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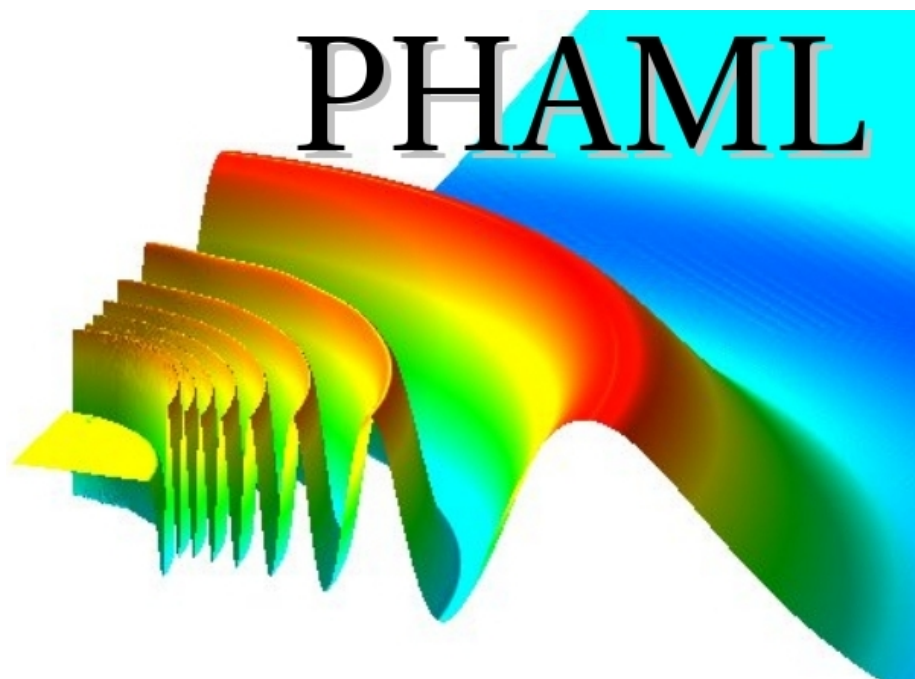
# **PHAML User's Guide**

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## PHAML User's Guide, Version 1.9.1

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## **Abstract**

PHAML (Parallel Hierarchical Adaptive MultiLevel) is a Fortran module for the solution of elliptic partial differential equations. It uses finite elements, adaptive grid refinement ( $h$ ,  $p$  or  $hp$ ) and multigrid solution techniques in a message passing parallel program. It has interactive graphics via OpenGL. This document is the user's guide for PHAML. The first part tells how to obtain any needed software, how to build and test the PHAML library, and how to compile and run the example programs. The second part explains the use of PHAML, including program structure and the various options that are available. The third part is a reference manual which describes the API (application programming interface) of PHAML. The reference manual begins with a 2 page Quick Start section for the impatient.

**Keywords:** adaptive grid refinement, dynamic load balancing, elliptic eigenvalue problems, elliptic partial differential equations, high order finite elements, hp-adaptivity, multigrid, parallel programming

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# Chapter 1

## Introduction

To start using PHAML immediately, see the Quick Start guide in Section 7.1.

Please note: This document changes with each release of PHAML. Changes made in the current release are printed in **red**. Changes made in a recent release are printed in **dark red**. Text that has been deleted is maintained for a few releases for comparison, but is printed in a tiny font size.

PHAML stands for Parallel Hierarchical Adaptive MultiLevel method. It solves systems of linear elliptic partial differential equations (PDEs) of the form

$$-\frac{\partial}{\partial x} \left( c_{xx} \frac{\partial u}{\partial x} \right) - \frac{\partial}{\partial x} \left( c_{xy} \frac{\partial u}{\partial y} \right) - \frac{\partial}{\partial y} \left( c_{yx} \frac{\partial u}{\partial x} \right) + c_x \frac{\partial u}{\partial x} + c_y \frac{\partial u}{\partial y} + c_u u = f \quad \text{in } \Omega \quad (1.1)$$

where  $c_{xx}$ ,  $c_{xy}$ ,  $c_{yy}$ ,  $c_x$ ,  $c_y$ ,  $c_u$  and  $f$  are functions of  $x$  and  $y$ , and the domain  $\Omega$  is a bounded, connected, region in  $R^2$ . The boundary conditions can be Dirichlet on part of the boundary,

$$u = g \quad \text{on } \partial\Omega_D \quad (1.2)$$

and natural or mixed on the remainder of the boundary,

$$\left( c_{xx} \frac{\partial u}{\partial x} + c_{xy} \frac{\partial u}{\partial y} \right) \frac{\partial y}{\partial s} - c_{yy} \frac{\partial u}{\partial y} \frac{\partial x}{\partial s} + c_{bc} u = g \quad \text{on } \partial\Omega_N \quad (1.3)$$

where  $g$  and  $c_{bc}$  are functions of  $x$  and  $y$ ,  $c_{xx}$ ,  $c_{xy}$  and  $c_{yy}$  are the same functions as in Equation 1.1, and the boundary  $\partial\Omega = \partial\Omega_D \cup \partial\Omega_N$  and  $\partial\Omega_D \cap \partial\Omega_N = \emptyset$ . Periodic boundary conditions are also supported.

For natural boundary conditions, differentiation with respect to  $s$  is with respect to a counterclockwise parameterization of the boundary  $(x(s), y(s))$  with  $\|(dx/ds, dy/ds)\| = 1$ . Note that when  $c_{xx} = c_{yy} = 1$  and  $c_{xy} = 0$  or the boundaries of the domain are parallel to the axes, the natural boundary condition reduces to the Neumann boundary condition

$$\frac{\partial u}{\partial n} = g$$

where  $n$  is the outward unit normal.

If the domain has curved boundaries, it is defined by subroutines that define the boundary parametrically. If it is polygonal, it can instead be defined by an initial triangulation given in data files created by the program Triangle.

PHAML also solves elliptic eigenvalue problems where the right hand side  $f(x, y)$  is  $\lambda u$ , and  $\lambda$  is an eigenvalue to be determined along with  $u$ . For eigenvalue problems, the boundary conditions must be homogeneous, i.e.,  $g = 0$ .

PHAML discretizes the PDE using the standard finite element method with piecewise polynomial functions over triangles. The approximating polynomial degrees may be a fixed constant or adaptive. The grid is generated by beginning with a very coarse grid of fixed degree and using uniform or adaptive h-refinement in the form of newest node bisection, uniform or adaptive p-refinement in which the polynomial degree is increased, or hp-adaptive refinement which combines both forms of refinement.

The standard process alternates between phases of grid adaptation and solution of the discrete equations. The default solution method is a hierarchical basis multigrid method. Other solution methods are available through other optional software packages. If the cross derivative or first order terms are included, i.e.  $c_{xy} \neq 0$ ,  $c_x \neq 0$  or  $c_y \neq 0$ , then one of the alternate solution methods for nonsymmetric matrices must be used. All this is performed in parallel using the full domain partition to minimize communication. The default load balancing method is a refinement-tree based partitioning algorithm. Other load balancing methods are available through other optional software packages.

PHAML can be run as a sequential program, a master/slave parallel program using MPI-2 or PVM, an SPMD parallel program using MPI-1, MPI-2, or PVM, **an SMP program using OpenMP, or a hybrid parallel program using both MPI and OpenMP.** Currently, the OpenMP code is under development and only a few parts of the code use it. See the release notes for the current status. PHAML optionally uses OpenGL to provide menu-driven interactive visualization.

## Chapter 2

# Software

### 2.1 Obtaining Software

PHAML and all required or optional auxiliary software can be obtained by freely downloading them from the web, although some have commercial alternatives. This section explains what software is used with PHAML, when you need that software, and where to obtain it. Most of this software is optional and you do not need to install it unless you are using the relevant features as described below.

#### 2.1.1 PHAML

PHAML is available for download at <http://math.nist.gov/phaml>. It is a gzipped tar file. When unpacked, it creates a directory called `phaml-x.x.x` where the x's are the version number. It has been tested under many Unix systems, but not under MS Windows. It is standard conforming Fortran 90, so (theoretically) it should work under Windows (at least sequentially) without too much effort. For the remainder of this document, we will assume that the operating system is some variant of Unix, including Linux.

#### 2.1.2 Fortran 90 and C Compilers

A Fortran 90 (or later Fortran standard) compiler is required to compile PHAML. In some cases, a C compiler will also be required to compile some wrapper routines to C libraries. Most Unix systems already have a C compiler installed, and many also have a Fortran 90 compiler installed. Check your local documentation to find out, or look for a command like `f90` or `f95`.

If you do not have a Fortran 90 compiler, many good commercial compilers are available for most (if not all) Unix systems. To identify what is available for your system, see Michael Metcalf's *Fortran 90/95/HPF Information File* at <http://www.fortran.com/fortran/metcalf.htm>.

There are currently two free Fortran 90 compilers, g95 at <http://www.g95.org> and GNU's gfortran at <http://gcc.gnu.org/fortran>.

### 2.1.3 BLAS and LAPACK

BLAS (Basic Linear Algebra Subroutines) and LAPACK (Linear Algebra PACKage) libraries are required for PHAML. The reference Fortran implementation for them can be obtained from Netlib at <http://www.netlib.org>. However, for best performance you should use a BLAS library that is optimized for your computer system. You may find that the BLAS and LAPACK libraries are already installed on your system, or that there are commercial optimized versions available for your system.

There are also freely available optimized BLAS packages such as ATLAS at <http://math-atlas.sourceforge.net> and GOTO BLAS at <http://www.tacc.utexas.edu/resources/software/>.

### 2.1.4 PVM and MPI

To run PHAML as a parallel program, you must have a message passing library (PHAML can be run as a sequential program, i.e. one processor, without a message passing library).

Message passing can be performed by either MPI (Message Passing Interface) or PVM (Parallel Virtual Machine). MPI is considered to be more of an industry standard, and is the recommended choice in most cases. PVM support in PHAML is maintained primarily for historical reasons, but PVM does have some features (for example, specifying the host on which to run the graphics process) that are not necessarily supported by an MPI implementation. Many of the other optional packages (e.g. PETSc, *hypr*, Zoltan, MUMPS, ARPACK) require MPI and cannot be used if PVM is chosen instead.

To run PHAML as a message passing parallel program, you must have an MPI library. Many computer systems will already have an MPI library installed. Check your local documentation. If not, there are three freely available implementations of MPI. The Open MPI implementation is available at <http://www.open-mpi.org>. The LAM implementation is available at <http://www.lam-mpi.org>. The MPICH2 implementation is available at <http://www-unix.mcs.anl.gov/mpi/mpich2>.

PVM is available at [http://www.csm.ornl.gov/pvm/pvm\\_home.html](http://www.csm.ornl.gov/pvm/pvm_home.html).

### 2.1.5 OpenGL (or Mesa), GLUT and f90gl

If you wish to use the visualization capabilities of PHAML, then you will need an OpenGL library (graphics library), GLUT (window, mouse, keyboard and menu management), and f90gl (the Fortran 90 interface to OpenGL and GLUT).

Some computers will already have an OpenGL library installed. If not, there are some commercial OpenGL libraries for some computer systems, and there is the freely available Mesa library. Mesa can be obtained from <http://www.mesa3d.org>, and further information about OpenGL can be found at <http://www.opengl.org>.

If your system already has an OpenGL library, then it is likely to also have GLUT already installed. However, many versions of GLUT are not compatible

with f90gl, so you may need to reinstall GLUT anyway. If you (re)install GLUT, you should get it from the software section of the f90gl web pages at <http://math.nist.gov/f90gl>. Alternatively, if you are installing Mesa, then also get MesaGLUT. This distribution works with f90gl since version 7.0.3.

It is highly unlikely that f90gl will already be installed on your system. You can obtain f90gl from <http://math.nist.gov/f90gl>.

### 2.1.6 Triangle

PHAML uses Jonathan Richard Shewchuk's mesh generator, Triangle [36], to generate the initial grid for arbitrary geometry. If you are only solving problems on the unit square, then you do not need Triangle – you can use the Triangle output files in the examples. You also do not need it if you have obtained Triangle data files elsewhere. Otherwise, you will need it. You also need it if you choose to define your domain through subroutines. Triangle is available at <http://www.cs.cmu.edu/~quake/triangle.html>.

### 2.1.7 ARPACK

To solve elliptic eigenvalue problems, PHAML needs an algebraic eigensolver. One of the supported eigensolvers is ARPACK [20]. You can get ARPACK and PARPACK from <http://www.caam.rice.edu/software/ARPACK>. For a sequential compilation you do not need PARPACK.

### 2.1.8 BLOPEX

Another supported eigensolver is BLOPEX [18, 19]. Currently, support for BLOPEX in PHAML is only available through PETSc 2.1.11. To include BLOPEX in PETSc, use `--download-blopex=1` when you configure PETSc for compilation.

### 2.1.9 *hypr*

*hypr* [13][14] is a package of iterative linear system solvers and preconditioners from the Lawrence Livermore National Laboratory. It is not required, but can be linked with PHAML to provide alternative linear system solvers. It is available at [http://www.llnl.gov/CASC/linear\\_solvers](http://www.llnl.gov/CASC/linear_solvers).

### 2.1.10 MUMPS

MUMPS [3][?] is a parallel direct linear system solver. It is not required, but can be linked with PHAML to provide an alternative linear system solver. It is available at <http://graal.ens-lyon.fr/MUMPS/index.html>.

### 2.1.11 PETSc

PETSc [6][7] is a package of iterative linear system solvers and preconditioners from Argonne National Laboratory. It is not required, but can be linked with PHAML to provide alternative linear system solvers. It is available at <http://www-unix.mcs.anl.gov/petsc/petsc-as>.

### 2.1.12 SuperLU

SuperLU [21] is a parallel direct linear system solver from the Lawrence Berkeley National Laboratory. It is not required, but can be linked with PHAML to provide an alternative linear system solver. It is available at <http://crd.lbl.gov/~xiaoye/SuperLU>.

### 2.1.13 Zoltan

Zoltan [11][12] is a library of dynamic load balancing methods. It is not required, but can be linked with PHAML to provide alternative methods for partitioning the grid for distribution over parallel processors. It can be obtained at <http://www.cs.sandia.gov/Zoltan>.

Zoltan may be compiled with support for certain third party libraries. Two of these are supported in PHAML through Zoltan. ParMETIS is a static partitioning library. Zoltan includes a complimentary copy of the ParMETIS software. DRUM is utility for load balancing on heterogeneous or dynamically changing parallel computers. It has not yet been officially released and can only be obtained by requesting it from the author.

## 2.2 Compiling PHAML

This section gives instructions for compiling the PHAML library. Throughout this section it is assumed that the current working directory is the PHAML root directory.

### 2.2.1 Creating the Makefiles

Before compiling you must create a **Makefile**. This is done by running the shell script **mkmkfile.sh**. **This script will make all the Makefiles in all the PHAML subdirectories, so you do not need to run any of the other mkmkfile.shs.**

Before running this script you must edit it to set system dependent values, like the paths to certain libraries, and specify the configuration you wish to build a **Makefile** for (what kind of computer, what message passing library, etc.). Instructions for modifying **mkmkfile.sh** are contained in the file, but are also summarized here. There are three steps to modifying **mkmkfile.sh**: 1) set the default system configuration, 2) set the paths, library names, etc., and 3) set values for particular systems.



In step 1, the default system configuration is set. If you will only be running in one environment, then you can set the configuration here and forget it. In fact, you can probably just make the `Makefile` once and forget it. If you will be using more than one environment, then you can set the defaults here, and they can be overridden by one of the methods described later in this section.

The configuration variables (e.g. `PHAML_ARCH`) and legitimate values (e.g. `origin`) can be found under Step 1 in `mkmkfile.sh`. You can also get a list of all the configuration variables, their legitimate values, and current defaults by executing

```
./mkmkfile.sh help
```

They are:

- `PHAML_ARCH` `origin` `rs6k` `sgi` `sun` `tflop` `x86`. This defines the type of computer architecture you have.
- `PHAML_OS` `aix` `cougar` `irixn32` `irix64` `linux` `solaris`. This defines the operating system running on your computer.
- `PHAML_F90` `absoft` `g95` `gfortran` `intel` `lahey` `nag` `pathscale` `pgi` `sgi` `sun` `xlf`. This defines the Fortran 90 compiler to use.
- `PHAML_C` `cc` `gcc`. This defines the C compiler to use.
- `PHAML_HASHSIZE` `1` `2`. This defines the size of hash key to use for global identifiers. `1` uses one integer and `2` uses two integers. Using `1` saves memory and reduces message sizes; using `2` allows smaller elements (more refinement levels).
- `PHAML_PARALLEL` `messpass_spawn` `messpass_nospawn` `openmp` `hybrid_spawn` `hybrid_nospawn` `sequential`. This defines the type of parallelism. You can select between running a sequential program (no parallelism), having a master process that spawns the slave and graphics processes, running an SPMD (single program multiple data) program in which all processes are identical and started from the command line, `running an SMP program using OpenMP`, and `running a hybrid OpenMP/MPI program either as master/slave or SPMD`. Note: `not all compilers support OpenMP`.
- `PHAML_PARLIB` `lam` `mpi` `mpich` `mpich2` `myrinet` `openmpi` `pvm` `none`. This defines the parallel library to use. `lam`, `mpich`, `mpich2` and `openmpi` and `pvm` refer to the packages by those names, `mpi` to vendor implementations of MPI, `myrinet` to a special implementation of MPICH for myrinet networks, and `none` should be used if `PHAML_PARALLEL` is `sequential` or `openmp`.
- `PHAML_GRAPHICS` `metro` `mesa` `none` `opengl`. This defines what OpenGL library should be used. `opengl` refers to vendor OpenGL libraries, `mesa` to the MESA library, `metro` to a commercial OpenGL library for Linux, and `none` indicates that the graphics program should not be built.

- `PHAML_BLAS atlas compiler goto source standard vendor`. This defines the BLAS library to use. `atlas` and `goto` are as mentioned in Section 2.1.3. `compiler` refers to a BLAS library provided with the Fortran compiler defined in `PHAML_F90`. `vendor` refers to a BLAS library supplied by the hardware vendor defined in `PHAML_ARCH`. `standard` refers to a BLAS library in a standard location like `/usr/lib`, and often is not a high performance library. As a last resort, `source` will use BLAS source code provided with PHAML.
- `PHAML_LAPACK atlas compiler source standard vendor`. This defines the LAPACK library to use, and is usually given the same value as `PHAML_BLAS`.
- `PHAML_ARPACK no yes`. This indicates whether or not to include ARPACK.
- `PHAML_BLOPEX no withpetsc`. This indicates whether or not to include BLOPEX, and what source of BLOPEX to use. `withpetsc` indicates that PETSc was configured with BLOPEX and a PETSc solver will be used with BLOPEX. Additional sources will be added in a future release of PHAML.
- `PHAML_HYPRE no yes`. This indicates whether or not to include *hypre*.
- `PHAML_MUMPS no yes`. This indicates whether or not to include MUMPS.
- `PHAML_PETSC no yes`. This indicates whether or not to include PETSc.
- `PHAML_SUPERLU no yes`. This indicates whether or not to include SuperLU.
- `PHAML_ZOLTAN no yes`. This indicates whether or not to use Zoltan. When Zoltan was compiled, it may or may not have included support for third party libraries. The next few parameters indicate which of them were built into Zoltan. These must agree exactly with how Zoltan was built, or you will get error messages from the linker.
- `PHAML_PARMETIS no yes`. Should be yes if and only if Zoltan was built with ParMETIS support.
- `PHAML_JOSTLE no yes`. Should be yes if and only if Zoltan was built with JOSTLE support.
- `PHAML_PATOH no yes`. Should be yes if and only if Zoltan was built with PaToH support.
- `PHAML_PARKWAY no yes`. Should be yes if and only if Zoltan was built with ParKway support.
- `PHAML_NEMESIS no yes`. Should be yes if and only if Zoltan was built with Nemesis support.

- `PHAML_DRUM` `no` `yes`. Should be `yes` if and only if Zoltan was built with DRUM support.
- `PHAML_SYSTEM` `none` `dragon` `looney` `pepe` `raritan` `speedy` `looneyjr` `sgis` `suns` `tflop`. This designates a specific computer system for overriding configuration variables in Step 3. Usually it is the hostname of the system.

If you don't find a value that matches your system, you can either (1) add that value and modify `mkmkfile.sh` to handle it by mimicking what is done for other values, or (2) use a similar value and edit `Makefile` to correct it for your system. However, if you edit `Makefile` then you will lose your changes the next time you run `mkmkfile.sh`.

In step 2, set the paths, library names, etc. Here you set shell variables in `mkmkfile.sh` with the names of commands, flags, paths, library names, etc., for the configurations you will be using. Many of these will already be correct, but examine them because some of them are specific to the systems on which PHAML was developed. In particular, many of the compiler command names have been changed to avoid conflicts between multiple compilers on one system, and nearly all of the paths will vary between systems.

In step 3, you can override values set in step 2 for particular system configurations. For example, suppose your MPI library has a different name on Linux than it has on all your other systems. Then in step 2 you set the name used on most systems, and in step 3 you override it if `PHAML_SYSTEM` is your linux box.

There are two other ways to override the default configuration variable values: 1) on the command line for `mkmkfile.sh` (described below), and 2) by setting environment variables in your shell. If an environment variable exists with the same name as a configuration variable, the value of that environment variable is used. Usually these variables are set in the shell startup file. For csh-type shell, an example is

```
setenv PHAML_OS linux
```

Once you finish modifying `mkmkfile.sh`, running it will create `Makefile` in the `src` directory and all the subdirectories under `example` and `testdir`. If a file named `Makefile` already exists, it is moved to `Makefile.bak` before the new file is created. To accept the default configuration, just run

```
./mkmkfile.sh
```

To override the default values and values given by environment variables, specify them on the command line by giving a space-separated list of configuration variables and values. Here, including the “PHAML\_” part of the configuration variable is optional. For example,

```
./mkmkfile.sh PARALLEL messpass.spawn PARLIB lam
```

## 2.2.2 Compiling the Library

Once `Makefile` has been built, just type `make` in either the PHAML root directory or `src` subdirectory to compile the PHAML library. If this shows that `Makefile` has errors in it, you can either edit `Makefile` to fix them (but then

they will appear again if you need to run `mkmkfile.sh` again), or fix the errors in `mkmkfile.sh` and rebuild `Makefile`. This will create the PHAML library `libphaml.a` and copy it to subdirectory `lib`. It also copies any module files (e.g., `*.mod`) to subdirectory `modules`, and creates a file `lib/CONFIG` that contains the values of the configuration variables used to create the library.

## 2.3 Testing the Library

After creating the PHAML library, you can test it by running the PHAML Test Suite. The suite contains over 100 short test programs to test nearly all aspects and options of PHAML. Tests that do not apply to the current compilation of the library are skipped. The tests have not yet been written for `PHAML_PARALLEL=messpass_nospawn` (i.e. SPMD programs). They can only be run with `sequential`, `messpass_spawn`, `openmp` or `hybrid_spawn` programs.

To run the full test suite on an interactive system, enter the command

```
make test
```

from either the PHAML root directory or the `testdir` subdirectory. If you are using a batch system like PBS or Torque/Maui, you can run the tests with one of the PBS scripts in the `testdir` directory. To delete all the files that were created by the tests, enter the command

```
make test what=clean
```

The tests can be grouped into three categories. The first tests consists of two very simple programs to verify that a program can be compiled, linked and run. The interactive tests test the use of the `pause` variables to `phaml_solve_pde`, spawning in a debug window, and graphics. The remaining tests are the non-interactive tests. Depending on how many optional auxiliary packages are included, the speed of your computer system, etc., these may take anywhere from a few minutes to a few hours to run.

You can run just the first tests, just the first and interactive tests, or just the first and noninteractive tests with the commands

```
make test what=first
```

```
make test what=interactive
```

```
make test what=noninteractive
```

If you are running the tests from a batch system where you cannot respond to interactive prompts, you should use `what=noninteractive`.

The individual interactive tests can be run by giving `what` the value `pause`, `debug`, or `graphics`. The noninteractive tests are in the directories `testdir/test_*`. You can run individual noninteractive tests by giving `what` the name of the subdirectory, with or without `test_`. For example, to run the ARPACK tests you can use either

```
make test what=test_arpack, or
```

```
make test what=arpack
```

The results of the tests are printed both to the screen and to the file `testdir/testresults`. The file only contains standard output, not standard error, so things like messages from the compiler will not appear in the file.

Each test consists of a short program that is compiled and run with the output directed to a file. The file is compared to an existing file that contains the expected output. Each test is identified as SUCCESS, FAILURE or WARNING. WARNING indicates that the output is not the same as the comparison file. Often this difference is just minor numerical differences (accumulated round off error) due to different processors, compilers, versions of auxiliary packages, etc. The warning message will direct you to a file that contains the differences, as identified by `diff`, which you should examine to see if the differences are significant.

## 2.4 Compiling and Running the Examples

Next you compile your application and link in the PHAML library. With most compilers you will need to specify the location of the module files when compiling program units that use `phaml`. Many compilers use the `-I` flag for this (e.g. `f90 -I $PHAML_HOME/modules myprog.f90`), but you should check your compiler's documentation. You also need to link with the PHAML library (e.g. `f90 myprog.f90 -L$PHAML_HOME/lib -lphaml`) and any other libraries your program needs (MPI, OpenGL, LAPACK, etc.). Your best start is to look at the examples in the `examples` directory, and their `Makefiles`. (which were created by `mkmkfile.sh`) These `Makefiles` were created when you ran `mkmkfile.sh` in the top PHAML directory. You should not run the `mkmkfile.sh` in the `examples` subdirectories.

With the examples, a successful `make` will create up to three executables:

1. the primary executable `phaml`. This is the program you run.
2. `phaml_slave` if `PHAML_PARALLEL` is `messpass_spawn`. This is spawned by `phaml`.
3. `phaml_graphics` if `PHAML_GRAPHICS` is not `none` and `PARALLEL` is not `messpass_nospawn`. This is spawned by `phaml` and `phaml_slave` if `PARALLEL` is `messpass_spawn`, or run from the command line if `PARALLEL` is `sequential`.

In the PHAML root directory, `make clean` will remove any files created by compilation (e.g. `*.o`) in `src`, `examples/*`, and `testdir/*`. `make reallyclean` will also remove everything in `lib` and `modules`, and the `Makefiles`.

In general, you run `phaml` as you would any parallel program that uses the message passing library you have selected. Some examples are:

1. A sequential compilation. Run the primary program.

```
phaml
```

If you requested graphics, you must also run the graphics program.

```
phaml_graphics
```

2. PVM, as a master/slave program. Start the PVM demon on the nodes of the virtual machine. Then just run the master program, which will spawn the slaves and graphics.

```
phaml
```

3. LAM MPI, with spawning. You only start the master program, which spawns the slaves and graphics, so specify the number of processors to be 1.

```
mpirun -w -np 1 phaml
```

I have also found that in this case you don't need to use `mpirun`, you can just execute the master.

```
phaml
```

4. MPICH MPI, as an SPMD program. Since this does not spawn the slaves and graphics, you must use `mpirun` to specify the right number of processes. If there are to be  $n$  slaves, this consists of the sum of the following: One for the master,  $n$  for the slaves, one if the master is doing graphics, and  $n$  if the slaves are doing graphics. For example, if there are four slaves and the master is doing graphics,

```
mpirun -machinefile <file with list of hosts> -np 6 phaml
```

5. For OpenMP or hybrid OpenMP/MPI programs, set the `OMP_NUM_THREADS` environment variable to the desired number of threads, probably equal to the number of cores on the SMP computer, or number of cores on each node of the distributed memory parallel computer, and run the same as a sequential or MPI program.
6. See also the file `doc/HINTS` for my notes on running under debuggers, using ssh, redirecting displays, etc.

For parallel code with PVM, you will need to have `phaml`, `phaml.slave` and `phaml.graphics` (if you are using graphics) in `$HOME/pvm3/bin/$PVM_ARCH`. I just keep a symbolic link in there for each one, which points to the executable in my working directory.

If you request graphics, a window should pop up with the graphical display. The following actions are defined by default:

- arrow keys - pan
- left mouse button - rotate
- middle mouse button - zoom
- right mouse button - a menu of actions

The menu contains a large number of ways to change the graphics. See Section 3.5 for further details.

## Chapter 3

# Scalar Linear Elliptic Boundary Value Problems

The primary function of PHAML is to solve second-order scalar linear self-adjoint elliptic partial differential equations of the form given in Equation 1.1 with Dirichlet, natural (often Neumann), or mixed boundary conditions given by Equations 1.2 and 1.3. This section explains how to write a program that uses PHAML to solve these problems. Other classes of problems that can be solved using PHAML are addressed in subsequent sections. The information here is organized by concept. For an organization by subroutine and parameters, see the reference guide in Section 7.

### 3.1 Main program

PHAML is a library of subroutines. The user must write a program that calls these subroutines to solve the application problem. Ordinarily this just requires a simple main program and subroutines that define the equations, but the program can be as complex as needed. Subroutines in the PHAML library can be called from either the main program or subroutines.

#### 3.1.1 Parallelism

There are **six** three models of parallel processing supported by PHAML. The programming model used must correspond to the one specified by `PHAML_PARALLEL` in `mkmkfile.sh` when the PHAML library was built (see Section 2.2.1). The correspondence is given in Table 3.1.

In the master/slave model, the parallelism is almost transparent to the user. The user only needs to specify `nproc` in `phaml_create`, and start the program in the manner specified by the message passing library, requesting 1 process. With some libraries, for example LAM and MPICH2, even this can be transparent, since you can just run the executable when you are only starting 1 process, unless

model	PHAML_PARALLEL
sequential	sequential
master/slave	messpass_spawn
SPMD	messpass_nospawn
OpenMP SMP	openmp
hybrid OpenMP/MPI master/slave	hybrid_spawn
hybrid OpenMP/MPI SPMD	hybrid_nospawn

Table 3.1: Correspondence between PHAML\_PARALLEL in `mkmkfile.sh` and the parallel programming model.

your system uses a queuing program like PBS. Sequential programs can use the same program as a master/slave program, and `nproc` will be ignored. Single Program Multiple Data (SPMD) programs require a little more knowledge of the parallelism. Each instance of the program must determine if it is to be the master, a slave, or a graphics server, and act accordingly. This is explained in the next section. Since SPMD PHAML programs behave as if they were a master/slave program, with process 0 acting as the master, this document will often make reference to the master and slaves, even though the program might be SPMD. With OpenMP, the parallelism is completely transparent to the user, except for setting the environment variable `OMP_NUM_THREADS` to the desired number of threads. Usually this is equal to the number of cores on an SMP computer, or, for hybrid OpenMP/MPI programs, the number of cores on each node of a distributed memory parallel computer or cluster. However, it is sometimes advantageous to use one less than the number of cores if the operating system is running a lot of background processes that would run on the unused core.

### 3.1.2 Program structure

#### 3.1.2.1 Master/slave and Sequential

In a master/slave program, the user writes a program for the master process. The same program can be used for sequential, OpenMP and hybrid OpenMP/MPI master/slave models. The slave and graphics programs are given in the `src` directory as `slave.f90` and `graphmain.f90`. To compile these programs, see the `Makefiles` in the `examples` directory.

Figure 3.1 illustrates a very simple main program for a master/slave model. More complicated examples can be found in the `examples` directory. This example illustrates the main steps in writing a PHAML program.

First it must use module `phaml`. This gives access to all the public subroutines, the defined constants that are used as values for subroutine arguments, the `phaml_solution_type` data structure, and the defined constant `my_real` which contains the kind number for reals in PHAML.

Second, there must be at least one variable of type `phaml_solution_type`.



```

program simple
use phaml
type(phaml_solution_type) :: sol
call phaml_create(sol,nproc=2)
call phaml_solve_pde(sol,
                     max_vert=100000,
                     print_grid_when=PHASES,
                     print_grid_who=MASTER,
                     print_error_when=PHASES,
                     print_error_what=LINF_ERR,
                     print_error_who=MASTER)
call phaml_destroy(sol)
end program simple

```

Figure 3.1: A simple main program.

Variables of this type contain all the information known about the solution of the equation, including the current grid, solution, parallel processes, etc. The type is a public entity in module `phaml`, but the internals of the type are private. This means you cannot directly manipulate components of the variable, you can only pass it to the PHAML subroutines that operate on it.

Next, the solution variable is initialized by subroutine `phaml_create`. This creates an initial grid, allocates and initializes other components of the variable, and, for master/slave programs, spawns the slave and graphics processes. Here we requested the use of 2 slave processors. Other arguments will be discussed in subsequent sections as they become relevant. A full list can be found in Section 7.4.4.

The PDE is solved by calling `phaml_solve_pde`. This subroutine requires that the first argument be a solution variable. There are over 100 more arguments to this subroutine, which will be discussed in subsequent sections and can be found in Section 7.4.15. To make this manageable, all arguments are optional, have reasonable defaults, and should be given as keyword arguments, i.e. *dummyarg=value*, as shown in the example.

Finally, the solution variable should be destroyed by `phaml_destroy` to free memory and terminate any processes that were spawned by `phaml_create`.

