging and in visualizing intermediate data, the functions writeFAB, writeFABname, writeLevel, and writeLevelname may be used. These functions are designed to output a single FArrayBox or a LevelData<FArrayBox> into a file which can then be read by Vislt. These functions may be called from debuggers such as gdb during the debugging process. The usual way these functions are used is as follows:

```
    place the line
#include ''FABView.H',1
in the file which contains the function main().
```

- 2. Run the code in the debugger, calling the needed IO function as needed.
- 3. in a shell environment, start Vislt to look at the output file

#### 6.3.1 Functions writeFAB and writeFABname

The functions writeFAB and writeFABname write a single FArrayBox into a file which uses the plotfile format of WriteAMRHierarchyHDF5, and which can then be viewed with Vislt (called separately). The writeFAB function writes the input FArrayBox into a file named fab.hdf5, while the writeFABname allows the user to specify the name of the output file. Note that the data is passed using a pointer to an FArrayBox.

```
void
writeFAB(const FArrayBox* a_data)
Arguments:
No Arguments are modified.
```

<sup>&</sup>lt;sup>1</sup>The actual function declarations are in AMRIO.H, but FABView.H contains fake "calls" to the various functions to ensure that they are all included in the executable file.

a\_data : data to be written

A file named fab.hdf5 is created if it doesn't already exist and overwritten if it does exist.

void

writeFABname(const FArrayBox\* a\_data, const char\* a\_filename)

Arguments:

No Arguments are modified.

- a\_data : data to be written
- a\_filename : name of file into which data is written

a\_filename is created if it doesn't already exist, and overwritten if it does exist.

#### 6.3.2 Functions writeLevel and writeLevelname

The functions writeLevel and writeLevelname write the data from a single LevelData<FArrayBox> into a file which uses the plotfile format of WriteAMRHierarchyHDF5, and which can then be viewed with Vislt (called separately). The writeLevel function writes the input LevelData<FArrayBox> into a file named LDF.hdf5, while the writeLevelname allows the user to specify the name of the output file.

Notes:

- Data is passed using a pointer to a LevelData<FArrayBox>.
- All data is written on an FArrayBox by FArrayBox basis, including any and all ghost cells.

void

writeLevel(const LevelData<FArrayBox>\* a\_data)

Arguments:

No Arguments are modified.

• a\_data : data to be written

A file named LDF.hdf5 is created if it doesn't already exist and overwritten if it does exist.

void

writeLevelname(const LevelData<FArrayBox>\* a\_data, const char\* a\_filename)

Arguments:

No Arguments are modified.

- a\_data : data to be written
- a\_filename : name of file into which data is written
- a\_filename is created if it doesn't already exist, and overwritten if it does exist.

#### 6.3.3 Functions writeDBL and writeDBLname

The functions writeDBL and writeDBLname write the data from a single DisjointBoxLayout into a file which uses the plotfile format of WriteAMRHierarchyHDF5, and which can then be viewed with Vislt (called separately). It does this by creating a LevelData<FArrayBox> using the input DisjointBoxLayout with the data initialized to the processor ID of each grid, and then calling the associated writeLevel functions. The writeDBL function writes the input DisjointBoxLayout into a file named DBL.hdf5, while the writeDBLname allows the user to specify the name of the output file.

Notes:

• Data is passed using a pointer to a DisjointBoxLayout.

#### void

writeDBL(const DisjointBoxLayout\* a\_data)

#### Arguments:

No Arguments are modified.

• a\_data : data to be written

A file named DBL.hdf5 is created if it doesn't already exist and overwritten if it does exist.

#### void

writeDBLname(const DisjointBoxLayout\* a\_data, const char\* a\_filename)

#### Arguments:

No Arguments are modified.

- a\_data : data to be written
- a\_filename : name of file into which data is written

a\_filename is created if it doesn't already exist, and overwritten if it does exist.

# Chapter 7

# Using PETSc in Chombo

PETSc (Portable, Extensible Toolkit for Scientific Computation) is a library of tools for computing with structured and unstructured grids. Its primary functionality is a time stepping class (TS), which has a nonlinear solver (SNES), which has a linear solver (KSP). Chombo provides support for constructing PETSc composite grid matrices (class PetscCompGrid) from Chombo's block structured hierarchy of grids. Applications can then use these sparse (unstructured "AIJ") matrices in PETSc's (time,nonlinear,linear) solvers.

This approach to solvers in Chombo is motivated by applications that have operators that are not solved well with Chombo's geometric multigrid solver ( $\S4.4.4$ ). The first application to use this functionality is the BISICLES ice sheet modeling code.

# 7.1 Building Chombo with PETSc

Follow the instructions on the PETSc web page and document to build PETSc if there is not a PETSc build on your machine. This involves setting the make variables PETSC\_DIR and PETSC\_ARCH in you environment or as an argument to make. Build your Chombo application with USE\_PETSC=TRUE.

# 7.2 PETSc Composite Grid Classes

Chombo provides a base class PetscCompGrid that provides a framework for constructing PETSc matrices from a Chombo AMR hierarchy of grids. Specific operators are derived from this base class (e.g., PetscCompGridPois). An example of the usage to solve a Laplacian is in lib/test/AMRElliptic/testPetscCompGrid.cpp.

# 7.3 PETSc Solver Usage

There are three components to building a PETSc solver in Chombo: 1) making a matrix  $(\S7.3.1)$ , 2) make a solver  $(\S7.3.2)$ , and 3) make the operator  $(\S7.3.3)$ .

### 7.3.1 Making a PETSc Matrix

Chombo currently has a constant coefficient Laplacian class (PetscCompGridPois) and a variable coefficient viscous tensor operator class PetscCompGridVTO. Figure 7.1 shows an example making a matrix for the constant coefficient Laplacian.

```
#include "PetscCompGridPois.H"
....
PetscCompGridPois petscop(0.,-1.,2); /* alpha, beta, and order */
RefCountedPtr<ConstDiriBC> bcfunc = RefCountedPtr<ConstDiriBC>(new ConstDiriBC(1,petscop.getGhostVect()));
BCHolder bc(bcfunc);
petscop.setCornerStencil(true);
petscop.define(cdomain,grids,refratios,bc,cdx*RealVect::Unit);
petscop.setVerbose(1);
Mat A; /* linear system matrix */
A = petscop.getMatrix();
```

Figure 7.1: Example of creating a matrix from testPetscCompGrid.cpp

This example has a simple operator "define" process:  $\alpha$  and  $\beta$ , in the matrix for the operator  $\alpha u + \nabla \cdot \beta \nabla u$ , and the order of the discretization are arguments to the constructor. The type of stencil is set with an operator specific method. The actual define method is a base class method and provides the grid hierarchy, domain, and boundary condition class. See lib/src/AMRElliptic/PetscCompGrid.H for the current version and Figure 7.2 for reference. The class ConstDiriBC implements homogeneous Dirichlet boundary

Figure 7.2: PetscCompGrid::define

conditions. Other boundary conditions can be implemented by deriving from the base class CompBC in lib/src/AMRElliptic/PetscCompGrid.H

### 7.3.2 Making a PETSc Solver

Figure 7.3 shows an example of a PETSc linear solver.

```
// solve A phi = rhs
Vec x, b; /* approx solution, RHS */
KSP ksp; /* linear solver context */
ierr = MatGetVecs(A,&x,&b); CHKERRQ(ierr);
ierr = petscop.putChomboInPetsc(rhs,b); CHKERRQ(ierr);
ierr = KSPCreate(PETSC_COMM_WORLD, &ksp); CHKERRQ(ierr);
ierr = KSPSetOperators(ksp, A, A, DIFFERENT_NONZERO_PATTERN);
CHKERRQ(ierr);
ierr = KSPSetFromOptions(ksp); CHKERRQ(ierr);
ierr = KSPSolve(ksp, b, x); CHKERRQ(ierr);
ierr = KSPDestroy(&ksp); CHKERRQ(ierr);
ierr = petscop.putPetscInChombo(x,phi); CHKERRQ(ierr);
```

Figure 7.3: Example of a linear solver from testPetscCompGrid.cpp

The PetscCompGrid class provides helper methods to move vector data between PETSc and Chombo data structures (putChomboInPetsc and putPetscInChombo). The rest of this solver code is standard PETSc and one can refer to the PETSc documentation for more information.

## 7.3.3 Making an operator

The PetscCompGrid class has one pure virtual method:

PetscErrorCode createOpStencil(IntVect,int,DataIterator,StencilTensor&) = 0;. Figure 7.4 shows an example of a simple 5 and 7 point stencil for the constant coefficient Laplacian. Note, the first two arguments are the components of a multilevel cell index, which can be used to construct the multilevel cell index object IndexML. The third argument is a data iterator that could be use to, for instance, index into a coefficient data structure; this example does not use this argument. Chombo provides some build-in operators and users are encouraged to implement their own as needed.

```
PetscErrorCode PetscCompGridPois::createOpStencil(IntVect a_iv,
                                                   int a_ilev,
                                                   DataIterator a_dit,
                                                   StencilTensor &a_sten)
 {
   Real dx=m_dxs[a_ilev][0],idx2=1./(dx*dx);
   PetscFunctionBeginUser;
   StencilTensorValue &v0 = a_sten[IndexML(a_iv,a_ilev)];
   v0.define(1);
   v0.setValue(0,0,m_alpha - m_beta*2.*SpaceDim*idx2);
   for (int dir=0; dir<CH_SPACEDIM; ++dir)</pre>
   {
     for (SideIterator sit; sit.ok(); ++sit)
        int isign = sign(sit());
        IntVect jiv(a_iv); jiv.shift(dir,isign);
        StencilTensorValue &v1 = a_sten[IndexML(jiv,a_ilev)];
        v1.define(1);
        v1.setValue(0,0,m_beta*idx2);
     }
     PetscFunctionReturn(0);
   }
```

 $Figure \ 7.4: \ Example \ of \ an \ operator \ from \ testPetscCompGridPois.cpp$ 

## Chapter 8

# **Using SUNDIALS in Chombo**

The SUite of Nonlinear and DIfferential/ALgebraic equation Solvers (SUNDIALS) [15, 16] provides robust and efficient time integrators and nonlinear solvers that can easily be incorporated into existing simulation codes. Within Chombo, it can be used as a time integrator for global time stepping, that is for a time step within a single level or time step which advances an entire AMR hierarchy. For details on SUNDIALS features and capabilities see the SUNDIALS project page at https://computing.llnl.gov/projects/sundials and the documentation at https://sundials.readthedocs.io.

Chombo interfaces are expressed through adaptors to the SUNDIALS NVector data structure. Because SUNDIALS doesn't have a concept of spatial mesh or discretization, the adaptors wrap actions using knowledge of "valid data," e.g. uncovered values in an AMR hierarchy, interior (not ghost cells), etc. The first application to use this functionality is the BISICLES ice sheet modeling code.

## 8.1 Building Chombo with SUNDIALS

Follow the instructions from the SUNDIALS documentation to build and install SUNDIALS if it is not already available on your machine. If you are compiling with the C++11 standard, we recommend using SUNDIALS version 6.3.0. Newer SUNDIALS versions can be used if C++14 or a later standard is selected. Next, set the make variable SUNDIALS\_DIR (which should point to the <code>install</code> directory with subdirectories containing header and library files) in your environment or as an argument to make. Then build your Chombo application with USE\_SUNDIALS=TRUE. The adaptors can be compiled for parallel execution with SUNDIALS using MPI=TRUE.

## 8.2 Chombo-SUNDIALS Adaptor Classes

Chombo provides a base class ChomboSundialsAdaptor that provides a virtual base class for constructing SUNDIALS NVector-compatible classes. Specific adaptors have

been derived from this base class; AmrAdaptor works with an AMR hierarchy, while SimpleAdaptor works with a single level. An example of the usage in a simple Laplacian heat equation model is in:

releasedExamples/Sundials/exec/testSundialsPDE.cpp .

### 8.3 SUNDIALS Adaptor Usage

The interface to use the SUNDIALS adaptor is straight-forward:

- Add SUNDIALS initialization to your driver, to create a SUNDIALS context object, SUNContext;
- Define and construct your data holder (such as LevelData<FArrrayBox>),
- Choose the appropriate adaptor class, and call its define with your Chombo data holder.
- Use the provided macros to create a SUNDIALS NVector;
- Define required "rhs" and "solver" functions for SUNDIALS; and finally
- Call SUNDIALS to perform the time integration.

#### **8.3.1** Define and construct adaptors

In the example code, you will see these lines:

```
#include "SimpleAdaptor.H"
...
LevelData<FArrayBox> phi(level0, ncomp, ghostVect);
SimpleAdaptor adaptor;
adaptor.define(&u, false);
...
N_Vector y0 = N_VNew_Ch(ctx, adaptor);
```

Figure 8.1: Example of creating an adaptor and NVector from a LevelData.

This simply hands the pointer to the SUNDIALS adaptor, and tells it ("false") to not delete the data pointer in the adaptor destructor. The function N\_VNew\_Ch is an NVector constructor which connects the SUNDIALS vector operations to the data and operations in the adaptor class.

