# hypre Documentation Release 2.31.0 

## Lawrence Livermore National Laboratory

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## INTRODUCTION

This manual describes hypre, a software library of high performance preconditioners and solvers for the solution of large, sparse linear systems of equations on massively parallel computers [FaJY2004]. The hypre library was created with the primary goal of providing users with advanced parallel preconditioners. The library features parallel multigrid solvers for both structured and unstructured grid problems. For ease of use, these solvers are accessed from the application code via hypre's conceptual linear system interfaces [FaJY2005] (abbreviated to conceptual interfaces throughout much of this manual), which allow a variety of natural problem descriptions.

This introductory chapter provides an overview of the various features in hypre, discusses further sources of information on hypre, and offers suggestions on how to get started.

### 1.1 Overview of Features

Scalable preconditioners provide efficient solution on today's and tomorrow's systems: hypre contains several families of preconditioner algorithms focused on the scalable solution of very large sparse linear systems. (Note that small linear systems, systems that are solvable on a sequential computer, and dense systems are all better addressed by other libraries that are designed specifically for them.) hypre includes "grey box" algorithms that use more than just the matrix to solve certain classes of problems more efficiently than general-purpose libraries. This includes algorithms such as structured multigrid.

Suite of common iterative methods provides options for a spectrum of problems: hypre provides several of the most commonly used Krylov-based iterative methods to be used in conjunction with its scalable preconditioners. This includes methods for nonsymmetric systems such as GMRES and methods for symmetric matrices such as Conjugate Gradient.

Intuitive grid-centric interfaces obviate need for complicated data structures and provide access to advanced solvers: hypre has made a major step forward in usability from earlier generations of sparse linear solver libraries in that users do not have to learn complicated sparse matrix data structures. Instead, hypre does the work of building these data structures for the user through a variety of conceptual interfaces, each appropriate to different classes of users. These include stencil-based structured/semi-structured interfaces most appropriate for finite-difference applications; a finite-element based unstructured interface; and a linear-algebra based interface. Each conceptual interface provides access to several solvers without the need to write new interface code.

User options accommodate beginners through experts: hypre allows a spectrum of expertise to be applied by users. The beginning user can get up and running with a minimal amount of effort. More expert users can take further control of the solution process through various parameters.

Configuration options to suit your computing system: hypre allows a simple and flexible installation on a wide variety of computing systems. Users can tailor the installation to match their computing system. Options include debug and optimized modes, the ability to change required libraries such as MPI and BLAS, a sequential mode, and modes enabling threads for certain solvers. On most systems, however, hypre can be built by simply typing configure followed by make, or by using CMake [CMakeWeb].

Interfaces in multiple languages provide greater flexibility for applications: hypre is written in C (with the exception of the FEI interface, which is written in $\mathrm{C}++$ ) and provides an interface for Fortran users.

### 1.2 Getting More Information

This user's manual consists of chapters describing each conceptual interface, a chapter detailing the various linear solver options available, detailed installation information, and the API reference. In addition to this manual, a number of other information sources for hypre are available.

- Reference Manual: This is equivalent to Chapter API in this user manual, but it can also be built as a separate document. The reference manual comprehensively lists all of the interface and solver functions available in hypre. It is ideal for determining the various options available for a particular solver or for viewing the functions provided to describe a problem for a particular interface.
- Example Problems: A suite of example problems is provided with the hypre installation. These examples reside in the examples subdirectory and demonstrate various features of the hypre library. Associated documentation may be accessed by viewing the README. html file in that same directory.
- Papers, Presentations, etc.: Articles and presentations related to the hypre software library and the solvers available in the library are available from the hypre web page at http://www.llnl.gov/CASC/hypre/.
- Mailing List: The mailing list hypre-announce can be subscribed to through the hypre web page at http: //www.llnl.gov/CASC/hypre/. The development team uses this list to announce new releases of hypre. It cannot be posted to by users.


### 1.3 How to get started

### 1.3.1 Installing hypre

As previously noted, on most systems hypre can be built by simply typing configure followed by make in the toplevel source directory. Alternatively, the CMake system [CMakeWeb] can be used, and is the best approach for building hypre on Windows systems in particular. For more detailed instructions, read the INSTALL file provided with the hypre distribution or the General Information section of this manual. Note the following requirements:

- To run in parallel, hypre requires an installation of MPI.
- Configuration of hypre with threads requires an implementation of OpenMP. Currently, only a subset of hypre is threaded.
- The hypre library currently does not directly support complex-valued systems.


### 1.3.2 Choosing a conceptual interface

An important decision to make before writing any code is to choose an appropriate conceptual interface. These conceptual interfaces are intended to represent the way that applications developers naturally think of their linear problem and to provide natural interfaces for them to pass the data that defines their linear system into hypre. Essentially, these conceptual interfaces can be considered convenient utilities for helping a user build a matrix data structure for hypre solvers and preconditioners. The top row of Figure 1 illustrates a number of conceptual interfaces. Generally, the conceptual interfaces are denoted by different types of computational grids, but other application features might also be used, such as geometrical information. For example, applications that use structured grids (such as in the left-most interface in Figure 1) typically view their linear problems in terms of stencils and grids. On the other hand, applications that use unstructured grids and finite elements typically view their linear problems in terms of elements and element stiffness
matrices. Finally, the right-most interface is the standard linear-algebraic (matrix rows/columns) way of viewing the linear problem.
The hypre library currently supports four conceptual interfaces, and typically the appropriate choice for a given problem is fairly obvious, e.g. a structured-grid interface is clearly inappropriate for an unstructured-grid application.