### 3.1.2.2 SPMD

An SPMD program is more complicated because the master, slave and graphics processes are identical. However, it still operates like a master/slave program, so each process must determine if it is the master, a slave, or a graphics server. This is illustrated in Figure 3.2 for a program that uses an MPI library. **The same program can be used for a hybrid OpenMP/MPI SPMD model.**

In addition to using module `phaml`, it must use module `mpif_mod` to get access to the MPI include file.

```

program simple_spmd
  use phaml
  use mpif_mod
  integer :: whodrawg
  integer :: jerr
  integer :: my_processor, total_nproc
  integer :: nslave, subtract, divide

  ! initialize MPI, find out how many processors and what my rank is

  call mpi_init(jerr)
  call mpi_comm_size(MPI_COMM_WORLD,total_nproc,jerr)
  call mpi_comm_rank(MPI_COMM_WORLD,my_processor,jerr)

  ! set the graphics options

  whodrawg = NO_ONE

  ! determine how many processors for slaves and graphics

  subtract = 1
  if (whodrawg == MASTER .or. whodrawg == EVERYONE) subtract = 2
  divide = 1
  if (whodrawg == SLAVES .or. whodrawg == EVERYONE) divide = 2
  nslave = (total_nproc-subtract)/divide

  ! call the master, slave or graphics program depending on my rank

  if (my_processor == 0) then
    call phaml_master(whodrawg,nslave)
  elseif (my_processor <= nslave) then
    call phaml_slave
  else
    call phaml_graphics
  endif
end program simple_spmd

subroutine phaml_master(whodrawg,nslave)
  use phaml
  integer, intent(in) :: whodrawg, nslave
  type(phaml_solution_type) :: sol
  call phaml_create(sol,nproc=nslave,draw_grid_who=whodrawg)
  call phaml_solve_pde(sol, ... )
  call phaml_destroy(sol)
end subroutine phaml_master

```

Figure 3.2: An SPMD main program using MPI.

```

subroutine pdecoefs(x,y,cxx,cxy,cyy,cx,cy,c,rs)
use phaml
real(my_real), intent(in) :: x,y
real(my_real), intent(out), dimension(:,:) :: cxx,cxy,cyy,cx,cy,c
real(my_real), intent(out), dimension(:) :: rs
cxx=1
cxy=0
cyy=1
cx=0
cy=0
c=0
rs=0
end subroutine pdecoefs

```

Figure 3.3: `pdecoefs` for Laplace's equation.

The first step is to initialize MPI, determine how many processes there are, and determine the rank of this process. This is needed to determine what type of process this will be, so it cannot be hidden inside `phaml_create` like it is for master/slave programs.

Then the number of slaves is computed based on the total number of processes (which is specified when the program is launched, for example by `mpirun`) and how many graphics processes are needed.

Each process determines its type based on its rank (`my_processor`) and the number of slaves, and calls the corresponding main subroutine. The slave and graphics main subroutines are in the PHAML library. The main subroutine for the master is nearly identical to the main program in the master/slave model, as shown in Figure 3.2.

## 3.2 Defining the problem

### 3.2.1 Defining the PDE

The PDE in Equation 1.1 must be defined in subroutine `pdecoefs`. For convenience, the equation is repeated here.

$$-\frac{\partial}{\partial x} \left( c_{xx} \frac{\partial u}{\partial x} \right) - \frac{\partial}{\partial x} \left( c_{xy} \frac{\partial u}{\partial y} \right) - \frac{\partial}{\partial y} \left( c_{yx} \frac{\partial u}{\partial x} \right) + c_x \frac{\partial u}{\partial x} + c_y \frac{\partial u}{\partial y} + c_u u = f \quad \text{in } \Omega$$

`pdecoefs` receives a point  $(x, y)$  and returns the value of the functions  $c_{xx}$ ,  $c_{xy}$ ,  $c_{yx}$ ,  $c_x$ ,  $c_y$ ,  $c_u$  and  $f$  at that point in the arguments `cxx`, `cxy`, `cyy`, `cx`, `cy`, `c` and `rs` respectively. The interface for this routine is given in Section 7.3.6. For a scalar PDE, the return variables are  $1 \times 1$  arrays (they have higher dimension for systems of equations; see Section 4.2) and can be assigned with a whole array

```

subroutine bconds(x,y,bmark,itype,c,rs)
use phaml
real(my_real), intent(in) :: x,y
integer, intent(in) :: bmark
integer, intent(out) :: itype
real(my_real), intent(out) :: c(:,:),rs(:)
if (bmark == 1) then
    itype = NATURAL
else
    itype = DIRICHLET
endif
c=0
rs=0
end subroutine bconds

```

Figure 3.4: `bconds` for homogeneous boundary conditions, natural on boundary piece 1 and Dirichlet elsewhere.

boundary condition	itype
Eq. 1.2	DIRICHLET
Eq. 1.3 with $c_{bc} = 0$	NATURAL
Eq. 1.3 with $c_{bc} \neq 0$	MIXED
periodic	PERIODIC

Table 3.2: Permitted values for `itype` to specify the type of boundary conditions.

assignment statement, making their “arrayness” transparent. Figure 3.3 shows how `pdecoefs` could be written for Laplace’s equation  $\nabla^2 u = 0$ .

### 3.2.2 Defining the boundary conditions

The boundary conditions are defined in subroutine `bconds`. The interface for this routine is given in Section 7.3.1. On each piece of the boundary (Section 3.2.3 explains how to define the boundary piecewise and send a piece ID number to `bconds`) the boundary can be Dirichlet as defined by Equation 1.2

$$u = g,$$

natural (usually Neumann) or mixed as defined by Equation 1.3

$$\left( c_{xx} \frac{\partial u}{\partial x} + c_{xy} \frac{\partial u}{\partial y} \right) \frac{\partial y}{\partial s} - c_{yy} \frac{\partial u}{\partial y} \frac{\partial x}{\partial s} + c_{bc} u = g$$

or periodic (usually on opposite sides of a rectangle).

**bconds** receives as input a point  $(x, y)$  at which to return the boundary conditions, and **bmark**, an integer ID number for the piece of the boundary that  $(x, y)$  is on. This ID is defined along with the boundary of the domain (Section 3.2.3).

The type of boundary condition for the indicated piece is returned in **itype** using a symbolic constant listed in Table 3.2. The functions  $c_{xx}$ ,  $c_{xy}$  and  $c_{yy}$  in Equation 1.3 are the same as those in the PDE, Equation 1.1, which are defined in subroutine **pdecoefs** (Section 3.2.1) and not repeated here. The function  $c_{bc}$  in Equation 1.3 is returned in the variable **c**. Finally, the function  $g$  in Equations 1.2 and 1.3 is returned in **rs**.

**c** and **rs** are arrays, but for scalar problems the dimensions are all 1, and they can be assigned with a whole array assignment statement. (They have higher dimension for systems of equations; see Section 4.2.)

Assuming the domain is the unit square with the left side assigned 1 for the ID, Figure 3.4 shows a subroutine for the boundary conditions

$$\partial u / \partial n = 0 \quad \text{on } x = 0 \quad (3.1)$$

$$u = 0 \quad \text{elsewhere} \quad (3.2)$$

Periodic boundary conditions say that the value of the solution on one piece of the boundary is the same as the value on another piece. Usually the two pieces are opposite sides of a rectangle, but for generality we need a means of indicating which two pieces are matched up. This is done by making the piece ID of the two pieces have the same absolute value, with one positive and the other negative. **c** and **rs** are not used with periodic boundary conditions, but it is prudent to set them to 0. For an example of periodic boundary conditions, see **examples/periodic**.

Natural and periodic boundary conditions require that there be at least one point with Dirichlet boundary conditions to make the solution unique.

### 3.2.3 Defining the domain and initial grid

The domain and initial grid can be defined in one of two ways. The first is to supply subroutines that define the boundary explicitly as a piecewise curve in  $R^2$ . This approach is required if the boundary is not polygonal, i.e. if it has curved pieces, but can also be used for polygonal domains. The second is to provide triangle data files that were produced by the program Triangle [36]. These files can be created by writing a **.poly** file and running Triangle, or by using the first approach once and saving the triangle data files that PHAML creates, or by using the triangle data files from one of the PHAML examples. In either case, this gives a starting triangulation, which is preprocessed to create the initial grid.

To define the domain with subroutines, you must write three subroutines similar to those used by the domain processor of Rice [34].

**subroutine boundary\_point(ipiece,s,x,y)** contains the definition of the boundary. Each piece of the boundary is given by a parametric curve  $(x(s), y(s))$

for  $s$  in some range of  $R$ . The ending point of one piece must be the same as the starting point of the next piece, including the last piece of the outer boundary connecting to the first piece of the outer boundary, and the last piece of each hole connecting to the first piece of the same hole. The input parameters `ipiece` and `s` indicate which piece of the boundary and parameter value to evaluate, and the output parameters `x` and `y` are the requested point on the boundary.

`ipiece` is also used as the boundary marker for the triangle data files, and will be passed to `bcond` (Section 7.3.1) to simplify evaluation of boundary conditions. The endpoints of the boundary pieces will always be vertices of the grid, and are assigned the boundary marker of the piece they start. To get a boundary marker that is unique to an individual vertex, define a boundary piece of length 0 at that vertex (i.e., starting and ending parameters are the same). The last boundary piece cannot be a single point. Instead, make that point be the first piece.

`function boundary_npiece(hole)` (Section 7.3.3) returns the number of pieces used to define the boundary. If `hole` is 0, it should return the number of pieces that define the outer boundary. Otherwise it should return the number of pieces that define the `holeth` hole. The holes are numbered consecutively starting with 1. `boundary_npiece(0)` must return a positive integer if and only if subroutines are used to define the boundary. If it returns 0 or a negative integer, the starting triangulation is read from triangle data files. If `hole` is larger than the number of holes in the domain, it should return 0.

`subroutine boundary_param(start,finish)` returns the range of parameter values for every piece of the boundary. The output variables `start` and `finish` are `real(my_real)` arrays of length equal to the total number of boundary pieces, and should be assigned the starting and ending parameters of each piece.

The starting grid is created by PHAML by creating a `.poly` file for input to Triangle, which is then run to create triangle data files. Running Triangle involves a call to the intrinsic subroutine `system`, which is not standard in Fortran 90. Most compilers provide this routine as an extension, and use the same behavior for it, but if you have trouble you may have to change the definition of `my_system` in `sysdep.f90`.

You can control the fineness of the starting grid with the parameter `max_blen` in `phaml.create`. No triangle boundary side in the starting triangulation will have length longer than `max_blen`, and Triangle is run with `-a max_blen2√3/4` to limit the area of the triangles to that of an equilateral triangle with side length `max_blen`.

The name of the `.poly` file is determined by the parameter `triangle_files` in `phaml.create`, which has the default value `"domain"`. The file name will be the character string with the added suffix `.poly`. Note that if the full path is not given in `triangle_files`, the location of the file may be compiler dependent.

The second method for defining the domain and initial grid is to use triangle data files. These are the edge (`.edge`), element (`.ele`), neighbor (`.neigh`), node (`.node`) and poly (`.poly`) files created by the program Triangle.

To indicate that the starting grid is given by triangle data files, have `function`

`boundary_npiece(0)` return 0 or a negative number. The parameter `triangle_files` in `phaml_create`, which has the default value "domain", then contains the root of the name of the five triangle data files. If Triangle inserted an iteration number (e.g. .1) into the name, then it should be included.

If you are solving a PDE on the unit square, or on one of the other domains in the examples, then you only need to copy the triangle data files from the example and, if necessary, specify the name of the files.

Otherwise you must define the polygonal domain with a `.poly` file, which is basically a list of vertices and boundary line segments. Holes are allowed. See the documentation for Triangle [37] for the exact definition of a `.poly` file. You should specify a boundary marker for each vertex and boundary segment. This boundary marker will be passed to function `bcond` to simplify the definition of boundary conditions. With a `.poly` file defined, create the triangle data files with the command

```
triangle -pnej root_of_filename.poly
```

The `-p` flag indicates the input is a poly file. The `-n`, and `-e` flags force the writing of neighbor and edge files. The `-q` flag requests a quality mesh, which imposes bounds on the angles. The `-j` flag says to jettison (remove) vertices that are not part of the triangulation. Also, the `-a` flag may be useful to impose a maximum size (area) of the triangles, and the `-I` flag can be used to prevent the insertion of the iteration number in the filename. See the documentation for Triangle [37].

Regardless of whether they are created by Triangle from a `.poly` file, created by PHAML from the boundary subroutines, or copied from somewhere else, the triangle data files define a starting triangulation. PHAML requires an initial grid that satisfies certain conditions. This initial grid is determined from the starting triangulation automatically by PHAML through a process of refinement and grid smoothing. The triangles of the initial grid are obtained by bisection or trisection of the starting triangles, followed by some node movement.

### 3.2.4 The true solution

If the true (a.k.a. exact or analytical) solution is known, it and its first derivatives can be provided through subroutines `true`s, `true`xs and `true`ys. If these are provided, then the program can print norms of the error (Section 3.4.2) or choose the error as the function to visualize with the graphics. If `true`s is defined, but `true`xs and/or `true`ys is not, then the energy norm of the error cannot be printed, but all other norms and the graphics are still valid.

The interfaces for these routines are given in Sections 7.3.9, 7.3.10 and 7.3.11. They are function subroutines that return the value of the solution,  $x$  derivative of the solution, and  $y$  derivative of the solution, respectively. The input variables give the point  $(x, y)$  at which to return the solution. There are also two input variables, `comp` and `eigen`, that are not relevant for scalar boundary value problems (they are used for systems of equations and eigenvalue problems).

### 3.3 Solution method

There are many arguments to `phaml_solve_pde` that affect the details of the solution method. This section explains the options available. It is organized with sections on discretization, refinement, the error indicator, the linear system solver, load balancing and termination. Each section briefly describes the methods used, but this is not intended to be a thorough treatment of parallel adaptive multilevel finite element methods. See the references for more details of the methods.

See also Section 7.4.15 which formally defines the arguments to `phaml_solve_pde`.

#### 3.3.1 Discretization

PHAML uses a standard Galerkin finite element method (see, for example [39][41]) to approximate the solution of Equations 1.1-1.3, which we briefly describe here. The domain is partitioned into a set of conforming triangles,  $T$ , (the grid or mesh) and the approximation space is defined as the Hilbert space of continuous functions that are polynomials over each triangle. The degree of the polynomial can be different over different triangles. The  $p$ -hierarchical basis,  $\Phi = \{\phi_i\}$ , of either Szabo & Babuška [41] or Carnevali et al. [9] spans the space, and gives a representation for the approximate solution

$$u_T = \sum_{\phi_i \in \Phi} \alpha_i \phi_i \quad (3.3)$$

Basis functions can be categorized in three groups. The linear ( $p = 1$ ) basis functions are in one-to-one correspondence with the vertices of the grid. Higher order basis functions are associated with edges or faces (triangles). There are  $p - 1$  edge bases associated with an edge of degree  $p$ , and  $(p - 1)(p - 2)/2$  face bases associated with a face of degree  $p$ .

The space has the energy inner product defined by

$$\langle u, v \rangle = \int_{\Omega} c_{xx} u_x v_x + c_{xy} u_y v_x + c_{yy} u_y v_y + c_x u_x v + c_y u_y + c_u uv \quad (3.4)$$

where  $c_{xx}$ ,  $c_{xy}$ ,  $c_{yy}$ ,  $c_x$ ,  $c_y$ , and  $c_u$  are from Equation 1.1, and the subordinate energy norm  $\|u\|^2 = \langle u, u \rangle$ . (Note: We need  $c_{xy} = c_x = c_y = 0$  for 3.4 to be an inner product because of symmetry, and  $c_u \geq 0$  for  $\langle u, u \rangle$  to be a norm because of positivity, but formally we proceed with these definitions knowing that some of the following statements are not strictly true.) The approximate solution is the function in the space that minimizes the energy norm of the error, i.e. the error is orthogonal to the space under the energy inner product. Thus the approximate solution satisfies

$$\langle u_T, \phi_i \rangle = \langle f, \phi_i \rangle_2 \quad \forall \phi_i \in \Phi \quad (3.5)$$

where  $\langle \cdot, \cdot \rangle_2$  is the  $L_2$  inner product  $\langle u, v \rangle_2 = \int_{\Omega} uv$ . Substituting Equation 3.3 into Equation 3.5 leads to the discretized form of the problem  $Ax = b$



with  $a_{ij} = \langle \phi_i, \phi_j \rangle$ ,  $b_i = \langle f, \phi_i \rangle_2$ , and  $x$  the vector of  $\alpha_i$ 's. Natural boundary conditions are imposed automatically by the energy inner product. Dirichlet boundary conditions are imposed by replacing the corresponding equations with equations that set the corresponding  $\alpha_i$ 's directly from the boundary condition.

There are only four arguments that affect how discretization is performed. First, you can select the degree of the polynomials in the approximation space with the argument `degree`. This sets all elements to start with the given degree, even if you use  $p$ - or  $hp$ -refinement. Second, the integrals of Equation 3.4 are computed by numerical quadrature, with the order of the quadrature rule determined such that integrals are exact for polynomials of the degree  $2(p-1)$  where  $p$  is the degree of the basis over each triangle. Sometimes this may not be accurate enough, in which case you can supply an increase of the order through `inc_quad_order`. Third, in triangles that are owned by a different processor (see Section 3.3.5), the quadrature is performed by the owner and communicated by message passing. If this were not done, then the integrals over those triangles would be less accurate than the same integrals on the processor that owns the triangle, because it will have refined the triangle into many smaller triangles. However, sometimes this doesn't matter, such as when solving Laplace's equation (all the integrals are exact) or when using a solver other than the (default) hierarchical basis multigrid solver 3.3.4.1, which don't use the unowned triangles. You can avoid the extra computation and communication with `ignore_quad_err=true`. Finally, you can select how the degree of an edge is determined when the triangles that share that edge have different degrees through `edge_rule`. If you select `MINIMUM_RULE` (the default), then the edge gets the smaller of the two triangle degrees. If you select `MAXIMUM_RULE`, then the edge gets the larger of the two triangle degrees.

### 3.3.2 Refinement

One of the major phases in `phaml_solve_pde` is refinement of the grid to increase the size of the approximation space, which reduces the norm of the discretization error.  $h$ -refinement refers to subdividing triangles by newest node bisection (see [24][26]).  $p$ -refinement refers to increasing the polynomial degree over a triangle. In either case the refinement can be undone, referred to as derefinement. Derefinement can be switched off with the argument `derefine=false`. Refinement can be done uniformly, i.e. refine all triangles in the grid, or adaptively, i.e. refine a selected subset of the triangles.

The type of refinement that is performed is determined by the argument `reftype`. This argument can have one of the values `H_UNIFORM`, `H_ADAPTIVE`, `P_UNIFORM`, `P_ADAPTIVE` or `HP_ADAPTIVE`. With `HP_ADAPTIVE` an element may be refined by either  $h$ - or  $p$ -refinement.  $hp$ -adaptive strategies are explained later in this section.

There are several ways to determine how much refinement to perform in one refinement phase. This is controlled by the argument `refterm`. The allowed values for this argument are given in Table 3.3.

DOUBLE_NVERT
DOUBLE_NVERT_SMOOTH
DOUBLE_NELEM
DOUBLE_NELEM_SMOOTH
DOUBLE_NEQ
DOUBLE_NEQ_SMOOTH
HALVE_ERREST
KEEP_NVERT
KEEP_NVERT_SMOOTH
KEEP_NELEM
KEEP_NELEM_SMOOTH
KEEP_NEQ
KEEP_NEQ_SMOOTH
ONE_REF
ONE_REF_HALF_ERRIND

Table 3.3: Permitted values for `refterm` to determine how much refinement to do.

“Double” means that the refinement should approximately double the number of vertices, elements or equations in the grid. The factor 2 was chosen because this corresponds to the increase that would occur with one uniform  $h$ -refinement of the grid. If a different factor is desired, it can be specified with the real valued argument `inc_factor`.

Strictly doubling the number of entities may lead to grids that are nonsymmetric, which may be undesirable if the solution is symmetric. This can be improved by using the values that contain `SMOOTH`. With these values, after the doubling is completed the refinement will continue by refining all elements with a similar error indicator (Section 3.3.3) to the last element refined.

“Keep” attempts to change the grid by derefining some elements and refining others while keeping the total number of entities approximately the same. This is useful, for example, with time dependent problems where the grid should track the movement of some feature of the solution. If the corresponding “max\_” value is supplied (for example, `max_vert` along with `KEEP_NVERT`), then the attempt is to match the number supplied. Otherwise, the attempt is to match the number of entities currently in the grid, which can cause a drift in the number of entities after several iterations.

`HALVE_ERREST` refines the grid until the maximum error indicator has been reduced by half. This corresponds to the expected reduction of error by a uniform  $h$ -refinement with `degree=1` and a smooth solution.

`ONE_REF` is a scheme in which no element gets refined more than once in a refinement phase. The argument `reftol` can provide a tolerance for which elements should be refined. All elements with an error estimate larger than  $\text{reftol}/\sqrt{n}$  are refined, where  $n$  is the starting number of elements.

ONE\_REF\_HALF\_ERRIND is also a scheme in which no element gets refined more than once in a refinement phase. Those elements with an error indicator larger than half (actually,  $1/\text{inc\_factor}$ ) of the maximum error indicator get refined.

A large number of strategies for *hp*-adaptive refinement have been implemented in PHAML. Selection of which strategy to use is controlled by the argument `hp_strategy`. Some strategies may override the values of some arguments, in particular `refterm`, `derefine`, `inc_factor`, and `error_estimator`. The strategies are briefly described here. For a more complete description, see the survey paper [30].

In all of the *hp* strategies, if an element that would be refined by *p* refinement has degree `max_deg`, then it is refined by *h* refinement (unless `stop_on_maxdeg` is `true`), and if an element that would be refined by *h* refinement has level `max_lev`, then it is refined by *p* refinement (unless `stop_on_maxlev` is `true`). If an element has both `max_deg` and `max_lev`, then it is not refined.

The following values are currently allowed for `hp_strategy`:

HP\_APRIORI is an extension of the second strategy in Ainsworth & Senior [2]. The basic approach is to refine elements by *p* refinement, except elements that contain a known point singularity are refined by *h* refinement. The extension is that we do not limit irregularities to point singularities, and you can specify the strength of that irregularity.

This strategy requires that the user provide a function subroutine that indicates where the solution has an irregularity (i.e. is singular or nonsmooth). See the end of `examples/L-domain/pde.f90` for an example of this subroutine. Also see Section 7.3.8. In theory, this routine should return the largest value of *m* such that the solution is in  $H^m(T)$ , i.e. the derivative up to order *m* are in  $L^2$ , where *T* is the triangle whose vertices are given as input to the function. For multicomponent solutions, it should return the worst (i.e. smallest) such *m* among the components.

In practice, it can be used to guide refinement in other *a priori* known trouble areas, such as sharp peaks, boundary layers and wave fronts. The actual use is that *p* refinement is performed if the current degree of the triangle is less than the returned value, and *h* refinement is performed otherwise. So, for example, if you know some region contains a boundary layer, you could return 3.1 for any triangle that intersects that region to perform *h*-refinement with cubic elements over the boundary layer.

Several *hp*-adaptive strategies use the same principle as the HP\_APRIORI strategy, but attempt to estimate the regularity in various ways. HP\_NEXT3P is an implementation of the first strategy given in Ainsworth & Senior [2]. Three error estimates are computed by the equilibrated residual method with spaces of degree  $p + 1$ ,  $p + 2$  and  $p + 3$ , where *p* is the current degree of the element. If  $\phi$  is the actual error over the element, the error estimates  $\eta_{p+q}$ ,  $q = 1, 2, 3$ , converge to  $\phi$  with error bounds

$$||\phi||^2 - \eta_{p+q}^2 \leq C(p+q)^{-2\alpha}.$$

The three error estimates are used to determine the three unknown constants  $||\phi||$ , *C* and  $\alpha$ . The regularity is then given by  $m = 1 + \alpha$ .

HP\_PRIOR2P\_E and HP\_PRIOR2P\_H1 are inspired by a strategy of Suli, Houston and Schwab [40]. Two error estimates are computed using the local  $p$  error estimator with spaces of degree  $p - 1$  and  $p - 2$ . The regularity is estimated by

$$m = 1 - \frac{\log(\eta_{p-1}/\eta_{p-2})}{\log((p-1)/(p-2))}$$

HP\_PRIOR2P\_E uses the energy norm of the error estimate, whereas HP\_PRIOR2P\_H1 uses the  $H^1$  norm.

HP\_TYPEPARAM selects the type parameter strategy of Gui and Babuška [16]. Perceived smoothness of the solution over element  $t$  is given by

$$R(t) = \begin{cases} \frac{e(t,p)}{e(t,p-1)} & e(t,p-1) \neq 0 \\ 0 & e(t,p-1) = 0 \end{cases}$$

where  $p$  is the degree of the element and  $e$  is the error indicator for the given degree. The type parameter,  $\gamma$ , determines the type of the element. If  $R(t) > \gamma$  then  $t$  is of  $h$ -type and will be  $h$  refined; otherwise it is  $p$ -type and will be  $p$  refined.  $\gamma$  is specified with `tp_gamma`.

Another way of determining if the solution is smooth is to compare the local *a posteriori* error estimate with a prediction of what the error estimate should be, after refinement, under the assumption that the solution is smooth and has the proper asymptotic convergence rate [23]. If  $\eta_E$  is the error estimate on element  $E$ , and  $E'$  is a child of  $E$  after refinement, then the predicted error indicator for  $E'$  is

$$\eta_{E'}^{\text{pred}} = \begin{cases} \frac{1}{2}\gamma_h \left(\frac{1}{\sqrt{2}}\right)^{p_E} \eta_E & \text{if } E \text{ was } h \text{ refined} \\ \sqrt{\gamma_p} \eta_E & \text{if } E \text{ was } p \text{ refined} \end{cases}$$

where  $p_E$  is the degree of element  $E$ , and  $\gamma_h$  and  $\gamma_p$  are control parameters given in `sp_gamma_h` and `sp_gamma_p`. Then, if  $E'$  is selected for refinement, it is refined by  $h$ -refinement if  $\eta_{E'} > \eta_{E'}^{\text{pred}}$ , and by  $p$ -refinement otherwise. This strategy is selected with `HP_SMOOTH_PRED`.

HP\_BIGGER\_ERRIND selects a strategy in which both the `LOCAL_PROBLEM_H` and `LOCAL_PROBLEM_P` error indicators are computed. It is similar to the strategy for 1D problems in [35]. The error indicator specified by `error_estimator` is used to determine which elements will be refined (see 3.3.3). For each element that is selected for refinement, it is  $h$  refined if `LOCAL_PROBLEM_H` gives the larger error indicator, and  $p$  refined if `LOCAL_PROBLEM_P` is larger. The premise is that the local problem error indicators approximate how much the solution will change if the refinement is performed, and one should perform the type of refinement that will cause the largest change, and hence reduce the error the most.

The Texas 3 Step strategy [32] is selected by `HP_T3S`. The three steps are 1) uniform  $h$  refinement, 2) adaptive  $h$  refinement, and 3) adaptive  $p$  refinement. Steps 2 and 3 are repeated until a termination criterion is met. The first step is intended to create a starting grid for which the rate of convergence has reached

the asymptotic region. The number of uniform refinements performed is controlled by `t3s_nunif`, which is 0 by default. If the method is performing poorly at the start, try increasing it.

Steps 2 and 3 attempt to reduce the error by a prescribed amount. Specifically, if the error estimate at the beginning of step 2 is  $\theta$ , then the adaptive  $h$  refinement attempts to reduce it to  $\gamma\eta\theta$ , and the adaptive  $p$  refinement attempts to reduce it to  $\eta\theta$ . The parameters  $\gamma$  and  $\eta$  are specified by `t3s_gamma` and `t3s_eta`, and currently default to 6 and 0.1. This is still experimental and may change. In each step a formula is used to determine how much to refine each element. The regularity of the solution is used in this formula, and is specified by subroutine `regularity` the same as the `HP_APRIORI` strategy. To avoid overrefining in one step, an upper bound is placed on the number of times an element can be refined in one step. These bounds are given by `t3s_maxref` for the  $h$  refinement of step 2, and `t3s_maxdeginc` for the  $p$  refinement of step 3. Currently they both default to 3, but this is still experimental and may change. Currently derefinement is not allowed with this strategy. You should use `derefine=.false.` to avoid a warning message.

`HP_ALTERNATE` gives another strategy that alternates between  $h$  and  $p$  adaptive refinement. It is similar to `HP_T3S`, but instead of using a formula to estimate the correct amount of refinement of each element, it simply performs adaptive refinement until the error estimate reaches the target value of  $\gamma\eta\theta$  for the  $h$  refinement step and  $\eta\theta$  for the  $p$  refinement step.  $\gamma$  and  $\eta$  are again specified with `t3s_gamma` and `t3s_eta`. Unlike the Texas 3 Step strategy, it does not begin with uniform refinements.

`HP_NLP` attempts to optimize the grid by solving a nonlinear programming optimization problem. It is based on a method proposed by Patra and Gupta [33]. In essence, the optimization problem says to determine a number of  $h$ -refinements and  $p$ -refinements of each element to minimize the number of degrees of freedom subject to the predicted error estimate being less than one fourth of the current error estimate, plus other constraints that keep the number of refinements within feasible bounds, such as the degree being positive. Among these constraints are use-specified limits on how much change can occur in one refinement stage. The parameters `nlp_max_h_dec`, `nlp_max_p_dec`, `nlp_max_h_inc`, and `nlp_max_p_inc` specify the maximum amount of  $h$  and  $p$  coarsening and refinement to use at once. The defaults are 1, 1, 5 and 2. Solution of the nonlinear program takes a long time, except for very coarse grids, so we do not encourage the use of this strategy. This strategy requires the use of a third party optimization routine which is not included in the PHAML distribution. If you feel the need to use this strategy, contact [phaml@nist.gov](mailto:phaml@nist.gov) for instructions on how to obtain and configure this routine. This strategy cannot be used in parallel, because the optimization routine is not parallel.

A strategy for 1D problems based on the coefficients of the Legendre polynomial expansion of the solution was given in [22]. If  $p$  refinement is optimal, then the coefficients should decrease exponentially. A least squares fit of the last four coefficients is made to  $ce^{-\sigma p_i}$ , where  $p_i$  are the degrees of the last four Legendre polynomials. If  $\sigma > 1$  then  $p$  refinement is performed, otherwise  $h$  refinement.

To extend this to 2D, we use the coefficients of the  $p$ -hierarchical basis. The  $i^{th}$  “coefficient” is given by the sum of the absolute values of all coefficients of bases of degree  $i$ . This strategy is selected with `HP_COEF_DECAY`.

Another strategy based on Legendre coefficients is given in [17]. This strategy, selected with `HP_COEF_ROOT`, uses a “root test” on the coefficients to estimate the regularity of the solution.  $p$  refinement is used where the solution is sufficiently smooth, and  $h$  refinement elsewhere. Again, PHAML uses the coefficients of the  $p$ -hierarchical basis instead of the Legendre coefficients.

A strategy based on that of Šolín et al. [38] is implemented as `HP_REFSOLN_ELEM`. This strategy has its own overall structure to the adaptive refinement algorithm, so most of the refinement-related parameters to PHAML have no effect, including the error indicator. Also note that the global energy norm error estimate is actually a relative  $H^1$  norm error estimate (which is equivalent), and the  $L_\infty$  and  $L_2$  error estimates are not available. However, all the usual error estimates are available through `phaml_query`. The method begins by computing a reference solution,  $u_{\text{ref}}$  on a grid that has been globally refined by both  $h$  and  $p$ . The error indicator for element  $E$  is computed as  $\|u_{\text{ref}} - u_T\|_{H^1(E)}$ . Elements for which the error indicator is larger than the largest error indicator divided by `inc_factor` are selected for refinement. For each element to be refined, 11 candidate refinements are considered:  $p$ -refine to degree  $p+1$ ,  $p$ -refine to degree  $p+2$ , and  $h$ -refine with the child degrees being all combinations of  $p_0$ ,  $p_0+1$  and  $p_0+2$  where  $p$  is the current degree of the element and  $p_0 = \lfloor (p+1)/\sqrt{2} \rfloor$ . For each candidate, the  $H^1$  projection of the reference solution onto the corresponding space,  $\Pi_{\text{candidate}}$ , is computed, and the projection error is computed as  $\|u_{\text{ref}} - \Pi_{\text{candidate}}\|_{H^1(E)}$ . The projection error onto the unrefined space is also computed. The winning candidate refinement is selected based on the amount by which the logarithm of the projection error is reduced relative to the increase in the number of degrees of freedom. Experiments have indicated that it is often beneficial to bias the refinement towards  $p$ -refinement, which can be done by setting `refsoln_pbias` to a number greater than 1. We have found that generally `refsoln_pbias=2` works best if the solution is singular or has exceptionally strong features, and `refsoln_pbias=4` is best for smooth solutions or moderately strong features. `HP_REFSOLN_ELEM` has not yet been implemented in parallel, so it currently requires `nproc=1`.