#### 8.3.2 Define the SUNDIALS functions

SUNDIALS works with C-style function and data pointers for its API, so that user-specific data has to be extracted and recast into user code types. Again, the example code provides a model for this for an explicit time integrator operator in Fig. 8.2. There is a macro "NV\_ADAP\_CH" that can be used to access the pointer to the adaptor, which can then expose Chombo data (using "getData()" here). Note that additional data needed in the rhs function can come from user-defined UserData, which can be an arbitrarily complex structure.

```
int f(Real t, N_Vector y, N_Vector yDot, void *data) {
  const UserData *ud = static_cast<UserData*>(data);
  const Real diffusionCoeff = ud->diffusionCoeff;
  const Real dx = ud->dx;

SimpleAdaptor* a_y = dynamic_cast<SimpleAdaptor*>(&NV_ADAP_CH(y));
  assert(a_y != NULL);
  SimpleAdaptor* a_yDot =
        dynamic_cast<SimpleAdaptor*>(&NV_ADAP_CH(yDot));
  assert(a_yDot != NULL);
  LevelData<FArrayBox>* ly = a_y->getData();
  LevelData<FArrayBox>* lyDot = a_yDot->getData();
  ...
  [update a_yDot data]
  return 0;
}
```

Figure 8.2: Example of defining a function for SUNDIALS.

## Chapter 9

# **Parallel Programming with Chombo**

## 9.1 Initialization and Scoping

Chombo provides no special MPI initialization function. This is done intentionally to make it easier for Chombo users to interface with other parallel packages (which may provide their own special initialization routines) in the same code that they use with Chombo.

If one is using Chombo in a parallel code, one must somehow call MPI\_Init before instantiating any Chombo data objects (whether that is in the context of some other package's initialization routines or not). One must also call MPI\_Finalize after all Chombo data objects have gone out of scope. To make sure these get done in the correct order, some care in scoping is required. Some Chombo classes, by necessity, call MPI functions in their destructors. One must be careful to make certain that all Chombo classes go out of scope before MPI\_Finalize gets called. A simple way to do this is shown here:

```
int main(int argc, char* argv[])
{
#ifdef MPI
MPI_Init(&argc, &argv);
#endif
//this sets the beginning of the scope of Chombo objects
{
LevelData<FArrayBox> phi, rhs;
...do a bunch of calculations...
//this sets the end of the scope of Chombo objects
}
#ifdef MPI
MPI_Finalize();
#endif
return(0);
```

}

In this example, MPI\_Init and MPI\_Finalize get used in the normal sense but the braces between them force Chombo destructors to be called before MPI\_Finalize is called.

#### 9.2 Overview of Chombo Data Parallelism

Our parallel model is box-based SPMD parallel programming. Distributed data always lives in containers (LayoutData, BoxLayoutData, and LevelData are the containers). All processors execute the same code. All processors have access to the unions of rectangles (the BoxLayouts and the DisjointBoxLayouts) and what processors each rectangle's data lives upon. The data associated with the rectangles is distributed among processors. We use smart iterators over our data objects which stop at the boxes which live on the current processor. To be complete, broadcast and gather functions are included for situations where parallelism cannot be hidden by iterators. As of release 3.2 it is also possible to use hybrid parallelism with OpenMP. When using Chombo's native iterators, it suffices to modify looping over the boxes. We use the same iterators, however, the boxes are indexed explicitly to let OpenMP know that they are parallelizable. The lower level code has been made thread-safe and examples of hybrid parallelism are available in AMRTimeDependent and AMRElliptic/AMRPoissonOp.\*. Note that the memory tracking feature of Chombo is not yet usable with hybrid parallelism.

As in the serial case, the class BoxLayout represents an arbitrary union of rectangles and LayoutData and BoxLayoutData both represent data built upon such a union. The class DisjointBoxLayout represents a disjoint union of rectangles and LevelData represents data built upon a disjoint union. The data in these data holders is distributed among processors according to the boxes that the data lives upon. A box's worth of data is considered atomic in this model.

Also, as in the serial case, the data holders have very different communication patterns even though all the holders distribute their data among processors. LayoutData is a distributed object that can not be involved in communication (it can be neither the source nor destination in copyTo or exchange). A BoxLayoutData object may be only the destination of a copyTo function and exchange is not defined for BoxLayoutData. A LevelData object, because it is built upon a disjoint layout, may be involved in any of our forms of data communication. Chombo also contains two templated communication functions that sometimes cannot be avoided in parallel applications: broadcast and gather. See section 9.5 for details.

## 9.3 Box-processor assignment

A BoxLayout is a set of boxes and processor assignments. We construct the layout with two matching lists: a Vector of boxes and a Vector of integers which represent

the processor into which data over the box will be distributed. Chombo does provide a load balancing function (see section 9.4 for details) which can generate these processor assignments. This function is not integrated into the BoxLayout class for the express purpose of providing a user the ability to use her own load balancing algorithm to generate processor assignments.

For now, assume we have a vbox = Vector<Box> which represents the grids from which we generate a BoxLayout and we have a vint = Vector<int> which represents the processor mapping we desire (we want data which resides on vbox[i] to reside on processor vint[i]). A BoxLayout can be constructed either incrementally:

```
BoxLayout boxlayout; //layout
Vector<Box> vbox; //grids
Vector<int> vint; //processor assignments
for(int ibox = 0; ibox < vint.size(); ibox++)
    boxlayout.addBox(vbox[ibox], vint[ibox]);
boxlayout.close();
or all at once:
Vector<Box> vbox; //grids
Vector<int> vint; //processor assignments
BoxLayout boxlayout(vbox, vint);
boxlayout.close();
```

Note that the close function must be called in either case after all the boxes are added. A DisjointBoxLayout is constructed in exactly the same way. If the boxes which are added to a DisjointBoxLayout are not disjoint (i.e. they have some nontrivial intersection) a runtime error is raised when close() is called.

### 9.4 LoadBalance

Chombo provides a load balancing function (called LoadBalance) to compute an assignment of boxes to processors for an AMR mesh hierarchy. The assignment is made to balance the computation workload on each processor (i.e., make it as even as possible). The meshes in the AMR hierarchy are represented using Vector< Vector< Box > >. The computational workload is a real number for each box in the hierarchy, represented as a Vector< Vector <Real> >. This is an input which the user may prescribe to her own needs. Load determination is far too application-specific to permit any kind of general solution. The resulting assignment is an integer for each box in the hierarchy, which gives the processor number (starting at zero) on which each box will reside. LoadBalance uses the Kernighan-Lin algorithm for solving knapsack problems. This algorithm has been used quite successfully for load balancing parallel AMR calculations [11]. The interface to LoadBalance is given by

```
void setGrids(Vector<DisjointBoxLayout>& vectGrids,
              const Vector<int>&
                                          vectRefRatio,
              const Vector<Vector<Box>& VVBoxNew,
              const int& numlevels)
{
  for(int ilev = 0; ilev < numlevels; ilev++)</pre>
  {
        const Vector<Box>& levelGrids = VVBoxNew[ilev];
        Vector<int> procAssign;
        int eekflag = LoadBalance(procAssign, levelGrids);
        assert(eekflag == 0);
        vectGrids[ilev].define(levelGrids, procAssign);
        vectGrids[ilev].close();
  }
}
```

Figure 9.1: Code snippet to show how LoadBalance is used to transform a list of boxes into DisjointBoxLayouts.

Here boxes are the input grids and procAssignments are the processor numbers that go with each box. The load for a given used by this version of LoadBalance is the number of points in the box. There is a more elaborate version of LoadBalance in Chombo which allows the user to input the loads for each box. See the reference manual for details. The return value of LoadBalance is an error code. If LoadBalance exited without error, 0 is returned. If anything other than zero is returned, the output values are undefined. An example of how to use LoadBalance is shown in 9.1.

#### 9.5 Broadcast and Gather

Chombo also contains two templated communication functions that sometimes cannot be avoided in parallel applications: broadcast and gather. Consider the following example: suppose one has a LevelData<FArrayBox> called resid and one wants to calculate the max norm of the data in this container. A naive way to do this would be:

```
//naive routine to calculate max norm of resid at varNum component
Real maxNorm(LevelData<FArrayBox>& resid, int varNum)
{
   Real maxnorm = 0;
   DataIterator dit = resid.iterator();
   for (dit.reset(); dit.ok(); ++dit)
   {
```

```
maxnorm = Max(maxnorm, resid[dit()].norm(0, varNum, 1);
}
return maxnorm;
}
```

This code is correct in serial and incorrect in parallel. In parallel, every processor will have a different value of maxnorm. To make this code correct, we must *gather* all the values of maxnorm, calculate the maximum value, and *broadcast* this value to all processors.

The interface to the templated gather function is as follows:

```
///gather a_input into a_outVec on a_dest
template <class T>
void gather(Vector<T>& a_outVec, const T& a_input, int a_dest);
```

This function gathers a\_input from every processor into Vector<T> a\_outVec on processor a\_dest. a\_outVec is a vector of length nProc() long with the value of a\_input on every processor in its elements.

The interface to the templated broadcast function is as follows:

```
///broadcast a_inAndOut to every processor from a_src
template <class T>
void broadcast(T& a_inAndOut, int a_src);
```

This function broadcasts a\_inAndOut from processor a\_src to all processors for both broadcast<T> and gather<T>. There are some restrictions on T, which are explained in section 9.5.1.

Here is how to make the previous example work in parallel:

```
//correct routine to calculate max norm of resid at varNum variable
Real maxNorm(LevelData<FArrayBox>& resid, int varNum)
{
   Real maxnormLocal = 0;
   DataIterator dit = resid.iterator();
   for (dit.reset(); dit.ok(); ++dit)
   {
      maxnormLocal = Max(maxnormLocal, resid[dit()].norm(0, varNum, 1);
   }
   //gather all maxnormLocals onto processor 0
   int srcProc = 0;
   Vector<Real> allMaxNorm(numProc());
   gather(allMaxNorm, maxnormLocal, srcProc);
   Real maxnorm = 0;
   if(procID() == srcProc)
   {
      for(int ivec = 0; ivec < numProc(); ivec++)
      maxnorm = Max(maxnorm, allMaxNorm[ivec]);</pre>
```

```
}

//broadcast the right answer to all procs
broadcast(maxnorm, srcProc);
return maxnorm;
}
```

This example will work in both the serial and parallel cases.

#### 9.5.1 linearIn, linearOut, linearSize

By "linearize," we mean "to convert a data structure into a contiguous block of memory." For either gather<T> or broadcast<T> to work, T must have the following template functions:

- int linearSize<T>(const T& inputT)
  Return the linear size of the object inputT in bytes.
- void linearIn<T>(T& outputT, const void\* const inBuf)
   Initialize the object outputT from the byte stream in inBuf.
- void linearOut<T>(void\* const outBuf, const T& inputT)

Output the object inputT into the byte stream outBuf. The memory for the buffer is assumed to be allocated elsewhere.

Chombo provides these functions for Box, IntVectSet, Real, int and a templated function for any Vector<T> (as long as T has the three functions itself).

## Chapter 10

## Chombo Fortran

#### 10.1 Introduction

The Chombo library is built with the ability to call Fortran routines from C++. There are many reasons to want to do this. For example, one many want to use the more complex data structures that C++ supports but may not want to forfeit the superior floating-point performance that Fortran offers. The details of mixed language programming, however, can be complex and both compiler and platform-dependent. Another complication is that C++ can be written in a dimension-independent form but the syntax of Fortran is intrinsically dimension-dependent. Array access, declaration and looping all require knowledge of the dimensionality of the problem. Chombo Fortran is designed to create abstractions which avoid these problems. Chombo Fortran allows the C++-Fortran programmer many advantages.

- The complicated data structures (classes) provided by Chombo in C++ can be passed to and used in Fortran routines.
- $\bullet$  The name-mangling differences between Fortran and C++ are handled automatically and cleanly.
- Type checking of arguments in calls to Fortran from C++ is handled automatically by the C++ compiler. This makes mixed language code far less error-prone.
- Dimension-independent Fortran code is made possible. This eliminates the maintenance problems associated with having to maintain separate Fortran kernels for simulation codes which differ only in the number of spatial dimensions.
- Very long Fortran argument lists and declarations (due to array specification) are greatly reduced by the Chombo Fortran macros. This makes Chombo Fortran less error-prone and easier to read.

The basic usage pattern is this. One uses Chombo Fortran to declare her subroutine argument lists and local floating point arguments. ChF interprets these macros in the

context of the input dimensionality and precision and creates a Fortran file. ChF also creates a prototype file to be included in the C++ calling file which unravels the compiler and platform-dependence of the Fortran name mangling (so C++ will be able to find the function).

#### 10.2 ChF Fortran macros

There are three classes of Fortran macros in ChF: array declaration, array access and dimension-handling. The array declaration macros are used to specify arguments to Fortran subroutines that will be called from C++. The array access macros are used to reference these arguments in the body of Fortran subroutines. The dimension-handling macros are used in the body of the Fortran subroutines to create dimension-independent code.