- Structured-Grid System Interface (Struct): This interface is appropriate for applications whose grids consist of unions of logically rectangular grids with a fixed stencil pattern of nonzeros at each grid point. This interface supports only a single unknown per grid point. See Chapter Structured-Grid System Interface (Struct) for details.
- Semi-Structured-Grid System Interface (SStruct): This interface is appropriate for applications whose grids are mostly structured, but with some unstructured features. Examples include block-structured grids, composite grids in structured adaptive mesh refinement (AMR) applications, and overset grids. This interface supports multiple unknowns per cell. See Chapter Semi-Structured-Grid System Interface (SStruct) for details.
- Finite Element Interface (FEI): This is appropriate for users who form their linear systems from a finite element discretization. The interface mirrors typical finite element data structures, including element stiffness matrices. Though this interface is provided in hypre, its definition was determined elsewhere (please send email to Alan Williams william@sandia.gov for more information). See Chapter Finite Element Interface for details.
- Linear-Algebraic System Interface (IJ): This is the traditional linear-algebraic interface. It can be used as a last resort by users for whom the other grid-based interfaces are not appropriate. It requires more work on the user's part, though still less than building parallel sparse data structures. General solvers and preconditioners are available through this interface, but not specialized solvers which need more information. Our experience is that users with legacy codes, in which they already have code for building matrices in particular formats, find the IJ interface relatively easy to use. See Chapter Linear-Algebraic System Interface (IJ) for details.

Linear System Interfaces


Fig. 1: Figure 1
Graphic illustrating the notion of conceptual linear system interfaces.
Generally, a user should choose the most specific interface that matches their application, because this will allow them to use specialized and more efficient solvers and preconditioners without losing access to more general solvers. For example, the second row of Figure Figure 1 is a set of linear solver algorithms. Each linear solver group requires different information from the user through the conceptual interfaces. So, the geometric multigrid algorithm (GMG) listed in the left-most box, for example, can only be used with the left-most conceptual interface. On the other hand, the ILU algorithm in the right-most box may be used with any conceptual interface. Matrix requirements for each
solver and preconditioner are provided in Chapter Solvers and Preconditioners and in Chapter API. Your desired solver strategy may influence your choice of conceptual interface. A typical user will select a single Krylov method and a single preconditioner to solve their system.
The third row of Figure Figure 1 is a list of data layouts or matrix/vector storage schemes. The relationship between linear solver and storage scheme is similar to that of the conceptual interface and linear solver. Note that some of the interfaces in hypre currently only support one matrix/vector storage scheme choice. The conceptual interface, the desired solvers and preconditioners, and the matrix storage class must all be compatible.

### 1.3.3 Writing your code

As discussed in the previous section, the following decisions should be made before writing any code:

- Choose a conceptual interface.
- Choose your desired solver strategy.
- Look up matrix requirements for each solver and preconditioner.
- Choose a matrix storage class that is compatible with your solvers and preconditioners and your conceptual interface.

Once the previous decisions have been made, it is time to code your application to call hypre. At this point, reviewing the previously mentioned example codes provided with the hypre library may prove very helpful. The example codes demonstrate the following general structure of the application calls to hypre:

- Build any necessary auxiliary structures for your chosen conceptual interface. This includes, e.g., the grid and stencil structures if you are using the structured-grid interface.
- Build the matrix, solution vector, and right-hand-side vector through your chosen conceptual interface. Each conceptual interface provides a series of calls for entering information about your problem into hypre.
- Build solvers and preconditioners and set solver parameters (optional). Some parameters like convergence tolerance are the same across solvers, while others are solver specific.
- Call the solve function for the solver.
- Retrieve desired information from solver. Depending on your application, there may be different things you may want to do with the solution vector. Also, performance information such as number of iterations is typically available, though it may differ from solver to solver.

The subsequent chapters of this User's Manual provide the details needed to more fully understand the function of each conceptual interface and each solver. Remember that a comprehensive list of all available functions is provided in Chapter API, and the provided example codes may prove helpful as templates for your specific application.

## STRUCTURED-GRID SYSTEM INTERFACE (STRUCT)

In order to get access to the most efficient and scalable solvers for scalar structured-grid applications, users should use the Struct interface described in this chapter. This interface will also provide access (this is not yet supported) to solvers in hypre that were designed for unstructured-grid applications and sparse linear systems in general. These additional solvers are usually provided via the unstructured-grid interface (FEI) or the linear-algebraic interface (IJ) described in Chapters Finite Element Interface and Linear-Algebraic System Interface (IJ).

Figure Structured Grid Example gives an example of the type of grid currently supported by the Struct interface. The interface uses a finite-difference or finite-volume style, and currently supports only scalar PDEs (i.e., one unknown per gridpoint).


Fig. 1: Structured Grid Example
An example 2D structured grid, distributed accross two processors.
There are four basic steps involved in setting up the linear system to be solved:

1. set up the grid,
2. set up the stencil,
3. set up the matrix,
4. set up the right-hand-side vector.

To describe each of these steps in more detail, consider solving the 2D Laplacian problem

$$
\begin{cases}\nabla^{2} u=f, & \text { in the domain, }  \tag{2.1}\\ u=0, & \text { on the boundary } .\end{cases}
$$

Assume (2.1) is discretized using standard 5-pt finite-volumes on the uniform grid pictured in Structured Grid Example, and assume that the problem data is distributed across two processes as depicted.

### 2.1 Setting Up the Struct Grid

The grid is described via a global index space, i.e., via integer singles in 1D, tuples in 2D, or triples in 3D (see Figure Boxes in Index Space).


Fig. 2: Boxes in Index Space
A box is a collection of abstract cell-centered indices, described by its minimum and maximum indices. Here, two boxes are illustrated.