Another strategy based on using a reference solution is that of Demkowicz [10]. `HP_REFSOLN_EDGE` is based on this strategy. It is similar to `HP_REFSOLN_ELEM`, but one first considers the refinement of edges by projecting the reference solution onto spaces corresponding to candidate refinements of the edge. Once it is determined how to refine the edge, the refinement of the element is determined by consistency of the  $h$ -refinement of the edge and the minimum edge rule for  $p$ . This method has not been implemented in parallel yet, so it currently requires `nproc=1`.

for all situations yet. In particular, it requires `system_size=1`, `num_eval=1` and `nproc=1`.

### 3.3.3 Error indicator

The heart of an adaptive refinement strategy is the error estimator, or more properly, error indicator. An error indicator is computed for each triangle, and those with the largest error indicators are refined. PHAML currently contains six error indicators. The error indicator is selected by the argument `error_estimator` which takes one of the values `EXPLICIT_ERRIND`, `LOCAL_PROBLEM_H`, `LOCAL_PROBLEM_P`, `EQUILIBRATED_RESIDUAL`, `HIERARCHICAL_COEFFICIENT` or `TRUE_DIFF` to select one of the error indicators. It can also take the value `INITIAL_CONDITION`.

Many error indicators are based on the interior residual within elements

$$r = f - \mathcal{L}u_T \quad (3.6)$$

and the boundary residual on element edges

$$R = \begin{cases} g - \mathcal{B}u_T & \text{if the edge is on } \partial\Omega_N \\ 0 & \text{if the edge is on } \partial\Omega_D \\ -[\frac{\partial u_T}{\partial n}] & \text{if the edge is interior} \end{cases} \quad (3.7)$$

where  $\mathcal{L}$  is the operator defined in Equation 1.1,  $\mathcal{B}$  is the operator defined in Equation 1.3,  $f$ ,  $g$ ,  $\partial\Omega_N$ , and  $\partial\Omega_D$  are defined in Equations 1.1-1.3,  $u_T$  is the approximate solution, and  $[\frac{\partial u_T}{\partial n}]$  is the jump in the normal derivative across the element boundary. See [1] for a more detailed treatment of error indicators.

`EXPLICIT_ERRIND` selects the explicit error indicator defined in Chapter 2 of [1], with slight modifications for the more general operator. The error indicator is based on the norm of the residual. The explicit error indicator,  $\eta_e$ , for an element  $e$  is given by

$$\eta_e^2 = h_e^2 \|r\|_{L_2(e)}^2 + h_e \|R\|_{L_2(\partial e)}^2 \quad (3.8)$$

where  $h_e$  is the longest edge length of the element. The error indicator is used to guide adaptive refinement. The energy norm error estimate is given by an unknown constant times the square root of the sum of the squares of the error indicators. For the  $L_2$  error estimate, the powers on the  $h$ 's are increased by 2. For the  $L_\infty$  error estimate, the  $L_\infty$  norm of the residuals is used and the maximum error indicator gives the error estimate. In PHAML the unknown constant has been chosen to be 1/20 for the energy norm and 1/100 for the  $L_2$  and  $L_\infty$  norms based on the results obtained with the example and test problems, although these constants work better with high order or  $hp$ -adaptive elements than linear elements. The integrals for the energy and  $L_2$  norms are approximated with a 4<sup>th</sup> order quadrature rule, and the  $L_\infty$  norm is approximated by values at the quadrature points of the same quadrature rule. The explicit error indicator is quite efficient to compute and generally provides good guidance for adaptive refinement, but one cannot be certain of the accuracy of the error estimates.

`LOCAL_PROBLEM_H` and `LOCAL_PROBLEM_P` are based on solving a small local problem. Equation 1.1 is modified by replacing  $f$  with the interior residual

*r*. For a linear PDE, the solution of this equation is the error. The modified equation is solved on a domain consisting of one or two triangles to get an estimate of the error over that small domain. To approximate the solution of this small problem with slightly higher accuracy than the current solution  $u_T$ , the triangle(s) are refined once either by  $h$ -refinement or  $p$ -refinement. The  $h$ -refinement form uses two triangles and refines them as a pair by bisection. Homogeneous Dirichlet boundary conditions are applied. The  $p$ -refinement form uses one triangle with polynomial degree one larger than that used for  $u_T$ . Natural boundary conditions are applied using the boundary residual  $R$ . Since the refinement used for the error indicator is exactly what would occur if the triangle(s) was (were) chosen for refinement by  $h$ - or  $p$ -refinement, this gives an estimate of how much the solution would change if this triangle(s) was (were) refined. The local problem error indicators are very accurate, but relatively expensive, because of the computation required to set up and solve the elemental matrices for each triangle.

The `EQUILIBRATED_RESIDUAL` error indicator is similar to the `LOCAL_PROBLEM_P` indicator, except that the residual is equilibrated so that the boundary fluxes are in equilibrium with the interior residual loads (see Chapter 6 in [1]). This insures that the discrete Neumann problem has a solution, which in turn allows the use of the full local approximation space of degree  $p$ , whereas the `LOCAL_PROBLEM_P` uses a reduced space spanned by the bases of exact degree  $p$ . It can be shown that this method produces an error estimate that is a guaranteed upper bound on the actual energy norm of the error, if the local problem was solved exactly. Of course, the local problem is solved approximately so the guarantee does not hold, but the PHAML examples typically find the ratio of error estimate to actual error to be about 1.25. This is the most accurate and stable error indicator in PHAML, but also the most computationally expensive.

A less expensive approach, given by `HIERARCHICAL_COEFFICIENT`, is to simply examine the hierarchical coefficients (see [24]) of  $u_T$  over each triangle, which is similar to computing the local problem indicator one refinement back. For linear elements, the coefficient of the  $h$ -hierarchical basis is used. For higher order elements, the coefficients of the  $p$ -hierarchical bases are used. This indicator can provide a very inexpensive way to guide adaptive refinement, but is not likely to give an accurate estimate of the norm of the error.

The `TRUE_DIFF` error indicator uses the difference between the true solution (if given, see 3.2.4) and  $u_T$ . The energy and  $L^2$  norms are estimated with a sixth order quadrature rule, and the  $L^\infty$  norm is estimated using the same quadrature points. This is primarily available for comparison with new error indicators and would not normally be used.

The `INITIAL_CONDITION` estimate uses the difference between the function given in subroutine `iconds` and  $u_T$ . Normally this is used to define an initial grid for time dependent or nonlinear problems (see Section 4.3), but it can also be used to provide a function that indicates where you believe the grid should be finer.



### 3.3.4 Linear system solver

Another major phase of `phaml_solve_pde` is the solution of the linear system of equations to get the coefficient vector of the solution. PHAML contains a hierarchical basis multigrid method as the primary linear system solver, along with two basic Krylov space solvers, conjugate gradients and GMRES. It also contains hooks into several freely available software packages of direct and iterative parallel solvers for comparative studies and for situations where the native solvers are not sufficient. For information on obtaining the optional software discussed in this section, see Section 2.1. The solver is selected by the arguments `solver` and `preconditioner`.

#### 3.3.4.1 Hierarchical basis multigrid solver

The default solver is the hierarchical basis multigrid solver (HBMG), which can be explicitly selected with `solver = MG_SOLVER`. (Note: this solver must NOT be used if the linear system is not symmetric (i.e.,  $c_{xy}$ ,  $c_x$  or  $c_y$  from Equation 1.1 is nonzero, or a coefficient matrix in a coupled system of equations is nonsymmetric) or indefinite (i.e.,  $c_u$  is negative with sufficiently large magnitude).) It combines the ideas of an  $hp$ -multigrid method [29]<sup>[31]</sup> with the  $h$ -hierarchical basis multigrid method for linear elements [25]. The equations corresponding to high order face basis functions 3.3.1 are first removed by static condensation [44]. A  $p$ -multigrid cycle is applied to the equations corresponding to vertex and edge basis functions as follows. The cycle is like a normal multigrid V-cycle, except the levels are given by the polynomial degree. Some number of Gauss-Seidel iterations are performed on all equations up to degree  $p_{\max}$  where  $p_{\max}$  is the maximum degree. Then the Gauss-Seidel iterations are applied to all equations up to degree  $p_{\max} - 1$ , then  $p_{\max} - 2$ , etc., until only the linear and quadratic equations are used. With the  $p$ -hierarchical basis, the “intergrid transfers” are automatic. Then the “coarse grid” equations (equations corresponding to the vertices, i.e. linear elements) are solved “exactly” by using a standard  $h$ -multigrid method. In PHAML, the  $h$ -hierarchical basis multigrid method is used, as follows. The residual is injected into the equations corresponding to the linear basis functions. The equations corresponding to the linear basis functions are relaxed by some red-black Gauss-Seidel iterations. The black equations are those corresponding to equations in the coarse grid, and the red equations are from the fine grid but not the coarse grid. A half iteration is allowed, which means relaxing the red equations but not the black. A basis change is then performed on those equations to convert from a nodal basis to a 2-level  $h$ -hierarchical basis, and the coarse grid equations are extracted. The process of relaxation, basis change and extraction is repeated until the grid consists only of the initial grid, i.e. all elements have refinement level 1. The coarsest grid problem is solved with a LAPACK direct solver. The process is then reversed by performing relaxation followed by conversion from 2-level  $h$ -hierarchical basis to nodal basis to get the next finer grid. After the finest level is reached, the second half of the  $p$ -multigrid cycle performs relaxations

on all equations up to degree 2, then up to degree 3, etc. This constitutes one multigrid V-cycle. Cycles are repeated until some termination criterion is met. Finally, the equations corresponding to face basis functions are solved directly.

There are several arguments to `phaml_solve_pde` that determine the specifics of the multigrid algorithm. `mg_prerelax_ho` and `mg_postrelax_ho` determine the number of Gauss-Seidel iterations to perform on each level of the  $p$ -multigrid cycle, before and after solving the linear basis equations. `mg_prerelax` and `mg_postrelax` give the number of half-red-black Gauss-Seidel iterations to perform before and after solving the coarse grid problem for the linear equations. A half iteration relaxes the red equations but not the black equations. So, for example, `mg_prerelax=2` specifies one red-black iteration.

The termination of the multigrid cycles can be specified as a fixed number of cycles or by a tolerance on the residual of the linear system. To terminate by a tolerance, specify the tolerance in `mg_tol`. Iterations continue until the  $\ell^2$  norm of the residual of the scaled linear system is less than `mg_tol`. There are two symbolic constants that provide special values for `mg_tol`. `MG_ERREST.TOL` says to cycle until the residual is reduced to some fraction of the global error estimate. This avoids excess computation from solving the system more accurately than is necessary relative to the discretization error. `MG_NO_TOL` says to use a fixed number of iterations rather than a tolerance. The argument `mg_cycles` gives the fixed number of cycles to perform. Used in conjunction with a tolerance, `mg_cycles` provides an upper bound on the number of cycles to guarantee the iteration will terminate.

The HBMG as implemented in PHAML uses the full domain partition approach to parallelism by default [27]. This approach allows the use of only two messages in each  $h$ -multigrid cycle, one at the coarsest grid and one at the finest grid, with a (usually very minor) reduction in the convergence rate of the  $h$ -hierarchical basis multigrid method, but still uses communication after each level of the  $p$ -multigrid cycle. PHAML also provides the option of using conventional parallelism with messages on each  $h$ -level, which gives exactly the same results as the sequential HBMG. This is selected by specifying `mg_comm = MGCOMM_CONVENTIONAL`.

### 3.3.4.2 Krylov space solvers

PHAML also contains two Krylov space solvers: conjugate gradients and GMRES. For a description of these methods, see [8]. The methods are selected by setting `solver` to be `CG_SOLVER` and `GMRES_SOLVER`, respectively. As with the HBMG method, PHAML begins by eliminating the equations associated with face basis function by static condensation. It also eliminates the equations associated with Dirichlet boundary conditions so that the working matrix is symmetric.

Either method can be used without preconditioning via `preconditioner=NO_PRECONDITION` or with HBMG as the preconditioner via `preconditioner=MG_PRECONDITION`. As a preconditioner, HBMG does not need to converge to the solution, just give an approximation. It is usually sufficient to use two iterations, so the default

HBMG parameters are set appropriately when HBMG is used as a preconditioner instead of a solver. Also, the HBMG preconditioner uses, by default, `ignore_quad_err=.true.` to avoid reducing the larger quadrature errors in un-owned elements, and `mg_comm = MGCOMM_NONE` to skip the communication steps in HBMG.

There are three parameters that control the Krylov space solvers. `krylov_iter` provides an upper bound on the number of iterations allowed, and `krylov_tol` gives a tolerance on the  $\ell^2$  norm of the residual. Convergence is declared when the residual is less than the tolerance. If the maximum number of iterations is achieved before convergence, the solution is accepted and a warning is printed. `krylov_restart` gives the number of restart vectors for GMRES.

### 3.3.4.3 Alternative direct solvers

#### 3.3.4.3.1 LAPACK

The LAPACK library [5] is always linked with a PHAML program. In limited situations it can be used as the linear system solver. It is specified by `solver=LAPACK_SPD_SOLVER` for most problems, which generate a symmetric positive definite matrix, or by `solver=LAPACK_INDEFINITE_SOLVER` for problems that generate an indefinite symmetric matrix. However, it can only be used as the solver for sequential programs because it is not a parallel library. And it should only be used for relatively small problems.

#### 3.3.4.3.2 MUMPS

MUMPS [3][4] is a parallel direct solver from the European project PARASOL. Currently the PHAML interface to MUMPS only supports double precision, i.e. `my_real=kind(0.0d0)` in `global.f90`. MUMPS is specified with `solver=MUMPS_SPD_SOLVER` for the symmetric positive definite solver, `solver=MUMPS_GEN_SOLVER` for the symmetric general solver, or `solver=MUMPS_NONSYM_SOLVER` for the non-symmetric solver.

#### 3.3.4.3.3 SuperLU

SuperLU [21] is a parallel direct solver from the Lawrence Berkley National Laboratories. It is specified by `solver=SUPERLU_SOLVER`.

### 3.3.4.4 Alternative iterative solvers

#### 3.3.4.4.1 PETSc

PETSc [6][7] is a parallel library of preconditioners and Krylov space iterative solvers from Argonne National Laboratories. PHAML provides access to most of the methods in PETSc and several parameters to those methods. Table 3.4 contains the available PETSc solvers as specified by `solver`. Table 3.5 contains the available preconditioners for the PETSc solvers specified by `preconditioner`.

PETSC_RICHARDSON_SOLVER	Richardson
PETSC_CHEBYCHEV_SOLVER	Chebyshev
PETSC_CG_SOLVER	conjugate gradients
PETSC_BICG_SOLVER	BiConjugate Gradients
PETSC_GMRES_SOLVER	generalized minimal residual
PETSC_BCGS_SOLVER	biconjugate gradients stabilized
PETSC_CGS_SOLVER	conjugate gradient squared
PETSC_TCQMR_SOLVER	transpose-free quasi-minimal residual
PETSC_TFQMR_SOLVER	transpose-free quasi-minimal residual
PETSC_CR_SOLVER	conjugate residual
PETSC_LSQR_SOLVER	least squares

Table 3.4: Available values of **solver** for the PETSc solvers.

NO_PRECONDITION	no preconditioning
MG_PRECONDITION	one hierarchical basis multigrid V-cycle
FUDOP_DD_PRECONDITION	an experimental domain decomposition
COARSE_GRID_PRECONDITION	precondition with solution on a coarse grid
PETSC_JACOBI_PRECONDITION	Jacobi
PETSC_BJACOBI_PRECONDITION	block Jacobi
PETSC_SOR_PRECONDITION	SOR and SSOR
PETSC_EISENSTAT_PRECONDITION	SOR with Eisenstat trick
PETSC_ICC_PRECONDITION	incomplete Cholesky
PETSC_ILU_PRECONDITION	incomplete LU
PETSC_ASM_PRECONDITION	additive Schwarz

Table 3.5: Available values of **precondition** for the PETSc solvers.

real(my_real)	petsc_richardson_damping_factor
real(my_real)	petsc_chebychev_emin
real(my_real)	petsc_chebychev_emax
integer	petsc_gmres_max_steps
real(my_real)	petsc_rtol
real(my_real)	petsc_atol
real(my_real)	petsc_dtol
integer	petsc_maxits
integer	petsc_ilu_levels
integer	petsc_icc_levels
real(my_real)	petsc_ilu_dt
real(my_real)	petsc_ilu_dtcol
integer	petsc_ilu_maxrowcount
real(my_real)	petsc_sor_omega
integer	petsc_sor_its
integer	petsc_sor_lits
logical	petsc_eisenstat_nodiagscaling
real(my_real)	petsc_eisenstat_omega
integer	petsc_asm_overlap

Table 3.6: Arguments that are passed to the PETSc solvers.

HYPRE_BOOMERAMG_SOLVER	algebraic multigrid
HYPRE_PCG_SOLVER	preconditioned conjugate gradients
HYPRE_GMRES_SOLVER	generalized minimal residual

Table 3.7: Available values of **solver** for the *hypre* solvers.

The FUDOP\_DD and COARSE\_GRID preconditioners are experimental and may be removed in a future release of PHAML. Their use is discouraged.

There are a number of arguments that are used as parameters to the PETSc methods. These are listed in Table 3.6. Refer to the PETSc User’s Manual [7] for explanations of these arguments.

If the preconditioner is not one of the ones starting with **PETSC**, then memory can be saved by not copying the matrix to the PETSc format. This is specified by **petsc\_matrix\_free=.true.**

If you use PETSc, you might have to make some changes to PHAML depending on what version of PETSc you have. See **mkmkfile.sh**, **petsc\_init.F90**, and **petsc\_interf.F90** and search for “before” to see if you need to make changes.

#### 3.3.4.4.2 *hypre*

*hypre* [13][14] is a package of parallel iterative solvers and preconditioners from

NO_PRECONDITION	no preconditioning
HYPRE_DS_PRECONDITION	diagonal scaling
HYPRE_BOOMERAMG_PRECONDITION	algebraic multigrid
HYPRE_PARASAILS_PRECONDITION	sparse approximate inverse (GMRES only)

Table 3.8: Available values of `precondition` for the *hypre* solvers.

integer	hypre_BoomerAMG_MaxLevels
integer	hypre_BoomerAMG_MaxIter
real(my_real)	hypre_BoomerAMG_Tol
real(my_real)	hypre_BoomerAMG_StrongThreshold
real(my_real)	hypre_BoomerAMG_MaxRowSum
integer	hypre_BoomerAMG_CoarsenType
integer	hypre_BoomerAMG_MeasureType
integer	hypre_BoomerAMG_CycleType
integer	hypre_BoomerAMG_NumGridSweeps(:)
integer	hypre_BoomerAMG_GridRelaxType(:)
integer	hypre_BoomerAMG_GridRelaxPoints(:,:)
real(my_real)	hypre_BoomerAMG_RelaxWeight(:)
integer	hypre_BoomerAMG_DebugFlag
real(my_real)	hypre_ParaSails_thresh
integer	hypre_ParaSails_nlevels
real(my_real)	hypre_ParaSails_filter
integer	hypre_ParaSails_sym
real(my_real)	hypre_ParaSails_loadbal
integer	hypre_ParaSails_reuse
integer	hypre_ParaSails_logging
real(my_real)	hypre_PCG_Tol
integer	hypre_PCG_MaxIter
integer	hypre_PCG_TwoNorm
integer	hypre_PCG_RelChange
integer	hypre_PCG_Logging
integer	hypre_GMRES_KDim
real(my_real)	hypre_GMRES_Tol
integer	hypre_GMRES_MaxIter
integer	hypre_GMRES_Logging

Table 3.9: Arguments that are passed to the *hypre* solvers.

Lawrence Livermore National Laboratories. The *hypr*e solvers are listed in Table 3.7 and preconditioners are listed in Table 3.8. Note that the BoomerAMG solver does not use a preconditioner, and the ParaSails preconditioner cannot be used with the PCG solver. There are also several arguments that are passed to the *hypr*e methods listed in Table 3.9. See the *hypr*e user’s guide (distributed with the software) for an explanation of these.

If you use *hypr*e, you might have to make some changes to PHAML depending on what version of *hypr*e you have. See `hypr_fix.c` (instructions are at the beginning of the file) and `mkmkfile.sh` (search for “hypr version”) to see if you need to make changes.

### 3.3.5 Load balancing

A parallel program with adaptive grid refinement must perform dynamic load balancing. When the grid is refined adaptively, some processors will perform more refinement than others, resulting in more grid elements, and hence more of the computational load, than others. Dynamic load balancing redistributes ownership of the elements to balance the load among the processors.

In PHAML, load balancing is performed by partitioning the grid into  $P$  sets where  $P$  is the number of slave processes. Each process is said to own the triangles in one of the partitions. Each vertex and edge has an associated triangle and is owned by the process that owns that triangle. PHAML uses a full domain partition (FuDoP) [26] in which each process also has additional coarse elements that cover the unowned part of the domain. Figure 3.5 illustrates an adaptively refined grid and the grid that would be seen by each of three processors, with the color indicating ownership.

One normally thinks of performing load balancing after refinement to redistribute the new grid. But it is also possible to perform predictive load balancing before refinement, which can reduce the amount of data to be redistributed. There also may be advantages to basing the balance on the number of elements, number of vertices or number of equations. PHAML provides the choice of balancing any of these entities before and/or after refinement. **The number of entities (elements, vertices or equations) that are associated with an element is used as the weight for the element in the partitioning algorithms.** For balancing after refinement, it divides up the number of entities as equally as possible. For balancing before refinement, it **estimates the number of entities that will be associated after refinement. If `refterm` is not `ONE_REF` or `ONE_REF_HALF_ERRIND`, it includes** the error indicator in the weight, so that those that are likely to be refined more times are given more weight. The selection of what to balance and when to balance is given by the arguments `prebalance` and `postbalance` which can take the value `BALANCE_NONE`, `BALANCE_ELEMENTS`, `BALANCE_VERTICES` or `BALANCE_EQUATIONS`. Usually, one of the two arguments is `BALANCE_NONE` so that either regular or predictive balancing is performed, but it is possible to perform balancing on both sides of refinement.

Grid partitioning algorithms have been the topic of extensive research, and there are several algorithms available. The method built into PHAML is the

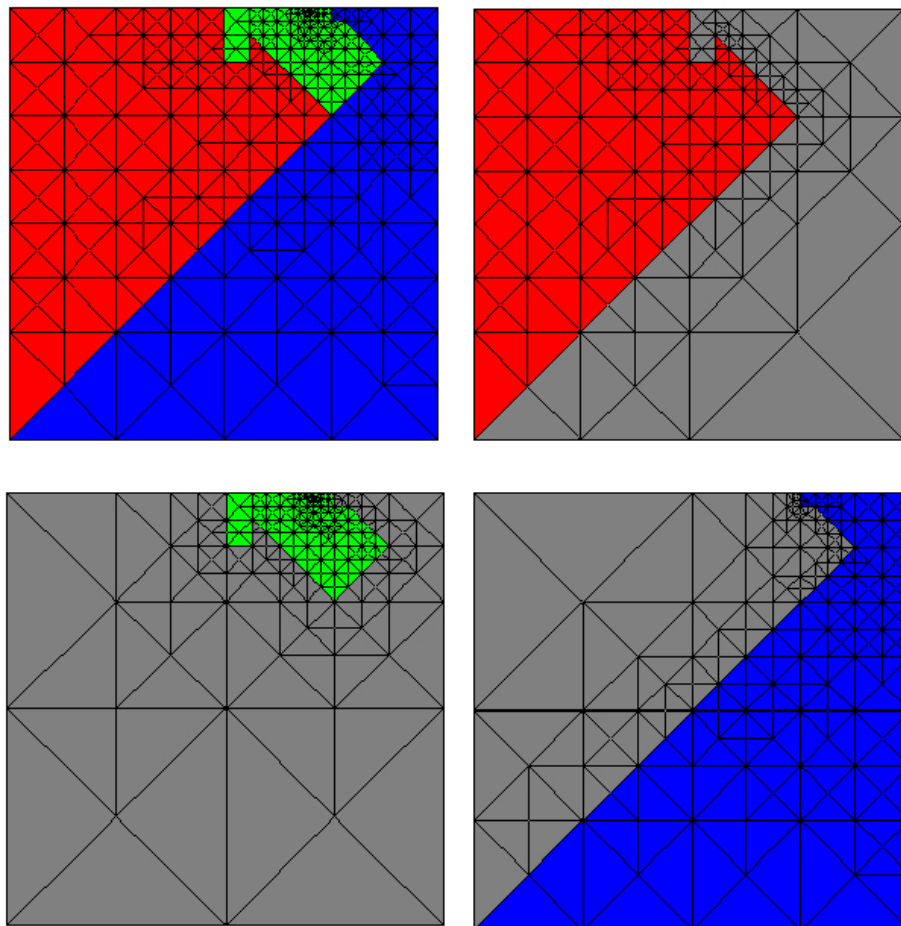


Figure 3.5: An adaptive grid partitioned for 3 processors, and the grid seen by each of the processors.



ZOLTAN_RCB	recursive coordinate bisection
ZOLTAN_OCT	RPI's Octree method
ZOLTAN_METIS	local diffusion method from ParMETIS
ZOLTAN_REFTREE	refinement tree
ZOLTAN_RIB	recursive inertial bisection
ZOLTAN_HSFC	Hilbert space filling curve
ZOLTAN_FILE	read Zoltan method and parameters from a file

Table 3.10: Available values of `partition_method` for Zoltan partitioners.

k-way refinement tree method [28]. This method uses the inherent refinement tree where the tree nodes correspond to triangles and the children of a tree node are the triangles created when a triangle is refined. Weights are attached to the leaf nodes and a tree traversal is performed to sum the weights. A second traversal is performed to partition the tree into  $P$  subtrees of equal summed weight. The children in the tree are ordered such that a tree traversal gives a space filling curve through the grid, and partitions are connected.

The choice of partitioning method is determined by the argument `partition_method`. The built in k-way refinement tree method is selected by the value `RTK`. Alternative partitioning methods are provided through the optional Zoltan dynamic load balancing library [11][12]. The values for selecting methods from Zoltan are given in Table 3.10. See the Zoltan User's Guide for descriptions of these methods, except `ZOLTAN_FILE`.

`ZOLTAN_FILE` lets you specify the method to use in Zoltan, and also to specify other Zoltan parameters, in a file. Basically, any parameter that can be set with `Zoltan_Set_Param` (see the Zoltan User's Guide) can be set by including a line containing the parameter name and value. In particular, you can specify the method with, for example, `LB_METHOD RCB`. For a full description of this file, see the `zoltanParams` web page [43]. The Zoltan parameter file is specified by the `phaml_solve_pde` argument `zoltan_param_file`, with a default value of `"zoltan.params"`.

DRUM (the Dynamic Resource Utilization Model) [15][42] supports resource-aware, large-scale scientific computing in heterogeneous and hierarchical parallel computing environments. It can improve the load balance on, for example, clusters containing different kinds of processors. Within PHAML, DRUM is accessed with the partitioning method `ZOLTAN_FILE`. The parameters for DRUM are specified in the Zoltan parameter file. See the DRUM documentation for a description of the DRUM parameters.

### 3.3.6 Termination

There are several ways to specify how long `phaml_solve_pde` should continue to work on the solution, i.e. when to terminate. This section presents them. At least one of them must be specified or else the subroutine will run forever.

Multiple termination criteria can be specified, and the routine will return when any of them are satisfied.

The argument `task` selects what task should be performed by `phaml_solve_pde`. Three of the values for `task` invoke a single pass of one part of the program, and no further termination criterion is needed. These are `BALANCE_ONLY`, `REFINE_ONLY`, and `SOLVE_ONLY`. The value `BALANCE_REFINE_SOLVE` invokes a loop over the three phases until one of the termination criteria is met. `SET_INITIAL` also loops over the three phases, but for the solve phase it interpolates the function in subroutine `iconds` (see Section 7.3.5). This is usually used for setting initial conditions for a time dependent problem, or an initial guess for a nonlinear problem (see Section 4.3).

The termination criteria are given by optional arguments and fall into two categories. The first is those that specify an upper bound on the number of something. These arguments all take an `integer` value. The arguments are `max_elem` (elements), `max_vert` (vertices), `max_eq` (equations in the linear system), and `max_refsolveloop` (number of times through the refine/solve loop). You can also specify `max_lev` (refinement levels) and `max_deg` (polynomial degree), but by default these are not termination criteria – if the refinement of an element would exceed the given value, then it is simply not performed. To use these as termination criteria, use the logical arguments `stop_on_maxlev` and `stop_on_maxdeg`.

The other category is to terminate when an error estimate is small enough. The global error estimate is computed from the local error indicator. The estimate can be an estimate of the energy norm,  $L^\infty$  norm, or  $L^2$  norm of the error. The arguments `term_energy_err`, `term_Linf_err`, and `term_L2_err` all take a `real(my_real)` value.

## 3.4 I/O

### 3.4.1 I/O files

PHAML writes printed output to two units, which can be specified through the integer arguments `output_unit` and `error_unit` of subroutine `phaml_create`. Error messages and warnings are written to `error_unit` and all other printed output is written to `output_unit`. By default, `output_unit` is 6 which most compilers provide as a pre-connected unit, often referred to as stdout. Also by default, `error_unit` is 0 which most compilers provide as a pre-connected unit, often referred to as stderr. If your compiler uses different unit numbers for pre-connected units, the correct units can be specified through these arguments. The two units can be the same if you want to have error messages and printed output intermixed. Warning messages from the master and slave processes can be suppressed by passing `print_warnings=.false.` to subroutine `phaml_solve_pde`. Error messages and warnings from graphics processes cannot be suppressed.

Usually stdout and stderr print to the terminal screen. In particular, they

usually print to the window from which the program was started. However, the processes of a parallel program might not have an associated window, and the destination of stdout and stderr is determined by the parallel library. For example, if using a master/slave model with LAM as the parallel library, output from the master process will go to the window from which it was started, but output from the slave processes will go to the window from which `lamboot` was run. Moreover, output from all the slaves is intermixed in this window. To get more control over the destination of output from the slaves you can use `spawn_form=DEBUG_SLAVE` in `phaml_create`. This will open a window for each slave process and normally the printed output goes to those windows.

If you prefer to have the printed output directed to a file, then you can open a unit and specify that unit in `output_unit` and/or `error_unit`. PHAML provides “parallel open” and “parallel close” subroutines for this purpose. Subroutine `phaml_popen` (Section 7.4.11) opens a unit with a different file for the master and each of the slaves. If you specify the name of the file to be `root.suffix` then the actual filenames are `rootXXXX.suffix` for slave number XXXX, where the number of digits in XXXX is the minimum needed for the number of slaves. The master process is number 0. Subroutine `phaml_pclose` (Section 7.4.10) closes the unit.

Thus the usual process for directing printed output (or error) to files is: 1) call `phaml_create` with some non-pre-connected unit number in `output_unit`, 2) call `phaml_popen` with that unit number and some root file name, 3) call `phaml_solve_pde`, 4) call `phaml_pclose` with that unit number. Note that if there are any error messages printed by `phaml_create`, they cannot be written to this file because the unit has not yet been opened. These messages are written to unit 0 by default. You can specify a different unit through the argument `output_now`, but it must be a pre-connected unit.

### 3.4.2 Printed I/O

PHAML provides for the printing of various quantities at various times during the execution of `phaml_solve_pde`. By default, the only printed output is a header containing a summary of the input variables to `phaml_solve_pde` and a trailer containing termination information, printed by the master process. Other output is controlled through a series of “when” and “who” arguments. The “when” arguments indicate how often to print something. They take the values `NEVER` (don’t print that quantity), `FINAL` (print it just before returning), `PHASES` (print it once in each refine/solve loop), and `FREQUENTLY` (print more often, usually used for debugging). The “who” arguments indicate which processes should print something. They take the values `NO_ONE` (don’t print that quantity), `MASTER` (the master prints that quantity for the entire grid), `SLAVES` (each slave prints that quantity for the part of the grid that it owns), `EVERYONE` (both the master and the slaves print), and `MASTER.ALL` (the master prints the quantity for each of the slaves).

`print_grid_when` and `print_grid_who` provide for printing a summary of the grid, such as the number of vertices, number of elements, etc. `print_linsys_when`

and `print_linsys_who` provide for printing a summary of the linear system, such as the number of equations, etc. `print_solver_when` and `print_solve_who` provide for printing a summary of the solver, such as the number of iterations used, the norm of the residual, etc.

`print_error_when` and `print_error_who` provide for printing certain norms of the error (if the true solution is provided) and error estimates. `print_error_what` indicates what global norm(s) to print, with the available norms being the energy norm,  $L^\infty$  norm, and  $L^2$  norm. The values for this argument are `NEVER`, `ENERGY_ERR`, `LINF_ERR`, `L2_ERR`, `ENERGY_LINF_ERR`, `ENERGY_L2_ERR`, `LINF_L2_ERR`, and `ENERGY_LINF_L2_ERR`. The energy and  $L^2$  errors are approximated using a sixth order quadrature rule for the integrals. The  $L^\infty$  norm is approximated using the quadrature points of a sixth order quadrature rule.