## 10.3 dimension-handling macros

The dimension-handling macros are:

- CHF\_DDECL and CHF\_AUTODECL for declaring variables and creating argument lists
- CHF\_DTERM for choosing multiple expressions or statements based on dimension
- CHF\_DINVTERM for choosing multiple expressions or statements based on dimension and reversing their order
- CHF\_DSELECT for choosing one expression or statement based on dimension
- CHF\_AUTOID for setting multiple variables based on dimension
- CHF\_MULTIDO and CHF\_AUTOMULTIDO for handling nested DO loops
- CHF\_ENDDO goes with CHF\_MULTIDO and CHF\_AUTOMULTIDO

CHF\_DDECL[arg0;arg1;arg2] translates to arg0, arg1, arg2 (in three dimensions). This is useful when one needs to declare variables that only exist in a dimension-dependent context. Say, for example, one has SpaceDim components of velocity called (u,v,w) in three dimensions. Since in two dimensions, the third component is not used in the code, one could declare these variables as

```
integer CHF_DDECL[ u; v; w ]
```

to avoid "unused variable" compiler warnings. This macro will respect carriage returns and other white space.

This is also used in creating argument lists for calling other routines. Using the previous example, to call a routine named F00 that expects SpaceDim arguments, one would write the call as

```
call FOO( CHF_DDECL[ u; v; w ] )
```

The CHF\_AUTODECL macro performs a similar function by expanding a root: CHF\_AUTODECL[arg] will expand to arg0, arg1, arg2 (in three dimensions). This can result in more compact code, especially for code intended to support higher dimensionality.

Similarly, CHF\_DTERM[arg0; arg1; arg2] translates to arg0arg1arg2 in three dimensions and arg0arg1 in two dimensions. This is useful if one has code that is dimension-dependent. For example:

```
integer CHF_DDECL[ii;jj;kk]
CHF_DTERM[
ii = CHF_ID(0,idir);
jj = CHF_ID(1,idir);
kk = CHF_ID(2,idir)]
```

This macro will respect carriage returns and other white space.

The CHF\_DINVTERM[arg0; arg1; arg2] macro is a variation on CHF\_DTERM which reverses the chosen arguments. It translates to arg2arg1arg0 in three dimensions and arg1arg0 in two dimensions. This is useful if one has indexing loops in code that is dimension-dependent. For example:

```
integer CHF_DDECL[ii;jj;kk]
CHF_DINVTERM[
do ii = 0,10;
do jj = 0,10;
do kk = 0,10]
```

Like CHF\_DTERM, this macro respects carriage returns and other white space.

The CHF\_DSELECT macro is a variation on CHF\_DTERM. Instead of choosing the arguments from 1 to SpaceDim, it chooses *only* the SpaceDim'th argument. This is useful for expressions that are different for each dimension. For example:

```
rho = CHF_DSELECT[ cos(x) ; sin(x*y) ; cos(x*z)*sin(y*z) ]
```

Like CHF\_DTERM, this macro respects carriage returns and other white space.

The macro CHF\_AUTOID[ii;idir] generates

```
CHF_DTERM[
ii0 = CHF_ID(0,idir);
ii1 = CHF_ID(1,idir);
ii2 = CHF_ID(2,idir)]
```

and can also be called with an additional optional argument, where CHF\_AUTOID[ii;idir;factor] generates

```
CHF_DTERM[
ii0 = factor*CHF_ID(0,idir);
ii1 = factor*CHF_ID(1,idir);
ii2 = factor*CHF_ID(2,idir)]
```

CHF\_MULTIDO and CHF\_AUTOMULTIDO are used to iterate over a box in a dimension-independent fashion by setting up nested Fortran DO loops. CHF\_ENDDO is used to terminate those DO loops correctly. Specifically, CHF\_MULTIDO[box;i;j;k] will generate a DO loop for i nested inside a DO loop for j and, in 3D, this will be nested inside a DO loop for k. The i loop will go from the first element of the low corner of box to the first element of the high corner of box. Similarly, the j loop will use the second element and, in 3D, the k loop will use the third element. CHF\_ENDDO will end all the DO loops set up by CHF\_MULTIDO.

CHF\_MULTIDO can also be used to iterate with a stride. The syntax for this is CHF\_MULTIDO[box;i;j;k;2], where the "2" could be any integer constant except 0. A negative stride will make the loop iterate backward in each dimension (from the high corner to the low corner). Be warned that using a variable name instead of an integer constant will not produce the desired result because ChomboFortran will just think you've coded a 4-dimensional loop so it will ignore the last variable.

Here is an example using these macros:

```
subroutine LOOP(CHF_FRA1[array],CHF_BOX[box])
integer CHF_DDECL[i;j;k]
integer productsum

productsum = 0
CHF_MULTIDO[box;i;j;k]
  productsum = productsum + CHF_DTERM[i;*j;*k]
  array(CHF_IX[i;j;k]) = productsum
CHF_ENDDO
return
end
```

The other sections contain exact definitions of the other macros used in this example.

The CHF\_AUTOMULTIDO macro also sets up nested loops, but constructs the indices of the loops based on a root. Specifically, CHF\_AUTOMULTIDO[box;i] is the same as CHF\_MULTIDO[box;i0;i1;i2]. Strides are also supported, so CHF\_AUTOMULTIDO[box;i;i2] is the same as CHF\_MULTIDO[box;i0;i1;i2;2] (note that the indices start with 0 instead of 1, which is consistent with the conventions elsewhere in ChomboFortran).

Here is the previous example written using the AUTO macros:

```
subroutine LOOP(CHF_FRA1[array],CHF_BOX[box])
```

```
integer CHF_AUTODECL[i]
integer productsum

productsum = 0
CHF_AUTOMULTIDO[box;i]
  productsum = productsum + CHF_DTERM[i0;*i1;*i2]
  array(CHF_AUTOIX[i]) = productsum
CHF_ENDDO

return
end
```

#### 10.4 Declaration macros

The declaration macros are used inside Fortran SUBROUTINE statements (in the argument list) to specify the types of the arguments to the subroutine.

The ChF system automatically generates type declaration statements for the variables named in ChF declaration macros so explicit declarations statements for these variables are unnecessary and will likely cause compilation errors.

The declaration macros can be used to declare variables of the basic data types (INTEGER and REAL\_T) and variables corresponding to Chombo C++ classes (Box, FArrayBox and IntVect, RealVect, Vector). Variables of the basic types can be scalars or 1D arrays (CHF\*1D macros). Variables of FArrayBox type can have single or multiple components (CHF\*F\* macros).

The macros automatically create and declare all the extra arguments related to array sizes that are needed. The ChF access macros can be used to access these variables. For example, the macro CHF\_LBOUND[ A;1 ] would return the lowest index of the array A in the second dimension (dimensions are counted starting at 0). As a special case, CHF\_UBOUND[ V ] is the same as CHF\_UBOUND[ V;0 ] and is used with Vectors and 1D arrays of basic data types.

The "\_CONST" qualifier in the macro names indicates that the variable named in the macro is not modified in the Fortran subroutine. This form of the macros should be used when the C++ variable is declared 'const'. This has no direct effect on the Fortran code or its execution, but it does affect the C++ code that calls the Fortran subroutine and the C++ prototype that is automatically-generated by ChF.

The following is the complete list of ChF Fortran declaration macros and their uses.

- CHF\_INT[ $\langle arq \rangle$ ] Declare a scalar integer argument.
- CHF\_CONST\_INT[ $\langle arg \rangle$ ] Declare a read-only scalar integer argument.
- CHF\_REAL [ $\langle arg \rangle$ ] Declare a scalar floating point argument.

- CHF\_CONST\_REAL [< arg>] Declare a read-only scalar floating point argument.
- CHF\_COMPLEX[< arg >] Declare a scalar complex argument.
- CHF\_CONST\_COMPLEX[<arg>] Declare a read-only scalar complex argument.
- CHF\_REALVECT[< arg>] Declare a real vector of SpaceDim length argument (indices go from 0 to SpaceDim-1).
- CHF\_CONST\_REALVECT[< arg>] Declare a constant real vector of SpaceDim length argument (").
- CHF\_INTVECT[< arg>] Declare an integer vector of SpaceDim length argument (").
- CHF\_CONST\_INTVECT[< arg>] Declare a constant integer vector of SpaceDim length argument (").
- CHF\_I1D[< arg >] Declare a C array of integers (indices go from 0 to CHF\_UBOUND[< arg >]).
- CHF\_CONST\_I1D[<arg>] Declare a read-only C array (").
- CHF\_R1D[< arg >] Declare a C array of reals (").
- CHF\_CONST\_R1D[<arg>] Declare a read-only C array of reals (").
- CHF\_VI[<arg>] Declare a Chombo Vector<int>.
- CHF\_CONST\_VI[<arg>] Declare a read-only Chombo Vector<int>.
- CHF\_VR[<arg>] Declare a Chombo Vector<Real>.
- CHF\_CONST\_VR[ $\langle arg \rangle$ ] Declare a read-only Chombo Vector $\langle$ Real $\rangle$ .
- CHF\_VC[<arg>] Declare a Chombo Vector<Complex>.
- CHF\_CONST\_VC[<arg>] Declare a read-only Chombo Vector<Complex>.
- CHF\_FIA[< arq>] Declare a multi-component integer C++ BaseFab argument.
- CHF\_CONST\_FIA[<arg>] Declare a read-only multi-component integer BaseFab argument.
- CHF\_FRA[<arq>] Declare a multi-component floating point BaseFab argument.
- $\bullet$  CHF\_CONST\_FRA [< arg>] Declare a read-only multi-component floating point Base-Fab argument.
- CHF\_FIA1[<arq>] Declare a single-component integer BaseFab argument.

- CHF\_CONST\_FIA1[<arg>) Declare a read-only single-component integer BaseFab argument.
- CHF\_FRA1[<arg>] Declare a single-component floating point BaseFab argument.
- CHF\_CONST\_FRA1[< arg>] Declare a read-only single-component floating point BaseFab argument.
- CHF\_BOX[< arg >] Declare a Box argument. Boxs are always read-only.

So a typical subroutine declaration would look like this:

```
subroutine TYPICAL(
& CHF_FRA[fab],
& CHF_CONST_FRA[constfab],
& CHF_BOX[region],
& CHF_CONST_REAL[dx],
& CHF_INT[intflag])
```

This routine takes two floating point BaseFabs (one constant), a box, a constant floating point scalar and an integer. Keep in mind that this is still Fortran. All arguments are still being sent as pointers so they can be changed in the Fortran code. The CONST modifier of the declaration just adds a const to the C++ prototype to allow the user to send const C++ variables without the C++ compiler complaining.

Chombo Fortran preprocessing of arguments can be disabled by adding the comment ! NO\_CHF to the end of the line with the subroutine statement. In a ".ChF" file, this should only be used where absolutely necessary (an example would be an internal procedure as shown in section 10.9.4). Ordinary Fortran subroutines should normally be placed in a separate ".F" file.

#### 10.5 Access macros

- CHF\_LBOUND [< arg>; < dim>] Access the lower bound of a BaseFab or Box < arg> in constant dimension < dim>. Returns an integer variable.
- CHF\_UBOUND [<arg>;<dim>] Access the upper bound of a BaseFab or Box <arg> in constant dimension <dim>. Returns an integer variable. Also used to access the upper bound of a 1D array or Chombo Vector, in which case <dim> need not be specified.<sup>1</sup>
- CHF\_NCOMP [< arg>] Access the number of components in the BaseFab < arg>.
   Returns an integer. Note that the components in Fortran code are numbered from 0 to CHF\_NCOMP(< arg>)-1 to be consistent with the requirements of C++.

 $<sup>^{1}</sup>$ the upper bound of a 1D array is always one less than the dimension specified in the C++ call to CHF\_I1D or CHF\_R1D.

- CHF\_IX[< index0>; < index1>; < index2>] Access an element of an array declared with one of the F\*A\* macros.
- CHF\_AUTOIX[<indexRoot>] Access an element of an array declared with one of the F\*A\* macros. Expands the indexRoot so that CHF\_AUTOIX[i] is the same as CHF\_IX[i0;i1;i2].
- CHF\_OFFSETIX[<indexRoot>;<offsetRoot>] Access an element of an array declared with one of the F\*A\* macros. Similar to CHF\_AUTOIX, but with an offset. For example, CHF\_OFFSETIX[i;-ii] expands to be the same as CHF\_IX[i0-ii0; i1-ii1; i2-ii2].
- CHF\_ID(<dim1>,<dim2>) Return 1 when the arguments have the same value. Used with CHF\_IX for accessing "nearby" array elements. Notice that CHF\_ID uses parentheses instead of square brackets and a comma instead of a semicolon. Simply put, CHF\_ID isn't really a macro—it is a 6x6 (to support up to 6D) identity matrix which gets declared in every subroutine. The parentheses are consistent with array access in Fortran,

#### Notes:

- The <arg> macro argument must be a variable that was declared with one of the BaseFab, Box, 1D array or Chombo Vector macros.
- The < dim> macro argument must be an integer constant in the range 0. . . CH\_SPACEDIM-1.
- The < dim1> and < dim2> macro arguments must be integer variables or constants in the range 0...CH\_SPACEDIM-1.
- Only SUBROUTINEs can be called from C++. FUNCTIONs are not supported.
- ullet The dimensions values are 0-based as in C++, not 1-based as is the default for Fortran.