The integers may have any value, negative or positive. The global indexes allow hypre to discern how data is related spatially, and how it is distributed across the parallel machine. The basic component of the grid is a box: a collection of abstract cell-centered indices in index space, described by its "lower" and "upper" corner indices. The scalar grid data is always associated with cell centers, unlike the more general SStruct interface which allows data to be associated with box indices in several different ways.

Each process describes that portion of the grid that it "owns", one box at a time. For example, the global grid in Figure Structured Grid Example can be described in terms of three boxes, two owned by process 0, and one owned by process 1. The following is the code (with visual annotations) for setting up the grid on process 0 (the code for process 1 is similar).


```
    HYPRE_StructGrid grid;
    int ndim = 2;
    int ilower[][2] = {{-3,1}, {0,1}};
    int iupper[][2] = {{-1,2}, {2,4}};
    /* Create the grid object */
1: HYPRE_StructGridCreate(MPI_COMM_WORLD, ndim, &grid);
```

```
    /* Set grid extents for the first box */
2: HYPRE_StructGridSetExtents(grid, ilower[0], iupper[0]);
    /* Set grid extents for the second box */
HYPRE_StructGridSetExtents(grid, ilower[1], iupper[1]);
    /* Assemble the grid */
    HYPRE_StructGridAssemble(grid);
```

The images along the top illustrate the result of the numbered lines of code. The Create() routine creates an empty 2D grid object that lives on the MPI_COMM_WORLD communicator. The SetExtents() routine adds a new box to the grid. The Assemble() routine is a collective call (i.e., must be called on all processes from a common synchronization point), and finalizes the grid assembly, making the grid "ready to use".

### 2.2 Setting Up the Struct Stencil

The geometry of the discretization stencil is described by an array of indexes, each representing a relative offset from any given gridpoint on the grid. For example, the geometry of the 5-pt stencil for the example problem being considered can be represented by the list of index offsets shown in Figure Figure $4 a$.


Fig. 3: Figure 4a
Representation of the 5-point discretization stencil for the example problem.
Here, the $(0,0)$ entry represents the "center" coefficient, and is the 0th stencil entry. The $(0,-1)$ entry represents the "south" coefficient, and is the 3rd stencil entry. And so on.
On process 0 or 1, the following code (with visual annotations) will set up the stencil in Figure Figure 4a. The stencil must be the same on all processes.


Fig. 4: Figure 4b
Need to combine this with 4 a .


```
HYPRE_StructStencil stencil;
int ndim
    = 2;
int size
    = 5;
```

(continues on next page)

```
int entry;
```

int entry;
int offsets[][2] = {{0,0}, {-1,0}, {1,0}, {0,-1}, {0,1}};
int offsets[][2] = {{0,0}, {-1,0}, {1,0}, {0,-1}, {0,1}};
/* Create the stencil object */
/* Create the stencil object */
1: HYPRE_StructStencilCreate(ndim, size, \&stencil);
1: HYPRE_StructStencilCreate(ndim, size, \&stencil);
/* Set stencil entries */
/* Set stencil entries */
for (entry = 0; entry < size; entry++)
for (entry = 0; entry < size; entry++)
{
{
HYPRE_StructStencilSetElement(stencil, entry, offsets[entry]);
HYPRE_StructStencilSetElement(stencil, entry, offsets[entry]);
}
}
/* Thats it! There is no assemble routine */

```
/* Thats it! There is no assemble routine */
```

The Create() routine creates an empty 2D, 5-pt stencil object. The SetElement () routine defines the geometry of the stencil and assigns the stencil numbers for each of the stencil entries. None of the calls are collective calls.

### 2.3 Setting Up the Struct Matrix

The matrix is set up in terms of the grid and stencil objects described in Sections Setting Up the Struct Grid and Setting Up the Struct Stencil. The coefficients associated with each stencil entry will typically vary from gridpoint to gridpoint, but in the example problem being considered, they are as follows over the entire grid (except at boundaries; see below):

$$
\left[\begin{array}{ccc} 
& -1 &  \tag{2.2}\\
-1 & 4 & -1 \\
& -1 &
\end{array}\right]
$$

On process 0 , the following code sets up matrix values associated with the center (entry 0 ) and south (entry 3 ) stencil entries as given by (2.2) and Figure Figure $4 a$ (boundaries are ignored here temporarily).

```
HYPRE_StructMatrix A;
double values[36];
int stencil_indices[2] = {0,3};
int i;
HYPRE_StructMatrixCreate(MPI_COMM_WORLD, grid, stencil, &A);
HYPRE_StructMatrixInitialize(A);
for (i = 0; i < 36; i += 2)
{
    values[i] = 4.0;
    values[i+1] = -1.0;
}
HYPRE_StructMatrixSetBoxValues(A, ilower[0], iupper[0], 2,
    stencil_indices, values);
HYPRE_StructMatrixSetBoxValues(A, ilower[1], iupper[1], 2,
    stencil_indices, values);
/* set boundary conditions */
...
```

```
HYPRE_StructMatrixAssemble(A);
```

The Create() routine creates an empty matrix object. The Initialize() routine indicates that the matrix coefficients (or values) are ready to be set. This routine may or may not involve the allocation of memory for the coefficient data, depending on the implementation. The optional Set routines mentioned later in this chapter and in Chapter API, should be called before this step. The SetBoxValues() routine sets the matrix coefficients for some set of stencil entries over the gridpoints in some box. Note that the box need not correspond to any of the boxes used to create the grid, but values should be set for all gridpoints that this process "owns". The Assemble() routine is a collective call, and finalizes the matrix assembly, making the matrix "ready to use".