`print_errest_what` controls which global error estimates are printed. It takes values that are similar to `print_error_what` except that “ERR” is replaced by `ERREST`.

The norms of the error and error estimates can be either absolute error (the default) or relative error. This is controlled by `errtype` which takes the values `ABSOLUTE_ERROR` and `RELATIVE_ERROR`. If the relative error is selected, then the printed norm of the error is divided by the norm of the true solution, and the printed error estimate is divided by the norm of the computed solution.

`print_error_when` also provides for monitoring the convergence rate of the iterative linear system solver, if the solver is the built-in hierarchical basis multi-grid, conjugate gradients, or GMRES method or one of the solvers from PETSc. If it has the value `FREQUENTLY` then the  $\ell^2$  norm of the residual is printed after each iteration. For GMRES, it is printed at the restarts. It also accepts the value `TOO_MUCH` which additionally sets the solution to 0.0 before starting the iterations, to avoid convergence in 1 iteration.

`print_time_when` and `print_time_who` provide for measuring execution time of the program. The execution time of each section of the program (refinement, reconciliation, load balancing, matrix assembly, linear system solution, communication) is printed. Each time the time is printed it prints the time for the most recent pass through the refine/solve loop and the total time so far. `clocks` determines how the time is measured. It can be `CLOCK_C` for the cpu clock, `CLOCK_W` for the wall clock, or `CLOCK_CW` for both.

`print_header_who` and `print_trailer_who` specify which processes should print the header and trailer, respectively.

### 3.4.3 Pausing

In order to examine printed or graphical output while a PHAML program is running, it is often useful to have the program pause until you indicate that it may continue. There are several “pause” arguments to `phaml.solve_pde` that provide for this. They are `logical` arguments. When the program pauses, the master prints “`press return to continue`” to its printed output unit. It then reads from standard input, which is usually associated with the window from which the master program was run. The pause arguments are `pause_at_start`

(pause upon entering `phaml_solve_pde`), `pause_at_end` (pause before leaving `phaml_solve_pde`), `pause_after_phases` (pause at the end of each refine/solve loop), and `pause_after_draw` (pause each time the graphics is updated).

## 3.5 Graphics

### 3.5.1 Overview

Graphics, or visualization, in PHAML is provided using OpenGL, which is a platform-independent specification of a graphics application programming interface, and GLUT, which supplies window, keyboard and mouse usage for OpenGL. There are libraries for both of these specifications available for nearly every computer. See Section 2.1.5 for URLs for more information on OpenGL and GLUT and how to obtain the software.

PHAML's graphics are interactive and menu driven. There may be graphics windows associated with the master process and/or with each of the slave processes. The master's graphics shows the grid and solution as a whole entity, whereas the slaves each show the grid and solution as they know it. The selection of which processes will have graphics is made with the argument `draw_grid_who` to `phaml_create`, which can have the value `MASTER`, `SLAVES`, `EVERYONE`, or `NO_ONE`. There is no option to have only a subset of the slaves do graphics.

The graphics servers, which are separate processes in the parallel program, receive messages from the associated master or slave whenever the grid or solution changes. For a program under a sequential compilation of PHAML, the messages are passed by writing files to `/tmp`. For a master/slave program, the graphics servers are spawned automatically during the execution of subroutine `phaml_create`. For an SPMD program, the graphics servers should be included in the number of processes launched (see Section 2.4). For a sequential program, you must start the graphics server, `phaml_graphics`, from the command line. If a sequential program terminates abnormally, you must terminate `phaml_graphics` by hand, and should check `/tmp` for leftover files `phaml_message` and `phaml_lock`, and remove them if they exist.

All interaction with the graphics is performed with the mouse and arrow keys. The left and middle mouse buttons and arrow keys can be assigned various operations to change the view, as discussed in Section 3.5.3. By default, the left button rotates, the middle button zooms and the arrow keys pan. The right button brings up a menu of actions you can take to modify the graphics. The main menu is shown in Table 3.11. Menu items with a right arrow ( $\Rightarrow$ ) bring up submenus.

The next section will show some of the visualizations that are available. The subsequent sections will discuss how to manipulate the graphics through the submenus.

view modifier	⇒
element edge color	⇒
element interior color	⇒
function	⇒
contour plots	⇒
preprocess function	⇒
subelement resolution	⇒
color scheme	⇒
toggle lights	⇒
element label	⇒
edge label	⇒
vertex label	⇒
associated element	⇒
eigenfunction to use	⇒
component to use	⇒
component scale	⇒
space filling curve	⇒
grid offset	⇒
crop (debug window)	
toggle axes	
toggle color key	
write postscript	⇒

Table 3.11: The main graphics menu.

### 3.5.2 Example visualizations

The PHAML graphics server has many options in the graphics menu. Primarily the graphics options are for different displays of the grid properties, functions over the grid, and the partitioning of the grid over the slaves. The functions that can be displayed include not only the computed solution, but also the true solution and error (if the true solution is given), and error estimates. In this section we present some examples of PHAML's graphics capability with very brief descriptions. The subsequent sections will discuss how to use the submenus to create these and other graphical displays.

Figure 3.6 shows several of the different ways you can display a function. Parts (a)–(f) show surface plots of the solution, which comes from using the menu to select `function` to be `solution`. Part (a) shows the surface with the triangle interiors colored by the solution value using the rainbow color scheme where blue represents small values and red represents large values. In part (b) the same function is shown with the grid added to the surface, drawn in black. Part (c) displays the grid in black on the surface, but selects `element interior color` to be `transparent`. Part (d) is the same but with hidden lines removed, which is achieved by selecting `element interior color` to be `white` so that the triangle faces hide the grid lines behind them. Part (e) also has the grid on the surface and hidden lines removed, but with the grid lines colored by solution value. Part (f) is a surface plot using the striped color scheme and placing contour lines on the surface. Parts (g)–(l) show 2D displays of the solution obtained by selecting `function` to be `no function` and viewing from straight above. Part (g) uses the striped color scheme with contour lines drawn in the  $x$ - $y$  plane. Part (h) shows the solution using the rainbow color scheme. Part (i) adds the grid to the previous image, and part (j) colors the triangle interiors transparent and the grid lines with the solution value. Part (k) is simply a contour plot with no grid lines and the triangle interiors transparent. Finally, part (l) adds the coloring of the triangle interiors by the solution value using the gray scale color scheme.

Figure 3.7 shows some additional visualizations of the grid. Part (a) simply shows the grid drawn in black. Part (b) shows the regions in which the grid is coarse or fine by coloring the interiors of the triangles by the triangle size, using blue for small triangles and red for large. In part (c) the triangle interiors are colored by the polynomial degree of the triangle, for visualizing the effect of  $p$ - and  $hp$ -adaptive refinement. And in part (d) we use `function`  $\Rightarrow$  `levels` to create a hierarchy of grids based on refinement level.

Figure 3.8 shows a few of the ways you can visualize how the grid has been partitioned for distribution over the slaves during the load balancing step. In part (a) the interior of the triangles is colored by owner and the grid lines are colored black. Part (b) colors the grid lines by owner and leaves the interior transparent. In part (c) the interiors are again colored by owner, but only the triangle edges that are on partition boundaries are drawn. Finally, part (d) shows an exploded grid where the partitions are physically separated.

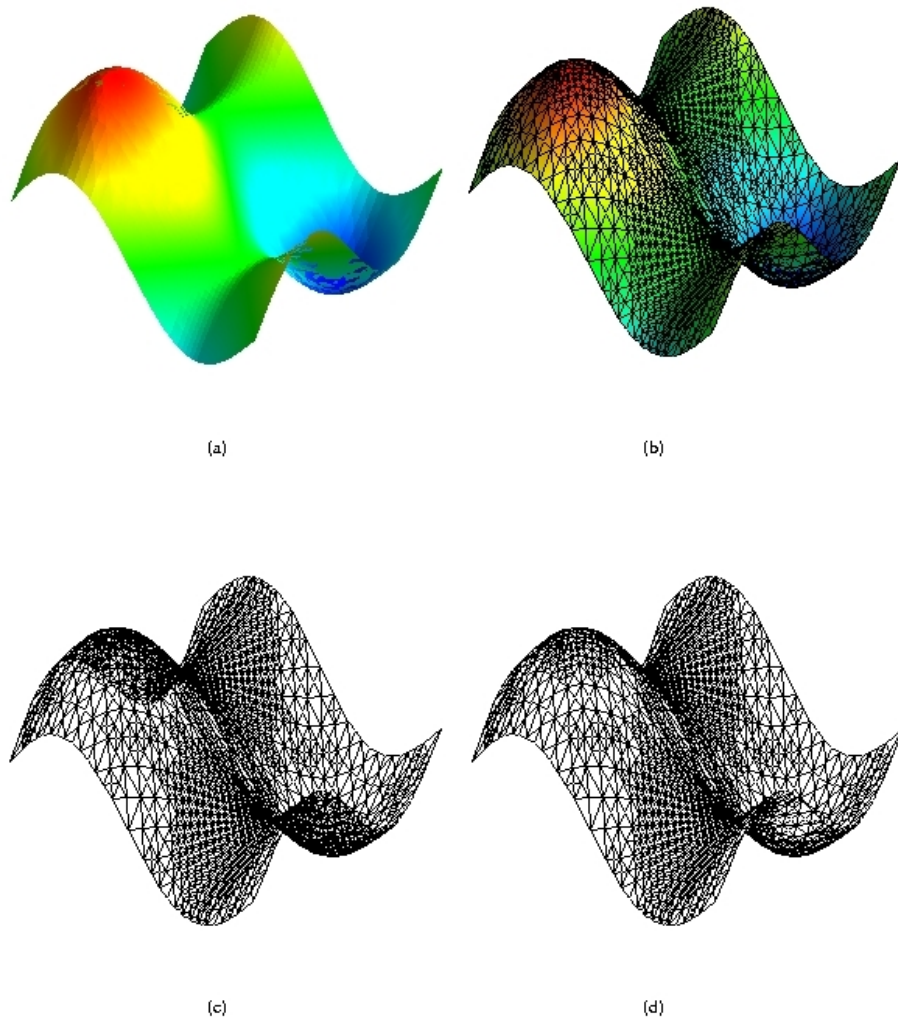
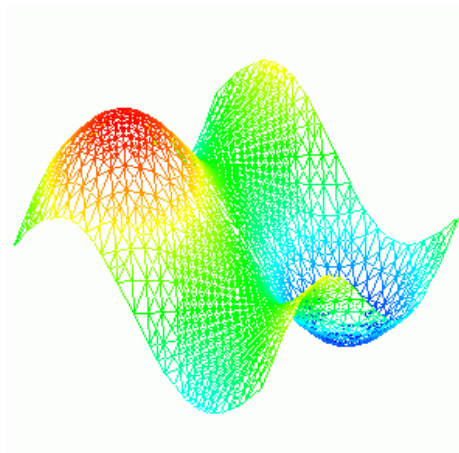
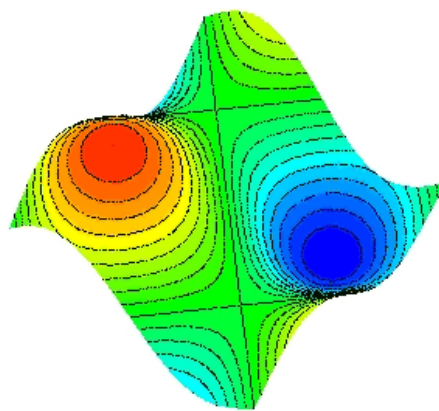


Figure 3.6: Visualizations of the solution.

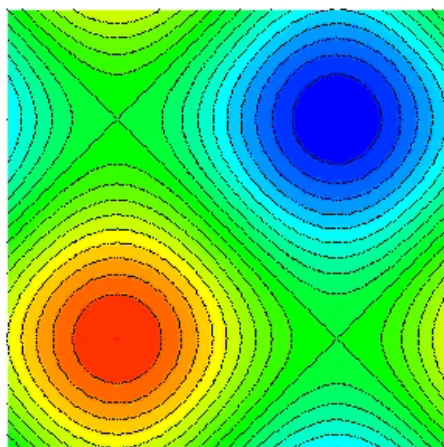




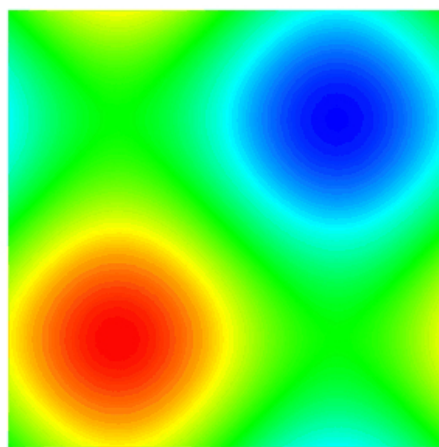
(e)



(f)



(g)



(h)

Figure 3.6: (continued) Visualizations of the solution.

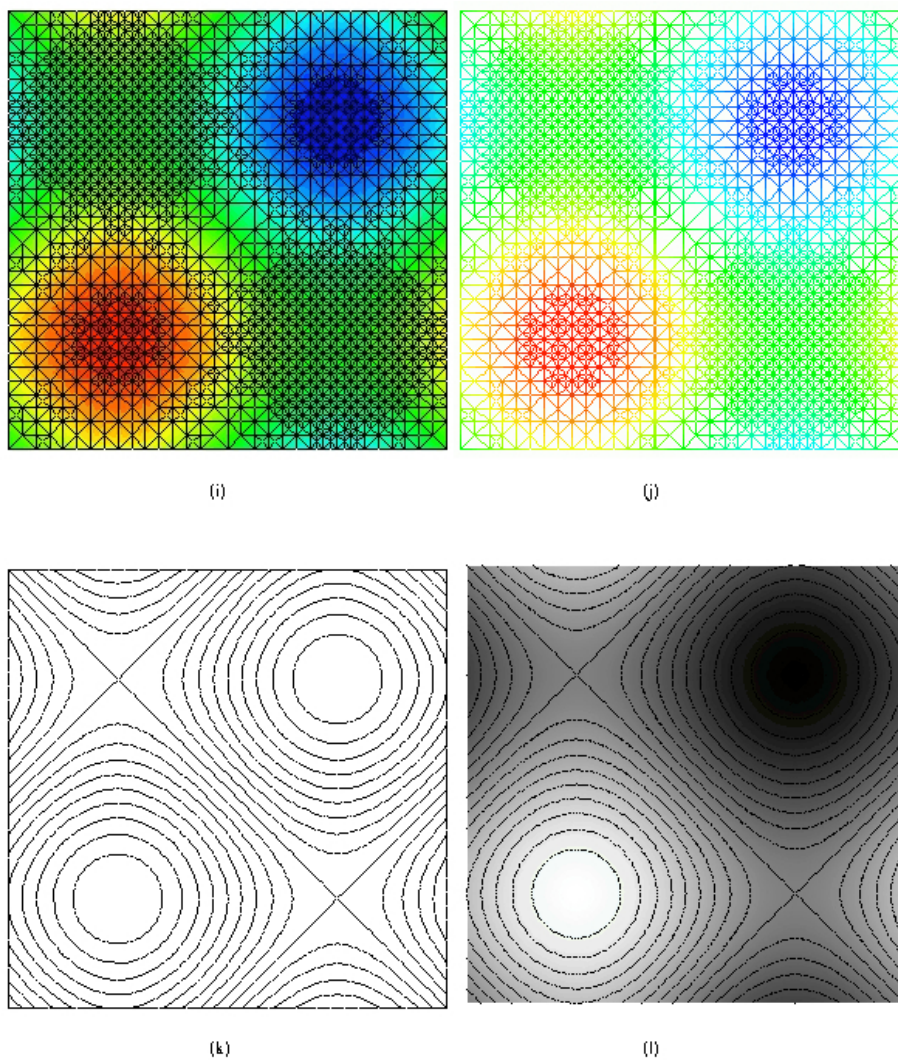


Figure 3.6: (continued) Visualizations of the solution.

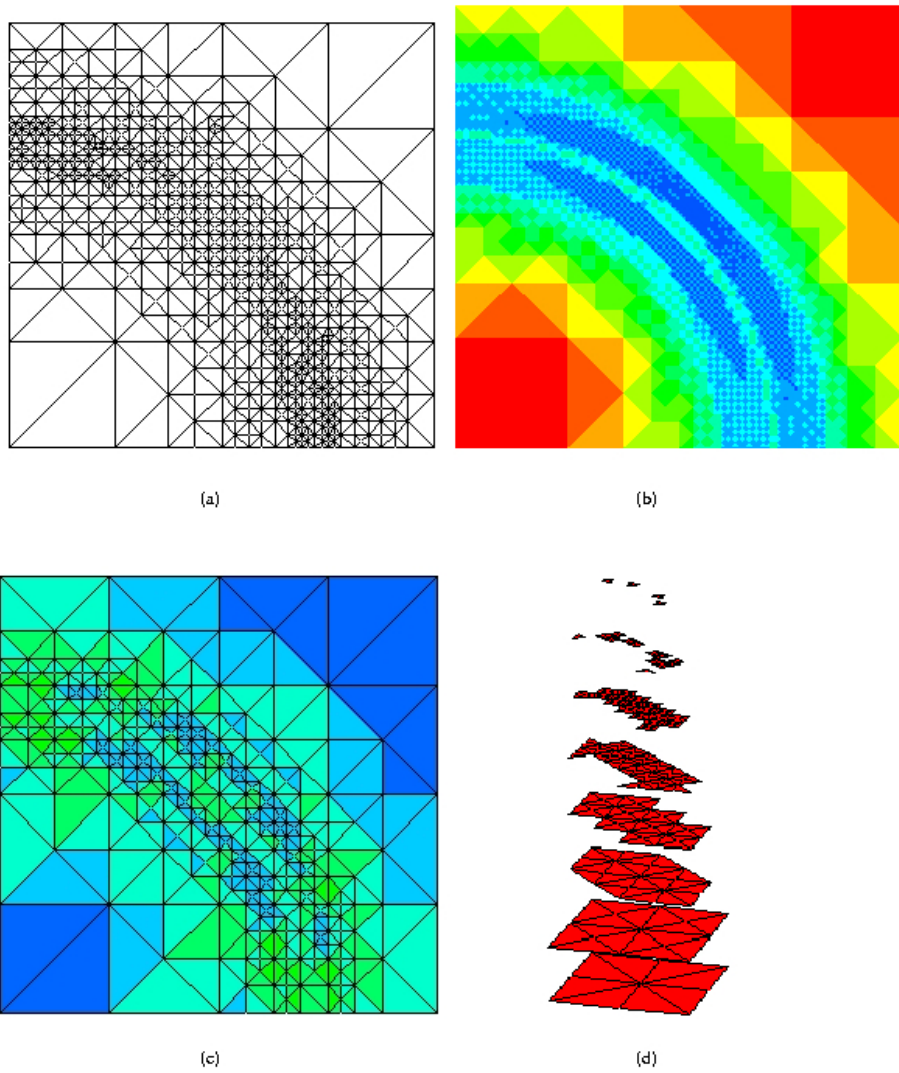


Figure 3.7: Visualizations of the grid.

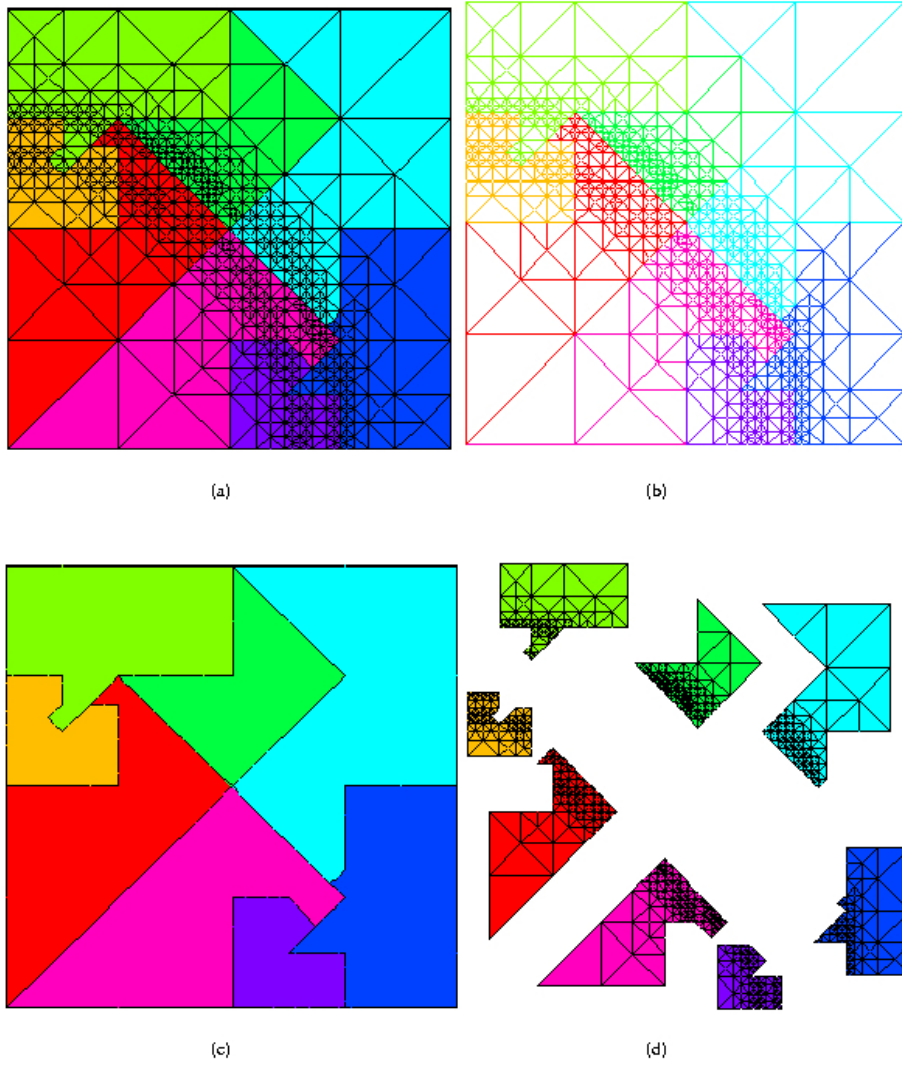


Figure 3.8: Visualizations of the partitions of the grid for load balancing.

left mouse button	⇒
middle mouse button	⇒
arrow keys	⇒
reset to initial view	
view from above	
view from above origin	
quit	

Table 3.12: The view modifier submenu.

### 3.5.3 View modifier

In this section, we discuss the view modifier. The **view modifier** submenu is shown in Table 3.12.

The first three items set the operation performed by the left mouse button, middle mouse button and arrow keys, respectively. The selected operation remains in effect until you select another one. The submenus below each of these are all the same. They contain **rotate**, **zoom**, **pan**, **scale x** (stretch or shrink along the  $x$  axis), **scale y**, **scale z**, **explode** (as in Figure 3.8(d)), and **move light** (see Section 3.5.6).

The next three entries are quick changes to a specific view of the grid. **reset to initial view** resets the rotation, zoom and pan settings to the original settings. **view from above** sets a view that looks straight down and at the center of the grid, with the (xmin,ymin) corner of the domain at the lower left corner of the image. This is normally used with **function** ⇒ **no function** for visualizations of the grid and contour plots. **view from above origin** looks at the point (0,0,0) from straight above. This is useful if you want to zoom in on the origin without having to pan repeatedly.

The final entry, **quit**, should not normally be used. If you terminate the graphics program using this menu item, it may or may not kill the master and/or slaves. The graphics program will terminate properly when **phaml.destroy** is called. Using the **quit** entry in the menu is only useful if the master or associated slave terminated early, and even then there might be a better way to terminate the graphics server (for example, with LAM the command **lamclean** will terminate all the processes).

### 3.5.4 Colors

By default, PHAML uses a rainbow color scheme. Colorization of items that have a continuous numerical value, like the solution, comes from a continuous spectrum with blue corresponding to small values and red to large values. Normally the spectrum is scaled to correspond to the minimum and maximum values of the item being drawn. Colorization of items that have a finite set of  $N$  discrete values, like the owner, comes from an equally spaced set of  $N$  colors in the spectrum from blue to red with blue assigned to the first item.

no lines
black
edge owner
vertex owner
computed solution
true solution
error
size
degree
partition boundary black

Table 3.13: The element edge color submenu.

There are four other color schemes available. The color scheme is selected in the submenu under **color scheme** which has the selections **rainbow**, **double rainbow**, **gray scale**, **striped**, and **stepped sequential**. The double rainbow passes through the colors twice, with the second time a little more pastel. This is useful when discrete colors need more separation, such as when coloring by owner and there are a lot of processors, or when coloring by element degree with an *hp*-adaptive grid. The gray scale scheme uses a continuous gray scale from black to white with black corresponding to small values and white to large values. The striped scheme is like the rainbow scheme except the colors are from a discrete set of equally space colors from the spectrum from blue to red. The number of colors in the scheme is one less than the number of contour lines used for contour plots, so that contour lines will fall on the boundary between colors. The stepped sequential scheme uses several steps of saturation/value within each of several hues ranging from red to violet. The number of steps and number of hues can be changed by use of a submenu. By default, **there are 4 steps of 6 hues for a total of 24 colors.** they are both 4 for a total of 16 colors. This scheme is particularly useful for discrete values such as owner and degree.

With a color scheme selected, you can choose how to color the grid lines (i.e. element edges) and triangles (i.e. element interiors) to get different information about the grid, partition, solution, error, etc. Table 3.13 shows the element edge color submenu for selecting how to color the element edges. **no lines** means don't draw the grid. **black** draws the edges black for a simple drawing of the grid. **edge owner** selects the color to indicate which slave process owns each edge. **vertex owner** selects the color to indicate the owner of the vertices at the ends of each edge. If the owners of the two endpoints are different, the color blends from one to the other along the length of the edge. **computed solution** uses the color corresponding to the value of the computed solution at the endpoints, and blends the colors along the length of the edge. **true solution** is similar but uses the true solution if it is given in function **trues**. Likewise, **error** uses the computed solution minus the true solution, if it is available. **size** colors the edges according to their length, using red for the largest edges and blue for

the smallest. This is useful for distinguishing element sizes when the grid is so fine that drawing the edges black would result in large black areas. **degree** colors the edges by the polynomial degree of the approximation space along that edge. Finally, **partition boundary black** draws only the edges that fall between two triangles with different owners, and edges on the domain boundary. This results in the partition boundary being drawn black.

The submenu for element interior color is similar to the one for element edge color, with just a few differences. The **no lines** entry is replaced by **transparent**, but has the same meaning of don't draw the element. **black** is replaced by **white**. Coloring an element interior white, which is the same as the graphics window background, has the effect of blocking whatever is behind the element, and can be used to draw hidden line plots. There is only one **owner** entry, used to indicate which slave owns each element. **computed solution**, **true solution**, **error**, **size**, and **degree** all have the same meaning as with edge color. There is no partition boundary entry. It has an additional entry for coloring each element by its error indicator.

### 3.5.5 Functions

The function submenu determines what function to draw as surface plots and contour plots. This menu has six entries. The first is **no function**, which simply draws the elements in the  $x$ - $y$  plane. This is usually used in conjunction with viewing from above. The next three entries are **computed solution**, **true solution**, and **error**. They draw the corresponding function. The fifth entry is **levels**. This draws the elements, including parent elements, in  $L$  discrete planes parallel to the  $x$ - $y$  plane, where  $L$  is the number of refinement levels in the grid (see Figure 3.7(d)). Each level contains the elements of that refinement level. You may find it useful to use **scale z** from the view modifier (Section 3.5.3) with this. The final entry is **error indicator**. This is a piecewise constant function with each element drawn at the height corresponding to its error indicator.

The functions (except levels) can be preprocessed in several ways with the submenu under **preprocess function**. The menu entries indicate what would be applied to a function  $f$ . They are **none** (no preprocessing, the default), **-f** (negate), **abs(f)** (absolute value), **f\*\*2** (square), and **log(abs(f))** (logarithm of the absolute value).

### 3.5.6 Lights

When the function is **no function**, **levels** or an error estimate, or the striped color scheme is used, the image is rendered in flat light. Otherwise, it is rendered with a light source resulting in a 3D appearance with shadows. There are five lights available in the PHAML graphics. Four of them are fixed in position: one to the right, one to the left, one above and one below. The fifth light is movable. The default is that only the light to the right is turned on. Lights can be turned on and off with the submenu under **toggle lights**. The movable light

can be moved with the mouse or arrow keys by selecting `move light` under the `view modifier` submenu.

### 3.5.7 Contour plots

In addition to being drawn as a surface or being represented by color, a function can be drawn as a contour plot. Contour plots are controlled by the submenu under `contour plots`. This submenu has four entries for selecting what function to plot, and three submenus for changing the properties of the contour plot.

The first four entries in the contour plot submenu select the function to plot. As usual, they are `no contour plot`, `computed solution`, `true solution`, and `error`. By default, no contour plot is drawn.

The next entry is `set number of uniform lines`. Initially PHAML uses 21 equally spaced contour values between the minimum and maximum values of the function. This submenu allows you to change the number of values while keeping them equally spaced. There are six entries that allow you to make the change directly from the menu: `increment by 1`, `decrement by 1`, `increment by 10`, `decrement by 10`, `double`, and `cut in half`. The final entry, `enter number in debug window`, lets you type in the number of contour lines you want. However, it prompts for this number from standard input, so the graphics process must have an associated window. This is achieved by using `spawn_form = DEBUG_GRAPHICS` or `spawn_form = DEBUG_BOTH` in the call to `phaml_create` (see Section 7.4.4).

You can also space the contour lines nonuniformly by using `set nonuniform lines`. This also requires a debug window. You will be prompted for the number of contour lines, and then to enter a comma separated list of the values for the contour lines.

The final menu entry gives two choices for the placement of the contours. They can either be placed on the  $x$ - $y$  plane, or be elevated by the function value, i.e., placed on the surface.

### 3.5.8 Multiple solutions

If you solve a coupled system of equations (multicomponent solution) or solve for more than one eigenvalue of an eigenvalue problem, then there is more than one solution to draw. Two submenus let you select which function to display. The default is to display the first one.

`eigenfunction to use` provides the selection of which eigenfunction to display. It has an entry of the form `eigenfunction 1`, `eigenfunction 2`, etc., up to the number of eigenvalues computed. If there are more than 9 eigenvalues, the last entry is `more  $\Rightarrow$` . This brings up a submenu containing `10's  $\Rightarrow$` , `20's  $\Rightarrow$` , etc. Under these submenus you will find the eigenfunctions with larger indices. PHAML sorts the eigenvalues from smallest to largest, and orders the corresponding eigenfunctions the same way.

`component to use` provides the selection of which component of a multicomponent solution to display. It contains entries of the form `component 1`,



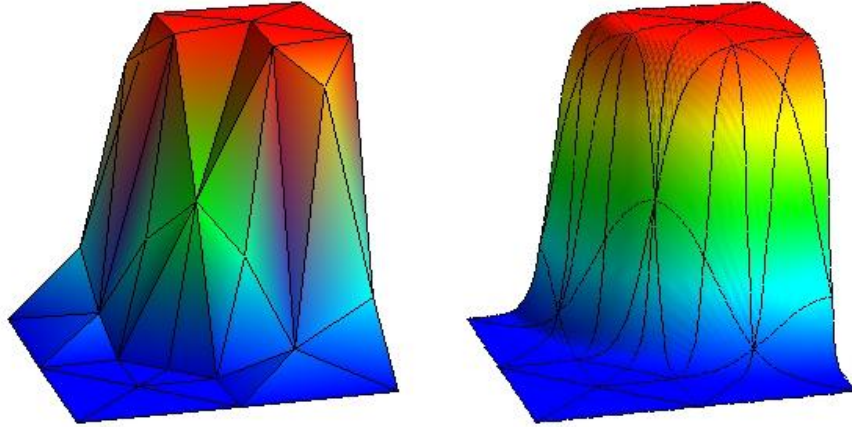


Figure 3.9: Improvement of a surface visualization by using subelement resolution.

component 2, etc. Like the eigenfunction selection, there are submenus for the higher indices if there are more than 9 components. The menu also contains two entries for displaying a composite of the components. If the components of the solution are  $u_1, u_2, \dots$ , then **L1 sum** will display  $|u_1| + |u_2| + \dots$ , and **L2 sum** will display  $u_1^2 + u_2^2 + \dots$ .

**component scale** provides the selection of how to scale the components when the function is computed solution, true solution or error. Normally the function is scaled by its maximum absolute value. For multicomponent solutions, you can choose to have each component scaled by its own maximum absolute value by selecting **individual**, or you can choose to have all components scaled the same using the maximum absolute value over all components by selecting **all the same**.

### 3.5.9 Miscellaneous features

This section addresses some menu items that are not big enough to warrant a section of their own.