## 10.6 C++ macros

The ChF C++ macros are intended to be used in C++ code that calls Fortran subroutines that have been declared using the ChF Fortran macros. The prototype header file that is automatically generated by the ChF Fortran macros must be #included in any file where the ChF C++ macros are used to call a Fortran subroutine. The name of this header file is of the form "<fortran\_file\_basename>\_F.H", where <fortran\_file\_basename> is the name of the Fortran source code file without the extension. Every Fortran subroutine that is called from C++ must appear in one and only one included prototype header file.

There are two aspects to using the ChF macros to call Fortran subroutines: specifying the name of the Fortran subroutine and specifying the arguments to the Fortran subroutine.

Fortran subroutines must be called from C++ by prefixing the name of the subroutine with FORT\_ and always using uppercase. For example, the Fortran subroutine named "F00" must be called from C++ using the name "F0RT\_F00". Attempts to access the Fortran name directly will fail on some systems because of compiler-dependent inter-language calling conventions.

The C++ prototypes for Fortran subroutines with no arguments will be generated with the keyword "void" in the argument list.

All arguments to a Fortran subroutine called from C++ must be specified in ChF declaration macros. The macro names indicate the data type of the argument and allow the ChF system to generate appropriate dimension-independent code. The macros used in C++ application code should match the macros that appear in the prototypes provided in the \*\_F.H header files, except that macros in application code should use the CHF\_ prefix where the macros used in the prototypes use the CHFp\_ prefix.<sup>2</sup>

Most of the declaration macros come in a CONST and non-CONST form. The CONST form should be used to declare arguments that are not modified by the Fortran subroutine. The Box macro does not have a CONST form because Boxes are assumed to be constant always.

The ChF C++ declaration macros are almost identical in syntax and usage to the Fortran declaration macros. The differences are:

- the C++ macros are case-sensitive.
- the single-component BaseFab macros (CHF\_\*F{I|R}1()) take 2 arguments (BaseFab, component\_number) in C++ and 1 in Fortran,
- the 1D array macros (CHF\_\*1D) take 2 arguments (array, length) in C++ and 1 in Fortran,
- for each Fortran subroutine < proc >, a C++ macro FORT\_< proc > is defined.

#### 10.7 Declaration macros

The C++ declaration macros are those that the application programmer uses to pass variables to Fortran routines from C++.

The following is the complete list of ChF C++ declaration macros and their uses.

- $CHF_INT(\langle arg \rangle)$  Pass a scalar int variable.
- CHF\_CONST\_INT( $\langle arg \rangle$ ) Pass a const scalar int variable.
- CHF\_REAL( $\langle arg \rangle$ ) Pass a scalar Real variable.
- CHF\_CONST\_REAL(< arg>) Pass a const scalar Real variable.

<sup>&</sup>lt;sup>2</sup>application code should never use the CHFp<sub>-</sub> macros directly.

- CHF\_COMPLEX( $\langle arg \rangle$ ) Pass a scalar Complex variable.
- CHF\_CONST\_COMPLEX( $\langle arg \rangle$ ) Pass a const scalar Complex variable.
- CHF\_REALVECT(<arg>) Pass a RealVect variable.
- CHF\_CONST\_REALVECT( $\langle arq \rangle$ ) Pass a constant RealVect variable.
- CHF\_INTVECT(<arg>) Pass a IntVect variable.
- CHF\_CONST\_INTVECT(<arg>) Pass a constant IntVect variable.
- CHF\_I1D( $\langle arq \rangle$ ,  $\langle len \rangle$ ) Pass a 1D array of ints of length  $\langle len \rangle$ .
- CHF\_CONST\_I1D( $\langle arg \rangle$ ,  $\langle len \rangle$ ) Pass a constant 1D array of ints of length  $\langle len \rangle$ .
- CHF\_R1D( $\langle arg \rangle$ ,  $\langle len \rangle$ ) Pass a 1D array of Reals of length  $\langle len \rangle$ .
- CHF\_CONST\_R1D(<arg>,<len>) Pass a constant 1D array of Reals of length <len>.
- CHF\_VI(<arg>) Pass a Vector<int>.
- CHF\_CONST\_VI(<arg>) Pass a constant Vector<int>.
- $CHF_VR(\langle arg \rangle)$  Pass a Vector $\langle Real \rangle$ .
- CHF\_CONST\_VR( $\langle arq \rangle$ ) Pass a constant Vector $\langle Real \rangle$ .
- CHF\_VC(<arg>) Pass a Vector<Complex>.
- CHF\_CONST\_VC(<arg>) Pass a constant Vector<Complex>.
- CHF\_FIA(<arq>) Pass a BaseFab<int>.
- $\bullet$  CHF\_CONST\_FIA(<arg>) Pass a const BaseFab<int> .
- $CHF\_FRA(\langle arg \rangle)$  Pass a BaseFab<Real>.
- CHF\_CONST\_FRA( $\langle arg \rangle$ ) Pass a const BaseFab<Real>.
- CHF\_FIA1( $\langle arg \rangle$ ,  $\langle comp \rangle$ ) Pass a single component of a BaseFab<int>.
- CHF\_CONST\_FIA1(<arg>, <comp>) Pass a single const component of a BaseFab<int>.
- CHF\_FRA1( $\langle arq \rangle$ ,  $\langle comp \rangle$ ) Pass a single component of a BaseFab<Real>.
- CHF\_CONST\_FRA1( $\langle arg \rangle$ ,  $\langle comp \rangle$ ) Pass a single const component of a BaseFab<Real>.

- CHF\_BOX( $\langle arg \rangle$ ) Pass a Box. Boxes are always const.
- FORT\_proc>(...) Call the Fortran subroutine c> with the arguments specified.

The macros CHF(\_CONST)\_FIA, CHF(\_CONST)\_FRA, CHF(\_CONST)\_FIA1, and CHF(\_CONST)\_FRA1 all come in a version with the appendix \_SHIFT and take an extra IntVect describing the amount to shift the associated box before passing the argument to Fortran. This can be used to align box data and/or change the reference frame. No change is required to the corresponding Fortran declaration macro and using the shift macro has no effect on the original data. In other words, there is no need to un-shift. A common use is to shift all boxes to the positive quadrant so that coarsening can be achieved by simple integer division. In the C++ code:

```
const int refRatio = 2;
 Box crBox (-IntVect::Unit, IntVect::Unit);
 Box fnBox = refine(crBox, refRatio);
 FArrayBox crFRA(crBox, 1);
  crFRA.setVal(0.);
 FArrayBox fnFRA(fnBox, 1);
  fnFRA.setVal(1.);
  const IntVect crShiftToZero = crBox.smallEnd();
  const IntVect fnShiftToZero = scale(crShiftToZero, refRatio);
  FORT_SUMFINE(CHF_FRA1_SHIFT(crFRA, 0, crShiftToZero),
               CHF_BOX_SHIFT(fnBox, fnShiftToZero),
               CHF_CONST_FRA1_SHIFT(fnFRA, 0, fnShiftToZero),
               CHF_CONST_INT(refRatio));
}
In the Fortran code:
      subroutine SUMFINE(
     &
       CHF_FRA1[crFRA],
      CHF_BOX[fnBox],
       CHF_CONST_FRA1[fnFRA],
         CHF_CONST_INT[refRatio])
      integer CHF_AUTODECL[iFn]
      integer CHF_AUTODECL[iCr]
      CHF_AUTOMULTIDO[fnBox;iFn]
С
         Coarsen iFn
         CHF_DTERM[
```

## 10.8 Language support

Chombo Fortran supports the Fortran standard language with a few exceptions. The exceptions include standard Fortran features that are not supported and an extension to the standard that is required.

Chombo Fortran does not support the following features of the Fortran standard:

- REAL, DOUBLE PRECISION, COMPLEX datatypes. The only floating point datatype
  that is supported is REAL\_T. REAL\_T is a Chombo Fortran extension to the Fortran
  standard.
- Appending "\*<*length*>" to a datatype is not supported. This is not standard Fortran, but is a common extension.
- Non-void functions are not supported by Chombo Fortran. Only subroutine statements are supported and those are only allowed with Chombo Fortran macros as arguments.

The code generated by the Chombo Fortran preprocessor conforms to the Fortran standard (ISO/IEC 1539:1991, ANSI X3.198-1992) with the following exceptions:

- The code produced by ChF may violate the Fortran standard maximum number of
  continuation lines in a statement (19). If this occurs, it will be necessary to provide
  a compiler option to increase the limit or change the original Fortran code so that
  it produces fewer continuation lines, usually by breaking a single statement into
  several separate statements.
- Chombo Fortran does not support input and output to the standard units (i.e., 5,6, "\*") on all combinations of C++ and Fortran compilers. Input and output to files should work correctly in all systems. This problem is a fundamental one of mixed-language programming and cannot be solved in any kind of a general way.

A special subroutine is provided which allows the Fortran code to print a special message and terminate execution of the program. This subroutine interfaces with the MayDay class in the Chombo C++ library. The subroutine has two versions, named MAYDAY\_ERROR and MAYDAY\_ABORT.

 The code generated for any ChomboFortran subroutine will contain an IMPLICIT NONE statement so this statement should not be used in the source code. As a result, all variables used in the subroutine must be explicitly declared else the code will not compile successfully.

### 10.9 Examples

#### 10.9.1 Dot Product Example

This routine multiplies each point of one BaseFab with the corresponding point the other BaseFab over the input Box and puts the result in the input Real.

```
subroutine DOTPRODUCT(
&
      CHF_REAL[dotprodout],
      CHF_CONST_FRA[afab],
&
&
      CHF_CONST_FRA[bfab],
      CHF_BOX[region])
integer CHF_DDECL[i;j;k]
 integer nv,ncomp
ncomp = CHF_NCOMP[afab]
 if(ncomp .ne. CHF_NCOMP[bfab]) then
     call MAYDAY_ERROR()
endif
dotprodout = zero
do nv = 0, ncomp-1
   CHF_MULTIDO[region; i; j; k]
   dotprodout = dotprodout +
         afab(CHF_IX[i;j;k],nv)*
&
         bfab(CHF_IX[i;j;k],nv)
&
    CHF_ENDDO
enddo
return
end
```

### 10.9.2 RealVect and IntVect Example

```
subroutine realVectTest(CHF_REALVECT[foo])

CHF_DTERM[
foo(0) = 1.0;
foo(1) = 2.0;
foo(2) = 3.0]

return
end
subroutine intVectTest(CHF_INTVECT[foo])

CHF_DTERM[
foo(0) = 1;
foo(1) = 2;
foo(2) = 3]

return
end
```

#### 10.9.3 Laplacian Example

This subroutine produces a standard (3 point in one dimension, 5 point in two dimensions, and 7 point in three dimensions) discrete Laplacian of the input BaseFab over the input box.

```
subroutine OPERATORLAP(
     CHF_FRA[lofphi],
&
&
      CHF_CONST_FRA[phi],
&
      CHF_BOX[region],
      CHF_CONST_REAL[dx])
&
REAL_T dxinv, lphi
 integer n,ncomp,idir
 integer CHF_DDECL[ii,i;jj,j;kk,k]
ncomp = CHF_NCOMP[phi]
 if(ncomp .ne. CHF_NCOMP[lofphi]) then
    call MAYDAY_ERROR()
 endif
dxinv = one/(dx*dx)
 do n = 0, ncomp-1
```

```
CHF_MULTIDO[region; i; j; k]
    lphi = zero
    do idir = 0, CH_SPACEDIM-1
    CHF_DTERM[
      ii = CHF_ID(idir, 0);
      jj = CHF_ID(idir, 1);
      kk = CHF_ID(idir, 2)]
      lphi = lphi +
&
           ( (phi(CHF_IX[i+ii;j+jj;k+kk],n)
&
           - phi(CHF_IX[i ;j ;k
                                       ],n))
           - (phi(CHF_IX[i
                           ;j ;k ],n)
&
           - phi(CHF_IX[i-ii;j-jj;k-kk],n))
&
           )*(dxinv)
&
    enddo
    lofphi(CHF_IX[i;j;k],n) = lphi
   CHF_ENDDO
 enddo
return
```

This can also be expressed in a simpler way making full use of the AUTO and OFFSET macros:

```
& CHF_FRA[lofphi],
& CHF_CONST_FRA[phi],
& CHF_BOX[region],
& CHF_CONST_REAL[dx])

REAL_T dxinv,lphi
integer n,ncomp,idir

integer CHF_AUTODECL[i], CHF_AUTODECL[ii]

ncomp = CHF_NCOMP[phi]
if(ncomp .ne. CHF_NCOMP[lofphi]) then
    call MAYDAY_ERROR()
endif

dxinv = one/(dx*dx)
```

subroutine OPERATORLAP(

```
do n = 0, ncomp-1
    CHF_AUTOMULTIDO[region; i]
    lphi = zero
    do idir = 0, CH_SPACEDIM-1
       CHF_AUTOID[ii;idir]
       lphi = lphi +
            ( (phi(CHF_OFFSETIX[i;+ii],n)
&
&
            - phi(CHF_AUTOIX[i],n))
&
            - (phi(CHF_AUTOIX[i],n)
            - phi(CHF_OFFSETIX[i;-ii],n))
&
&
            )*(dxinv)
    enddo
    lofphi(CHF_AUTOIX[i],n) = lphi
    CHF_ENDDO
 enddo
 return
 end
```

#### 10.9.4 Internal Procedure Example

This example demonstrates use of an internal procedure in a "\*.ChF" file. Ordinary Fortran arguments could have also been declared for the internal procedure.

```
subroutine testNoChFCall(
    CHF_BOX[box])

integer CHF_AUTODECL[i]
integer c

c = 0
call countCells
write(*,*) c
call countCells
write(*,*) c
return

contains
subroutine countCells ! NO_CHF
```

```
CHF_AUTOMULTIDO[box;i]

c = c+1

CHF_ENDDO

end subroutine

end
```

#### 10.10 Landmines

This section is intended to point out some known uses of Chombo Fortran that will result in errors.