Matrix coefficients that reach outside of the boundary should be set to zero. For efficiency reasons, hypre does not do this automatically. The most natural time to insure this is when the boundary conditions are being set, and this is most naturally done after the coefficients on the grid's interior have been set. For example, during the implementation of the Dirichlet boundary condition on the lower boundary of the grid in Figure Structured Grid Example, the south coefficient must be set to zero. To do this on process 0 , the following code could be used:

```
int ilower[2] = {-3, 1};
int iupper[2] = { 2, 1};
/* create matrix and set interior coefficients */
...
/* implement boundary conditions */
...
for (i = 0; i < 12; i++)
{
    values[i] = 0.0;
}
i = 3;
HYPRE_StructMatrixSetBoxValues(A, ilower, iupper, 1, &i, values);
/* complete implementation of boundary conditions */
...
```


### 2.4 Setting Up the Struct Right-Hand-Side Vector

The right-hand-side vector is set up similarly to the matrix set up described in Section Setting Up the Struct Matrix above. The main difference is that there is no stencil (note that a stencil currently does appear in the interface, but this will eventually be removed).

On process 0 , the following code sets up the right-hand-side vector values.

```
HYPRE_StructVector b;
double values[18];
int i;
HYPRE_StructVectorCreate(MPI_COMM_WORLD, grid, &b);
HYPRE_StructVectorInitialize(b);
```

(continues on next page)

```
for (i = 0; i < 18; i++)
{
    values[i] = 0.0;
}
HYPRE_StructVectorSetBoxValues(b, ilower[0], iupper[0], values);
HYPRE_StructVectorSetBoxValues(b, ilower[1], iupper[1], values);
HYPRE_StructVectorAssemble(b);
```

The Create() routine creates an empty vector object. The Initialize() routine indicates that the vector coefficients (or values) are ready to be set. This routine follows the same rules as its corresponding Matrix routine. The SetBoxValues() routine sets the vector coefficients over the gridpoints in some box, and again, follows the same rules as its corresponding Matrix routine. The Assemble() routine is a collective call, and finalizes the vector assembly, making the vector "ready to use".

### 2.5 Symmetric Matrices

Some solvers and matrix storage schemes provide capabilities for significantly reducing memory usage when the coefficient matrix is symmetric. In this situation, each off-diagonal coefficient appears twice in the matrix, but only one copy needs to be stored. The Struct interface provides support for matrix and solver implementations that use symmetric storage via the SetSymmetric() routine.

To describe this in more detail, consider again the 5-pt finite-volume discretization of (2.1) on the grid pictured in Figure Structured Grid Example. Because the discretization is symmetric, only half of the off-diagonal coefficients need to be stored. To turn symmetric storage on, the following line of code needs to be inserted somewhere between the Create() and Initialize() calls.

```
HYPRE_StructMatrixSetSymmetric(A, 1);
```

The coefficients for the entire stencil can be passed in as before. Note that symmetric storage may or may not actually be used, depending on the underlying storage scheme. Currently in hypre, the Struct interface always uses symmetric storage.

To most efficiently utilize the Struct interface for symmetric matrices, notice that only half of the off-diagonal coefficients need to be set. To do this for the example being considered, we simply need to redefine the 5-pt stencil of Section Setting Up the Struct Stencil to an "appropriate" 3-pt stencil, then set matrix coefficients (as in Section Setting Up the Struct Matrix) for these three stencil elements only. For example, we could use the following stencil

$$
\left[\begin{array}{ll}
(0,1) &  \tag{2.3}\\
(0,0) & (1,0)
\end{array}\right] .
$$

This 3-pt stencil provides enough information to recover the full 5-pt stencil geometry and associated matrix coefficients.

## SEMI-STRUCTURED-GRID SYSTEM INTERFACE (SSTRUCT)

The SStruct interface is appropriate for applications with grids that are mostly—but not entirely—structured, e.g. block-structured grids (see Figure Figure $6 a$ ), composite grids in structured adaptive mesh refinement (AMR) applications (see Figure Figure 9), and overset grids. In addition, it supports more general PDEs than the Struct interface by allowing multiple variables (system PDEs) and multiple variable types (e.g. cell-centered, face-centered, etc.). The interface provides access to data structures and linear solvers in hypre that are designed for semi-structured grid problems, but also to the most general data structures and solvers.

The SStruct grid is composed out of a number of structured grid parts, where the physical inter-relationship between the parts is arbitrary. Each part is constructed out of two basic components: boxes (see Figure Boxes in Index Space) and variables. Variables represent the actual unknown quantities in the grid, and are associated with the box indices in a variety of ways, depending on their types. In hypre, variables may be cell-centered, node-centered, face-centered, or edge-centered. Face-centered variables are split into x-face, y-face, and z-face, and edge-centered variables are split into x-edge, y-edge, and z-edge. See Figure Figure 5 for an illustration in 2D.


Fig. 1: Figure 5
Grid variables in hypre are referenced by the abstract cell-centered index to the left and down in 2D (analogously in 3D). In the figure, index $(i, j)$ is used to reference the variables in black. The variables in grey-although contained in the pictured cell-are not referenced by the $(i, j)$ index.

The SStruct interface uses a graph to allow nearly arbitrary relationships between part data. The graph is constructed from stencils or finite element stiffness matrices plus some additional data-coupling information set by the GraphAddEntries() routine. Two other methods for relating part data are the GridSetNeighborPart() and GridSetSharedPart () routines, which are particularly well suited for block-structured grid problems. The latter is useful for finite element codes.

There are five basic steps involved in setting up the linear system to be solved:

1. set up the grid,
2. set up the stencils (if needed),
3. set up the graph,
4. set up the matrix,
5. set up the right-hand-side vector.

### 3.1 Block-Structured Grids with Stencils

In this section, we describe how to use the SStruct interface to define block-structured grid problems. We do this primarily by example, paying particular attention to the construction of stencils and the use of the GridSetNeighborPart () interface routine.