The **subelement resolution** submenu is useful when high order elements are used. By default, a triangle is drawn as a piece of a plane defined by the three vertices of the triangle. Color blending, contour lines, etc., are also limited by this definition. This applies both to 3D surfaces and 2D drawings in the  $x$ - $y$  plane. This is fine for piecewise linear elements since the solution is a plane over each triangle. But with higher order element, any detail on the subelement level is lost. This menu entry defines how much subelement resolution to use. The drawing is still done by drawing pieces of a plane, i.e. triangles, but multiple triangles are drawn within a grid element. With subelement level 0, the element is drawn as a single triangle (the default). With subelement level 1, the element

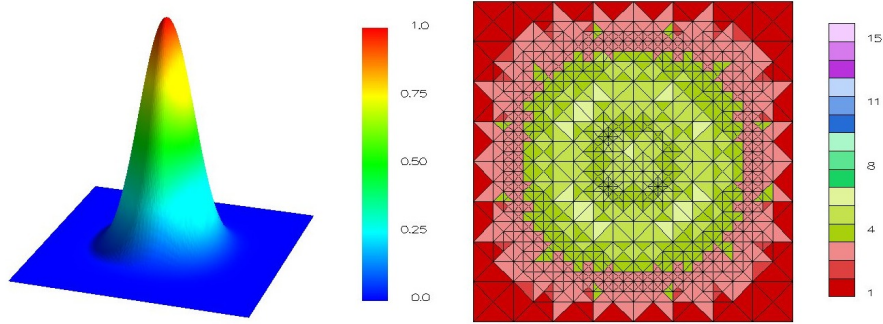


Figure 3.10: Examples of the color key.

is drawn as 4 triangles, formed by connecting the midpoints of the element edges. At subelement level 2, each of those 4 triangles are drawn as 4 triangles by connecting the midpoints of their sides, resulting in 16 triangles in a grid element. In general, subelement level  $\ell$  results in drawing  $4^\ell$  triangles in each element. The effect of using a high level of subelement resolution can be seen in Figure 3.9.

`subelement resolution` has entries for 0, 1, 2 and 3 levels of subelement resolution. Since the number of triangles drawn grows exponentially with the number of subelement levels, using higher values of subelement resolution can be very slow. However, if needed, values larger than 3 are obtainable with the submenu entries `increase`, which adds 1, and `decrease`, which subtracts 1.

`x`, `y` and `z` axes can be added and removed from the plot with the `toggle axes` entry. The axes are fairly primitive with tic marks and values only at the ends and midpoint.

A color key can be added and removed with the `toggle color key` entry. The key gives the correspondence between the color of the interior of the elements and numeric values. Figure 3.10 illustrates two uses of the color key: a continuous color key for the function value using the rainbow color scheme, and a discrete color key showing the element degree using the stepped sequential color scheme.

`crop (debug window)` provides a means of restricting how much of the domain is used in the display. Since it requires input from standard input, the graphics process must have an associated window. This is achieved by using `spawn_form = DEBUG_GRAPHICS` or `spawn_form = DEBUG_BOTH` in the call to `phaml.create` (see Section 7.4.4). When this menu entry is selected, it will prompt for the crop region to be entered as `xmin`, `xmax`, `ymin`, `ymax`.

The `grid offset` submenu helps to solve a potential problem with the rendering of the graphics. The elements are drawn as triangles and the grid lines are drawn as lines separately, but they occupy the same space. This can cause a problem in determining which should show when the image is rendered. Even if they

are separated very slightly, there can be a problem because of machine roundoff error. This submenu allows you to change how far the grid lines are offset from the triangles. If you find that grid lines are disappearing, you should increase the offset. If you find that the grid lines appear to be separated from the surface, you should decrease the offset. The submenu entries let you increase or decrease by 1 or 10 at a time.

### 3.5.10 Development aids

Some of the graphics options were created as aids in the development and debugging of PHAML. These are probably not much use to the end user. They are the **element label**, **edge label**, and **vertex label**, which label each entity with its index in PHAML's data structure, **associated element** which displays the edge-element and vertex-element correspondence used to determine the owner of edges and vertices, and **space filling curve** which displays the space filling curve associated with the refinement tree partition method.

### 3.5.11 Postscript

The **write postscript** submenu lets you save the current visualization to an encapsulated postscript file. It writes vector graphics in the postscript language. This means the saved image is high quality and scalable, but it creates very large files. If you want smaller files you can use a screen capturing program and save the image in a raster graphics format like JPEG.

**write postscript** has two options in the submenu: **unsorted** and **sorted**. The **sorted** option sorts the entities of the image by distance from the viewer before writing the postscript file. There isn't really any reason to use the **unsorted** option, and it will probably be removed in the future.

When **write postscript** is selected, it creates a file called **renderX.eps** where X is 0 for the master's graphics processor or the slave number for the slaves' graphics processors (note there cannot be more than 9 slaves for this to work properly). The location of this file is compiler dependent, but it usually ends up in the directory where the master program was started or in the user's home directory. You may have to modify this file slightly in two ways. First, rename it to something more meaningful! Second, some printers need to have **showpage** added as the last command in the file, but that confuses some viewers. If you try to print the file and nothing happens, add this line.

## 3.6 Post-solution utilities

### 3.6.1 Store and Restore

There are times when it may be useful to save a PHAML solution and use it in some subsequent program. For example, one might solve a problem on a batch-only system and save the solution, and then later run a program on an interactive system that reads the solution and displays it with PHAML's

graphics capability. PHAML provides a routine that writes the entire contents of a `phaml_solution_type` variable to a set of files (one file for the master and one for each slave), and a routine that reads those files into a `phaml_solution_type` variable. The files should be connected to a unit number using `phaml_popen` and `phaml_pclose` as described in section 3.4.1.

`phaml_store` takes two arguments: a `phaml_solution_type` variable and an integer specifying the unit to write to. `phaml_restore` also takes a `phaml_solution_type` variable (which should first be created with `phaml_create`, but otherwise be empty) and a unit to read from. It also takes two optional logical arguments. `do_draw_grid` indicates whether or not you want to invoke graphical output immediately after reading the file, and `pause` indicates whether or not to pause after the graphics. A program that reads files with `phaml_restore` must have the same number of slaves as the program that created the files with `phaml_store`.

### 3.6.2 Store Matrix

One might wish to store the linear system that represents the discretized PDE in a file for processing external to PHAML, for example solving the linear system with Matlab. Subroutine `phaml_store_matrix` does this. This routine stores the discretized PDE in `phaml_solution` so that it can be processed by an external program. In addition to a `phaml_solution_type` variable, it takes four optional arguments, `stiffness_unit`, `rhs_unit`, `mass_unit`, and `inc_quad_order`. `inc_quad_order` increases the order of the quadrature rule used for computing the matrix and right hand side entries, the same as in `phaml_solve_pde`.

For an elliptic boundary value problem, the discrete problem is a linear system of equations,  $Ax = b$ . `phaml_store_matrix` can store the stiffness matrix,  $A$ , and/or the right hand side,  $b$ . The presence of the optional arguments `stiffness_unit` and `rhs_unit` determines whether or not the stiffness matrix and right hand side will be stored, respectively. Either or both can be present. `mass_unit` should not be present.

For an elliptic eigenvalue problem, the discrete problem is a generalized eigenvalue problem  $Ax = \lambda Mx$ . `phaml_store_matrix` can store the stiffness matrix,  $A$ , and/or the mass matrix,  $M$ . The presence of the optional arguments `stiffness_unit` and `mass_unit` determines whether or not the stiffness matrix and mass matrix will be stored, respectively. Either or both can be present. `rhs_unit` should not be present.

When present, `stiffness_unit`, `rhs_unit` and `mass_unit` should contain the I/O unit of a file. It is the responsibility of the calling program to open the unit for formatted, sequential writing before calling `phaml_store_matrix` and closing it after. This is done with the usual Fortran `open` and `close` statements, not the PHAML `phaml_popen` and `phaml_pclose` statements. If more than one is present, they must not be the same file.

The matrices are written in the Matrix Market exchange format. See <http://math.nist.gov/MatrixMarket/> for a description of the format and routines for reading the format in Fortran, C and Matlab. They are written in the real,

general, coordinate format. The right hand side vector is written as an  $N \times 1$  matrix in the real, general, coordinate Matrix Market format.

### 3.6.3 Query

PHAML provides for the recovery of many quantities of interest through subroutine `phaml_query`, which can be called after returning from `phaml_solve_pde`. This includes information about the grid, such as number of elements, number of vertices, minimum and maximum polynomial degree, etc. Most of them can be for the whole grid or for the grid that each slave has. A second category includes error estimates in various norms, and, if the true solution is provided, the norms of the error and true solution. The norms of the error are absolute error; you can get the relative error by dividing by the norm of the solution. A third category provides interesting quantities related to eigenvalue problems.

For a complete list of the quantities that can be recovered through `phaml_query`, see Section 7.4.12.

### 3.6.4 Solution evaluation

PHAML provides for evaluating the computed solution and its derivatives at a point in the domain through subroutine `phaml_evaluate` (see Section 7.4.6). After returning from `phaml_solve_pde`, you can pass the `phaml_solution_type` variable to `phaml_evaluate` to obtain the solution and/or first and second derivatives. You can request any subset of them. You also pass two arrays, `x` and `y`, containing the points at which to evaluate the solution. The solution is returned in the arrays `u`, `ux`, `uy`, `uxx` and `uyy`, which must have the same dimension as `x` and `y`. These are all optional arguments, and which ones are present determines which ones are evaluated. If you are solving a system of equations or an eigenvalue problems, there are optional arguments to specify which component or eigenfunction you want to evaluate. If a point outside the domain is given, the solution is returned as 0.

Another way to obtain the solution for post processing is through the subroutine `phaml_get_grid_soln` (see Section 7.4.7). This routine returns the vertices of the triangles in the grid in two arrays, `x` and `y`, and the solution at the vertices in the array `u`. The vertices are not returned in any particular order relative to their placement in the domain. If you are solving a system of equations or an eigenvalue problem, there are optional arguments to specify which component or eigenfunction you want to evaluate.

There is also a subroutine to evaluate an “old” solution, which can be used for time dependent and nonlinear problems. See Section 4.3 for information on this routine.

### 3.6.5 Functionals

In this context, a functional of the solution is some form of integral of the computed solution. Subroutine **phaml\_integrate** (see Section 7.4.9) computes

$$\iint_{\Omega} k(x, y) u_i^p(x, y) u_j^q(x, y) dx dy.$$

$u_i$  and  $u_j$  are two components of the solution of a system of equations, or two eigenfunctions of an eigenvalue problem, or  $u_i$  is a solution and  $u_j$  is 1.  $p$  and  $q$  are integer powers.  $k$  is a kernel function defined by the user in function **phaml\_integral\_kernel** (Section 7.3.7). In addition to a point at which to evaluate the kernel, the function takes an integer, **kernel**, which allows for the definition of several kernel functions in the same subroutine.

## Chapter 4

# Problem Extensions

### 4.1 Eigenvalue Problems

In addition to elliptic boundary value problems, PHAML can solve elliptic eigenvalue problems of the form

$$-\frac{\partial}{\partial x} \left( c_{xx} \frac{\partial u}{\partial x} \right) - \frac{\partial}{\partial x} \left( c_{xy} \frac{\partial u}{\partial y} \right) - \frac{\partial}{\partial y} \left( c_{yx} \frac{\partial u}{\partial x} \right) + c_x \frac{\partial u}{\partial x} + c_y \frac{\partial u}{\partial y} + c_u u = \lambda f u \text{ in } \Omega \quad (4.1)$$

$$u = 0 \text{ on } \partial\Omega_D \quad (4.2)$$

$$\left( c_{xx} \frac{\partial u}{\partial x} + c_{xy} \frac{\partial u}{\partial y} \right) \frac{\partial y}{\partial s} - c_{yy} \frac{\partial u}{\partial y} \frac{\partial x}{\partial s} + c_{bc} u = 0 \text{ on } \partial\Omega_N \quad (4.3)$$

Note that the boundary conditions must be homogeneous. Usually the function  $f$  is identically 1. However there are some occasions where other  $f$  are useful. For example, to put the Laplacian operator in polar coordinates into the form of Equation 4.1, one may multiply the canonical form of the equation by  $x$  (a.k.a.  $r$  in polar coordinates) which results in  $f = x$ .

You indicate that an eigenvalue problem is being solved by specifying `eq_type = EIGENVALUE` in subroutine `phaml_create`. The solution returns eigenfunctions,  $u$ , and the corresponding eigenvalues  $\lambda$ . More than one eigenpair can be computed. The number of desired eigenpairs is specified through the argument `num_eval` to subroutine `phaml_solve_pde`. Subroutines `phaml_evaluate`, `phaml_evaluate_old`, `phaml_integrate`, `phaml_query`, and `phaml_scale` take an optional integer argument, `eigen`, which specifies which eigenfunction to use. Subroutine `phaml_query` returns the entire set of eigenvalues in the `real(my_real)` array argument `eigenvalues`. Printed output prints all of the eigenpairs.

By default PHAML computes the smallest eigenvalue(s). You can compute eigenvalues in the interior of the spectrum by using the argument `lambda0` to `phaml_solve_pde`. PHAML will compute the eigenvalues that are closest to `lambda0`. The argument `lambda0_side` determines the position of the computed

eigenvalues relative to `lambda0`. It can be `EIGEN_LEFT` (eigenvalues less than `lambda0`), `EIGEN_RIGHT` (eigenvalues greater than `lambda0`), or `EIGEN_BOTH`. To obtain interior eigenvalues, a transformation that moves the desired eigenvalues to the ends of the spectrum must be used. Two transformations are supported in PHAML, specified by the argument `transformation` to `phaml_solve_pde`. `SHIFT_INVERT` uses the shift and invert transformation which, in part, uses  $(A - \lambda_0 M)^{-1}$ . `SHIFT_SQUARE` uses the shift and square transformation which, in part, uses  $(A - \lambda_0 M)^2$ . Currently, `SHIFT_SQUARE` is only supported for BLOPEX and only with `EIGEN_BOTH`. Also BLOPEX with `SHIFT_INVERT` does not currently support `EIGEN_BOTH`.

If  $u$  is a solution of Equation 4.1, then so is  $\alpha u$  for any scalar number  $\alpha$ . Thus the solution must be scaled to meet some condition to be unique. PHAML provides three options for how to scale the eigenfunctions through the argument `scale_evec` to subroutine `phaml_solve_pde`. `SCALE_LINF` requests that the  $\ell^\infty$  norm of the eigenvector,  $x$ , of the discrete problem be 1. With linear elements, this is the same as the  $L^\infty$  norm of the eigenfunction  $u$ . `SCALE_L2` scales such that the  $\ell^2$  norm of  $x$  is 1. `SCALE_M` scales such that the  $M$  norm of the eigenvector,  $\sqrt{x^T M x}$  where  $M$  is the mass matrix, or equivalently the  $L^2$  norm of the eigenfunction, is 1.

PHAML uses either ARPACK or BLOPEX to solve the discrete eigenproblem, so you must have PHAML configured with one of these optional packages (see Sections 2.1.7, 2.1.8 and 2.2.1). There are three arguments to `phaml_solve_pde` that affect ARPACK's behavior: `arpack_ncv` sets the number of Lanczos basis vectors, `arpack_maxit` sets the maximum number of IRLM iterations, and `arpack_tol` sets the relative accuracy of the eigenvalues. See the ARPACK User's Guide [20] for a deeper explanation of these arguments. There are also three arguments that affect BLOPEX's behavior: `blopex_maxit` sets the maximum number of iterations, `blopex_atol` sets a tolerance on the absolute residual, and `blopex_rtol` sets a relative tolerance.

## 4.2 Coupled Systems or Multicomponent Solutions

PHAML provides for the solution of certain types of coupled systems of elliptic PDEs, either boundary value problems or eigenvalue problems. These problems are also referred to as multicomponent problems if you prefer to think of it as a vector equation, rather than a system of equations, with a multicomponent solution, rather than multiple solutions. For these problems, Equations 1.1 - 1.3 are the same except the functions  $c_{xx}$ ,  $c_{xy}$ ,  $c_{yy}$ ,  $c_x$ ,  $c_y$ ,  $c_u$  and  $c_{bc}$  are  $n \times n$  arrays and the functions  $u$ ,  $f$  and  $g$  are vectors of length  $n$ , where  $n$  is the number of equations in the coupled system. An example of subroutine `pdecoefs` for a coupled system is given in Figure 4.1 for the equations

$$-\nabla^2 u + v = f_1$$



```

subroutine pdecoefs(x,y,cxx,cxy,cyy,cx,cy,c,rs)
use phaml
real(my_real), intent(in) :: x,y
real(my_real), intent(out), dimension(:,:) :: cxx,cxy,cyy,cx,cy,c
real(my_real), intent(out), dimension(:) :: rs

cxx(1,1) = 1.0_my_real; cxx(1,2) = 0.0_my_real
cxx(2,1) = 0.0_my_real; cxx(2,2) = 1.0_my_real

cyy(1,1) = 1.0_my_real; cyy(1,2) = 0.0_my_real
cyy(2,1) = 0.0_my_real; cyy(2,2) = 1.0_my_real

c(1,1) = 0.0_my_real; c(1,2) = 1.0_my_real
c(2,1) = 1.0_my_real; c(2,2) = 0.0_my_real

rs(1) = -(2.0_my_real*exp(x-y) - (x+y)**4/8.0_my_real)
rs(2) = -(3.0_my_real*(x+y)**2 - exp(x-y))

cxy=0
cx=0
cy=0
end subroutine pdecoefs

```

Figure 4.1: `pdecoefs` for a coupled system.

$$-\nabla^2 v + u = f_2$$

The number of equations is specified by the argument `system_size` to subroutine `phaml_create`. Subroutines `phaml_evaluate`, `phaml_evaluate_old`, `phaml_integrate`, `phaml_query`, and `phaml_scale` take an optional integer argument `comp` which specifies which component to use.

If any of the matrices  $c_{xx}$ ,  $c_{yy}$ ,  $c_u$  or  $c_{bc}$  is nonsymmetric, or  $c_{xy}$ ,  $c_x$  or  $c_y$  is nonzero, then you must use a nonsymmetric solver like `LAPACK_INDEFINITE_SOLVER` (single processor only), `MUMPS_NONSYM_SOLVER` or a nonsymmetric solver from PETSC. Do not use the built-in hierarchical basis multigrid method (the default) as a solver or as a preconditioner.

### 4.3 Parabolic, Nonlinear, Etc. Problems

PHAML can be used to solve parabolic (time dependent) and nonlinear PDEs, but the user has to provide the iteration control in the main program. See the example `examples/parabolic` for an example of solving a time dependent parabolic problem using an implicit finite difference scheme in time, and

`examples/nonlinear` for solving a nonlinear PDE using a simple Picard iteration. Other approaches to handling the time dimension or nonlinearity can be implemented, but they must be single step methods, i.e., must only require one previous solution in the iterative step.

The feature of PHAML that facilitates these problems is the storage of an “old” solution. You create an old solution with subroutine `phaml_copy_soln_to_old` (Section 7.4.3). This copies the current solution component of the `phaml_solution_type` variable to another component called the old solution. This would normally be done at the beginning or end of each iteration. To use the old solution, call `phaml_evaluate_old` (Section 7.4.8). You can evaluate the solution and/or the first derivatives and/or the second derivatives. This would normally be called from subroutine `pdecoefs`.

PHAML only provides for saving one “old” solution. You would think this would prohibit the use of methods that require more than one old solution. However, in many cases such problems can be reformulated as a coupled system of equations. See `examples/waveeq` for an example of solving a problem with a second order time derivative, which requires two old solutions, as a coupled system of two first order equations. An example of a second order backward difference scheme, which requires two old solutions, implemented as a coupled system of two equations is given in `examples/bdf2`. And `examples/irk4` contains a fourth order 6-stage implicit Runge Kutta method implemented as a coupled system of six equations.

A time dependent problem requires setting an initial condition to start the solution. Similarly, a nonlinear problem requires an initial guess of the solution. These initial functions are set by calling `phaml_solve_pde` with `task=SET_INITIAL` and an appropriate termination criterion. The solution will be set to be the function in subroutine `iconds` (Section 7.3.5). It should then be copied to the old solution before starting the iteration.

## Chapter 5

# Examples

Several examples of using PHAML are provided in directory `phaml-x.x.x/examples`. These can be used as tutorials to understand how PHAML works, or as templates for starting the implementation of your application. Depending on your set up, you might have to edit `master.f90` (or `spmd.f90` if you are using the SPMD model) slightly before running them, to change the termination criterion, number of processors, graphics choice, etc. The examples include:

`simple` – a trivial program that should be the first one you try.

`elliptic` – contains several linear scalar elliptic boundary value problems selected by a case statement.

`rectangle` – defines a rectangular domain with an  $N \times M$  grid by writing a Triangle `.poly` file.

`L-domain` – the classic L shaped domain with a singular solution commonly used for testing adaptive refinement codes.

`domains` – contains several domains defined by Triangle `.poly` files.

`curved` – defines a domain with curved boundaries.

`periodic` – periodic boundary conditions.

`eigenvalue` – an elliptic eigenvalue problem.

`parabolic` – solves a time-dependent parabolic PDE by an implicit finite difference scheme in  $t$ .

`nonlinear` – solves a nonlinear equation by Picard iteration.

`system` – solves a coupled system of 2 elliptic PDEs.

`all` – illustrates the use of all of the `phaml_*` routines.

`runtimepde` – determine user provided routines at run time

**waveeq** – solves the second order wave equations as a coupled system of two first order equations.

**bdf2** – solves a parabolic PDE by second order backward differences implemented as a coupled system of two equations.

**irk4** – solves a parabolic PDE by a 4<sup>th</sup> order, 6-stage implicit Runge Kutta method implemented as a coupled system of six equations.

There are also several programs under the **testdir** directory, but they do not contain comments to help you understand them.

# Chapter 6

## Release notes

This section is reserved for listing the changes with each new release of PHAML. See also the files `doc/HISTORY` and `doc/UPGRADING`. Changes in other parts of this document are indicated by font color and size. Changes made in the current release are printed in **red**. Changes made in a recent release are printed in **dark red**. Text that has been deleted is maintained for a few releases, but is printed in a tiny size.

### 6.1 Version 1.9.1

Released April 25, 2011

#### 6.1.1 Summary of changes

- `mark_reftype_one` returning “n” is not a fatal error in `mark_for_refinement`
- bug: exceeded dimension of `multi_p` in equilibrated residual error indicator with multiple eigenvalues
- bug: used `system_size` instead of `num_eigen` for dimension of energy error estimate in bigger\_errind *hp* strategy
- bug: master not getting any\_changes in `reconcile_requests` and `pred_load_balance`
- bug: use `alltoall` instead of `send/recv` in `enforce_overlap`

#### 6.1.2 Major changes

This is a bug fix release.

#### 6.1.3 Upgrading

No changes compared to 1.9.0.

## 6.2 Version 1.9.0

Released April 13, 2011

### 6.2.1 Summary of changes

- Additional OpenMP sections:
  - `all_error_indicators`
  - most cases of adaptive refinement
  - multigrid, except basis change
- Changed the way `reterm=KEEP_*` is handled when the corresponding `max_` is supplied, so it doesn't drift.
- Use the usual error indicators with the reference solution strategies.
- Moved printing of linear system information to before solving it.
- Added option of printing solver information.
- Added a user callable routine to return the vertices of the grid and solution at the vertices.
- Added `waveeq`, `bdf2` and `irk4` examples to illustrate how to use coupled systems in place of multiple old solutions.
- Set order of quadrature rule for `norm_error` based on element degree.
- Modification to T3S hp-adaptive strategy to avoid stalling in some cases.
- Quit using `assoc_elem` for temporary storage of `vert_master` in `init_grid`.
- Faster, less careful barycentric coordinates for basis functions if quad precision not selected.
- Remove support for PETSc ILU drop tolerance (removed from PETSc 3.1)
- Change in PETSc interface `#include` because of change in PETSc 3.1 .h files.
- Added an option in `mkmkfile.sh` for PETSc (3.1 and later) as a single library.
- Change in BLOPEX support for changes in BLOPEX 1.1.
- Added jump term for periodic boundary conditions in explicit error indicator.
- Added subsections for `phaml_solve_pde` arguments in the User's Guide.

- Dropped PVM support.
- Check allocation status on all allocate statements.
- Replaced `my_real` with explicit single or double precision in arguments to blas and lapack routines.
- Allow reference solution to have level `max_lev+1` in `REFSOLN_EDGE`.
- Added OpenMP ordered construct and dynamic schedule to matrix assembly and static condensation.
- Moved subroutines from `gridtype.f90` to `grid_util.f90`.
- Changed factor for `MG_ERREST_TOL` to `1/1000`.
- Print warning if multigrid doesn't reach tolerance.
- Increased `mg_cycles` in tests as necessary to meet convergence test.
- Don't force balancing in predictive load balance.
- Changed MCSD to ACMD in header comments.
- bug: doubly periodic boundary conditions didn't work.
- Workaround for PGI compiler bug in `explicit_ei`.
- bugs in `resid_natural_bconds`: used outer do loop variable in implied do, double use of `rs`, didn't evaluate `pdecoefs` for natural boundary condition.
- bug: inconsistency in allocation/use of `oldsoln`
- bug in the handling of additional space for fill-in in MUMPS.
- bug: initial elements in reverse order when drawing space filling curve.
- bug: shortened some subroutine names that were too long in `hp_strategies.f90`.
- bug: for printed relative error estimate, dividing by the norm of the true solution instead of the computed solution when still sequential.
- bug: made initialization of quadrature rules a critical section in OpenMP.
- bug: equilibrated residual error indicator did not work with periodic boundary conditions.
- bug: `save_matrix` did not zero out Dirichlet columns in non-Dirichlet rows.
- bug: With conventional communication, don't relax unowned equations.
- bug in `unbisect_triangle_pair` with periodic boundary conditions
- bug: miscounted number of nonzero entries in a row with periodic boundary conditions on high order element

## 6.2.2 Major changes

Most of PHAML has now been parallelized with OpenMP. The major additions in this release were adaptive refinement and multigrid. This required changes to both of those algorithms to group entities into independent sets that can be operated on in a data parallel sense.

Printed output associated with solving the linear system has been enhanced. The output from `print_linsys` now occurs between assembly and solution, which helps to monitor the program progress. A new facility, `print_solve`, provides information about the solver, such as how many iterations it used, after the solve phase is complete.

A new utility routine was added to return the vertices of the triangles and the solution at the vertices. See `phaml_get_grid_soln` in Section 7.4.7.

Three new examples were added to illustrate how to formulate algorithms that require more than one “old” solution as a system of equations. These examples illustrate a second order backward difference, fourth order implicit Runge-Kutta, and solution of the wave equation.

Support for PVM was removed.

## 6.2.3 Upgrading

Existing programs shouldn’t require any changes. However, due to the major restructuring of adaptive refinement and multigrid, you cannot expect to get results that are identical to those of the previous version. If you really need to use the previous adaptive refinement algorithm, you can set `use_old_refinement=.true.` in `refine_adapt.f90`. However, this will be removed in a future release.

There are changes in `mkmkfile.sh`. You will need to edit it with your system specifics.

The multigrid algorithm now prints a warning if it doesn’t reach the desired residual tolerance in the allowed number of iterations. If you get this warning, you might want to increase the number of iterations via `mg_cycles` or shut off the warnings via `print_warnings`.

## 6.3 Version 1.8.0

Released April 9, 2010

### 6.3.1 Summary of changes

- Improved weights for load balancing high order and hp-adaptive elements.
- When determining if the distribution is sufficiently out of balance to do load balancing, check the balance of the entities that are being balanced instead of the balance of the elements.
- Additional OpenMP sections:



- Static condensation
- Elemental matrix computation and assembly
- Extend REFSOLN\_EDGE to handle multicomponent solutions.
- Estimate Sobolev regularity for NLP using PRIOR2P\_H1 instead of NEXT3P.
- When insuring  $\tau$  permits a feasible solution for NLP, don't scale the error estimate by  $\tau$ .
- Use inc\_factor to determine which edges get refined in REFSOLN\_EDGE instead of hard coded 10, and set default inc\_factor to 10 for REFSOLN\_EDGE and REFSOLN\_ELEM.
- Moved compute\_global\_max\_errind to grid\_util.f90.
- Removed unused numhref and numpref from mark\_reftype\_one.
- Removed unused edge\_block.
- Don't load balance with one processor.
- Scale graphics perspective by domain size.
- REFSOLN\_ELEM singular projection workaround
- Immediate return from static\_condensation if there are no face equations.
- bug: delta\_dof not set on early return from p\_refine\_element\_interior.
- bug: times not set in print\_error\_info with sequential and save\_convergence.
- bug: not including system\_size in count\_dof face dof.
- bug: LOCAL\_P error estimate was not using off diagonal coefficients for coupled systems of PDEs.

### 6.3.2 Major changes

The weights used in the load balancing algorithm were changed, mainly to include the additional work associated with higher degree elements. Tests typically showed a factor of 2-3 increase in speed on 8 processors with high order elements or hp-adaptive grids.

Additional parts of the program have been parallelized with OpenMP, but still not enough to really be useful.

Control over how much refinement occurs in one phase with the REFSOLN\_EDGE hp strategy was added, and REFSOLN\_EDGE was extended to handle systems of equations.

### 6.3.3 Upgrading

The default `inc_factor` for `REFSOLN_ELEM` is now 10. To get the old behavior use `inc_factor = 2.0`.

The changes in the load balancing weights will likely change your results slightly, due to changes in the refinement, which come from the different partition of the grid. There is no way to get the old behavior.

The `mkmkfile.sh` from the previous version should still work.

## 6.4 Version 1.7.0

Released January 26, 2010

### 6.4.1 Summary of changes

- Added `HP_REFSOLN_EDGE` *hp*-adaptive strategy.
- Broke grid module into several smaller modules.
- Began adding support for OpenMP:
  - Added `openmp`, `hybrid_spawn` and `hybrid_nospawn` options in `mkmkfile.sh`.
  - Created OpenMP stub library.
  - Support for h-uniform refinement.
  - Support for p-uniform refinement.
- Added a toggleable color key to the graphics.
- Removed unnecessary assignments in `init_guess_h`.
- If stopping on max lev or deg, don't switch type when max is reached.
- Removed possible p refinement of mate during h refinement.
- Cleaned up parts of `mkmkfile.sh`.
- Added a note in `examples/*/mkmkfile.sh` about not running that file.
- Changed the ParMETIS approach in Zoltan to the new recommendation.
- Changed default stepped sequential color scheme to 4 steps of 6 hues.
- Graphics: corrected computation of minimum and maximum element size.
- Removed `doc/HISTORY` and `doc/UPGRADE`
- Bug: `det_one_opt_ref_elem`, array shape mismatch for subset.
- Bug: `init_guess_p` assumed edge rule is maximum rule.

- Bug: `init_guess_h` used `elem` instead of `mate` for `elemental_matrix` call.
- Bug: grid degrees of freedom count did not include multiple components

### 6.4.2 Major changes

`mkmkfile.sh` has substantial changes. You should modify the new version with your system information rather than using your old one.

A new *hp*-adaptive strategy, `HP_REFSOLN_EDGE` was added.

A color key was added to the graphics to indicate the numeric range that the colors represent. In the menu that the right mouse button brings up, look for “toggle color key”.

Support for OpenMP has begun, to provide shared memory parallelism on multicore machines, and hybrid parallelism on clusters of multicore machines. However, there is not much useful at this point. Look for further support to be added in future versions.

### 6.4.3 Upgrading

`mkmkfile.sh` has substantial changes. You should modify the new version with your system information rather than using your old one.

The number of steps and hues in the stepped sequential color scheme has changed to (4,6). Use the color scheme submenu to change it if you want the old (4,4).

## 6.5 Version 1.6.2

Released September 28, 2009

### 6.5.1 Summary of changes

- Added second interface to PHAML public routines in `runtimepde`.
- Bug in `krylov.f90`, deallocating unallocated rhs.

### 6.5.2 Major changes

There are no significant changes.

### 6.5.3 Upgrading

Fully compatible.

## 6.6 Version 1.6.1

Released September 18, 2009

### 6.6.1 Summary of changes

- Added a new example, runtimepde, that allows changing the user provided routines at run time.
- Modified mkmkfile.sh for normal usage of LAM and Open MPI compiler wrappers.
- Changed graphics window size from 300 pixels to 500 pixels.
- Bug in phaml\_slave, rpos too small.
- Bug in refsoln\_errest, sqrt of negative number.
- Bug in static condensation with nonsymmetric matrix; assumed symmetric.
- Bug in make\_linsys, not setting extra\_bcns with loc\_bconds\_s.

### 6.6.2 Major changes

There are no really significant changes.

### 6.6.3 Upgrading

mkmkfile.sh has substantial changes. You should modify the new version with your system information rather than using your old one.

## 6.7 Version 1.6.0

Released August 4, 2009

### 6.7.1 Summary of changes

- Added cross derivative and first order terms to the PDE.
- Added choice between minimum rule and maximum rule for edge degree, and made minimum rule the default.
- Added HP\_REFSOLN\_ELEM hp strategy.
- Added HP\_NLP hp strategy.
- Added stepped sequential color scheme.
- Replace exchange\_neigh\_vect with send\_neigh\_vect and rcv\_neigh\_vect in some places to allow overlapping communication with computation, and only send to nearest neighbors.
- Skip final communication in matrix\_times\_vector when not needed.