- Be aware that using C++ and Fortran together defeats most bounds checkers. If you step out of bounds in a Fortran, as a rule, your bounds checker will not save you. This holds for both Fortran and Chombo Fortran.
- Combining Fortran and Chombo Fortran in the same file is a bad idea, with the exception of internal procedures as shown in the example of section 10.9.4. The Chombo Fortran parser keys on the word "subroutine," and dissects the argument list as described above. If ordinary Fortran subroutines are put into a Chombo Fortran file, the parser will fail to produce correct code. To use both Fortran and Chombo Fortran in the same application, put them into separate files. The Chombo makefile system recognizes files with ".F" extensions as Fortran and files with ".ChF" extensions as Chombo Fortran files.
- Send constants to Chombo Fortran (or plain Fortran, for that matter) using temporary variables. The C++ macros in Chombo Fortran are precisely that-macros. If you insert an explicit constant in a Chombo Fortran call, the macro will simply try to take the address of the explicit constant, resulting in undefined behavior. Say you want to send the number four to a Chombo Fortran routine. Here are both the incorrect and correct ways to do so.

```
//error! gets tranlated to senseless:
//myfunc_(&4);
FORT_MYFUNC(CHF_CONST_INT(4));

//correct. address sent to Fortran is legal. This gets translated to
//myfunc_(&ivar);
int ivar = 4;
FORT_MYFUNC(CHF_CONST_INT(ivar));
```

The exception to this is the 2nd argument to the CHF\_\*1D macros, which can be a literal constant.

- The arguments of a ChF Fortran macro must be enclosed in square brackets and separated by semicolons. Commas between the brackets will pass through to Fortran, as in the example in section 10.9.3 where CHF\_DDECL[ii,i;jj,j;kk,k] translates to ii,i,jj,j,kk,k or ii,i,jj,j. The one apparent exception is CHF\_ID(<dim1>,<dim2>), but as noted above, CHF\_ID is a matrix, not a macro.
- Chombo Vectors and 1D arrays always start at index 0. You cannot call CHF\_LBOUND on a Vector or 1D array. The value returned from CHF\_UBOUND on a 1D array will always be one less than the length value passed as the 2nd argument in the C++ call to CHF\_\*1D.

## Chapter 11

## Multidimensional Chombo

This section describes the Chombo support for solving multidimensional systems of equations. "Multidimensional" in this sense means a system in which part of the solution approach is in an N-dimensional space, while another part is in an M-dimensional space, where  $N \neq M$ .

## 11.1 Namespace implementation

To implement multidimensional Chombo, we make use of namespaces to encapsulate each dimensionality. If Chombo is compiled with multidimensional support, then a version of the Chombo libraries is complied for each dimension and placed in a corresponding namespace. This allows the different dimensionalities to co-exist in a single build while avoiding naming collisions. Note that in order for Chombo's multidimensional build system to work correctly with your application code, you *must* include the namespace headers and footers in your source and header files (see Section 1.2.8). The lone exception is for code in which inter-dimensional transfers take place, which must explicitly reference each dimensionality by name.

### 11.2 Transdimensional utilities

To move between the different dimensionalities, we will need slicing and injection capability. This functionality is found in /Chombo/lib/src/MultiDim

## 11.2.1 struct SliceSpec

A SliceSpec specifies a slice in space, using a direction d and a position p. The SliceSpec class is actually a dimensional object, and is a part of the BoxTools library. A DIM-dimensional SliceSpec defines a (DIM-1)-dimensional slice through DIM-Dimensional space, with the d-th coordinate index set to p. When referring to an IntVect,

the d index defines which element will be removed. For a Box and related objects, d refers to the dimension which will be removed during a slicing operation.

#### 11.2.2 Slicing

To move to a lower-dimensional space, we use the slicing functionality found in Slicing.H.transdim, which supports moving from the D space to the D-1 space. The Slicing code is heavily templated to work in a multidimensional setting. If DLo is the lower-dimension namespace and DHi is the higher dimension namespace (for example, DLo = D2 and DHi = D3), then the following functions are available.

Makes a\_to the indicated (by a\_spec) slice of a\_from; a\_from must be of dimension one larger then a\_to.

Defines a\_to as a\_from with its a\_slicespec.direction-th coordinate missing; a\_to is one dimension lower, e.g. if a\_from is a D3::Box, then a\_to is a D2::Box. If a\_outofbounds isn't NULL, we set it to true if a\_slicespec.position is outside of a\_from, along the a\_slicespec.direction-th axis.

• DLo::ProblemDomain

Functions similarly to the sliceBox function, but also maintains periodicity information appropriately.

• template<T>

Sets a\_to to be a slice of a\_from, along the a\_slicespec.direction—th axis at the a\_slicespec.position—th position. Data from the slice of a\_from is copied to the destination BaseFab> a\_to. As usual, a\_to is one dimension lower than a\_from; if a\_from is a D3::BaseFab, a\_to is a D2::BaseFab. It is an error if a\_slicespec.position is outside of a\_from.

• void

template<class T> void

• void

Template specialization for LevelData<FluxBox> Makes a\_from a slice of a\_to at a\_slicespec.position along the a\_slicespec.direction-th axis. If a\_to is defined, we use its DisjointBoxLayout, otherwise we create an appropriate new DisjointBoxLayout for it.

#### 11.2.3 Injection

To move from the D space to the D+1 space, we use the injection functionality in Injection.H.transdim. In general, the inject functions simply define a higher-dimensional version of the lower-dimensional input, with the "new" dimension defined by the SliceSpec::direction argument d. For the case of an IntVect, the value of the newly added dimension is set to SliceSpec::position p. For a Box and containers defined on a Box or DisjointBoxLayout, the higher-dimensional destination is defined to be one-cell wide in the new direction, and is located at  $i_d = p$ .

Makes a\_to like a\_from, only one dimension higher. The "new" transverse dimension is a\_spec.direction, and the value at that dimension is a\_spec.position.

Sets a\_to to be a\_from with an extra dimension - a\_slicespec.direction - in which it's just one cell thick and has coordinate value a\_slicespec.position.

• template<typename T>

Sets a\_to to be a\_from with an extra dimension - a\_slicespec.direction - in which it's just one cell thick and has coordinate value a\_slicespec.position.

 void injectDisjointBoxLayout(DHi::DisjointBoxLayout& a\_to, const DLo::DisjointBoxLayout& a\_from, const DHi::SliceSpec& a\_slicespec)

Sets a\_to to be a\_from with an extra dimension - a\_slicespec.direction - in which it's just one cell thick and has coordinate value a\_slicespec.position. Data values are copied from a\_from to fill a\_to.

• template <class T>

Sets a\_to to be a\_from with an extra dimension - a\_slicespec.direction - in which it's just one cell thick and has coordinate value a\_slicespec.position. If a\_to is defined, we use its DisjointBoxLayout, otherwise we give it a DisjointBoxLayout with the same assignment-to-processors as a\_from has. Data values are copied from a\_from to fill a\_to.

Version of injection for LevelData<FluxBox>. Sets a\_to to be a\_from with an extra dimension - a\_slicespec.direction - in which it's just one cell thick and has coordinate value a\_slicespec.position. If a\_to is defined, we use its DisjointBoxLayout, otherwise we give it a DisjointBoxLayout with the same assignment-to-processors as a\_from has. Data values are copied from a\_from to fill a\_to.

#### 11.2.4 The ReductionCopier class

The ReductionCopier is a specialized Copier designed to support a reduction from a higher-dimensional DisjointBoxLayout to a smaller-dimensional one by copying all of the data in the transverse direction to the destination boxLayout. It is assumed that this will be used with a different sort of LDOperator (such as the SumOp described below), since a simple copy operation wouldn't make much sense.

Essentially, the ReductionCopier computes and stores the intersection list required to do a copyTo operation which copies all of the data in src into dest, assuming that an intersection can be found by shifting the location of a data point in src in the transverse direction.

Note that this class operates entirely in a single dimensional namespace. The usage pattern is to perform the reduction operation in the higher-dimensional space, then use a slicing operation to move to a lower-dimensional space. A simple usage example is shown in figure 11.1, with a corresponding code example in figure 11.2.

Full constructor and define functions (constructor simply calls define)

- a\_level source DisjointBoxLayout
- a\_dest destination boxes
- a\_destGhost number of ghost cells to be filled around the destination grids
- a\_transverseDir the direction of the reduction operation
- a\_exchange if true, this copier is being defined for an exchange, rather than a copyTo operation.

### 11.2.5 The SpreadingCopier class

The SpreadingCopier is a specialized Copier designed to support a reduction from a lower-dimensional DisjointBoxLayout to a higher-dimensional one by copying all of the data in the transverse direction to the destination boxLayout. It is assumed that this will be used with an appropriate LDOperator such as the SpreadingOp described below.

Essentially, the SpreadingCopier computes and stores the intersection list required to do a copyTo operation which copies the data in src into dest, assuming that an intersection can be found by shifting the location of a data point in src in the transverse direction.

Like the ReductionCopier (only in reverse) this class operates entirely in a single dimensional namespace. The usage pattern is to use an injection operation to move the

data from the lower-dimensional space to the higher-dimensional space, then perform the spreading operation in the higher-dimensional space.

Full constructor and define functions (constructor simply calls define)

- a\_level source DisjointBoxLayout
- a\_dest destination boxes
- a\_destGhost number of ghost cells to be filled around the destination grids
- a\_transverseDir the direction of the reduction operation
- a\_exchange if true, this copier is being defined for an exchange, rather than a copyTo operation.

#### 11.2.6 The SumOp class

The SumOp class is an instance of LDOperator<FarrayBox> which performs a summing operation of the data in src in the summingDir direction, multiplies by the scale, and places the sum in the corresponding location in dest. Note that scale is a public member function, and so may be set without an access function.

```
    SumOp(int a_summingDir);
    Constructor - sets scale to one.
    - a_summingDir - direction in which to sum
    void op(FarrayBox& dest,
        const Box& RegionFrom,
        const Interval& Cdest,
        const Box& RegionTo,
        const FArrayBox& src,
```

const Interval& Csrc) const;

Computes componentwise sum of src over the RegionFrom in the summingDir direction, multiplies by scale and places the result in dest in the RegionTo.

- dest destination FAB
- RegionFrom Region in src over which to compute sum
- Cdest Interval in dest into which sum will be computed.
- RegionTo Region in dest into which the sum will be placed.
- src source FAB
- Csrc interval in src over which sum will be computed (must be the same size as Cdest).
- virtual void linearIn(FArrayBox& arg, void\* buf, const Box& R, const Interval& comps) const;

Linearization function.

#### 11.2.7 The SpreadingOp class

The SpreadingOp class spreads the data in src along the summingDir direction, multiplying by scale, and placing the resulting values in the corresponding locations in dest. Essentially the inverse of the SumOp class. Like the SumOp, scale is a public data member.

- SpreadingOp(int a\_spreadingDir);
   Defining constructor. Sets default value for scale to 1.0.
  - a\_spreadingDir direction in which to spread data.
- void op(FArrayBox& dest, const Box& RegionFrom, const Interval& Cdest, const Box& RegionTo, const FArrayBox& src, const Interval& Csrc) const;

Perform spreading operation — for each IntVect in RegionFrom, scale value in src by scale and then place scaled value into dest for each IntVect in RegionTo which corresponds to a translation from the source location along the spreadingDir direction. This is a componentwise operation, so the scaled values of the first component of Csrc will be placed in the first component of Cdest, etc.

- dest
- RegionFrom
- Cdest Interval in dest over which operation is performed.