Consider the solution of the diffusion equation

$$
\begin{equation*}
-\nabla \cdot(D \nabla u)+\sigma u=f \tag{3.1}
\end{equation*}
$$

on the block-structured grid in Figure Figure $6 a$, where $D$ is a scalar diffusion coefficient, and $\sigma \geq 0$. The discretization [MoRS1998] introduces three different types of variables: cell-centered, $x$-face, and $y$-face. The three discretization stencils that couple these variables are also given in the figure. The information in this figure is essentially all that is needed to describe the nonzero structure of the linear system we wish to solve.


Fig. 2: Figure 6a
Example of a block-structured grid with five logically-rectangular blocks and three variables types: cell-centered, $x$-face, and $y$-face. Discretization stencils for the cell-centered (left), $x$-face (middle), and $y$-face (right) variables are also pictured.


Fig. 3: Figure 6b
Need to combine this with 6a.


Fig. 4: Figure 7
One possible labeling of the grid in Figure Figure $6 a$.

The grid in Figure Figure $6 a$ is defined in terms of five separate logically-rectangular parts as shown in Figure Figure 7 , and each part is given a unique label between 0 and 4 . Each part consists of a single box with lower index $(1,1)$ and upper index $(4,4)$ (see Section Setting Up the Struct Grid), and the grid data is distributed on five processes such that data associated with part $p$ lives on process $p$. Note that in general, parts may be composed out of arbitrary unions of boxes, and indices may consist of non-positive integers (see Figure Boxes in Index Space). Also note that the SStruct interface expects a domain-based data distribution by boxes, but the actual distribution is determined by the user and simply described (in parallel) through the interface.


```
    HYPRE_SStructGrid grid;
    int ndim = 2, nparts = 5, nvars = 3, part = 3;
    int extents[][2] = {{1,1}, {4,4}};
    int vartypes[] = {HYPRE_SSTRUCT_VARIABLE_CELL,
        HYPRE_SSTRUCT_VARIABLE_XFACE,
        HYPRE_SSTRUCT_VARIABLE_YFACE};
    int nb2_n_part = 2, nb4_n_part = 4;
    int nb2_exts[][2] = {{1,0}, {4,0}}, nb4_exts[][2] = {{0,1}, {0,4}};
    int nb2_n_exts[][2] = {{1,1}, {1,4}}, nb4_n_exts[][2] = {{4,1}, {4,4}};
    int nb2_map[2] = {1,0}, nb4_map[2] = {0,1};
    int nb2_dir[2] = {1,-1}, nb4_dir[2] = {1,1};
1: HYPRE_SStructGridCreate(MPI_COMM_WORLD, ndim, nparts, &grid);
    /* Set grid extents and grid variables for part 3 */
    HYPRE_SStructGridSetExtents(grid, part, extents[0], extents[1]);
    HYPRE_SStructGridSetVariables(grid, part, nvars, vartypes);
    /* Set spatial relationship between parts 3 and 2, then parts 3 and 4 */
HYPRE_SStructGridSetNeighborPart(grid, part, nb2_exts[0], nb2_exts[1],
    nb2_n_part, nb2_n_exts[0], nb2_n_exts[1], nb2_map, nb2_dir);
: HYPRE_SStructGridSetNeighborPart(grid, part, nb4_exts[0], nb4_exts[1],
    nb4_n_part, nb4_n_exts[0], nb4_n_exts[1], nb4_map, nb4_dir);
6: HYPRE_SStructGridAssemble(grid);
```

Code on process 3 for setting up the grid in Figure fig-sstruct-example\}.

As with the Struct interface, each process describes that portion of the grid that it "owns", one box at a time. Figure fig-sstruct-grid shows the code for setting up the grid on process 3 (the code for the other processes is similar). The "icons" at the top of the figure illustrate the result of the numbered lines of code. Process 3 needs to describe the data pictured in the bottom-right of the figure. That is, it needs to describe part 3 plus some additional neighbor information that ties part 3 together with the rest of the grid. The Create() routine creates an empty 2D grid object with five parts that lives on the MPI_COMM_WORLD communicator. The SetExtents() routine adds a new box to the grid. The SetVariables() routine associates three variables of type cell-centered, $x$-face, and $y$-face with part 3 .

At this stage, the description of the data on part 3 is complete. However, the spatial relationship between this data and the data on neighboring parts is not yet defined. To do this, we need to relate the index space for part 3 with the index spaces of parts 2 and 4 . More specifically, we need to tell the interface that the two grey boxes neighboring part 3 in the bottom-right of Figure fig-sstruct-grid also correspond to boxes on parts 2 and 4 . This is done through the two calls to the SetNeighborPart () routine. We discuss only the first call, which describes the grey box on the right of the figure. Note that this grey box lives outside of the box extents for the grid on part 3, but it can still be described using the index-space for part 3 (recall Figure Boxes in Index Space). That is, the grey box has extents $(1,0)$ and $(4,0)$ on part 3's index-space, which is outside of part 3's grid. The arguments for the SetNeighborPart () call are simply the lower and upper indices on part 3 and the corresponding indices on part 2 . The final two arguments to the routine indicate that the positive $x$-direction on part 3 (i.e., the $i$ component of the tuple $(i, j)$ ) corresponds to the positive $y$-direction on part 2 and that the positive $y$-direction on part 3 corresponds to the positive $x$-direction on part 2 .

The Assemble() routine is a collective call (i.e., must be called on all processes from a common synchronization point), and finalizes the grid assembly, making the grid "ready to use".