- Reduced communication in `sum_fudop_vect`.
- Changed BLOPEX dummy routine from C to Fortran.
- Removed restriction on neighboring degrees in derefinement.
- Immediate return from exchange routines if only one processor.
- In `make_elist`, don't scale `reftol` by `normsoln` with T3S or ALTERNATE.
- Made minimum value 1 instead of 0 for scaling for color by degree.
- Added cross derivative to p-hierarchical basis function and local evaluate.
- Added relaxation at unowned vertices with `conventional_comm`.
- In high order relaxation, group equations of the same degree together, order the relaxations from high to low degree, and use local black relaxation.
- Slight modification to NEXT3P.
- Inconsequential coding change in `HP_ERRIND`.
- Changed the way `LAMHF77` is defined in `mkmkfiles`.
- Bug in allocation of `element_errind` in `phaml_restore`.
- Bug in order of subscripts in `u_K` in equilibrated residual `errest` with mixed bc.
- Bug; use quadrature rule that stays in the triangle in equilibrated residual.
- Bug in matching dimensions in `where` statement in `krylov`.
- Bug; checking size of unallocated allocatable array in `check_stall_special`.
- Bug; Texas 3 Step might not set work at end of `bisect_triangle_pair` resulting in divide by zero.
- Bug in setting edge next in `compress`.
- Bug in `eigen`; not summing `linsys` residual over processors.
- Bug; not measuring communication time in `phaml_cg`.
- Bug; `skip_matrix` in `basis_change` didn't work.
- Bug in second derivatives of basis functions.
- Bug in explicit error estimate with high order bases.
- Bug in explicit error estimate with mixed b.c.; wrong sign on `u` term.

### 6.7.2 Major changes

The class of problems that can be solved has been extended by adding the cross derivative and first derivative terms to the operator. Note that the discretization of these terms results in a nonsymmetric matrix, so you need to use a nonsymmetric solver like those in PETSc or *hypre*, or MUMPS\_NONSYM\_SOLVER.

Two more *hp*-adaptive strategies have been added.

A new color scheme has been added which provides more distinct separation of neighboring colors. This is useful when you need a large number of discrete colors, such as the element degree in *p*-adaptive or *hp*-adaptive methods, and the element owner when there are a lot of processors.

### 6.7.3 Upgrading

The minimum rule for determining the degree of an edge between two triangles of different degrees was added and made the default. To get the previous behaviour, use `edge_rule=MAXIMUM_RULE`.

The color scheme for color-by-owner was changed to be the same color scheme used by everything else. One consequence of this is that the direction of the rainbow has been reversed, i.e., processor 1 is blue and processor `nproc` is red.

There are changes that will cause slightly different answers in some cases.

`mkmkfile.sh` has substantial changes. You should modify the new version with your system information rather than using your old one.

## 6.8 Version 1.5.0

Released October 9, 2008

### 6.8.1 Summary of changes

- Added COEF\_DECAY *hp* strategy.
- Added COEF\_ROOT *hp* strategy.
- Added SMOOTH\_PRED *hp* strategy.
- Added NEXT3P *hp* strategy.
- Added EQUILIBRATED\_RESIDUAL error estimator.
- Added the option of whether or not to solve on the initial grid.
- Added the ability to use `max_lev` and `max_deg` as termination criteria.
- Check for not stalling when element  $p = 1$  with all three edges  $p > 1$ .
- If `errtype` is RELATIVE, use relative error for error estimate termination criteria.

- When doing  $h$  refinement, if the mate is scheduled for  $p$  refinement, do that refinement first.
- Do not compute error indicator of refined elements if not needed.
- Do not determine if refinement is  $h$  or  $p$  if not needed.
- Reduced default `max_deg` by one if `error_estimator` is `LOCAL_PROBLEM_P`.
- Increased quadrature rule order by one in `phaml.integrate`.
- Multiply  $p$  indicator by two in `HP_BIGGER_ERRIND` strategy.
- Finished `LOCAL_PROBLEM_H` and `LOCAL_PROBLEM_P` error indicators for nonconstant coefficients and multiple eigenvalues.
- Removed dead code associated with static condensation, pairing of triangles in the initial grid, and finding the `bparam` of a point on the boundary.
- Changed `mkmkfile.sh` to allow sequential MUMPS.
- Fixed documentation error concerning `reftol` and `ONE_REF`.
- Fixed documentation error and header concerning `LAST`.
- Updated the Quickstart section of the user's guide to include Open MPI.
- Bug in `norm_solution`: 2nd and higher eigenvectors not available.
- Bug in selection of elements to refine with `ONE_REF*` and high order elements.
- Bug: duplicate send tag when printing initial error `FREQUENTLY`.
- Bug in `LOCAL_PROBLEM_H` error indicator with Neumann boundary conditions.
- Bug in allocation size of message to graphics when `PARALLEL` is sequential.

### 6.8.2 Major changes

The major change is the addition of the equilibrated residual error estimator and four more  $hp$ -adaptive strategies. The option of not solving the discrete problem on the initial grid can be useful if `phaml.solve_pde` is called more than once.

### 6.8.3 Upgrading

None of the changes in this version require any changes in application codes, however some of the changes may cause the grid and/or computed solution to be slightly different in some cases. The `mkmkfile.sh` from the previous version should still work, unless you are using the sequential version of MUMPS.

## 6.9 Version 1.4.0

Released April 25, 2008

### 6.9.1 Summary of changes

- Added T3S hp strategy.
- Added ALTERNATE hp strategy.
- Added TYPEPARAM hp strategy.
- Added option of conventional communication for multigrid.
- Changed HBMG to an hp-multigrid cycle.
- Can run high order HBMG in parallel.
- Added support for BLOPEX (only via PETSc so far).
- Added option to compute eigenvalues on the left or right of  $\lambda_0$ .
- Solve on the initial grid and print error and draw.
- Use number of elements instead of degrees of freedom for ONE\_REF criterion.
- Use error estimate instead of error indicator for ONE\_REF criterion.
- Send mindeg and maxdeg to master in `get_grid_info`.
- Reconcile edge degree.
- Declare CG convergence with relative residual instead of absolute.
- Use sum of exports instead of max to see if redistribution is needed.
- Added `max_lev` to `exchange_*_vect` and `no_soln` to `exchange_fudop_soln_resid`.
- Improve efficiency of face basis functions.
- Improve efficiency of `elem_exact`.
- Tweaked the determination of the final error code.
- Discontinue maintenance of HISTORY and UPGRADE in doc/
- Fix spacing in PETSc related parameters in header.
- Bug in `gmres` when linear system size is 1 or 2.
- Bug in `phaml.f90`; not setting `nvert1` if refinement not requested.
- Bug; no `tol=tol/100` in `CG_SOLVER`.
- Bug; missing `no_master=.true.` in check for stalled refinement.
- Bug in `linsys_resid`; sending message to master when it's not there.



### 6.9.2 Major changes

The option of using conventional communication with the hierarchical basis multigrid method was added as an alternative to the full domain partition reduced communication. With conventional communication the multigrid method gives the exact same answers as a sequential implementation. It is selected by giving `mg_comm = MGCOMM_CONVENTIONAL` to `phaml.solve_pde`.

The high order bases part of the HBMG multigrid solver was changed from several Gauss-Seidel iterations ( $p_{\max}$  by default) to a  $p$ -multigrid cycle. See Section 3.3.4.1. Correspondingly, the default for `mg_prerelax_ho` and `mg_postrelax_ho` was changed to 1.

Partial support for the eigensolver BLOPEX was added. Currently it can only be accessed through PETSc, i.e., you must have PHAML configured with PETSc, you must have PETSc configured with BLOPEX, and you must use a PETSc solver (but you can use the multigrid preconditioner). Support for BLOPEX through *hypre* and as a stand-alone package will be added in a future release.

Support was added to request eigenvalues on either side of  $\lambda_0$ , rather than the closest eigenvalues on both sides. To get eigenvalues less than  $\lambda_0$ , use `lambda0_side = EIGEN_LEFT`. To get eigenvalues greater than  $\lambda_0$ , use `lambda0_side = EIGEN_RIGHT`.

Three more *hp*-adaptive strategies were added: the Texas 3 Step strategy, a similar strategy that alternates between  $h$  and  $p$  refinement, and the type parameter strategy. These are still experimental.

### 6.9.3 Upgrading

The logical parameter `mg_nocomm` was replaced by integer `mg_comm` since there are now three options for the multigrid communication. If you used `mg_nocomm=.true.`, replace it with `mg_comm=MGCOMM_NONE`.

Changes in the multigrid solver may cause slightly different answers in some cases.

`doc/HISTORY` and `doc/UPGRADING` are no longer maintained. That information is contained in this section of the User's Guide.

The `mkmkfile.sh` has important changes. You must modify the new version for your system.

## 6.10 Version 1.3.1

Released January 28, 2008

### 6.10.1 Summary of changes

- Bug in `linsys.io.f90`; mixed kinds in `max`.

### 6.10.2 Major changes

None. This was a bug fix release.

### 6.10.3 Upgrading

Fully compatible.

## 6.11 Version 1.3.0

Released January 24, 2008

### 6.11.1 Summary of changes

- Added quadrature rules of order 21 to 45 for triangles.
- Changed default max\_deg to 22.
- Replaced singular\_points with function regularity.
- Renamed HP\_AS2 to HP\_APRIORI
- Renamed HP\_RAS1E and HP\_RAS1H1 to HP\_PRIOR2P\_E and HP\_PRIOR2P\_H1.
- Removed degree  $\geq 3$  restriction on PRIOR2P strategies.
- New parallel implementation of conjugate gradients.
- Added option of printing relative error instead of absolute error.
- Added norms of true solution to phaml\_query.
- Added MUMPS nonsymmetric solver.
- Added second derivatives uxx and uyy to phaml\_evaluate\_old.
- Added first and second derivatives to phaml\_evaluate.
- Reorganized refinement loop.
- Added missing special case to LOCAL\_PROBLEM\_P error indicator.
- Slight change in the order of high order relaxation in hierarchical basis multigrid.
- Use quadrature order high enough for exact integrals in phaml\_integrate.
- Increased quadrature order for elemental matrices to be exact for mass matrix.
- More extensive use of double precision in graphics.

- In graphics, scale true by `maxabstrue` instead of `maxabssolut`.
- Improve vertical scaling in graphics.
- Removed extraneous choices for displaying error indicator.
- Added processor number to graphics postscript file name.
- Removed printing of error before first solution.
- Removed extra relaxation pass for linear elements in `hbmng`.
- Removed old farg kludge for MPICH.
- Removed `GRAPHICS_TREE` from `messpass` modules.
- In `run_tests` scripts under `testdir`, don't assume "." is in the path.
- Reduce roundoff error in computing barycentric coordinates.
- Minor bug in `petsc_interf.f90`.
- Allocation bug in `superlu_interf.f90`.
- Bug in `evaluate_oldsoln_local`; using unallocated pointer.
- Bug in graphics; contour plot of error.
- Fixed a floating point incompatibility between LAPACK source code and `gfortran`.
- Bug in graphics from `phaml_restore`; initialized twice.
- Bug in printing convergence history of conjugate gradients.

### 6.11.2 Major changes

The *hp*-adaptive strategy AS2 was renamed APRIORI, and the array `singular_points` was replaced by subroutine `regularity`. This allows more flexibility in specifying *a priori* knowledge about solution irregularities. See section 3.3.2. Other *hp*-adaptive strategies were also renamed for consistency with the new strategies.

### 6.11.3 Upgrading

You must add a new subroutine to the file where you define your problem, usually `pde.f90`. The function subroutine `regularity` was added for guiding *hp*-adaptive refinement by the APRIORI strategy. You can copy a dummy version from the end of `examples/simple/pde.f90`.

The symbolic constants for `hp_strategy` have been changed. If you are using *hp*-adaptive refinement, you will have to change this. `HP_AS2` is now `HP_APRIORI`. `HP_RAS1E` is now `HP_PRIOR2P_E`. `HP_RAS1H1` is now `HP_PRIOR2P_H1`.

If you are using the AS2 (now APRIORI) *hp*-adaptive strategy, the means for specifying irregularity has been changed. The argument `singular_points` has been removed. Instead, you must write a function subroutine called `regularity` to indicate that a triangle contains an irregular part of the solution. See the end of `examples/L-domain/pde.f90` for a working example of this subroutine. To duplicate prior behavior, `regularity` should return a value less than 1.0 if one of the given triangle vertices is a point that you would have given in `singular_points`.

The default value for `max_deg` was changed to 22, which is the maximum for the new higher order quadrature rules. To duplicate the previous behavior, use `max_deg=10`.

The order of the quadrature rule for elemental matrices was increased to be exact for the mass matrix instead of the stiffness matrix. If you have a *u* term in your PDE or your right hand side is not a polynomial of sufficiently small degree, this will cause a slight change in your results. To duplicate the previous behavior, use `inc_quad_order=-1` for linear elements, or -2 for high order elements.

Optional arguments for second derivatives were added to `phaml_evaluate_old`, but not at the end of the list. If you used this routine with `comp` and/or `eigen` given as non-keyword arguments, you will need to change that call.

In `phaml_evaluate`, the returned solution `soln` was replaced by `u`, `ux`, `uy`, `uxx`, `uyy`. If you designated `soln` as a keyword argument, it will have to be changed to `u`. If you used this routine with `comp` and/or `eigen` given as non-keyword arguments, you will need to change that call.

Many of the changes in this release will cause slight changes in the numerical results. Your computed solution should have roughly the same accuracy, but don't expect it to be exactly the same as that computed by the previous release.

Your `mkmkfile.sh` from 1.2.0 should still work.

## 6.12 Version 1.2.0

Released October 11, 2007

### 6.12.1 Summary of changes

- Added explicit error indicator.
- Made explicit error indicator the default.
- Added optional argument `inc_quad_order` to `phaml_store_matrix`.
- Added optional argument `error_estimator` to `phaml_query`.
- Added second derivatives to basis and evaluation routines.
- Added an estimate of the error in the eigenvalues.

- Changed how error indicators are computed.
- Changed how initial solution is set during refinement with SET\_INITIAL.
- Removed option to compute two error indicators.
- Added a warning when the ARPACK eigenvalue is not close to the Raleigh quotient.
- Removed the restriction that neighboring elements differ in degree by at most 1.
- Removed some dead code related to error indicators.
- Extended hierarchical coefficient error indicator to high order elements.
- Keep quadrature points inside triangles when necessary.
- Faster finding of element that contains a given point.
- Upgraded PETSc to version 2.3.3.
- Moved control of *hypr* version from mkmkfile.sh to source code.
- Removed BoomerAMG\_IOutDat.
- Removed some unused variables.
- Added PBS scripts for running tests on batch systems with PBS or Torque/Maui.
- Removed doc/USER\_GUIDE.
- Bug in deallocation in PetscInterf.F90.
- Bug in assignment of vertices to edge with periodic boundary conditions.
- Bug in src/mkmkfile.sh with zoltanP\_r\_f\_dum.f90.
- Bug: uninitialized loc\_Linf\_norm in norm\_error.

### 6.12.2 Major changes

Most of the major changes concern the error indicators.

A new error indicator was added. This is the explicit error indicator defined in Chapter 2 of [1], which is based on computing the norm of the residual (see Section 3.3.3). It is much more efficient than the local problem error indicators, but less accurate. It should guide adaptive refinement well, but if you want a good estimate of the error you should use one of the local problem error indicators. The new error indicator is specified with `error_estimator=EXPLICIT_ERRIND`. It is also now the default error indicator.

The HIERARCHICAL\_COEFFICIENT error indicator has been extended to high order elements. Formerly, it could only be used with linear elements.

The option to compute two error indicators was removed. This also means the LOCAL\_PROBLEM error indicator is removed, because it computed two error indicators, but the individual LOCAL\_PROBLEM\_H and LOCAL\_PROBLEM\_P error indicators remain.

### 6.12.3 Upgrading

There were slight changes to how most of the error indicators are computed. You might see a small difference in your results.

The option to compute two error indicators was removed. If you were using this, you might need to change some of the arguments to `phaml.solve_pde` and `phaml.query` by removing the 2 or 12 from the end of the symbolic constant.

The default error indicator was changed to `EXPLICIT_ERRIND` which is much faster. This will cause a small change in your results. To (nearly) get the previous results, use `error_estimator=LOCAL_PROBLEM_H`.

There was a change in how the initial condition is assigned from function icons. If you use this, you might see a slight change in your results.

The restriction that neighboring elements differ in degree by at most 1 was removed. If you use  $p$  or  $hp$  refinement, you might see a slight change in your results.

The interface to PETSc was upgraded to version 2.3.3. If you are using an older version of PETSc then you may need to make some changes in `mkmkfile.sh`, `petsc_init.F90` and `petsc_interf.F90`. Search for "before" to find the changes.

The control over selecting what version of *hypre* you use (if you use *hypre*) was removed from `mkmkfile.sh`. Instead, it is controlled by commenting out some lines of `hypre_fix.c`. If you use *hypre*, check that file to see if you need to make any changes. Also check `mkmkfile.sh` and search for "hypre version".

There are changes in `mkmkfile.sh` that might effect you if you use PETSc or *hypre*. Otherwise, `mkmkfile.sh` from Version 1.1.0 should still work.

## 6.13 Version 1.1.0

Released July 3, 2007.

### 6.13.1 Summary of changes

- Added native conjugate gradients and GMRES solvers.
- Added `ZOLTAN_FILE` to partition methods.
- Added support for DRUM.
- Added a routine to store the matrix as a Matrix Market file.
- Changed HBMG convergence test to use relative residual instead of absolute residual.

- Changed tests with high order elements to use CG\_SOLVER and run in parallel.
- Changed mkmkfile.sh handling of Zoltan optional third party libraries.
- Changed the tests where the solution was a polynomial of degree less than or equal to the degree of the approximation space.
- Removed multigrid F cycle.
- Bug in eigen.f90; static condensation needs contribution of other processors.
- Bug; uninitialized variables in graphics with SEQUENTIAL.
- Bug in maintaining high order oldsln during derefinement in parallel.

### 6.13.2 Major changes

Native Conjugate Gradients and GMRES solvers were added. They can be used without preconditioning, or with the hierarchical basis multigrid preconditioner. This now provides a native solver for high order bases in parallel. See Section 3.3.4.2.

A new way of specifying a partitioning method from Zoltan was added. It uses Jim Teresco's zoltanParams library [43] to read Zoltan parameters from a file. This allows for not only selecting the partitioning method in a file, but also for setting any other Zoltan parameter. This method is selected with `partition_method=ZOLTAN_FILE` and the file is specified by the argument `zoltan_param_file`. See Section 3.3.5.

Support was added for DRUM, the Dynamic Resource Utilization Model [15][42]. This is useful to improve load balancing in heterogeneous and hierarchical parallel computing environments. DRUM is accessed through Zoltan, with the partition method `ZOLTAN_FILE` and the DRUM parameters given in the Zoltan parameter file. See Section 3.3.5.

A user-callable utility routine was added to store the stiffness matrix and/or right hand side in a file in Matrix Market format. For eigenvalue problems, the mass matrix can also be stored. This allows for computations on the linear systems to be performed outside of PHAML, for example with Matlab. See Sections 3.6.2 and 7.4.17.

### 6.13.3 Upgrading

The multigrid F cycle was removed, including FMG\_PRECONDITION. If you used the FMG preconditioner, replace it with MG\_PRECONDITION with a sufficient number of cycles.

The test for convergence of the hierarchical basis multigrid method was changed from using the absolute residual to using the relative residual. It is possible that this may change your answers slightly, but they should agree to

several significant digits. In rare cases, the grid may be slightly different, if the error estimate for the element was very close to the cutoff value. There is no way to recover the previous behavior.

The `mknkfile.sh` has important changes. You must modify the new version for your system.

## **6.14 Version 1.0.0**

Released May 4, 2007.

The first non-beta release. Beta users should see the files `HISTORY` and `UPGRADING` for changes since the last beta release.



# Chapter 7

## Reference Manual

### 7.1 Quick Start

This section provides a minimum of information about getting PHAML up and running the example programs. For many people, this will be sufficient to get started using PHAML with your application, as many applications may be implemented by modifying one of the examples. If you encounter problems, refer to the appropriate section of the User's Guide for further details. Also read the remainder of the User's Guide for a better understanding of how PHAML works and what options may be useful to you.

The four steps in getting started with PHAML are

1. obtaining the software,
2. compiling the PHAML library,
3. compiling an example, and
4. running the example.

For this brief tutorial, we will build and run the `simple` example as a master/slave message passing MPI program with the master spawning the slave processes, and without graphics.

#### 7.1.1 Obtaining the software

PHAML can be obtained from the PHAML web page <http://math.nist.gov/phaml> by following the Software link. It comes as a gzipped tar file for Unix-like systems. (It has not yet been tested on MS Windows systems, but since it is written in standard Fortran 90, the adventurous may find that it works on MS Windows, too.) When unpacked, it will place everything in a new directory called `phaml-x.x.x`, where `x.x.x` is the current version number.

PHAML requires the BLAS and LAPACK libraries. You will probably find these are already installed on your computer, but if not, see Section 2.1.3.

For parallelism, you need an MPI library. You will probably find that Open MPI, LAM, MPICH, or a commercial MPI library is already installed on your computer, but if not see Section 2.1.4.

### 7.1.2 Compiling the PHAML library

The first step in compiling the PHAML library is to create the **Makefile**. This is done with the shell script `mkmkfile.sh` in the top PHAML directory. Since PHAML allows so many options in terms of what compilers and libraries to use, it would be difficult to auto-locate these files. So, you must modify `mkmkfile.sh` to specify some paths, command names, and defaults for your computer system. Instructions for modifying it can be found within the file.

Now create the **Makefile** with

```
./mkmkfile.sh PARALLEL messpass.spawn PARLIB mpi GRAPHICS none
```

You should replace `mpi` with `mpich`, `mpich2`, `openmpi`, or `lam` if you use an MPICH, Open MPI, or LAM library. You may omit some of the arguments if your defaults are already set to these values; defaults can be determined with `mkmkfiles.sh help`.

`make` should now compile the library and place it in the `lib` subdirectory.

### 7.1.3 Compiling an Example

Go to the directory `examples/simple` and type `make`. (The **Makefiles** for the examples were also created by `mkmkfile.sh`.) This should create the executables `phaml` and `phaml_slave`.

### 7.1.4 Running the Example

The details of running an MPI program vary with the different MPI libraries. You may need to check your MPI documentation to find the correct command(s). It may also require starting some daemon before running the execution command.

Note that you should specify *one* process, because you are running the master processes which will spawn the slave processes. The number of slaves is specified in the main program, `master.f90`.

If you are using LAM, try

```
lamboot
```

```
mpirun -np 1 phaml
```

If you are using MPICH, try

```
mpirun -np 1 phaml
```

If you are using MPICH2, try

```
mpiexec -n 1 phaml
```

If you are using Open MPI, try

```
mpiexec -np 1 phaml
```

### 7.1.5 Now what?

If you have successfully run the first example, you are ready to install the graphics and any other optional software you desire (Section 2.1), run the other examples, and begin working on your own application!

## 7.2 Public Entities in PHAML

The statement `use phaml` in a program unit provides access to the public entities in PHAML. These consist of a derived type, variables for the user to use, symbolic constants, and procedures. The procedures are described in Section 7.4. The other entities are described in this section.

### 7.2.1 `phaml_solution_type`

`phaml_solution_type` is a type that contains all the data used for solving the PDE (grid, etc.). The type is public, but the contents are private. You can declare one or more variables of this type and pass them to the PHAML procedures.

### 7.2.2 `my_real`

`my_real` is a symbolic constant that determines the kind of real numbers used in PHAML. This is defined in `global.f90`. You can change that definition there to select the kind to use, either single or double precision. Other kinds of real, if supported by your Fortran compiler, are not supported by PHAML because of the reliance on BLAS and LAPACK routines, which only come in single and double precision. Currently `my_real` is set for default double precision. To be sure that your program is using the same `kind` values, you should declare your variables as `real(kind=my_real)` and attach the kind to constants, e.g. `0.0_my_real`.

### 7.2.3 `pde` and `my_pde_id`

`pde` is a rank 1 allocatable array of type `phaml_solution_type`. If you are going to use more than one `pde` and they must communicate with each other, then you must use this array for your `phaml_solution` variables. It must be allocated before using `phaml_create`, and should be deallocated when you are done with it.

`my_pde_id` is an identifier for which `pde` a `phaml_solution` is associated with, usually the index into the array `pde` described above. It is usually used as the case statement variable in the user provided routines that define the PDE and boundary conditions.

See `examples/system.ss` for an example of the use of `pde` and `my_pde_id`. However, having multiple `phaml_solution_type` variables communicate with each other is very slow, so use of this capability is discouraged.

### 7.2.4 symbolic constants

Except for `my_real`, the symbolic constants are used as values for the arguments of the procedures. In this document and in the PHAML source code they are written in capital letters, but Fortran is not case sensitive so this is not necessary. They are defined as they arise in the description of the PHAML procedures. You can find a list of them by looking at the `public` statements at the beginning of `phaml.f90`. If you have a problem with one of the names conflicting with an entity in your code, you can use Fortran's renaming capability to circumvent it, for example

```
use phaml, MY_EVAL_NAME => EIGENVALUE
```

## 7.3 User Provided Routines

The specifics of the problem to be solved are defined through a set of external subroutines that the user must provide. They must be external subroutines (i.e., not module or internal subroutines) because these are called from within PHAML and are compiled after the PHAML library has been built. They should contain a `use phaml` statement to get access to `my_real` and possibly other entities from PHAML.

This section describes the purpose of each of these routines and defines the interface of each. Also see `pde.f90` in each subdirectory of the PHAML `examples` directory for examples of these routines.

### 7.3.1 bconds

Subroutine `bconds` returns the boundary conditions at a given point. At each point, the boundary conditions are of the form

$$u = g$$

or

$$\left( c_{xx} \frac{\partial u}{\partial x} + c_{xy} \frac{\partial u}{\partial y} \right) \frac{\partial y}{\partial s} - c_{yy} \frac{\partial u}{\partial y} \frac{\partial x}{\partial s} + c_{bc} u = g$$

as described in Section 3.2.2.

```
subroutine bconds(x,y,bmark,itype,c,rs)
```

`real(my_real), intent(in) :: x,y` – the point at which to evaluate the boundary conditions.

`integer, intent(in) :: bmark` – the boundary marker assigned in the Triangle data files.

`integer, intent(out) :: itype(:)` – the type of boundary condition at this point, given by symbolic constants from module `phaml`. It must be one of `DIRICHLET` (the first form above), `NATURAL` (the second form with  $c$  identically 0.0), or `MIXED` (the second form with  $c$  nonzero). The dimension

of `itype` is `system_size`. The  $i^{th}$  value is the type of boundary condition for the  $i^{th}$  component.

`real(my_real), intent(out) :: c(:, :)` – the function  $c_{bc}$  in the boundary condition. The dimension is `system_size` by `system_size`. The  $(i, j)^{th}$  entry is the coefficient of the  $j^{th}$  component in the boundary condition for the  $i^{th}$  component.

`real(my_real), intent(out) :: rs(:)` – the function  $g$  in the boundary condition. The dimension is `system_size`. The  $i^{th}$  entry is the right side of the boundary condition for the  $i^{th}$  component.

### 7.3.2 boundary\_point

Subroutine `boundary_point` defines the boundary of the domain, if it is given by subroutines rather than triangle data files.

```
subroutine boundary_point(ipiece,s,x,y)
```

`integer, intent(in) :: ipiece` – the piece of the boundary from which to return a point.

`real(my_real), intent(in) :: s` – the parameter for the point to be determined.

`real(my_real), intent(out) :: x,y` – the point  $(x(s), y(s))$  on piece `ipiece` of the boundary.

### 7.3.3 boundary\_npiece

If the boundary of the domain is defined by the boundary subroutines, function `boundary_npiece` returns the number of boundary pieces in the definition. If the domain is defined by triangle data files, it returns 0 or a negative number. If the domain contains holes, it returns the number of pieces in the requested hole.

```
function boundary_npiece(hole)
```

`integer, intent(in) :: hole` – the hole for which to return the number of pieces, or 0 for the outer boundary. Holes are numbered consecutively starting with 1.

`integer :: boundary_npiece` – the number of boundary pieces in the requested hole or in the outer boundary if `hole` is 0.

### 7.3.4 boundary\_param

Subroutine `boundary_param` gives the range of parameter values for each boundary piece.

```
subroutine boundary_param(start,finish)
```

`real(my_real), intent(out) :: start(:), finish(:) – start(i) and finish(i) are the beginning and ending parameter values for the  $i^{th}$  piece of the boundary, i=1,number of boundary pieces.`

### 7.3.5 iconsds

Subroutine `iconsds` is used for setting the solution to a given function. For example, it can be used for initial conditions for a time dependent problem or the initial guess for a nonlinear problem.

`function iconsds(x,y,comp,eigen)`

`real(my_real), intent(in) :: x,y – the point at which to evaluate the function.`

`integer, intent(in) :: comp – for multicomponent solutions, which component to return.`

`integer, intent(in) :: eigen – for eigenvalue problems in which more than one eigenvalue is computed, which eigenfunction to return.`

`real(my_real) :: iconsds – the function value.`

### 7.3.6 pdecoefs

Subroutine `pdecoefs` returns the values of the PDE coefficient functions and right side. The PDE is given by

$$-\frac{\partial}{\partial x} \left( c_{xx} \frac{\partial u}{\partial x} \right) - \frac{\partial}{\partial x} \left( c_{xy} \frac{\partial u}{\partial y} \right) - \frac{\partial}{\partial y} \left( c_{yy} \frac{\partial u}{\partial y} \right) + c_x \frac{\partial u}{\partial x} + c_y \frac{\partial u}{\partial y} + c_u u = f \quad \text{in } \Omega$$

as described in Section 3.2.1.

`subroutine pdecoefs(x,y,cxx,cxy,cyy,cx,cy,c,rs)`

`real(my_real), intent(in) :: x,y – the point at which to evaluate the PDE.`

`real(my_real), intent(out) :: cxx(:, :), cxy(:, :), cyy(:, :), cx(:, :), cy(:, :), c(:, :)` – the functions  $c_{xx}$ ,  $c_{xy}$ ,  $c_{yy}$ ,  $c_x$ ,  $c_y$  and  $c_u$  respectively. The dimension is `system_size` by `system_size`. The  $(i, j)^{th}$  entry is the coefficient of the  $j^{th}$  component in the PDE for the  $i^{th}$  component.

`real(my_real), intent(out) :: rs(:)` – the function  $f$ . The dimension is `system_size`. The  $i^{th}$  value is the right side of the PDE for the  $i^{th}$  component.

### 7.3.7 phaml\_integral\_kernel

Subroutine `phaml_integrate` described in Sections 3.6.5 and 7.4.9 computes functionals of the computed solution of the form

$$\iint_{\Omega} k(x, y) u_i^p(x, y) u_j^q(x, y) dx dy$$

where  $k$  is a kernel function defined in `phaml_integral_kernel`. In addition to a point at which to evaluate the kernel, the function takes an integer, `kernel`, which allows for the definition of several kernel functions in the same subroutine.

```
function phaml_integral_kernel(kernel,x,y)
```

```
integer, intent(in) :: kernel – an integer passed through phaml_integrate
to allow a choice of different kernels.
```

```
real(my_real), intent(in) :: x,y – the point at which to evaluate the
kernel.
```

```
real(my_real) :: phaml_integral_kernel – the kernel value,  $k(x, y)$ .
```

### 7.3.8 regularity

One strategy for *hp*-adaptive refinement is to provide *a priori* knowledge about the singular nature of the solution. Then  $h$  refinement is done near singularities and  $p$  refinement is done where the solution is smooth. This strategy is selected with `HP_APRIORI`, and the function `regularity` provides the user's *a priori* knowledge. This function is also used by the Texas 3 Step strategy selected with `HP_T3S`.

In theory, this routine should return the largest value of  $m$  such that the solution is in  $H^m(T)$ , i.e. the derivatives up to order  $m$  are in  $L^2$ , where  $T$  is the triangle whose vertices are given as input to the function. For multicomponent solutions, it should return the worst (i.e. smallest) such  $m$  among the components.