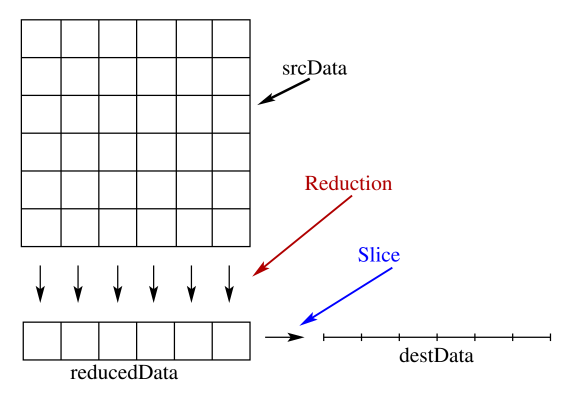


Figure 11.1: schematic of reduction operations carried out by the code in Figure 11.2

- RegionTo
- src
- Csrc Interval in src over which operation is performed. Must have the same size as Cdest.
- virtual void linearIn(FArrayBox& arg, void\* buf, const Box& R, const Interval& comps) const;

Linearization function.

```
// function to reduce a 2D LevelData<FArrayBox> to 1D by
// summing over the y-direction
void reduce2DTo1D(D1::LevelData<D1::FArrayBox> >& destData,
                D2::LevelData<D2::FarrayBox> >& srcData)
{
 // create 2D version of 1D boxes
 D2::LevelData<D2::FarrayBox> sliceGrids;
 D2::SliceSpec slice(1,0);
 injectDisjointBoxLayout(sliceGrids, srcData.getBoxes(), slice);
 // define sliced data holder
 D2::LevelData<D2::FArrayBox> reducedData(sliceGrids, srcData.nComp);
 // define ReductionCopier to compute intersections (sum in the y-direction)
 int transverseDir = 1;
 ReductionCopier reduceCopier(srcData.getBoxes(), sliceGrids, 1);
 SumOp op(transverseDir);
 op.scale = 1.0;
 // do summing operation -- sums data in srcData along lines in the
 // y-direction and places sum in reducedData
 srcData.copyTo(srcData.interval(),
                 reducedData, reducedData.interval(),
                 reduceCopier, op);
 // finally, take the data in reducedData (which is a 2D object)
 // and slice to 1D
 sliceLevelData(destData, reducedData, slice);
}
```

Figure 11.2: Code fragment illustrating use of ReductionCopier

# 11.3 Phase field example

As a simple example of a multidimensional application, we solve the phase space example from [31]:

$$\frac{\partial f}{\partial t} + \frac{\partial vf}{\partial x} + \frac{\partial af}{\partial v} = 0$$

$$a = -\frac{\partial \phi}{\partial x},$$
(11.1)

$$\frac{\partial^2 \phi}{\partial x^2}(x,t) = 4\pi G \int f(x,v,t)dv, \qquad (11.2)$$

where G is the gravitational constant ( $\pi$  for the example in [31]). This example may be found in releasedExamples/MultiDimPhase in the Chombo distribution.

#### 11.3.1 Algorithm

At time t we have  $f_{ij}^n = f(x_i, v_j, t^n)$ .

#### 1. Compute phase velocity a

- (a) Compute rhs for Poisson solve:  $\rho_i = \sum_{j=0}^{j < N_j} f_{i,j}^n \Delta v$
- (b) do 1D Poisson solve for  $\phi(x,t)$ : Solve  $L_i^{1D}\phi=\rho_i$
- (c) Compute a in 1D:  $a_i = G_i^{1D,CC} \phi$
- (d) Spread a to 2D mesh:  $a_{ij} = a_i$ .

#### 2. Advect f

- (a) predict  $f_{face}^{n+\frac{1}{2}}$  using AMRGodunovUnsplit predictor
- (b) update f:

$$f_{ij}^{n+1} = f_{ij}^{n} - \frac{\Delta t}{\Delta x} \left( v_j f_{i+\frac{1}{2},j}^{n+\frac{1}{2}} - v_j f_{i-\frac{1}{2},j}^{n+\frac{1}{2}} \right) - \frac{\Delta t}{\Delta v} \left( a_i f_{i,j+\frac{1}{2}}^{n+\frac{1}{2}} - a_i f_{i,j-\frac{1}{2}}^{n+\frac{1}{2}} \right)$$

# 11.4 Building the multidim example

Building a multidim Chombo application like the MultiDimPhase example is somewhat different from the standard Chombo build process, since it relies on a build shell script to orchestrate the build. This script has been incorporated into the build system rules, so building is fairly straightforward, but does require two separate GNUmakefiles, as may be found in releasedExamples/MultiDimPhase/exec. The primary makefile (GNUmakefile in the MultiDimPhase example) merely specifies the location of CHOMBO\_HOME, the min and max dimensions used by the given compilation, and the name of the second makefile.

The second makefile (GNUmakefile.multidim in the MultiDimPhase example, specifies which directories should be compiled in which dimensionalities (including the "multi-Dim" dimensionality which is where any inter-dimensional transfers happen and which is compiled without a particular SpaceDim) are defined in the 1dsrc\_dirs, 2dsrc\_dirs, 3dsrc\_dirs, 4dsrc\_dirs, 5dsrc\_dirs, 6dsrc\_dirs and mdsrc\_dirs (for the multi-dimensional source files) variables in the local GNUmakefile.multidim in the exec directory. At this point, the only real difference when invoking the makefile is that no dimensionality should be specified (you can, but it won't do anything):

```
obtain Chombo
edit Chombo/lib/mk/Make.defs.local as usual
cd Chombo/releasedExamples/MultiDimPhase/exec
gmake all
(go get a cup of coffee, and maybe some lunch...)
./phase<config>.ex inputs
```

Note that the executable doesn't have a "<DIM>d" in its name – just look for a phase\*.ex executable.

#### 11.5 Some Pointers That Could Save You Some Heartache

Here a few non-obvious things that you can check to make compiling a multi-dimensional example somewhat less painful.

- Have your code organized by directories that you can easily categorized as dimensional or non-dimensional.
- In all your files that are NOT your main driver, be sure have (after all the includes and before all the code) #include "NamespaceHeader.H" and (after all the code) #include "NamespaceFooter.H" (and remember that sis never capitalized in these declarations).
- In your file that includes main, do not use the namespace headers and footers. Instead use #include" UsingNamespace.H".
- Do not try to link with Chombo Fortran functions directly from the library. It messes with the dependencies and gives helpful errors such as "cannotfind a rule to generate...". Either write a c-code wrapper (in the library) or copy the relevant subroutines to your application and change the names.
- Once the libraries are compiled (and you have stopped changing them) use "make noLibs" to save yourself a lot of time in compiling.
- You might be tempted to do something sensible like make a class with mixeddimension member data. Don't do that. You will never get it to compile. Make sure your interfaces are single dimension.

# Chapter 12

# Particles in Chombo

The previous sections of this manual focus on tools for solving PDEs using Eulerian discretizations of space, in which the underlying grids on which the solution is represented are stationary. Many problems, however, are better suited to Lagrangian - or Particle - methods, in which the mesh points themselves move with time. Each of these points carries with it a position coordinate, as well as any number of other properties, which may or may not also change as the simulation advances. The simplest example of a problem amenable to solution by particle methods is the gravitational N-body problem, in which a collection of N self-gravitating particles evolves under Newton's laws of classical mechanics. Other applications of particle methods include vortex methods for incompressible fluid flow, molecular dynamics, Lagrangian flow tracers, and Particle-in-Cell methods for solving the Vlasov-Poisson equation for plasma physics or cosmology.

Chombo provides a number of tools for implementing these methods. These tools have been designed so that, as far as practical, Chombo code for doing particle simulations looks like Chombo code for doing Eulerian simulations. At the data container level or higher, all the particle classes have a straightforward analog to a class in traditional, rectangular array-based Chombo. Included are classes that represent single particles, container classes analogous to FArrayBox for storing and spatially sorting particle data, and distributed data containers analogous to BoxLayoutData to facilitate SPMD-based parallel programming. As with BoxLayoutData, these containers have been designed so that all the necessary data communication happens internally, so that the user rarely has to invoke MPI calls directly. In in this section, we describe these tools and explain how to use them in your own simulations. Only the basic components of each class's API are described here, for the full interface, please consult the Doxygen documentation.

## 12.1 Particle Datatypes

The base class for particle data is the BinItem - an item capable of being assigned to a bin based on its spatial location. The BinItem class has only one data member - a SpaceDimdimensional RealVect that stores the item's physical position. The most important parts

of the public interface for BinItem are described below. The full constructor looks like:

BinItem(const RealVect& a\_position)
 Full constructor. Places the BinItem object in a usable state.

#### **Arguments:**

a\_position The position coordinate of this object.

It also includes methods for getting and setting the position:

void setPosition(const RealVect& a\_position)
 Change the position attribute.

#### **Arguments:**

- a\_position The new position coordinate of this object.
- void setPosition(const Real a\_position, const int a\_dimension)

  Change one component of the position attribute.

#### **Arguments:**

- a\_position The new position component.
- a\_dimension The dimension to change.
- RealVect& position()
   Return a reference to the position.
- const RealVect& position()
   Return a const reference to the position.
- Real position(const int a\_dir)
   Get one component of the position.

#### **Arguments:**

a\_dir The dimension to get.

and equality and inequality operators:

- virtual bool operator==(const BinItem& a\_p) const Check whether two BinItems have equal data.
- virtual bool operator==(const BinItem\* a\_p) const Same as the above, defined for a pointer.
- virtual bool operator!=(const BinItem& a\_p) const
   Check whether two BinItems don't have equal data.

Finally, it's also necessary to define *linearization* functions, so that each particle knows how to write a serial representation of itself into a chunk of memory. These functions won't be used commonly in application code, but are used internally for IO and data communication. The linearization routines are:

- virtual int size() const
   Return the number of bytes in the linear representation of this object.
- virtual linearOut(void\* a\_buf) const
   Write out a serialized representation of this object into the memory address pointed to by a\_buf.
- virtual linearIn(void\* a\_buf)
   Set the data for this object by reading in the serialized representation from the memory address pointed to by a\_buf.

It is expected that users will need to define their own derived particle types based on BinItem that are better suited to their application domain. One example of how to do this provided in the Particle class. In addition to a position, Particle also contains RealVects that store the Particle's velocity and acceleration, as well as a Real attribute that stores the mass. The full constructor for Particle looks like:

 Particle(const Real& a\_mass, const RealVect& a\_position, const RealVect& a\_velocity=RealVect::Zero)

Full constructor. Places the Particle object in a usable state.

#### **Arguments:**

- a\_mass The mass of this particle.
- a\_position The position coordinate of this particle.
- a\_velocity The velocity of this particle. Defaults to zero.

The are also additional methods for getting and setting the mass, acceleration, and so on. For example:

void setMass(const Real a\_mass)
 Set / change the mass attribute.

#### **Arguments:**

- a\_mass The new mass of this object.
- Real& mass()

Retrieve a reference to the mass attribute.

• const Real& mass()

Return a const reference to the mass attribute.

These additional attributes make Particle appropriate for gravitational N-body calculations, or for Vlasov-Poisson simulations in which the forces are gravitational-only, as in cosmological dark matter simulations. Other possibilities include adding a charge attribute for plasma simulations, or giving each particle a unique integer id. Whenever you define a new particle type, the equality / inequality operators and the linearization routines will need to be modified to account for the new data. The Particle class shows one example of how to do this.

#### 12.2 Particle Containers

We also provide templated container classes for storing and sorting particle items based on their position. There are several reasons you might want to do this; for example, to divide the particles among processors based on a spatial domain decomposition, or to compute which particles are "nearby" in a molecular dynamics calculation. Currently, Chombo provides two different types of containers, those based on BinFab<T> and those based on ListBox<T>. We describe each of these, as well as the advantages of each, below.

#### **12.2.1** BinFab<T>

BinFab<T> is a class for storing particle data on the Box level. It works just like an FArrayBox, except that instead of a floating point number, each cell in the Box contains a List<T>. Each of these cells in the BinFab<T> is associated with a region in space, and T-objects belong in the associated List if and only if their position lie within that region. The constructor for BinFab<T> looks like this:

 BinFab(const Box& a\_domain, const RealVect& a\_mesh\_spacing, const RealVect& a\_origin)

Full constructor. Constructs an empty BinFab<T> on the  $Box\ a\_domain$ .

#### **Arguments:**

- a\_domain The Box on which this BinFab<T> is defined.
- a\_mesh\_spacing The physical size of the cells in this BinFab<T>, used for binning.
- a\_origin The origin of the coordinate system, also used for binning.

Since each BinItem or derived particle datatype stores its own physical position, the BinFab<T> must have a coordinate system origin in order to interpret this information properly. BinFab<T> also contains methods for adding items to the container. When one of these is called, the items automatically gets sorted into the proper cell. For example:

virtual void addItem(const T& a\_item)
 Copy one item into the appropriate bin.

#### **Arguments:**

a\_item The T-object that will be copied.

virtual void addItems(const List<T>& a\_list)
 Copy multiple items into the appropriate bins.