With the neighbor information, it is now possible to determine where off-part stencil entries couple. Take, for example, any shared part boundary such as the boundary between parts 2 and 3. Along these boundaries, some stencil entries reach outside of the part. If no neighbor information is given, these entries are effectively zeroed out, i.e., they don't participate in the discretization. However, with the additional neighbor information, when a stencil entry reaches into a neighbor box it is then coupled to the part described by that neighbor box information.
Another important consequence of the use of the SetNeighborPart() routine is that it can declare variables on different parts as being the same. For example, the face variables on the boundary of parts 2 and 3 are recognized as being shared by both parts (prior to the SetNeighborPart () call, there were two distinct sets of variables). Note also that these variables are of different types on the two parts; on part 2 they are $x$-face variables, but on part 3 they are $y$-face variables.

For brevity, we consider only the description of the $y$-face stencil in Figure Figure $6 a$, i.e. the third stencil in the figure. To do this, the stencil entries are assigned unique labels between 0 and 8 and their "offsets" are described relative to the "center" of the stencil. This process is illustrated in Figure Figure 7a. Nine calls are made to the routine HYPRE_SStructStencilSetEntry (). As an example, the call that describes stencil entry 5 in the figure is given the entry number 5 , the offset $(-1,0)$, and the identifier for the $x$-face variable (the variable to which this entry couples). Recall from Figure Figure 5 the convention used for referencing variables of different types. The geometry description uses the same convention, but with indices numbered relative to the referencing index $(0,0)$ for the stencil's center. Figure fig-sstruct-graph shows the code for setting up the graph .


HYPRE_SStructGraph graph;
HYPRE_SStructStencil c_stencil, x_stencil, y_stencil;


Fig. 5: Figure 7a
Assignment of labels and geometries to the $y$-face stencil in Figure fig-sstruct-example\}.


Fig. 6: Figure 7b
Need to combine this with 7a.

```
int c_var = 0, x_var = 1, y_var = 2;
int part;
HYPRE_SStructGraphCreate(MPI_COMM_WORLD, grid, &graph);
    /* Set the cell-centered, x-face, and y-face stencils for each part */
    for (part = 0; part < 5; part++)
{
    HYPRE_SStructGraphSetStencil(graph, part, c_var, c_stencil);
    HYPRE_SStructGraphSetStencil(graph, part, x_var, x_stencil);
    HYPRE_SStructGraphSetStencil(graph, part, y_var, y_stencil);
}
3: HYPRE_SStructGraphAssemble(graph);
```

Code on process 3 for setting up the graph for Figure fig-sstruct-example \}.
With the above, we now have a complete description of the nonzero structure for the matrix. The matrix coefficients are then easily set in a manner similar to what is described in Section Setting Up the Struct Matrix using routines MatrixSetValues() and MatrixSetBoxValues() in the SStruct interface. As before, there are also AddTo variants of these routines. Likewise, setting up the right-hand-side is similar to what is described in Section Setting Up the Struct Right-Hand-Side Vector. See the hypre reference manual for details.

An alternative approach for describing the above problem through the interface is to use the GraphAddEntries() routine instead of the GridSetNeighborPart() routine. In this approach, the five parts would be explicitly "sewn" together by adding non-stencil couplings to the matrix graph. The main downside to this approach for block-structured grid problems is that variables along block boundaries are no longer considered to be the same variables on the corresponding parts that share these boundaries. For example, any face variable along the boundary between parts 2 and 3 in Figure Figure $6 a$ would represent two different variables that live on different parts. To "sew" the parts together correctly, we would need to explicitly select one of these variables as the representative that participates in the discretization, and make the other variable a dummy variable that is decoupled from the discretization by zeroing out appropriate entries in the matrix. All of these complications are avoided by using the GridSetNeighborPart() for this example.

### 3.2 Block-Structured Grids with Finite Elements

In this section, we describe how to use the SStruct interface to define block-structured grid problems with finite elements. We again do this by example, paying particular attention to the use of the FEM interface routines and the GridSetSharedPart() routine. See example code ex14.c for a complete implementation.

Consider a nodal finite element (FEM) discretization of the Laplace equation on the star-shaped grid in Figure Figure $8 a$. The local FEM stiffness matrix in the figure describes the coupling between the grid variables. Although we could still describe this problem using stencils as in Section Block-Structured Grids with Stencils, an FEM-based approach (available in hypre version 2.6.0b and later) is a more natural alternative.
The grid in Figure Figure $8 a$ is defined in terms of six separate logically-rectangular parts, and each part is given a unique label between 0 and 5 . Each part consists of a single box with lower index $(1,1)$ and upper index $(9,9)$, and the grid data is distributed on six processes such that data associated with part $p$ lives on process $p$.


Fig. 7: Figure 8a
Example of a star-shaped grid with six logically-rectangular blocks and one nodal variable. Each block has an angle at the origin given by $\gamma=\pi / 3$. The finite element stiffness matrix (right) is given in terms of the pictured variable ordering (left).

$$
\begin{aligned}
& 0 \\
& 1 \\
& 2 \\
& 3
\end{aligned}\left(\begin{array}{cccc}
0 & 1 & 2 & 3 \\
4-k & -1 & -2+k & -1 \\
-1 & 4+k & -1 & -2-k \\
-2+k & -1 & 4-k & -1 \\
-1 & -2-k & -1 & 4+k
\end{array}\right) \alpha, \begin{aligned}
& \alpha=(6 \sin (\gamma))^{-1}, \quad k=3 \cos (\gamma), \quad \gamma=\pi / 3
\end{aligned}
$$