In practice, it can be used to guide refinement in other *a priori* known trouble areas, such as sharp peaks, boundary layers and wave fronts. The actual use is that  $p$  refinement is performed if the current degree of the triangle is less than the returned value, and  $h$  refinement is performed otherwise. So, for example, if you know some region contains a boundary layer, you could return 3.1 for any triangle that intersects that region to perform  $h$ -refinement with cubic elements over the boundary layer.

```
function regularity(x,y)
```

```
real(my_real), intent(in) :: x(3),y(3) – the  $x$  and  $y$  coordinates of
the three vertices of the triangle.
```

```
real(my_real) :: regularity – the returned regularity.
```

### 7.3.9 trues

If the true solution of the PDE is known, you can provide it in function **trues**. This is used for printing norms of the error and for graphical display of the error and true solution. If you do not know the true solution, returning **huge(0.0\_my\_real)** will prevent printing and plotting of a bogus error.

```
function trues(x,y,comp,eigen)

real(my_real), intent(in) :: x,y – the point at which to evaluate the
    true solution

integer, intent(in) :: comp – for multicomponent solutions, which com-
    ponent to return.

integer, intent(in) :: eigen – for eigenvalue problems in which more
    than one eigenvalue is computed, which eigenfunction to return.

real(my_real) :: trues – the return value.
```

### 7.3.10 truexs

If the true solution of the PDE is known, you can provide the  $x$  derivative of it in function **truexs**. This is used for printing the energy norm of the error. If **trues** does not return **huge(0.0\_my\_real)** and you request the energy norm of the error, then you must provide the  $x$  derivative of the true solution in **truexs** or an incorrect value of the energy norm of the error will be printed.

```
function truexs(x,y,comp,eigen)

real(my_real), intent(in) :: x,y – the point at which to evaluate the
    derivative of the true solution

integer, intent(in) :: comp – for multicomponent solutions, which com-
    ponent to return.

integer, intent(in) :: eigen – for eigenvalue problems in which more
    than one eigenvalue is computed, which eigenfunction to return.

real(my_real) :: truexs – the return value.
```

### 7.3.11 trueys

If the true solution of the PDE is known, you can provide the  $y$  derivative of it in function **trueys**. This is used for printing the energy norm of the error. If **trues** does not return **huge(0.0\_my\_real)** and you request the energy norm of the error, then you must provide the  $y$  derivative of the true solution in **trueys** or an incorrect value of the energy norm of the error will be printed.

```
function trueys(x,y,comp,eigen)

real(my_real), intent(in) :: x,y – the point at which to evaluate the
    derivative of the true solution
```



`integer, intent(in) :: comp` – for multicomponent solutions, which component to return.

`integer, intent(in) :: eigen` – for eigenvalue problems in which more than one eigenvalue is computed, which eigenfunction to return.

`real(my_real) :: trueys` – the return value.

### 7.3.12 `update_usermod`

The application can include a module called `phaml_user_mod` to provide access to global variables in the user provided subroutines. However, values assigned to these variables in the main program are only available in the master process. Subroutine `update_usermod` passes the current value of these variables from the master to the slaves. For an example use of this facility, see `examples/parabolic` where it is used to pass the time step to the slaves. A working version of this subroutine is shown in Figure 7.1, or can be copied from `examples/parabolic/pde.f90`. In the example, the `integer` variable `ivar` and the `real(my_real)` variables `rvar1` and `rvar2` are declared in module `phaml_user_mod`. If you do not use this facility, you can copy a dummy version of the subroutine from `examples/simple/pde.f90`.

## 7.4 PHAML procedures

### 7.4.1 `phaml_compress`

`phaml_compress` compresses the `phaml_solution_type` data by moving all the unused elements, edges and vertices after the used ones. If considerable derefinement has occurred, this can significantly reduce the size of files created by `phaml_store`.

```
subroutine phaml_compress(phaml_solution)
  type(phaml_solution_type), intent(inout) :: phaml_solution – the solution to compress.
```

### 7.4.2 `phaml_connect`

`phaml_connect` connects two `phaml_solution_type` variables so they can communicate. If one will request that another evaluate its solution, then they must be connected. See `examples/system_ss` and Section 7.2.3. However, having multiple `phaml_solution_type` variables communicate with each other is very slow, so use of this capability is discouraged.

```
subroutine phaml_connect(pde1, pde2)
  integer, intent(in) :: pde1, pde2 – indices into the pde array in module phaml.
```

```

subroutine update_usermod(phaml_solution)

!-----
! This routine updates the module variables on the slave processes by
! sending them from the master process
!-----

use phaml
use phaml_user_mod

!-----
! Dummy arguments

type(phaml_solution_type), intent(in) :: phaml_solution

!-----
! Local variables:

! Declare these arrays big enough to hold the variables to be sent

integer :: iparam(1)
real(my_real) :: rparam(2)

!-----
! Begin executable code

! Copy the module variables into the arrays, putting integer variables
! into iparam and real variables into rparam.

    iparam(1) = ivar
    rparam(1) = rvar1
    rparam(2) = rvar2

! Call the routine that performs the actual exchange. Don't change this line.

    call master_to_slaves(phaml_solution,iparam,rparam)

! Copy the arrays into the module variables, using the same correspondence
! between module variable and array index as was used above.

    ivar = iparam(1)
    rvar1 = rparam(1)
    rvar2 = rparam(2)

end subroutine update_usermod

```

Figure 7.1: An example of subroutine update\_usermod.

### 7.4.3 phaml\_copy\_soln\_to\_old

`phaml_copy_soln_to_old` makes a copy of the solution in `phaml_solution` which can be evaluated by `phaml_evaluate_old` (Section 7.4.8). The main purpose of this is to provide the “old” solution and derivatives for time stepping in time dependent problems and iterating in nonlinear problems.

```
subroutine phaml_copy_soln_to_old(phaml_solution)
```

```
type(phaml_solution_type), intent(inout) :: phaml_solution – the phaml  
solution in which to make a copy of the solution.
```

### 7.4.4 phaml\_create

`phaml_create` creates a variable to contain a `phaml_solution`, i.e., it initializes the internals of a variable of type `phaml_solution_type`. It also spawns the associated slave and graphics processes. This should be called with each variable of type `phaml_solution_type` in your program.

All arguments except `phaml_solution` are optional.

```
subroutine phaml_create(phaml_solution, nproc, draw_grid_who, spawn_form,  
debug_command, display, graphics_host, output_unit, error_unit, output_now,  
id, system_size, eq_type, max_blen, triangle_files, update_umod)
```

```
type (phaml_solution_type) phaml_solution – the variable to initialize.
```

```
integer nproc – the number of processes working in parallel. The default is  
1.
```

`draw_grid_who` – which processes should display grid graphics. The grid graphics are not just the grid, but also the computed solution and many other useful displays. Valid values are:

**MASTER** – the graphics process associated with the master process draws a grid that is a consolidation of the slave’s grids.

**SLAVES** – each slave has an associated graphics process, which draws the grid as known to that slave. Note that there will be a graphics window for each slave.

**EVERYONE** – both MASTER and SLAVES.

**NO\_ONE** – no graphics are drawn.

The default is **NO\_ONE**.

`spawn_form` – whether or not to provide debugging capability with spawned processes. For MPI, An xterm running the slave under the debugger given by `debug_command` is brought up. For PVM the debugger is determined by the PVM environment (see the file doc/HINTS). Valid values are:

NORMAL\_SPAWN – no debugging.

DEBUG\_SLAVE – spawn the slave processes under the debugger.

DEBUG\_GRAPHICS – spawn the graphics processes under the debugger.

DEBUG\_BOTH – spawn both slaves and graphics under the debugger.

The default is NORMAL\_SPAWN.

**character(len=\*) debug\_command** – the command name of the debugger to use in an xterm when **spawn\_form** is not NORMAL\_SPAWN and MPI is used. Limited to 64 characters. The default is "gdb".

**character(len=\*) display** – in some environments, the **-display** argument is needed for xterm when **spawn\_form** is not NORMAL\_SPAWN. This value is used for that argument. If it is "default" then the **-display** argument is not used for xterm. Limited to 64 characters. The default is "default".

**character(len=\*) graphics\_host** – the name of the host on which to spawn the graphics processes. This is useful if you want to force the graphics processes to run on the workstation whose display is in front of you, or if you want graphics to run on a particular architecture. PVM will obey this request. For MPI, this is a “hint” which may be obeyed or ignored depending on the implementation of MPI. ignores the request.

**integer output\_unit** – the unit number for printed output. It should either be a pre-connected unit (e.g. standard output) or be opened as **FORMATTED** with a call to **phaml\_popen** immediately after **call phaml\_create**. The default is 6.

**integer error\_unit** – the unit number for error messages. It should either be a pre-connected unit (e.g. standard error) or be opened as **FORMATTED** with a call to **phaml\_popen** immediately after **call phaml\_create**. The default is 0.

**integer output\_now** – since **output\_unit** and **error\_unit** may not be available until after **phaml\_create** is complete, a unit must be provided for any output (i.e. error messages) that occurs from subroutine **phaml\_create**. This should be a pre-connected unit. The default is 6.

**integer id** – an identifier for the PDE, simply for use by the user in subroutines **pdecoefs**, **bconds**, etc. It is available as **my\_pde\_id** in module **phaml**. See Section 7.2.3. The default is 0.

**integer system\_size** – number of equations in a coupled system of PDEs, or equivalently, number of components in a multicomponent solution. The default is 1.

**integer eq\_type** – type of equation to create. Valid values are:

ELLIPTIC – solve an elliptic boundary value PDE.

**EIGENVALUE** – solve an elliptic eigenvalue PDE.

The default is **ELLIPTIC**.

**real(my\_real) max.blen** – maximum length of a boundary segment if the domain is defined by the boundary subroutines.

The default is  $\infty$ , defined to be **huge(0.0\_my\_real)**.

**character(len=\*) triangle\_files** – the root name of the **.node**, **.ele**, **.neigh**, **.poly** and **.edge** triangle data files. If the domain is defined by triangle data files and the file names include Triangle's iteration number (usually **.1**), include the iteration number. If the domain is defined by the boundary subroutines, this is used as the root of the created triangle data files.

The default is **"domain"**.

**logical update\_umod** – run **update\_usermod** as soon as the communication has been initialized. This is needed if **phaml\_user\_mod** has any parameters that affect the definition of the domain, or need to be set before the first solution on the initial grid.

The default is **.false.**

#### 7.4.5 phaml\_destroy

**phaml\_destroy** destroys a variable of type **phaml\_solution\_type**, i.e., frees the memory contained in it. It also terminates the slave and graphics processes. If you have multiple **phaml\_solution\_type** variables that are not needed simultaneously, you should destroy those no longer needed before creating those not yet used, to reduce the possibility of running out of memory. Also, if a **phaml\_solution\_type** variable is used more than once and is to be recreated to give it a fresh start, it should be destroyed before being created again to avoid a memory leak.

subroutine **phaml\_destroy**(**phaml\_solution**, **finalize\_mpi**)

type (**phaml\_solution\_type**) **phaml\_solution** – the variable to destroy

logical **finalize\_mpi** – if **.false.**, do not call **mpi\_finalize** from the master. It should be **.true.** if and only if this is the last call to **phaml\_destroy**.

The default is **.true.**

#### 7.4.6 phaml\_evaluate

**phaml\_evaluate** evaluates the computed solution and/or derivatives of it at the given points.

subroutine **phaml\_evaluate**(**phaml\_solution**, **x**, **y**, **u**, **ux**, **uy**, **uxx**, **uyy**, **comp**, **eigen**)

`type(phaml_solution_type), intent(in) :: phaml_solution` – the solution to evaluate.

`real(my_real), intent(in) :: x(:), y(:)` – array of x and y coordinates at which to evaluate the solution.

`real(my_real), intent(out), optional :: u(:), ux(:), uy(:), uxx(:), uyy(:)` – returned as the computed solution and derivatives at the given points. Those that are present determine what is evaluated. Must have `size(u) == size(x)`, etc.

`integer, intent(in), optional :: comp` – for multicomponent solutions, which component to evaluate. The default is 1.

`integer, intent(in), optional :: eigen` – for eigenvalue problems, which eigenfunction to evaluate. The default is 1.

#### 7.4.7 `phaml_get_grid_soln`

`phaml_get_grid_soln` returns the vertices of the grid and the solution at those vertices.

`subroutine phaml_get_grid_soln(phaml_solution, x, y, u, comp, eigen)`  
`type(phaml_solution_type), intent(in) :: phaml_solution` – the solution from which to get the grid and solution.

`real(my_real), pointer :: x(:), y(:), u(:)` – x and y are returned as the coordinates of the vertices of the triangles of the grid; u is the computed solution at those vertices. They should be disassociated on input and will be allocated to the correct size. You should deallocate them when done with them. To determine the number of points returned, use `size(x)`.

`integer, intent(in), optional :: comp` – for multicomponent solutions, which component to return. The default is 1.

`integer, intent(in), optional :: eigen` – for eigenvalue problems, which eigenfunction to return. The default is 1.

#### 7.4.8 `phaml_evaluate_old`

`phaml_evaluate_old` evaluates the “old” solution and/or derivatives at the given point(s). Unlike all the other `phaml` subroutines, it should not be called from the main program. It is intended to be called from the user routines that define the problem (`pdecoefs`, etc.) to provide the “old” solution and derivatives for time stepping in time dependent problems and iterating in nonlinear problems.

`subroutine phaml_evaluate_old(x, y, u, ux, uy, uxx, uyy, comp, eigen)`

`real(my_real), intent(in) :: x, y` – x and y coordinates of the point at which to evaluate the solution.

`real(my_real), intent(out), optional :: u, ux, uy, uxx, uyy` – returned as the old solution, x derivative, y derivative, second x derivative and second y derivative at the given point. Any combination of the five may be present.

`integer, intent(in), optional :: comp` – for multicomponent solutions, which component to evaluate. The default is 1.

`integer, intent(in), optional :: eigen` – for eigenvalue problems, which eigenfunction to evaluate. The default is 1.

#### 7.4.9 phaml\_integrate

`phaml_integrate` returns a functional of the computed solution, i.e. an integral of a computed solution or product of two computed solutions or powers of computed solutions, weighted by a kernel function. It computes

$$\iint_{\Omega} k(x, y) u_{comp1, eigen1}^p(x, y) u_{comp2, eigen2}^q(x, y) dx dy$$

where  $k$  is a kernel function defined in `phaml_integral_kernel` (see Section 7.3.7).

`function phaml_integrate(phaml_solution, kernel, comp1, eigen1, comp2, eigen2, p, q)`

`type(phaml_solution_type), intent(in) :: phaml_solution` – the solution to integrate.

`integer, intent(in) :: kernel` – allows you to select among different kernel functions. It is passed to `phaml_integral_kernel` where it can be used, for example, in a case statement to determine the kernel to use.

`integer, intent(in), optional :: comp1, eigen1, comp2, eigen2` – which component(s) of a multicomponent solution and which eigenfunction(s) of an eigenvalue problem to use. All of them default to 1. If `comp1` is omitted then `comp2` must also be omitted, and likewise for `eigen1` and `eigen2`. If `comp2` and `eigen2` are both omitted, then  $u_{comp2, eigen2}$  is omitted from the integral.

`integer, intent(in), optional :: p, q` – the powers to which to raise the computed solutions in the integral. The default is 1.

#### 7.4.10 phaml\_pclose

`phaml_pclose` closes unit number `unit` on all processors in `phaml_solution`. See also `phaml_popen` in Section 7.4.11.

```
subroutine phaml_pclose(phaml_solution, unit)

type (phaml_solution_type), intent(in) :: phaml_solution – the so-
    lution for which the files are being closed.

integer, intent(in) :: unit – the unit number to close.
```

#### 7.4.11 phaml\_popen

`phaml_popen` opens unit number `unit` on all processors in `phaml_solution`. If `file` is of the form “root.suffix” then the actual filenames are `rootXXXX.suffix` for processor number `XXXX`, where the number of digits in `XXXX` is the minimum needed for the number of processors in `phaml_solution`. The master processor is number 0. If there is no “.” in `file`, then there is no suffix and `XXXX` is appended to the filename. The filename is limited to 128 characters and the number of processors is limited to 9999. If the filename does not contain the full path, the location of the file is compiler dependent. Good places to look for it are the current working directory and the user’s home directory.

```
subroutine phaml_popen(phaml_solution, unit, file, form)

type (phaml_solution_type), intent(in) :: phaml_solution – the so-
    lution for which the files are being opened.

integer, intent(in) :: unit – unit number to open.

character(len=*), intent(in) :: file – base file name for the files.

character(len=*), intent(in), optional :: form – must take the value
    "FORMATTED" or "UNFORMATTED" to determine the type of file to open.
    Default is "FORMATTED".
```

#### 7.4.12 phaml\_query

`phaml_query` returns values of interest about the grid and solution. All arguments except `phaml_solution`, `comp`, `eigen` and `error_estimator` are optional and `intent(out)`. Include those you wish to receive as keyword arguments. `phaml_solution` is mandatory and `intent(in)`. `comp`, `eigen` and `error_estimator` are optional and `intent(in)`.

```
subroutine phaml_query(phaml_solution, nvert, nvert_proc, nvert_own,
nelem, nelem_proc, nelem_own, neq, neq_proc, neq_own, nlev, min_degree,
max_degree, linf_error, energy_error, l2_error, max_error_indicator,
```



linf\_error\_estimate, energy\_error\_estimate, l2\_error\_estimate, linf\_solution,  
l2\_solution, energy\_solution, linf\_u, l2\_u, energy\_u, linf\_true, l2\_true,  
energy\_true, eigenvalues, eigenvalue\_error\_estimate, max\_linsys\_resid,  
ave\_linsys\_resid, eigen\_l2\_resid, arpack\_iter, arpack\_nconv, arpack\_numop,  
arpack\_numopb, arpack\_numreo, arpack\_info, comp, eigen, error\_estimator)

type(phaml\_solution\_type) :: phaml\_solution – the solution to query

integer :: nvert – the number of vertices in the grid

integer, dimension(nproc) :: nvert\_proc – the number of vertices in the  
grid of each processor

integer, dimension(nproc) :: nvert\_own – the number of vertices owned  
by each processor

integer :: nelem – the number of elements in the grid

integer, dimension(nproc) :: nelem\_proc – the number of elements in  
the grid of each processor

integer, dimension(nproc) :: nelem\_own – the number of elements owned  
by each processor

integer :: neq – the number of equations in the linear system

integer, dimension(nproc) :: neq\_proc – the number of equations in the  
linear system of each processor

integer, dimension(nproc) :: neq\_own – the number of equations owned  
by each processor

integer :: nlev – the number of levels of refinement

integer :: min\_degree – the smallest polynomial degree of an element

integer :: max\_degree – the largest polynomial degree of an element

real(my\_real) :: linf\_error – the  $L^\infty$  norm of the error (at the vertices  
and quadrature points of a sixth order quadrature rule), if the true solution  
is known

real(my\_real) :: energy\_error – the energy norm of the error (approx-  
imated by a sixth order quadrature rule), if the true solution and its  
derivatives are known

real(my\_real) :: l2\_error – the  $L^2$  norm of the error (approximated by a  
sixth order quadrature rule), if the true solution is known

real(my\_real) :: max\_error\_indicator – the largest error indicator

`real(my_real) :: linf_error_estimate` – an estimate of the  $L^\infty$  norm of the error  
`real(my_real) :: energy_error_estimate` – an estimate of the energy norm of the error  
`real(my_real) :: l2_error_estimate` – an estimate of the  $L^2$  norm of the error  
`real(my_real) :: linf_solution` – the discrete  $\ell^\infty$  norm of the solution vector  
`real(my_real) :: l2_solution` – the discrete  $\ell^2$  norm of the solution vector  
`real(my_real) :: energy_solution` – the discrete energy norm of the solution vector  
`real(my_real) :: linf_u` – the continuous  $L^\infty$  norm of the solution, approximated using the vertices and the quadrature points of a sixth order quadrature rule  
`real(my_real) :: l2_u` – the continuous  $L^2$  norm of the solution, approximated with a sixth order quadrature rule  
`real(my_real) :: energy_u` – the continuous energy norm of the solution, approximated with a sixth order quadrature rule  
`real(my_real) :: linf_true` – the continuous  $L^\infty$  norm of the true solution, approximated using the vertices and the quadrature points of a sixth order quadrature rule  
`real(my_real) :: l2_true` – the continuous  $L^2$  norm of the true solution, approximated with a sixth order quadrature rule  
`real(my_real) :: energy_true` – the continuous energy norm of the true solution, approximated with a sixth order quadrature rule  
`real(my_real), dimension(num_eval) :: eigenvalues` – the computed eigenvalues, for eigenvalue problems  
`real(my_real), dimension(num_eval) :: eigenvalue_error_estimate` – an estimate of the error in each eigenvalue  
`real(my_real) :: max_linsys_resid` – for eigenvalue problems, the largest  $\ell^2$  norm of the residual of the linear systems solved (scaled by the norm of the right hand side)  
`real(my_real) :: ave_linsys_resid` – for eigenvalue problems, the average of the  $\ell^2$  norm of the residuals of the linear systems solved (scaled by the norm of the right hand side)

`real(my_real), dimension(num_eval) :: eigen_l2_resid` – the  $\ell^2$  norm of the residual of the eigensystem, for each eigenvalue  $\|Ax - \lambda Mx\|/\|\lambda Mx\|$

`integer :: arpack_iter` – from ARPACK, number of iterations used

`integer :: arpack_nconv` – from ARPACK, number of converged Ritz values

`integer :: arpack_numop` – from ARPACK, number of  $OP \cdot x$  operations

`integer :: arpack_numopb` – from ARPACK, number of  $B \cdot x$  operations

`integer :: arpack_numreo` – from ARPACK, number of reorthogonalizations

`integer :: arpack_info` – from ARPACK, info (error flag)

`integer :: comp` – for multicomponent solutions, which component to report. Energy norms cover all components in a single norm.  $L^\infty$  and  $L^2$  norms cover each component in individual norms. The default is 1.

`integer :: eigen` – for eigenvalue problems, which eigenfunction to report. The default is 1.

`integer :: error_estimator` – which error indicator to use for error estimates and error indicators. The default is `EXPLICIT_ERRIND`.

### 7.4.13 phaml\_restore

`phaml_restore` restores information for `phaml_solution` from files created by subroutine `phaml_store` (see Section 7.4.16).

```

subroutine phaml_restore(phaml_solution, unit, do_draw_grid, pause)

type (phaml_solution_type), intent(inout) :: phaml_solution – the
    solution into which the data will be read. If phaml_solution was pre-
    viously used, it should be destroyed first with phaml_destroy (see Section
    7.4.5) to avoid a memory leak. Whether or not it was previously used, it
    must be created with phaml_create (see Section 7.4.4) before passing it
    to phaml_restore.

integer, intent(in) :: unit – the unit number to read from, which should
    have been opened with phaml_popen using the same form as was used when
    phaml_store created the files.

logical, intent(in), optional :: do_draw_grid – whether or not to draw
    the solution immediately after reading it.

logical, intent(in), optional :: pause – if do_draw_grid is true, whether
    or not to pause after drawing the grid.
    
```

#### 7.4.14 phaml\_scale

phaml\_scale scales the computed solution by multiplying by factor.

```
subroutine phaml_scale(phaml_solution, factor, comp, eigen)

type(phaml_solution_type), intent(inout) :: phaml_solution – the so-
    lution to scale.

real(my_real), intent(in) :: factor – the factor by which to multiply.

integer, intent(in), optional :: comp – for multicomponent solutions,
    which component to scale. The default is 1.

integer, intent(in), optional :: eigen – for eigenvalue problems, which
    eigenfunction to scale. The default is 1.
```

#### 7.4.15 phaml\_solve\_pde

phaml\_solve\_pde solves the PDE. All arguments are optional, except phaml\_solution, so you only need to provide those for which you do not want to take the default. All arguments are intent(in) except phaml\_solution which is intent(inout) and item which is intent(out). It is recommended that the call use keyword arguments for all arguments except possibly phaml\_solution. Since there is such a large number of arguments to this routine, they are placed into subsections by category.

```
subroutine phaml_solve_pde(phaml_solution, item, max_elem, max_vert,
max_eq, max_lev, max_deg, stop_on_maxlev, stop_on_maxdeg, max_refsolveloop,
term_energy_err, term_Linf_err, term_L2_err, task, print_grid_when, print_grid_who,
print_error_when, print_error_who, print_error_what, print_errest_what,
print_linsys_when, print_linsys_who, print_solver_when, print_solver_who,
print_time_when, print_time_who, print_eval_when, print_eval_who, print_header_who,
print_trailer_who, print_warnings, clocks, draw_grid_when, pause_after_draw,
pause_after_phases, pause_at_start, pause_at_end, solve_init, sequential_vert,
inc_factor, error_estimator, errtype, reftype, refterm, reftol, hp_strategy,
t3s_gamma, t3s_eta, t3s_nunif, t3s_maxref, t3s_maxdeginc, tp_gamma, sp_gamma_h,
sp_gamma_p, nlp_max_h_dec, nlp_max_h_inc, nlp_max_p_dec, nlp_max_p_inc,
refsoln_pbias, derefine, partition_method, edge_rule, zoltan_param_file,
prebalance, postbalance, petsc_matrix_free, solver, preconditioner,
mg_cycles, mg_tol, mg_prerelax, mg_postrelax, mg_prerelax_ho, mg_postrelax_ho,
dd_iterations, krylov_iter, krylov_restart, krylov_tol, mg_comm, ignore_quad_err,
eigensolver, num_eval, lambda0, lambda0_side, transformation, scale_evec,
arpack_ncv, arpack_maxit, arpack_tol, blopex_maxit, blopex_atol, blopex_rtol,
degree, inc_quad_order, hypre_BoomerAMG_MaxLevels, hypre_BoomerAMG_MaxIter,
hypre_BoomerAMG_Tol, hypre_BoomerAMG_StrongThreshold, hypre_BoomerAMG_MaxRowSum,
hypre_BoomerAMG_CoarsenType, hypre_BoomerAMG_MeasureType, hypre_BoomerAMG_CycleType,
hypre_BoomerAMG_NumGridSweeps, hypre_BoomerAMG_GridRelaxType, hypre_BoomerAMG_GridRelaxPoints
```

hypre\_BoomerAMG\_RelaxWeight, hypre\_BoomerAMG\_DebugFlag, hypre\_ParaSails\_thresh,  
hypre\_ParaSails\_nlevels, hypre\_ParaSails\_filter, hypre\_ParaSails\_sym,  
hypre\_ParaSails\_loadbal, hypre\_ParaSails\_reuse, hypre\_ParaSails\_logging,  
hypre\_PCG\_Tol, hypre\_PCG\_MaxIter, hypre\_PCG\_TwoNorm, hypre\_PCG\_RelChange,  
hypre\_PCG\_Logging, hypre\_GMRES\_KDim, hypre\_GMRES\_Tol, hypre\_GMRES\_MaxIter,  
hypre\_GMRES\_Logging, petsc\_richardson\_damping\_factor, petsc\_chebychev\_emin,  
petsc\_chebychev\_emax, petsc\_gmres\_max\_steps, petsc\_rtol, petsc\_atol,  
petsc\_dtol, petsc\_maxits, petsc\_ilu\_levels, petsc\_icc\_levels, petsc\_ilu\_dt,  
petsc\_ilu\_dtcol, petsc\_ilu\_maxrowcount, petsc\_sor\_omega, petsc\_sor\_its,  
petsc\_sor\_lits, petsc\_eisenstat\_nodiagscaling, petsc\_eisenstat\_omega,  
petsc\_asm\_overlap, coarse\_size, coarse\_method)

#### 7.4.15.1 Miscellaneous arguments

`type (phaml_solution_type) phaml_solution` – the variable that contains the main data structures. It must be created by subroutine `phaml_create` before passing it to `phaml_solve_pde`.

`integer task` – what task to perform. Valid values are:

`BALANCE_REFINE_SOLVE` – go through a loop repeatedly doing load balance, refinement and solution phases.

`SET_INITIAL` – loop through the phases, but for the solution phase use interpolation of the function in `iconds`.

`BALANCE_ONLY` – just do one load balancing phase.

`REFINE_ONLY` – just do one refinement phase.

`SOLVE_ONLY` – just do one solution phase.

The default is `BALANCE_REFINE_SOLVE`.

`logical solve_init` – determines whether or not to compute the solution on the initial grid. Sometimes you might not want to if the grid comes from a previous call.

The default is `.true.`

#### 7.4.15.2 Termination arguments

`integer iterm` – termination code. If positive, a termination criterion was met. If negative, an error occurred. For the current meaning of the codes, see `global.f90` and look for the string `termination`.

`integer max_elem` – the maximum number of elements to use (termination criterion).

The default is  $\infty$ , defined to be `huge(0)`.

**integer max\_vert** – maximum number of vertices to use (termination criterion).

The default is  $\infty$ , defined to be `huge(0)`.

**integer max\_eq** – maximum number of equations in the linear system (a.k.a. degrees of freedom) to use (termination criterion).

The default is  $\infty$ , defined to be `huge(0)`.

**integer max\_lev** – the maximum number of  $h$  refinement levels to use. If an element is flagged for  $h$  refinement and the number of levels would exceed `max_lev`, then the element is quietly not refined, except for some  $hp$  strategies where it is  $p$ -refined instead. The number of levels is also limited by the size of the hash keys, which is set at the time the PHAML library is compiled (see Section 2.2.1) and the number of element in the initial grid. For 32 bit integers and a very coarse initial grid the maximum is about 25-30 for `PHAML_HASHSIZE=1` and 55-60 for `PHAML_HASHSIZE=2`. If  $h$  refinement would cause the hash to overflow, the element is quietly not refined, except for some  $hp$  strategies where it is  $p$ -refined instead.

The default is  $\infty$ , defined to be `huge(0)`.

**integer max\_deg** – the maximum polynomial degree for the approximation space. If an element is flagged for  $p$  refinement and the degree would exceed `max_deg`, then the element is quietly not refined, except for some  $hp$  strategies where it is  $h$ -refined instead.

In most cases the default is 22, which corresponds to the maximum degree for which the currently implemented quadrature rules will give the exact solution if the solution is a polynomial of degree `max_deg`. If `reftype` is `HP_ADAPTIVE` and `hp_strategy` is `HP_NEXT3P`, the default is 19. Otherwise, if `error_estimator` is `LOCAL_PROBLEM_P` or `EQUILIBRATED_RESIDUAL`, or `reftype` is `HP_ADAPTIVE` and `hp_strategy` is `HP_REFSOLN_ELEM` or `HP_REFSOLN_EDGE`, the default is 21.

**logical stop\_on\_maxlev** – if `.true.`, then `max_lev` is used as one of the termination criteria.

The default is `.false.`

**logical stop\_on\_maxdeg** – if `.true.`, then `max_deg` is used as one of the termination criteria.

The default is `.false.`

**integer max\_refsolveloop** – number of times to go through the refine/solve loop (termination criteria).

The default is  $\infty$ , defined to be `huge(0)`.

**real(my\_real) term\_energy\_err** – terminate when the energy error estimate is less than this value (termination criteria).

The default is 0.0.

`real(my_real) term_Linf_err` – terminate when the  $L^\infty$  error estimate is less than this value (termination criteria).

The default is 0.0.

`real(my_real) term_L2_err` – terminate when the  $L^2$  error estimate is less than this value (termination criteria).

The default is 0.0.

#### 7.4.15.3 Output control arguments

`integer print_grid_when` – how often to produce a printed summary of the grid (number of vertices and elements, etc.). Valid values are:

NEVER – don't print.

PHASES – after each refinement phase.

FINAL – only at the end.

FREQUENTLY – possibly more often than PHASES (used for debugging).

The default is NEVER.

`integer print_grid_who` – which processes should print the summary of the grid. Valid values are:

NO\_ONE – don't print.

SLAVES – slave processes print a summary of the grid as they know it.

MASTER – master process prints the composite grid.

EVERYONE – both SLAVES and MASTER.

MASTER\_ALL – master prints the individual grids of each slave process.

The default is NO\_ONE.

`integer print_linsys_when` – how often to produce a printed summary of the linear system (number of equations, sparsity, etc.). Valid values are:

NEVER – don't print.

PHASES – after each linear system **assembly**. solution phase.

FREQUENTLY – possibly more often than PHASES (used for debugging).

The default is NEVER.

`integer print_linsys_who` – which processes should print the summary of the linear system. Valid values are:

NO\_ONE – don't print.

SLAVES – slave processes print a summary of the system as they know it.

MASTER – master process prints the composite linear system.

EVERYONE – both SLAVES and MASTER.

MASTER\_ALL – master prints the individual systems of each slave process.

The default is NO\_ONE.

**integer print\_solver\_when** – how often to produce a printed summary of the solver performance (number of iterations, residual, etc.). Valid values are:

NEVER – don't print.

PHASES – after each linear system solution phase.

FREQUENTLY – possibly more often than PHASES (used for debugging).

The default is NEVER.

**integer print\_solver\_who** – which processes should print the summary of the solver performance. Valid values are:

NO\_ONE – don't print.

SLAVES – slave processes.

MASTER – master process.

EVERYONE – both SLAVES and MASTER.

MASTER\_ALL – master process.

The default is NO\_ONE.

**integer print\_error\_when** – how often to print the norms of the error (defined to be the difference between the computed solution and the function defined in function **true**) and error estimates. What norms of the error and error estimates are printed is determined by **print\_error\_what** and **print\_errest\_what**. It also prints the factor by which they have been reduced since the last time they were printed, and the effectivity index of the error estimate (ratio of the error estimate to the error) if both the error and error estimate are printed. If **true** returns **huge(0.0\_my\_real)** to indicate that the true solution is not known, then the error is not printed. If **truex** or **truey** return **huge(0.0\_my\_real)** and **true** does not, then the energy norm of the error (if requested) is printed as **huge(0.0\_my\_real)**. Valid values are:

NEVER – don't print.

PHASES – after each solution phase.

FINAL – only at the end.

FREQUENTLY – possibly more often than PHASES (used for debugging).