#### **Arguments:**

- a\_list The List of T-objects that will be copied.
- virtual void addItemsDestructive(List<T>& a\_list)
   Copy multiple items into the appropriate bins, and remove them for the input list.

#### **Arguments:**

a\_list The List of T-objects that will be transferred.

If any of the particle positions change after them to the container (which they probably will in a time-dependent calculation), there are also reBin() functions to put them all back in the right Lists:

virtual void reBin()
 Sort all the items into the appropriate bins. This version loses all the items not in this BinFab<T>'s domain.

Sort all the items into the appropriate bins. Items that leave the domain defined by a\_valid go into a\_list.

#### **Arguments:**

- a\_lost The List of T-objects lost items will go into.
- a\_valid The Box that defines the valid domain. If this is omitted, the Box used to define the BinFab<T> will be used instead.
- a\_in If this is false, this will put \*valid\* objects into a\_lost, instead.

Note that indexing the BinFab<T> with an IntVect index will allow one to access the List<T> associated with that cell. For example, the code snippet in 12.1 demonstrates how to move all the particles in a BinFab<T> by some amount, and then call the reBin function to make sure they are in the proper cells.

Finally, BinFab<T> can also be used to define Derived classes. For instance, you might want a class that bins particles in velocity space instead of position space. This can be accomplished by deriving from BinFab and re-implementing the reBin and addItem methods as necessary.

```
void moveParticles(BinFab<BinItem>& a_particles,
                   const Real a_dt)
{
 BoxIterator bit(a_particles.box());
 RealVect velocity(D_DECL(1.0,0.5,0.0));
 RealVect amountToMove = a_dt*velocity;
  for (bit.reset(); bit.ok(); ++bit)
   {
      CH_XD::List<BinItem>& thisList = a_particles(bit(),0);
      if (thisList.length() > 0)
        {
          // now loop over the items in the list and move them
          ListIterator<BinItem> lit(thisList);
          for (lit.rewind(); lit; ++lit)
              BinItem& thisParticle = thisList[lit];
              thisParticle.position() = thisParticle.position()
                                      + amountToMove;
            }
        }
   }
  // finally, do rebinning
 a_particles.reBin();
}
```

Figure 12.1: Code snippet to show how move all the particles in a BinFab<T> and re-bin them.

Figure 12.2: Code snippet to show how move all the particles in a BinFab and re-bin them.

#### 12.2.2 Distributed BinFab<T>s

BinFab<T> implements the proper linearization functions so that it can be used with the LevelData and BoxLayoutData containers to store data that is distributed accross processes. That is, you can have a LevelData<BinFab<BinItem> >, and the copyTo and exchange() functions all work as expected, as does indexing into the container using a DataIndex. The code snippet in 12.2 shows how to use the moveParticles function from 12.1 to advance an entire domain's worth of particles in parallel. For simplicity, we assume that the DisjointBoxLayout used to define the LevelData<BinFab<T> > covers the entire domain - i.e. this is a unigrid, domain decomposed calculation. This example also assumes that the LevelData<BinFab<T> > has been defined with one row of ghost cells, and that the time step is limited such that the particles will not move more than one cell in a single step.

#### **12.2.3** ListBox<T>

The BinFab<T> data structure is appropriate when there are many particles in every cell of every box. However, in many applications, this will not be the case. Furthermore, we often do not care where individual particles are found within a single Box, but merely want a convenient way to distribute the particles among the MPI processes. In this case, a ListBox<T> may be more appropriate. a ListBox<T> is like a BinFab<T>, except

that there is only one List<T> for the entire box instead of one for every cell. The ListBox<T> has methods for adding, accessing, and removing Ts from the List, as well as for checking whether the Ts lie within the domain of the defining Box. The (partial) public interface for ListBox<T> is as follows:

#### • BinFab()

Default constructor. The ListBox<T> is left unusable, and define() must be called later.

 BinFab(const Box& a\_domain, const RealVect& a\_mesh\_spacing, const RealVect& a\_origin)

Full constructor. Constructs an empty BinFab<T> on the Box a\_domain.

#### **Arguments:**

- a\_domain The Box on which this BinFab<T> is defined.
- a\_mesh\_spacing The physical size of the cells in this BinFab<T>, used for binning.
- a\_origin The origin of the coordinate system, also used for binning.
- define(const Box& a\_domain, const RealVect& a\_mesh\_spacing, const RealVect& a\_origin)

Define function - same as full constructor.

#### **Arguments:**

- a\_domain The Box on which this BinFab<T> is defined.
- a\_mesh\_spacing The physical size of the cells in this BinFab<T>, used for binning.
- a\_origin The origin of the coordinate system, also used for binning.

undefine()

Free memory, return object to unusable state.

inline List<T>& listItems()

Get a reference to the list of items.

inline const List<T>& listItems() const

Get a reference to the list of items, const version.

 virtual void getItems(List<T>& a\_list, const Box& a\_valid) const

Copy items from this ListBox<T> and put them into a\_list only if they are contained within a\_valid.

#### **Arguments:**

- a\_list The List<T> into which to copy the items.
- a\_valid The Box that defines the valid region.
- virtual void getItemsDestructive(List<T>& a\_list, const Box& a\_valid) const

Copy items from this ListBox<T> and put them into a\_list only if they are contained within a\_valid. This operation removes the items from this ListBox<T>.

#### **Arguments:**

- a\_list The List<T> into which to copy the items.
- a\_valid The Box that defines the valid region.
- virtual void getInvalid(List<T>& a\_list,
   const Box& a\_valid) const

Copy items from this ListBox<T> and put them into a\_list only if they are NOT contained within a\_valid.

#### **Arguments:**

- a\_list The List<T> into which to copy the items.
- a\_valid The Box that defines the valid region.

virtual void getInvalidDestructive(List<T>& a\_list,
 const Box& a\_valid) const

Copy items from this ListBox<T> and put them into a\_list only if they are NOT contained within a\_valid. This operations removes the items from this ListBox.

#### **Arguments:**

- a\_list The List<T> into which to copy the items.
- a\_valid The Box that defines the valid region.
- virtual void addItem(const T& a\_item)

Add an item to this ListBox if it is contained within the defining Box.

#### **Arguments:**

- a\_item The item to add.
- virtual void addItems(const List<T>& a\_list)

Add all the items in a\_list to this ListBox if it is contained within the defining Box.

#### **Arguments:**

- a\_item The item to add.
- virtual void addItemsDestructive(const List<T>& a\_list)

Add all the items in a\_list to this ListBox if it is contained within the defining Box. This operation removes this items from a\_list.

#### **Arguments:**

- a\_item The item to add.
- virtual size\_t numItems()

Returns the number of items in the container.

virtual RealVect& meshSpacing()
 Returns the mesh spacing used for binning.

- virtual const Box& box()
   Returns the Box that defines this ListBox<T>.
- virtual void clear()
   Delete all the items in this ListBox<T> and return to an undefined state.

#### 12.2.4 ParticleData<T>

Internally, data communication must be handled somewhat differently for particle data defined on a set of ListBox<T>s. To facilitate this, we have provided a new data container class: the ParticleData<T>. In addition to representing a collection of ListBox<T>es on a single AMR level, a ParticleData<T> has an *outcast* list for storing particles that do not belong on any of the constituent Boxes, and methods for redistributing these particles to the proper ListBox<T> and MPI process. These methods can be used to perform parallel rebinning on a set of particles after their positions have been modified in a simulation.

ParticleData<T> behaves quite similarly to a LevelData<FarrayBox>, with a few exceptions. One major difference is that the BoxLayout on which the ParticleData<T> is defined must used fixed-size Boxes. This simplifies the interal implementation of ParticleData<T>, because the right Box for a given particle can be computed using a simple binning operation. Another difference is that you are not restricted to defining a ParticleData<T> on a DisjointBoxLayout. This is done to support load-balancing schemes in which particles that lay on the same spatial bin are handled by different processors. Other than that the use of the two classes is quite similar. For example, a ParticleData<T> can be indexed with a DataIndex, allowing data-parallel access to distributed ListBox<T>s. It also provides methods for handling ghost particles, as well as methods analagous to BinFab<T>::reBin() for shuffling particles between Boxes and processes. Some the most important functions in the API for ParticleData<T> are:

ParticleData()
 Default constructor. The ParticleData<T> is left unusable, and define() must be called later.

 const RealVect& a\_meshSpacing,
const RealVect& a\_origin);

Full constructor. Constructs an empty ParticleData<T> on the BoxLayout a\_dp. **Arguments:** 

- a\_dp The BoxLayout on which this ParticleData<T> is defined. This layout must used fixed-size Boxes.
- a\_domain The problem domain of the simulation on this level. Used for handling periodic boundaries.
- a\_fixedBoxSize The size of the Boxes in the layout.
- a\_meshSpacing The mesh spacing on this level.
- a\_origin The origin of the coordinate system, used to bin the particle positions.

```
    void define(const BoxLayout& a_dp,
        const ProblemDomain& a_domain,
        const int& a_fixedBoxSize,
        const RealVect& a_meshSpacing,
        const RealVect& a_origin);
```

Define function - same as full constructor.

#### **Arguments:**

- a\_dp The BoxLayout on which this ParticleData<T> is defined. This layout must used fixed-size Boxes.
- a\_domain The problem domain of the simulation on this level. Used for handling periodic boundaries.
- a\_fixedBoxSize The size of the Boxes in the layout.
- a\_meshSpacing The mesh spacing on this level.
- a\_origin The origin of the coordinate system, used to bin the particle positions.
- const BoxLayout& getBoxes() const;

Get the BoxLayout on which this ParticleData<T> is defined.

void gatherOutcast();

Collect the particles that are not in the right Boxes anymore and put them in the outcast list.

- void remapOutcast();
   Redistribute the particles in the outcast list to the right ListBox<T>. The can involve parallel communication.
- void fillGhosts(ParticleData<P>& a\_particlesWithGhosts, const int a\_numGhost);

A routine for grabbing copies of "ghost particles." On exit, each ListBox<T> in a\_particlesWithGhosts will contain all the particles within a\_numGhost of its Box. Periodic boundaries will be respected.

#### **Arguments:**

- a\_particlesWithGhosts A ParticleData<T> like this one, except that the grids have all been grown by a number of ghost cells.
- a\_numGhosts The number of cells the Boxes have been grown by.

For example, to move a ParticleData worth of particles, rebinning every time step, you would do something like the code snippet in figure 12.3

## 12.3 AMR with particles

For calculations with AMR, we provide the following tools for transferring particles between AMR levels: the ParticleValidRegion, for defining the region of space a given AMR level "owns", and the collectValidParticles function, for transferring particles to/from a given level based on its ParticleValidRegion. To see these tools in action, please see the AMRParticleMesh released example.

## 12.3.1 ParticleValidRegion

This is a class for defining the spatial region on which particle data is valid for an AMR level - that is, if a particle lives on an level's valid region, then it will be stored in that level's ParticleData<T>. In principle, this is could the same as the "valid region" of the AMR hierarchy, but for some applications it is convenient to shrink the valid region by some number of buffer cells around the coarse/fine boundaries. For example, for Particle-in-Cell calculations, it's convenient to arrange the particles onto AMR levels in such a way that particles on the fine level are guaranteed to have their entire interpolation kernels on the fine level for some number of time steps.

To create a ParticleValidRegion, you use the following constructor:

Figure 12.3: Code snippet to show how move all the particles in a BinFab and re-bin them.

```
    ParticleValidRegion(const DisjointBoxLayout& a_domain,
    const LevelData<BaseFab<bool> >* a_crseValid,
    const int a_refRatio,
    const int a_buffer);
```

Routine for defining an AMR level's valid region for particles.

#### **Arguments:**

- a\_domain The grids on this AMR level.
- a\_crseValid A boolean mask that defines the valid region of the coarser AMR level.
- a\_refRatio The AMR refinement ratio.
- a\_buffer The number of buffer cells used to define the AMR level.

You can also use the following method to get a pointer to the boolean mask the defines a ParticleValidRegion:

const LevelData<BaseFab<bool> >\* mask() const
 Get a pointer to the valid region. The mask will be True if the cell belongs on this region, False if not.

#### 12.3.2 collectValidParticles

The following function is used to transfer particles on to or off of a ParticleData<T>, based on whether the particles lives on a given ParticleValidRegion. Usage:

Routine for transferring particles between different levels.

#### **Arguments:**

- a\_partValid The destination to which particles will be sent if they meet the requirements.
- a\_domain The source from which the particles will be transferred.
- a\_mask This defined the valid region.

- a\_meshSpacing The mesh spacing used for binning particles. Should be for the same level as a\_mask.
- a\_refRatio The src will be coarsened or refined by this amount before it is compared to the mask.
- a\_flip If false, particles will be transferred if they live in cells where the mask is true. If true, they will be transferred if the mask is false.
- a\_origin The origin of the coordinate system, used to bin the particle positions. Default is (0.0, 0.0, 0.0)

#### 12.4 Particle IO

For data analysis and plotting, we provide the following HDF5 IO functions, analogous to writeAMRHierarchyHDF5 and readAMRHierarchyHDF5. These functions read or write a single ParticleData to or from a given HDF5 file. For an example of how to incorporate this into an AMR program, please see the AMRParticleMesh released Example.

```
    writeParticlesToHDF(HDF5Handle& a_handle,
const ParticleData<P>& a_particles,
const std::string& a_dataType);
```

Write the particles in the ParticleData to an HDF5 file.