Fig. 8: Figure 8b
Need to combine this with 8 a.


```
    HYPRE_SStructGrid grid;
    int ndim = 2, nparts = 6, nvars = 1, part = 0;
    int ilower[2] = {1,1}, iupper[2] = {9,9};
    int vartypes[] = {HYPRE_SSTRUCT_VARIABLE_NODE};
    int ordering[12] = {0,-1,-1, 0,+1,-1, 0,+1,+1, 0,-1,+1};
    int s_part = 2;
    int ilo[2] = {1,1}, iup[2] = {1,9}, offset[2] = {-1,0};
    int s_ilo[2] = {1,1}, s_iup[2] = {9,1}, s_offset[2] = {0,-1};
    int map[2] = {1,0};
    int dir[2] = {-1,1};
1: HYPRE_SStructGridCreate(MPI_COMM_WORLD, ndim, nparts, &grid);
    /* Set grid extents, grid variables, and FEM ordering for part 0 %/
    HYPRE_SStructGridSetExtents(grid, part, ilower, iupper);
    HYPRE_SStructGridSetVariables(grid, part, nvars, vartypes);
HYPRE_SStructGridSetFEMOrdering(grid, part, ordering);
    /* Set shared variables for parts 0 and 1 (0 and 2/3/4/5 not shown) */
5: HYPRE_SStructGridSetSharedPart(grid, part, ilo, iup, offset,
    s_part, s_ilo, s_iup, s_offset, map, dir);
6: HYPRE_SStructGridAssemble(grid);
```

Code on process 0 for setting up the grid in Figure Figure $8 a$.
As in Section Block-Structured Grids with Stencils, each process describes that portion of the grid that it "owns", one box at a time. Figure fig-sstruct-fem-grid shows the code for setting up the grid on process 0 (the code for the other processes is similar). The "icons" at the top of the figure illustrate the result of the numbered lines of code. Process 0 needs to describe the data pictured in the bottom-right of the figure. That is, it needs to describe part 0 plus some additional information about shared data with other parts on the grid. The SetFEMOrdering () routine sets the ordering of the unknowns in an element (an element is always a grid cell in hypre). This determines the ordering of the data passed into the routines MatrixAddFEMValues () and VectorAddFEMValues() discussed later.

At this point, the layout of the data on part 0 is complete, but there is no relationship to the rest of the grid. To couple the parts, we need to tell hypre that some of the boundary variables on part 0 are shared with other parts, i.e., they are the same as some of the variables on other parts. This is done through five calls to the SetSharedPart () routine. Only the first call is shown in the figure; the other four calls are similar. The arguments to this routine are the same as

SetNeighborPart () with the addition of two new offset arguments, named offset and s_offset in the figure. Each offset represents a pointer from the cell center to one of the following: all variables in the cell (no nonzeros in offset); all variables on a face (only 1 nonzero); all variables on an edge ( 2 nonzeros); all variables at a point ( 3 nonzeros). The two offsets must be consistent with each other.

The graph is set up similarly to Figure fig-sstruct-graph, except that the stencil calls are replaced by calls to GraphSetFEM(). The nonzero pattern of the stiffness matrix can also be set by calling the optional routine GraphSetFEMSparsity().

Matrix and vector values are set one element at a time. For the example in this section, calls on part 0 would have the following form:

```
int part = 0;
int index[2] = {i,j};
double m_values[16] = {...};
double v_values[4] = {...};
HYPRE_SStructMatrixAddFEMValues(A, part, index, m_values);
HYPRE_SStructVectorAddFEMValues(v, part, index, v_values);
```

Here, m_values contains local stiffness matrix values and v_values contains local variable values. The global matrix and vector are assembled internally by hypre, using the shared variables to couple the parts.

### 3.3 Structured Adaptive Mesh Refinement

We now briefly discuss how to use the SStruct interface in a structured AMR application. Consider Poisson's equation on the simple cell-centered example grid illustrated in Figure Figure 9. For structured AMR applications, each refinement level should be defined as a unique part. There are two parts in this example: part 0 is the global coarse grid and part 1 is the single refinement patch. Note that the coarse unknowns underneath the refinement patch (gray dots in Figure Figure 9) are not real physical unknowns; the solution in this region is given by the values on the refinement patch. In setting up the composite grid matrix [McCo1989] for hypre the equations for these "dummy" unknowns should be uncoupled from the other unknowns (this can easily be done by setting all off-diagonal couplings to zero in this region).

In the example, parts are distributed across the same two processes with process 0 having the "left" half of both parts. The composite grid is then set up part-by-part by making calls to GridSetExtents() just as was done in Section BlockStructured Grids with Stencils and Figure fig-sstruct-grid (no SetNeighborPart calls are made in this example). Note that in the interface there is no required rule relating the indexing on the refinement patch to that on the global coarse grid; they are separate parts and thus each has its own index space. In this example, we have chosen the indexing such that refinement cell $(2 i, 2 j)$ lies in the lower left quadrant of coarse cell $(i, j)$. Then the stencil is set up. In this example we are using a finite volume approach resulting in the standard 5-point stencil in Section Setting Up the Struct Grid in both parts.

The grid and stencil are used to define all intra-part coupling in the graph, the non-zero pattern of the composite grid matrix. The inter-part coupling at the coarse-fine interface is described by GraphAddEntries() calls. This coupling in the composite grid matrix is typically the composition of an interpolation rule and a discretization formula. In this example, we use a simple piecewise constant interpolation, i.e. the solution value in a coarse cell is equal to the solution value at the cell center. Then the flux across a portion of the coarse-fine interface is approximated by a difference of the solution values on each side. As an example, consider approximating the flux across the left interface of cell $(6,6)$ in Figure Figure 2. Let $h$ be the coarse grid mesh size, and consider a local coordinate system with the origin at the

part 0

Fig. 9: Figure 9
Structured AMR grid example. Shaded regions correspond to process 0 , unshaded to process 1 . The grey dots are dummy variables.
center of cell $(6,6)$. We approximate the flux as follows

$$
\begin{aligned}
\int_{-h / 4}^{h / 4} u_{x}(-h / 4, s) d s & \approx \frac{h}{2} u_{x}(-h / 4,0) \approx \frac{h}{2} \frac{u(0,0)-u(-3 h / 4,0)}{3 h / 4} \\
& \approx \frac{2}{3}\left(u_{6,6}-u_{2,3}\right) .
\end{aligned}
$$