With the hierarchical basis multigrid solver, built in conjugate gradient and GMRES solvers, and the PETSc solvers, this causes an estimate of the  $\ell^2$  norm of the residual to be printed after each iteration. See **KSPDefaultMonitor** in the PETSc documentation. For **GMRES\_SOLVER** it is only printed at the restarts.



TOO\_MUCH – possibly more often than FREQUENTLY. Also this sets the solution to 0.0 before the solution phase as well as printing the residual after each iteration of some solvers as with FREQUENTLY.

The default is NEVER.

`integer print_error_who` – which processes should print the error. Individual processor energy norms of the error cannot be computed, so the energy norm of the error is printed only by the master. Valid values are:

NO\_ONE – don't print.

SLAVES – slave processes print the error over the grid as they know it.

MASTER – master process prints the error over the composite grid.

EVERYONE – both SLAVES and MASTER.

MASTER\_ALL – master prints the individual errors of each slave process.

The default is NO\_ONE.

`integer print_error_what` – what norms of the error to print, selected from energy,  $L^\infty$ , and  $L^2$ . The  $L^\infty$  norm of the error is an approximation given by the maximum error at the vertices and the quadrature points of a sixth order quadrature rule. The  $L^2$  norm and energy norm are approximated using a sixth order quadrature rule over the triangles of the grid. Valid values are:

NEVER – don't print any.

ENERGY\_ERR – energy norm.

LINF\_ERR –  $L^\infty$  norm.

L2\_ERR –  $L^2$  norm.

ENERGY\_LINF\_ERR – energy and  $L^\infty$  norms.

ENERGY\_L2\_ERR – energy and  $L^2$  norms.

LINF\_L2\_ERR –  $L^\infty$  and  $L^2$  norms.

ENERGY\_LINF\_L2\_ERR – all three norms.

The default is NEVER.

`integer print_errest_what` – what norms of the error estimate to print, selected from energy,  $L^\infty$ , and  $L^2$ . Valid values are:

NEVER – don't print any.

ENERGY\_ERREST – energy norm.

LINF\_ERREST –  $L^\infty$  norm.

L2\_ERREST –  $L^2$  norm.

ENERGY\_LINF\_ERREST – energy and  $L^\infty$  norms.

ENERGY\_L2\_ERREST – energy and  $L^2$  norms.

LINF\_L2\_ERREST –  $L^\infty$  and  $L^2$  norms.

ENERGY\_LINF\_L2\_ERREST – all three norms.

The default is NEVER.

**integer errtype** – type of error and error estimates to use in printed output.

Valid values are:

ABSOLUTE\_ERROR – absolute error.

RELATIVE\_ERROR – relative error. Norms of the error are scaled by the norm of the true solution. Error estimates are scaled by the norm of the computed solution.

The default is ABSOLUTE\_ERROR.

**integer print\_time\_when** – how often to print the amount of time used by the program. Valid values are:

NEVER – don't print.

PHASES – after each refinement/solve loop.

LAST – the time for the last cycle.

FINAL – only at the end.

LAST\_AND\_FINAL – the time of the last cycle, and the total time.

FREQUENTLY – possibly more often than PHASES (used for debugging).

The default is NEVER.

**integer print\_time\_who** – which processes should print the time. Valid values are:

NO\_ONE – don't print.

SLAVES – slave processes print their own times.

MASTER – master process prints maximum time over all slaves.

EVERYONE – both SLAVES and MASTER.

MASTER\_ALL – master prints the individual times of each slave process.

The default is NO\_ONE.

**integer clocks** – which clock(s) (cpu and/or wall) to use for timing. Valid values are:

CLOCK\_C – cpu clock.

CLOCK\_W – wall clock.

CLOCK\_CW – both cpu and wall clock.

The default is `CLOCK_W`.

`integer print_eval_when` – for eigenvalue problems, how often to print the eigenvalues. Valid values are:

`NEVER` – don't print.

`PHASES` – after each refinement/solve loop.

`FINAL` – only at the end.

The default is `NEVER`.

`integer print_eval_who` – for eigenvalue problems, which processes should print the eigenvalues. Valid values are:

`NO_ONE` – don't print.

`SLAVES` – slaves print the eigenvalues.

`MASTER` – master process prints the eigenvalues.

`EVERYONE` – both `SLAVES` and `MASTER`.

The default is `NO_ONE`.

`integer print_header_who` – which processes should print a header message and the values of the parameters when the subroutine starts. Valid values are:

`NO_ONE` – don't print.

`SLAVES` – slaves print the header.

`MASTER` – master prints the header.

`EVERYONE` – both `SLAVES` and `MASTER`.

The default is `MASTER`.

`integer print_trailer_who` – which processes should print a trailer message when the subroutine completes. Valid values are:

`NO_ONE` – don't print.

`SLAVES` – slaves print the trailer.

`MASTER` – master prints the trailer.

`EVERYONE` – both `SLAVES` and `MASTER`.

The default is `MASTER`.

`logical print_warnings` – if `.false.`, warning messages are not printed.

The default is `.true.`

`integer draw_grid_when` – how often to update the graphics. Valid values are:

NEVER – don't draw.

PHASES – after each refinement and solve phase.

FINAL – only at the end.

FREQUENTLY – possibly more often than PHASES (used for debugging).

The default is NEVER.

`logical pause_after_draw` – if `.true.`, the program will prompt for keyboard input after updating the graphics.

The default is `.false.`

`logical pause_after_phases` – if `.true.`, the program will prompt for keyboard input after each refinement/solve loop.

The default is `.false.`

`logical pause_at_start` – if `.true.`, the program will prompt for keyboard input before starting subroutine `phaml_solve_pde`.

The default is `.false.`

`logical pause_at_end` – if `.true.`, the program will prompt for keyboard input before returning from subroutine `phaml_solve_pde`.

The default is `.false.`

#### 7.4.15.4 Refinement arguments

`integer degree` – sets the initial degree of the polynomials in the finite element space. If no  $p$  refinement is performed, it is the fixed degree of the space.

The default is to use the existing degree in each element. In `phaml_create`, all elements are initialized to have degree 1.

`logical derefine` – if `.true.`, perform derefinement as well as refinement during adaptive refinement.

The default is `.true.`

`integer error_estimator` – select what to use for an error estimate to guide adaptive refinement and estimate global norms of the error. Valid values are:

EXPLICIT\_ERRIND – error indicator based on the norm of the residual and jump.

LOCAL\_PROBLEM\_H – computes an estimate over a pair of elements by performing one  $h$  refinement and solving a local Diriclet residual problem.

LOCAL\_PROBLEM\_P – computes an estimate over a triangle by performing one  $p$  refinement and solving a local Neumann residual problem.

**EQUILIBRATED\_RESIDUAL** – computes an estimate over a triangle by performing one  $p$  refinement and solving a local Neumann residual problem with an equilibrated residual.

**HIERARCHICAL\_COEFFICIENT** – use the coefficient of the  $h$ -hierarchical basis for linear elements, or the highest order  $p$ -hierarchical bases for high order elements. It is significantly faster than the local problem estimates, but of lower quality.

**TRUE\_DIFF** – use the difference between the true solution at the newest vertex of the element and surrounding vertices of the parent element. Can only be used if the true solution is known and supplied, and only for  $h$  refinement of linear elements.

**INITIAL\_CONDITION** – an error estimate based on interpolation of the function in subroutine **iconds** (Section 7.3.5). If **task** is **SET\_INITIAL** then it must be **INITIAL\_CONDITION**.

The default is **EXPLICIT\_ERRIND**, but it is always set to **INITIAL\_CONDITION** if **task** is **SET\_INITIAL**.

**integer reftype** – type of refinement to perform. Valid values are:

**H\_UNIFORM** – uniform  $h$  refinement.

**H\_ADAPTIVE** – adaptive  $h$  refinement.

**P\_UNIFORM** – uniform  $p$  refinement.

**P\_ADAPTIVE** – adaptive  $p$  refinement.

**HP\_ADAPTIVE** – adaptive  $h$  and  $p$  refinement.

The default is **H\_ADAPTIVE**.

**integer refterm** – termination criteria for a refinement phase (**DOUBLE** or **HALVE** really mean to multiply or divide by **inc\_factor**). Valid values are:

**DOUBLE\_NVERT** – double the number of vertices.

**DOUBLE\_NVERT\_SMOOTH** – double the number of vertices, and then refine all remaining elements in the same error estimate bin (helps to maintain symmetries in the grid).

**DOUBLE\_NELEM** – double the number of elements.

**DOUBLE\_NELEM\_SMOOTH** – double the number of elements, and then refine all remaining elements in the same error estimate bin.

**DOUBLE\_NEQ** – double the number of equations.

**DOUBLE\_NEQ\_SMOOTH** – double the number of equations, and then refine all remaining elements in the same error estimate bin.

**HALVE\_ERREST** – reduce the maximum error indicator by half.

KEEP\_NVERT – keep the same number of vertices or, if `max_vert` is present, try to make it close to `max_vert`. reduce number to `max_vert`, and adjust the grid through derefinement and refinement.

KEEP\_NVERT\_SMOOTH – keep the same number of vertices or `match` reduce number to `max_vert`, but then refine all remaining elements in the same error estimate bin.

KEEP\_NELEM – keep the same number of elements or, if `max_elem` is present, try to make it close to `max_elem`. reduce number to `max_elem`.

KEEP\_NELEM\_SMOOTH – keep the same number of elements or `match` reduce number to `max_elem`, but then refine all remaining elements in the same error estimate bin.

KEEP\_NEQ – keep the same number of equations or, if `max_eq` is present, try to make it close to `max_eq`. reduce number to `max_eq`.

KEEP\_NEQ\_SMOOTH – keep the same number of equations or `match` reduce number to `max_eq`, but then refine all remaining elements in the same error estimate bin.

KEEP\_ERREST – keep the same maximum error indicator. Currently not supported.

ONE\_REF – refine all elements with an error indicator larger than  $\text{reftol}/\sqrt{n}$  where  $n$  is the number of elements, but refine each element only once (just  $h$  or  $p$ , and don't refine children).

ONE\_REF\_HALF\_ERRIND – perform one refinement of all elements with an error indicator larger than half (or  $1/\text{inc\_factor}$ ) of the maximum error indicator.

The default is DOUBLE\_NEQ\_SMOOTH.

`real(my_real) inc_factor` – usually the factor by which to increase the size of the grid during one refinement phase. For *hp*-adaptive refinement with REFSOLN\_EDGE or REFSOLN\_ELEM, it is the factor by which the maximum error indicator is divided to get the error indicator cut-off that determines which elements are refined.

The default is usually 2.0. For *hp*-adaptive refinement with REFSOLN\_EDGE or REFSOLN\_ELEM it is 10.0.

`real(my_real) reftol` – tolerance for refining elements if `refterm` is ONE\_REF. If `refterm` is ONE\_REF, then at least one of `reftol`, or `term_energy_err` must be given.

The default is `term_energy_err/2`.

`integer hp_strategy` – select strategy for `reftype=HP_ADAPTIVE`. See Section 3.3.2. Valid values are:

HP\_APRIORI – use  $h$  refinement around *a priori* known trouble spots, and  $p$  refinement elsewhere. The regularity,  $m$ , of the solution is given by the user in function subroutine **regularity** (Sect. 7.3.8).  $h$  refinement is used if the degree of the element is larger than  $m$ .

HP\_NEXT3P –  $hp$  strategy in which the regularity is estimated using error estimators computed with spaces of degree  $p + 1$ ,  $p + 2$ , and  $p + 3$ .

HP\_PRIOR2P\_E –  $hp$  strategy in which the regularity is estimated using energy norm error estimators computed with spaces of degree  $p - 1$  and  $p - 2$ .

HP\_PRIOR2P\_H1 –  $hp$  strategy in which the regularity is estimated using  $H^1$  error estimators computed with spaces of degree  $p - 1$  and  $p - 2$ .

HP\_TYPEPARAM – the type parameter strategy.

HP\_BIGGER\_ERRIND – compute both LOCAL\_PROBLEM\_H and LOCAL\_PROBLEM\_P error indicators and refine an element by  $h$  or  $p$  depending on which is bigger.

HP\_T3S – the Texas 3 Step strategy.

HP\_ALTERNATE – alternate between  $h$  and  $p$  adaptive refinement, reducing the error estimate to a prescribed level at each step.

HP\_NLP – determine what refinements to do using nonlinear programming.

HP\_COEF\_DECAY – strategy based on the decay rate of the  $p$ -hierarchical coefficients.

HP\_COEF\_ROOT – strategy that estimates the regularity using a root test with the  $p$ -hierarchical coefficients.

HP\_SMOOTH\_PRED – a strategy that compares the local *a posteriori* error estimate with a predicted error estimate under the assumption that the solution is smooth.

HP\_REFSOLN\_ELEM – a strategy that considers several possible refinements of an element and picks one that reduces the difference from a reference solution the most.

HP\_REFSOLN\_EDGE – a strategy that considers several possible refinements of an edge and picks one that reduces the difference from a reference solution the most, and refines elements to be consistent with the chosen edge refinement.

The default is HP\_PRIOR2P\_H1.

`real(my_real) tp_gamma` – the type parameter  $\gamma$  for HP\_TYPEPARAM.

The default is 0.2.

`integer t3s_nunif` – number of uniform  $h$  refinements to start HP\_T3S.

The default is 0.

`real(my_real) t3s_gamma` – the parameter  $\gamma$  for HP\_T3S and HP\_ALTERNATE.  
The default is 6.0.

`real(my_real) t3s_eta` – the parameter  $\eta$  for HP\_T3S and HP\_ALTERNATE.  
The default is 0.1.

`integer t3s_maxref` – upper bound on the number of  $h$  refinements to perform in one step of HP\_T3S.  
The default is 3.

`integer t3s_maxdeginc` – upper bound on the number of  $p$  refinements to perform in one step of HP\_T3S.  
The default is 3.

`real(my_real) sp_gamma_h` – parameter  $\gamma_h$  for SMOOTH\_PRED.  
The default is 4.0.

`real(my_real) sp_gamma_p` – parameter  $\gamma_p$  for SMOOTH\_PRED.  
The default is 0.4.

`integer nlp_max_h_dec` – maximum number of  $h$  coarsenings at one time in HP\_NLP.  
The default is 1.

`integer nlp_max_p_dec` – maximum number of  $p$  coarsenings at one time in HP\_NLP.  
The default is 1.

`integer nlp_max_h_inc` – maximum number of  $h$  refinements at one time in HP\_NLP.  
The default is 5.

`integer nlp_max_p_inc` – maximum number of  $p$  refinements at one time in HP\_NLP.  
The default is 2.

`real(my_real) refsoln_pbias` – bias the HP\_REFSOLN\_ELEM strategy toward  $p$ -refinement. Recommended values are 2.0 for singular or nearly singular solutions and 4.0 for smooth solutions or moderately strong features. 1.0 is no bias.  
The default is 4.0.

`integer edge_rule` – the rule for determining how to assign the polynomial degree for edges. Valid values are:

MINIMUM\_RULE – use the minimum of the degrees of the triangles sharing the edge.



MAXIMUM\_RULE – use the maximum of the degrees of the triangles sharing the edge.

The default is MINIMUM\_RULE.

#### 7.4.15.5 Load balancing arguments

integer sequential\_vert – number of vertices in the grid before it starts running in parallel.

The default is 100.

integer prebalance – what to balance when partitioning the grid before refinement. Valid values are:

BALANCE\_NONE – do not partition the grid before refinement.

BALANCE\_ELEMENTS – equal number of elements in each partition.

BALANCE\_VERTICES – equal number of vertices in each partition.

BALANCE\_EQUATIONS – equal number of equations in each partition.

The default is BALANCE\_ELEMENTS.

integer postbalance – what to balance when partitioning the grid after refinement. Valid values are:

BALANCE\_NONE – do not partition the grid after refinement.

BALANCE\_ELEMENTS – equal number of elements in each partition.

BALANCE\_VERTICES – equal number of vertices in each partition.

BALANCE\_EQUATIONS – equal number of equations in each partition.

The default is BALANCE\_NONE.

integer partition\_method – what method to use for partitioning the grid. Valid values are:

RTK – the k-way refinement tree method implemented in PHAML.

ZOLTAN\_RCB – recursive coordinate bisection, from Zoltan.

ZOLTAN\_OCT – RPI's Octree method, from Zoltan.

ZOLTAN\_METIS – the local diffusion method from ParMETIS via Zoltan.

ZOLTAN\_REFTREE – the refinement tree method, from Zoltan.

ZOLTAN\_RIB – recursive inertial bisection, from Zoltan.

ZOLTAN\_HSFC – Hilbert space filling curve, from Zoltan.

ZOLTAN\_FILE – give Zoltan method and parameters in a file.

The default is RTK.

character(len=\*) zoltan\_param\_file – name of the file containing parameters for ZOLTAN\_FILE. The default is zoltan.params.

#### 7.4.15.6 Assembly arguments

`integer inc_quad_order` – increment the order of the quadrature rules by this amount.

The default is 0.

`logical ignore_quad_err` – if `.true.`, when setting up the linear system, ignore the quadrature errors in the large triangles outside the owned region. This is acceptable (and reduces computation and communication) when the PDE coefficients and right hand side are constant (for example, Laplace’s equation), or when using a solver other than HBMG, but will reduce the convergence of the discretization error otherwise.

The default is `.false.` if the solver is HBMG and `.true.` otherwise.

#### 7.4.15.7 Solver arguments

`integer solver` – what method to use as the solver. Valid values are:

`MG_SOLVER` – the hierarchical basis multigrid method implemented in PHAML.

`CG_SOLVER` – conjugate gradient solver implemented in PHAML.

`GMRES_SOLVER` – GMRES solver implemented in PHAML.

`PETSC_RICHARDSON_SOLVER` – Richardson solver from PETSc.

`PETSC_CHEBYCHEV_SOLVER` – Chebychev solver from PETSc.

`PETSC_CG_SOLVER` – Conjugate Gradients from PETSc.

`PETSC_GMRES_SOLVER` – Generalized Minimal Residual (GMRES) from PETSc.

`PETSC_TCQMR_SOLVER` – Transpose-Free Quasi-Minimal Residual (QMR) from PETSc.

`PETSC_BCGS_SOLVER` – BiConjugate Gradients Stabilized (BiCGSTAB) from PETSc.

`PETSC_CGS_SOLVER` – Conjugate Gradient Squared from PETSc.

`PETSC_TFQMR_SOLVER` – Transpose-Free Quasi-Minimal Residual (QMR) from PETSc.

`PETSC_CR_SOLVER` – Conjugate Residual from PETSc.

`PETSC_LSQR_SOLVER` – Least Squares from PETSc.

`PETSC_BICG_SOLVER` – BiConjugate Gradients from PETSc.

`HYPRE_BOOMERAMG_SOLVER` – BoomerAMG algebraic multigrid from *hypre*.

`HYPRE_PCG_SOLVER` – Preconditioned Conjugate Gradients from *hypre*.

`HYPRE_GMRES_SOLVER` – GMRES from *hypre*.

MUMPS\_SPD\_SOLVER – Symmetric positive definite solver from MUMPS (a parallel sparse direct solver). Only double precision is supported, i.e. `my_real=kind(0.0d0)` in `global.f90`.

MUMPS\_GEN\_SOLVER – General symmetric solver from MUMPS, for when the matrix might not be positive definite, such as large negative  $c_u$ . Only double precision is supported, i.e. `my_real=kind(0.0d0)` in `global.f90`.

MUMPS\_NONSYM\_SOLVER – Nonsymmetric solver from MUMPS, for multi-component problems with nonsymmetric coupling, and problems in which  $c_{xy}$ ,  $c_x$ , or  $c_y$  is nonzero. Only double precision is supported, i.e. `my_real=kind(0.0d0)` in `global.f90`.

SUPERLU\_SOLVER – parallel sparse direct solver SuperLU.

LAPACK\_INDEFINITE\_SOLVER – the indefinite solver from LAPACK. This is available for certain debugging and testing operations, and can only be used for very small problem sizes and only with one processor.

LAPACK\_SPD\_SOLVER – the LAPACK solver for symmetric positive definite band matrices. This is available for certain debugging and testing operations, and can only be used for very small problem sizes and only with one processor.

Note: when the number of vertices is less than `sequential_vert`, the *hypre* and SuperLU solvers are replaced by a LAPACK solver for the whole matrix on each processor.)

The default is `MG_SOLVER`.

`integer preconditioner` – what method to use as a preconditioner for the Krylov methods. Valid values are:

`NO_PRECONDITION` – no preconditioning.

`MG_PRECONDITION` – a hierarchical basis multigrid V-cycle preconditioner.

`FUDOP_DD_PRECONDITION` – a domain decomposition preconditioner with FuDoP.

`COARSE_GRID_PRECONDITION` – exact solver on a coarse grid as preconditioner.

`PETSC_JACOBI_PRECONDITION` – Jacobi preconditioner from PETSc.

`PETSC_BJACOBI_PRECONDITION` – Block Jacobi preconditioner from PETSc.

`PETSC_SOR_PRECONDITION` – SOR (and SSOR) preconditioner from PETSc.

`PETSC_EISENSTAT_PRECONDITION` – SOR with Eisenstat trick from PETSc.

`PETSC_ICC_PRECONDITION` – Incomplete Cholesky preconditioner from PETSc.

`PETSC_ILU_PRECONDITION` – Incomplete LU preconditioner from PETSc.

`PETSC_ASM_PRECONDITION` – Additive Schwarz preconditioner from PETSc.

HYPRE\_DS\_PRECONDITION – Diagonal scaling preconditioner from *hypre*.  
HYPRE\_BOOMERAMG\_PRECONDITION – BoomerAMG algebraic multigrid from *hypre*.  
HYPRE\_PARASAILS\_PRECONDITION – ParaSails sparse approximate inverse from *hypre*.

Note: For `solver=CG_SOLVER` and `solver=GMRES_SOLVER`, the preconditioner must be one of `NO_PRECONDITION` or `MG_PRECONDITION`. For `solver=HYPRE_PCG_SOLVER`, the preconditioner must be one of `NO_PRECONDITION`, `HYPRE_DS_PRECONDITION`, or `HYPRE_BOOMERAMG_PRECONDITION`. For `solver=HYPRE_GMRES_SOLVER`, the preconditioner must be one of `NO_PRECONDITION`, `HYPRE_DS_PRECONDITION`, `HYPRE_BOOMERAMG_PRECONDITION`, or `HYPRE_PARASAILS_PRECONDITION`. The *hypre* preconditioners can only be used with the *hypre* PCG and GMRES solvers.

The default is:

`NO_PRECONDITION` – when not applicable  
`HYPRE_BOOMERAMG_PRECONDITION` – for the *hypre* solvers  
`MG_PRECONDITION` – otherwise

`integer mg_cycles` – number of multigrid V-cycles to use in one solution phase or as a preconditioner.

If HBMG is the solver, then the default is 1 if `mg_tol` is `MG_NO_TOL` and infinite otherwise. If HBMG is the preconditioner, then the default is 2.

`real(my_real) mg_tol` – perform multigrid cycles until the  $\ell^2$  norm of the scaled linear system residual is less than `mg_tol` (up to a maximum of `mg_cycles` cycles). In addition to positive real numbers, it can have the following values:

`MG_NO_TOL` – do not use the tolerance test for ending the multigrid cycles, use a fixed number of cycles given by `mg_cycles` instead.  
`MG_ERREST_TOL` – reduce the residual to some fraction of the error estimate, currently  $1/1000$   $1/100$ .

The default is `MG_ERREST_TOL` if HBMG is the solver, and `MG_NO_TOL` if HBMG is the preconditioner.

`integer mg_prerelax` – number of half red-black relaxation sweeps to make before coarse grid correction in the *h*-hierarchical linear bases.

The default is 1.

`integer mg_postrelax` – number of half red-black relaxation sweeps to make after coarse grid correction in the *h*-hierarchical linear bases.

The default is 2.

**integer mg\_prerelax\_ho** – number of relaxation sweeps to make on each level of the high order  $p$ -multigrid cycle before coarse grid correction.

The default is 1.

**integer mg\_postrelax\_ho** – number of relaxation sweeps to make on each level of the high order  $p$ -multigrid cycle after coarse grid correction.

The default is 1.

**integer mg\_comm** – selects how much communication to do in the parallel hierarchical basis multigrid method. Permitted values are

**MGCOMM\_FUDOP** – the full domain partition approach with two messages per cycle.

**MGCOMM\_CONVENTIONAL** – conventional communication at each level.

**MGCOMM\_NONE** – no communication. Should only be used as a preconditioner.

The default is **MGCOMM\_FUDOP** if **HBMG** is the solver and **MGCOMM\_NONE** if **HBMG** is the preconditioner.

**integer krylov\_iter** – the maximum number of iterations to use with the native Krylov space solvers **CG\_SOLVER** and **GMRES\_SOLVER**.

The default is 100.

**integer krylov\_restart** – the GMRES restart parameter for **GMRES\_SOLVER**.

The default is 20.

**real(my\_real) krylov\_tol** – convergence tolerance on the  $\ell^2$  norm of the linear system residual for the native Krylov space solvers. In addition to positive real numbers, it can have the following values:

**KRYLOV\_ERREST\_TOL** – reduce the residual to some fraction of the error estimate, currently 1/100.

The default is **KRYLOV\_ERREST\_TOL**.

**integer dd\_iterations** – number of iterations to use for the FuDoP domain decomposition preconditioner.

The default is 1.

**integer coarse\_size** – for the coarse grid preconditioner, maximum size of the coarsened grid.

The default is 5000.

**integer coarse\_method** – for the coarse grid preconditioner, the method to use to solve the coarse grid problem. Permitted values are

**LAPACK\_INDEFINITE\_SOLVER** – the indefinite solver from LAPACK.

MUMPS\_GEN\_SOLVER – general symmetric solver from MUMPS.

SUPERLU\_SOLVER – parallel sparse direct solver SuperLU.

The default is LAPACK\_INDEFINITE\_SOLVER.

logical `petsc_matrix_free` – if the solver is one of the PETSc methods and this parameter is `.true.`, memory is saved by using a matrix-free approach that does not copy the matrix to a PETSc data structure. If `.true.`, then you cannot use the PETSc preconditioners.

The default is `.false.`

The following parameters are passed to subroutines in PETSc. See the documentation for PETSc for explanations and default values.

real(my_real)	<code>petsc_richardson_damping_factor</code>
real(my_real)	<code>petsc_chebychev_emin</code>
real(my_real)	<code>petsc_chebychev_emax</code>
integer	<code>petsc_gmres_max_steps</code>
real(my_real)	<code>petsc_rtol</code>
real(my_real)	<code>petsc_atol</code>
real(my_real)	<code>petsc_dtol</code>
integer	<code>petsc_maxits</code>
integer	<code>petsc_ilu_levels</code>
integer	<code>petsc_icc_levels</code>
real(my_real)	<code>petsc_ilu_dt</code>
real(my_real)	<code>petsc_ilu_dtcol</code>
integer	<code>petsc_ilu_maxrowcount</code>
real(my_real)	<code>petsc_sor_omega</code>
integer	<code>petsc_sor_its</code>
integer	<code>petsc_sor_lits</code>
logical	<code>petsc_eisenstat_nodiagscaling</code>
real(my_real)	<code>petsc_eisenstat_omega</code>
integer	<code>petsc_asm_overlap</code>

The following parameters are passed to subroutines in *hypr*. See the documentation for *hypr* for explanations and default values.

integer	hypre_BoomerAMG_MaxLevels
integer	hypre_BoomerAMG_MaxIter
real(my_real)	hypre_BoomerAMG_Tol
real(my_real)	hypre_BoomerAMG_StrongThreshold
real(my_real)	hypre_BoomerAMG_MaxRowSum
integer	hypre_BoomerAMG_CoarsenType
integer	hypre_BoomerAMG_MeasureType
integer	hypre_BoomerAMG_CycleType
integer	hypre_BoomerAMG_NumGridSweeps(:)
integer	hypre_BoomerAMG_GridRelaxType(:)
integer	hypre_BoomerAMG_GridRelaxPoints(:, :)
real(my_real)	hypre_BoomerAMG_RelaxWeight(:)
integer	hypre_BoomerAMG_DebugFlag
real(my_real)	hypre_ParaSails_thresh
integer	hypre_ParaSails_nlevels
real(my_real)	hypre_ParaSails_filter
integer	hypre_ParaSails_sym
real(my_real)	hypre_ParaSails_loadbal
integer	hypre_ParaSails_reuse
integer	hypre_ParaSails_logging
real(my_real)	hypre_PCG_Tol
integer	hypre_PCG_MaxIter
integer	hypre_PCG_TwoNorm
integer	hypre_PCG_RelChange
integer	hypre_PCG_Logging
integer	hypre_GMRES_KDim
real(my_real)	hypre_GMRES_Tol
integer	hypre_GMRES_MaxIter
integer	hypre_GMRES_Logging

#### 7.4.15.8 Eigenvalue arguments

`integer eigensolver` – For eigenvalue problems, what software package to use to solve the discrete eigenproblem. Valid values are:

ARPACK\_SOLVER – ARPACK

BLOPEX\_SOLVER – BLOPEX

The default is ARPACK\_SOLVER.

`integer num_eval` – For eigenvalue problems, the number of eigenvalues to find.

The default is 1.

`real(my_real) lambda0` – For eigenvalue problems, find the eigenvalues closest to lambda0.

The default is  $-\infty$  defined as `-huge(0.0_my_real)`, i.e., find the smallest eigenvalues.

**integer lambda0\_side** – Which side of `lambda0` to compute eigenvalues on.  
Valid values are:

**EIGEN\_LEFT** – eigenvalues less than `lambda0`

**EIGEN\_RIGHT** – eigenvalues greater than `lambda0`

**EIGEN\_BOTH** – eigenvalues on both sides of `lambda0`

If `eigensolver` is **BLOPEX\_SOLVER** then 1) if `transformation` is **SHIFT\_INVERT** it must not be **EIGEN\_BOTH**, and 2) if `transformation` is **SHIFT\_SQUARE** it must be **EIGEN\_BOTH**.

The default is **EIGEN\_RIGHT** if `eigensolver` is **BLOPEX\_SOLVER** and `transformation` is **SHIFT\_INVERT**, and **EIGEN\_BOTH** otherwise.

**integer transformation** – What spectral transformation to use for interior eigenvalues. Valid values are:

**SHIFT\_INVERT** – shift and invert

**SHIFT\_SQUARE** – shift and square

If `eigensolver` is **ARPACK\_SOLVER** it must not be **SHIFT\_SQUARE**.

The default is **SHIFT\_INVERT**.

**integer scale\_evec** – For eigenvalue problems, the eigenvectors can be multiplied (scaled) by an arbitrary constant. These are the choices for scaling:

**SCALE\_LINF** – scale so the (discrete)  $\ell^\infty$  norm is 1. For linear elements, this is also the  $L^\infty$  norm.

**SCALE\_L2** – scale so the (discrete)  $\ell^2$  norm is 1.

**SCALE\_M** – scale so the  $M$  norm,  $\sqrt{x^T M x}$ , is 1, where  $M$  is the mass matrix. The  $M$  norm is also the (continuous)  $L^2$  norm.

The default is **SCALE\_LINF**.

The following parameters are passed to ARPACK. See the ARPACK documentation for more information on them.

**integer arpack\_ncv** – number of Lanczos basis vectors to use

The default is 20.

**integer arpack\_maxit** – maximum number of IRLM iterations

The default is 100.

**real(my\_real) arpack\_tol** – relative accuracy to which eigenvalues are computed

The default is  $10^{-10}$ .



The following parameters are passed to BLOPEX.

`integer blopex_maxit` – maximum number of iterations

The default is 100.

`real(my_real) blopex_atol` – tolerance on the absolute residual

The default is  $10^{-6}$ .

`real(my_real) blopex_rtol` – relative tolerance

The default is  $10^{-6}$ .

#### 7.4.16 `phaml_store`

`phaml_store` stores information from `phaml_solution` into files for later use.

```
subroutine phaml_store(phaml_solution, unit)
```

`type (phaml_solution_type), intent(in) :: phaml_solution` – the solution to store.

`integer, intent(in) :: unit` – the unit number to write to, which should have been opened as either FORMATTED or UNFORMATTED with subroutine `phaml_popen`. UNFORMATTED is likely to be faster and create smaller data files. FORMATTED may be required if you will be restoring with a different compiler or architecture, and might not reproduce floating point numbers exactly.

#### 7.4.17 `phaml_store_matrix`

`phaml_store_matrix` stores the stiffness matrix, right hand side, and/or mass matrix in a file in Matrix Market exchange format (<http://math.nist.gov/MatrixMarket/formats.html>).

```
subroutine phaml_store_matrix(phaml_solution, stiffness_unit, rhs_unit,  
mass_unit, inc_quad_order)
```

`type (phaml_solution_type), intent(in) :: phaml_solution` – the solution containing the linear system to store.

`integer, intent(in), optional :: stiffness_unit, rhs_unit, mass_unit` – the I/O unit numbers for writing the stiffness matrix, right hand side, and mass matrix, respectively. They should be open for formatted sequential writing prior to calling `phaml_store_matrix` and closed afterwards. Which ones are present determines which are written.

`integer, intent(in), optional :: inc_quad_order` – increase the order of the quadrature rule by this amount. Default is 0.

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