#### **Arguments:**

- a\_handle The previously opened HDF5 filehandle to which the particles will be written.
- a\_particles The particles that will be written out.
- a\_dataType A string labelling these particles; useful when you have multiple particle types.

```
    readParticlesFromHDF(HDF5Handle& a_handle,
ParticleData<P>& a_particles,
const std::string& a_dataType);
```

Read the particles from the HDF5 file and used them to initialize the ParticleData. **Arguments:** 

- a\_handle The previously opened HDF5 filehandle from which the particles will be read.
- a\_particles The ParticleData that the particles will be placed into.
- a\_dataType A string labelling these particles; useful when you have multiple particle types.

These routines output the particles in a format that can be recognized by the yt analysis code.

# 12.5 Example code

To demonstrate the use of the Chombo's particle tools in a relatively simple application, we have provided the AMRParticleMesh released example. This example implements the collisionless fluid and self-gravity components of the hybrid AMR scheme for cosmology calulculations described in [25], i.e. there is no hyperbolic component, just particles and an elliptic solve. W use a kick-drift-kick leapfrog time integrator to advance the particles and AMR multigrid to solve Poisson's equation with periodic boundary conditions.

The code can be found in releasedExamples/AMRParticleMesh in the Chombo distribution. This example program uses the full AMRTimeDependent class structure, and demonstrates how to work with ListBox<T> and ParticleData<T> in an AMR context. We provide initial conditions for a "Zel'dovich Pancake" collapse calculation. To build the program, navigate to Chombo/releasedExamples/AMRParticleMesh/exec, and then type make with the desired configuration flags. For instance:

```
make -j8 DIM=2 MPI=TRUE DEBUG=FALSE OPT=TRUE amrPM
```

To run the program in parallel, use:

```
mpirun -np 4 ./amrPM2d.Linux.64.mpiCC.gfortran.OPT.MPI.ex inputs
```

Your configuration string might be slighly different, depending on your platform and the flags you set for compilation.

In addition to the full example program, there are also a number test programs in Chombo/lib/test/ParticleTools. These programs test the basic functionality of the ParticleTools classes in isolation, and can also be referenced to see basic usage of the tools in practice.

# Chapter 13

# **Chombo Debugging and Performance Tools**

# 13.1 Overview of Chombo Debugging Tips

Chombo contains many complex datatypes and it can be difficult to look at one's data with simple gdb print statements. This chapter is intended to provide some tools for the user to be able to debug her applications with more facility.

Here are some general points.

- All the examples provided use gdb.
- To avoid ugly printouts while in gdb, one puts in her ~/.gdbinit :

```
set print static-members off set print pretty on
```

• To stop in a Fortran function, one usually has to add an underscore to the end of the name. To stop in Fortran subroutine myfortranfunc,

```
<gdb prompt> break myfortranfunc_
```

• gdb is most useful when run within emacs. To do this, type

```
Meta-x gdb
```

while the buffer is in a file in the same directory as the executable (the input file or the makefile are popular choices). The advantages of this are many:

- One gets emacs's nifty color scheme.

- One interacts with the code directly. This means that when her code segfaults at a particular point, she will be looking at the offending line of code.
- One can set breakpoints by just being at the line she wants to stop at and typing control-x spacebar.
- Avoid using more than one gdb session in an emacs session. It gets confusing. One should fire up another emacs if one needs to debug two codes at once.
- If one wants to view some complex datatype (say class BagODonuts) for which Chombo does not yet have a nifty print function, she can write her own print function and call it using

```
<gdb prompt> call printBagODonuts(&myBagODonuts)
```

All such functions will work most reliably if she always writes them to take pointers and define them using extern ''C''. Then gdb will not get confused about copy constructors and demangling. Chombo provides many examples of such functions. They live in

Chombo/lib/src/BoxTools/DebugOut.H

- These functions are not "safe." There is no type checking in gdb. Anything one sends to the function will be interpreted as a pointer and it will try to run with it. It is very easy to seg fault one's gdb session if she
  - Forgets the & so the address you are sending is nonsense.
  - Mismatches the call with the data type.

# 13.2 Chombo Print Utilities

Chombo provides a bunch of functions to print out various datatypes in the debugger. To use these functions, one includes <code>DebugDump.H</code> in her code. The prototypes for the functions are in are in <code>Chombo/lib/src/BoxTools/DebugOut.H</code>. The functions are used in gdb as follows:

```
<gdb prompt> call dumpIVS(&myIVS).
```

A list of some of them and what they print out follows:

void dumpLDF(const LevelData<FArrayBox>\* memLDF);
 Dump a level's worth of data to standard out. Only use this for small data sets.

- void dumpDBL(const DisjointBoxLayout\* a\_dblInPtr);
   Dump a DisjointBoxLayout to standard out.
- void dumpIVS(const IntVectSet\* a\_ivsInPtr)
   Dump the points of an IntVectSet to standard out.

# 13.3 Viewing data objects with Vislt from gdb

There are two ways to use Vislt to help examine data during a debugging session. One may use the writeFAB, writeFABname, writeLevel, and writeLevelname functions described in section 6.3 to write data in a FArrayBox or LevelData<FArrayBox> to a file, and then call VisIt from a shell to view the data. Alternatively, the viewFAB and viewLevel functions allow VisIt to be called directly from the gdb session.

If one has an FArrayBox fab and a LevelData<FArrayBox> ldf, one may do the following:

```
<gdb prompt> call viewFAB(&fab)
<gdb prompt> call viewLevel(&ldf)
```

which will result in fab and ldf being written to separate temporary files, and then one Vislt processes being started with two windows controlling the different data objects.

# 13.4 pout()

In Chombo, the pout() function is used in place of the std::cout output stream object. defined in header parstream H

In serial this function returns std::cout. In parallel, this creates a file called pout.n where n is the procID() of the given processor. Output is then directed to these files. This keeps the output from different processors from getting all jumbled up. Used just as one would use a standard output stream.

# 13.5 Memory Tracking

```
Chombo provides a simply internal memory tracking facility.
    from memtrack.H:

class Memtrack
{
    public:
        static void ReportUnfreedMemory(ostream& os);

/// calls ReportUnfreedMemory with pout()
        static void UnfreedMemory();

    static void memTrackingOn();

    static void memtrackingOff();

    static void overallMemoryUsage(long long& currentTotal, long long& peak);
};
```

This is compiled into Chombo and turned on by default when the compiler macro ENABLE\_MEMORY\_TRACKING is defined.

ReportUnfreedMemory produces a breakdown of current memory usage broken down by Chombo class. This does not include the system image of the program itself, or stack usage. This will also not include memory allocated by the user, just memory allocated by Chombo functions. RTTI is used in some places to identify the type of objects held in a Chombo Vector template container.

example output of an UnfreedMemory() call:

```
Vector 3Box: 56 bytes (0 Mb)

Vector 5Entry: 252 bytes (0 Mb)

Vector Ui: 28 bytes (0 Mb)

Vector i: 16 bytes (0 Mb)

Total Unfreed: 352 bytes (0 Mb)

peak memory usage: 360 bytes (0 Mb)
```

## 13.6 TraceTimer

TraceTimer class is a self-tracing code instrumentation system for Chombo (or any other package really). The user interface is specified by a small set of macros. The usage

model is that you just leave these timers in the code, for good. Initially, your application will have 'main' and a few heavy functions instrumented, and the lower level Chombo library instrumentation. As your tool or application matures, it will garner a larger set of instrumentation giving clear views of your code performance. After a routine has been cleverly and lovingly optimized, the timers are left in place to spot when some later bug fix or \*improvement\* undoes your previous labors.

You should never need to use or interact with the the classes TraceTimer or AutoStart. Use the macros. They call the right functions and classes for you.

The first macro is what people will use the most:

```
CH_TIME("label");
```

This is the simplest interface for TraceTimer. You place this macro call in a function you wish to be timed. It handles making the timer, calling 'start' when you enter the function, and calling 'stop' when you leave the function. A good idea is to use a 'label' specific enough to be unambiguous without being overwhelming. for instance:

In this case, we have a class with many constructors and define functions that all funnel into a single general function. We can just call this 'define' and not worry about naming/instrumenting all the different overloaded instances. If you slip up and use the same label twice, that is not a real problem, the two locations will be timed and tracked properly (even if one is a sibling or parent of the other). The only place it will make things a little harder is in the output where you might have the same name show up and look confusing.

In serial, you will see a file called time.table (in parallel, you will get a time.table.n (where n is the rank number) files).

By default, Chombo compiles in the instructions for the timers wherever the macros appear. If the compiler macro CH\_TIMER is defined, then all the CH\_TIME\* macros evaluate to empty expressions at compile time.

So, you put some CH\_TIME calls in your code and ran it, and nothing happened: Chombo looks for the environment variable **CH\_TIMER**. If it is set to anything (even if it is set to 'false' or 'no' or whatever) then the timers will be active and reporting will happen. If this environment variable is not set, then all the timers check a bool and return after doing nothing.

One point of interest with using the environment variable: In parallel jobs using mpich, only processor 0 inherits the environment variables from the shell where you invoke 'mpirun', the rest read your .cshrc (.bashrc, etc.) file to get their environment. To time all your processes, you need to make sure the **CH\_TIMER** environment variable gets to all your processes.

#### 13.6.1 Auto hierarchy

The timers automatically figure out their parent/child relationships. They also can be placed in template code. This has some consequences. First, if you have a low level function instrumented that has no timers near it in the code call stack, you will see it show up as a child of a high level timer. the root timer "main" will catch all orphaned timers. So, even though you might make no call to, say, 'exchange' in your 'main' function, you might very well call a function, that calls a function, that calls 'exchange'. Since no code in between was instrumented, this exchange is accounted for at 'main'. This might look strange, but it should prove very powerful. An expensive orphan is exactly where you should consider some more timers, or reconsidering code design.

For performance reasons, child timers have only one parent. As a consequence each CH\_TIME("label") label can show up at multiple places in your output. Each instance has it's own timer. So, each path through the call graph that arrives at a low-level function has a unique lineage, with it's own counter and time. Thus, I can instrument LevelData::copyTo once, but copyTo can appear in many places in the time.table file.

#### 13.6.2 Finer control

The next level up in complexity is the set of \*four\* macros for when you want sub-function resolution in your timers. For instance, in a really huge function that you have not figured out how to re-factor, or built with lots of bad cut n paste code 're-use'.

```
CH_TIMERS("parent");
CH_TIMER("child1", t1);
CH_TIMER("child2", t2);
CH_START(t1);
//some code go here
CH_STOP(t1);
CH_START(t2);
//some other code go here
CH_STOP(t2);
CH_START(t1);
//can start something here again
CH_STOP(t1);
```

One very good place to use the more sophisticated API is within loops. START and STOP are very fast compared to the timer declaration:

CH\_TIMERS has the same semantic as CH\_TIME, except that you can declare an arbitrary number of children after it in the same function scope. The children here do not autostart and autostop, you have to tell them where to start and stop timing. The children can themselves be parents for timers in called functions, of course. The children obey a set of mutual exclusions. The following generate run time errors:

- double start called
- double stop called
- start called when another child is also started
- you leave the function with a child not stopped the following will generate compile time errors:
- more than one CH\_TIME macro in a function
- invoking CH\_TIMER("child", t) without having first invoked CH\_TIMERS
- re-using the timer handle ie. CH\_TIMER("bobby", t1); CH\_TIMER("sally", t1)
- mixing CH\_TIME macro with CH\_TIMER
- mixing CH\_TIME macro with CH\_TIMERS

You do not have to put any calls in your main routine to activate the clocks or generate a report at completion, this is handled with static initialization and an atexit function. The exception to this is for parallel reporting. Since atexit and MPI\_Finalize() do not interact in an agreeable fashion, you need to explicitly call the Chombo macro CH\_TIMER\_REPORT() before the code reaches MPI\_Finalize

There is a larger argument about manual instrumentation being detrimental to clean software. Profiling the code is supposed to tell you where to expend your optimization effort. Manual instrumentation opens the door to people wasting time \*assuming\* what parts of the code are going to take up lots of time and instrumenting them, before seeing any real performance data. Good judgment is needed. We have a body of knowledge about Chombo that will inform us about a good minimal first set of functions to instrument.

# Chapter 14

# **Troubleshooting**

- error: 'H5Pset\_fapl\_mpio' was not declared in this scope
  - This is a compilation error. The user is trying to build the parallel version of Chombo but is not linking against an HDF5 build that has used --enable-parallel when it was configured
  - Most Chombo users need access to both a serial and a parallel version of their HDF5 libraries. These are distinguished in the Chombo makefiles as HDFINCFLAGS and HDFMPIINCFLAGS (these values should be different).

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