The first approximation uses the midpoint rule for the edge integral, the second uses a finite difference formula for the derivative, and the third the piecewise constant interpolation to the solution in the coarse cell. This means that the equation for the variable at cell $(6,6)$ involves not only the stencil couplings to $(6,7)$ and $(7,6)$ on part 1 but also nonstencil couplings to $(2,3)$ and $(3,2)$ on part 0 . These non-stencil couplings are described by GraphAddEntries() calls. The syntax for this call is simply the part and index for both the variable whose equation is being defined and the variable to which it couples. After these calls, the non-zero pattern of the matrix (and the graph) is complete. Note that the "west" and "south" stencil couplings simply "drop off" the part, and are effectively zeroed out (currently, this is only supported for the HYPRE_PARCSR object type, and these values must be manually zeroed out for other object types; see MatrixSetObjectType() in the reference manual).

The remaining step is to define the actual numerical values for the composite grid matrix. This can be done by either MatrixSetValues() calls to set entries in a single equation, or by MatrixSetBoxValues() calls to set entries for a box of equations in a single call. The syntax for the MatrixSetValues() call is a part and index for the variable whose equation is being set and an array of entry numbers identifying which entries in that equation are being set. The entry numbers may correspond to stencil entries or non-stencil entries.


Fig. 10: Figure 2
Coupling for equation at corner of refinement patch. Black lines (solid and broken) are stencil couplings. Gray line are non-stencil couplings.

## FINITE ELEMENT INTERFACE

Warning: FEI is not actively supported by the hypre development team. For similar functionality, we recommend using Block-Structured Grids with Finite Elements, which allows the representation of block-structured grid problems via hypre's SStruct interface.

### 4.1 Introduction

Many application codes use unstructured finite element meshes. This section describes an interface for finite element problems, called the FEI, which is supported in hypre.


Fig. 1: Example of an unstructured mesh.

FEI refers to a specific interface for black-box finite element solvers, originally developed in Sandia National Lab, see [CIEA1999]. It differs from the rest of the conceptual interfaces in hypre in two important aspects: it is written in C++, and it does not separate the construction of the linear system matrix from the solution process. A complete description of Sandia's FEI implementation can be obtained by contacting Alan Williams at Sandia (william@sandia.gov). A simplified version of the FEI has been implemented at LLNL and is included in hypre. More details about this implementation can be found in the header files of the FEI_mv/fei-base and FEI_mv/fei-hypre directories.

### 4.2 A Brief Description of the Finite Element Interface

Typically, finite element codes contain data structures storing element connectivities, element stiffness matrices, element loads, boundary conditions, nodal coordinates, etc. One of the purposes of the FEI is to assemble the global linear system in parallel based on such local element data. We illustrate this in the rest of the section and refer to example 10 (in the examples directory) for more implementation details.

In hypre, one creates an instance of the FEI as follows:

```
LLNL_FEI_Impl *feiPtr = new LLNL_FEI_Impl(mpiComm);
```

Here mpiComm is an MPI communicator (e.g. MPI\_COMM $\backslash$ WORLD). If Sandia's FEI package is to be used, one needs to define a hypre solver object first:

```
LinearSystemCore *solver = HYPRE_base_create(mpiComm);
FEI_Implementation *feiPtr = FEI_Implementation(solver,mpiComm,rank);
```

where rank is the number of the master processor (used only to identify which processor will produce the screen outputs). The LinearSystemCore class is the part of the FEI that interfaces with the linear solver library. It will be discussed later in Sections FEI Solvers and Using HYPRE in External FEI Implementations.

Local finite element information is passed to the FEI using several methods of the feiPtr object. The first entity to be submitted is the field information. A field has an identifier called fieldID and a rank or fieldSize (number of degree of freedom). For example, a discretization of the Navier Stokes equations in 3D can consist of velocity vector having 3 degrees of freedom in every node (vertex) of the mesh and a scalar pressure variable, which is constant over each element. If these are the only variables, and if we assign fieldID 7 and 8 to them, respectively, then the finite element field information can be set up by

```
nFields = 2; /* number of unknown fields */
fieldID = new int[nFields]; /* field identifiers */
fieldSize = new int[nFields]; /* vector dimension of each field */
/* velocity (a 3D vector) */
fieldID[0] = 7;
fieldSize[0] = 3;
/* pressure (a scalar function) */
fieldID[1] = 8;
fieldSize[1] = 1;
feiPtr -> initFields(nFields, fieldSize, fieldID);
```

Once the field information has been established, we are ready to initialize an element block. An element block is characterized by the block identifier, the number of elements, the number of nodes per element, the nodal fields and the element fields (fields that have been defined previously). Suppose we use 1000 hexahedral elements in the element block 0 , the setup consists of

```
elemBlkID = 0; /* identifier for a block of elements */
nElems = 1000; /* number of elements in the block */
elemNNodes = 8; /* number of nodes per element */
/* nodal-based field for the velocity */
nodeNFields = 1;
nodeFieldIDs = new[nodeNFields];
nodeFieldIDs[0] = fieldID[0];
/* element-based field for the pressure */
elemNFields = 1;
elemFieldIDs = new[elemNFields];
elemFieldIDs[0] = fieldID[1];
feiPtr -> initElemBlock(elemBlkID, nElems, elemNNodes, nodeNFields,
    nodeFieldIDs, elemNFields, elemFieldIDs, 0);
```

The last argument above specifies how the dependent variables are arranged in the element matrices. A value of 0 indicates that each variable is to be arranged in a separate block (as opposed to interleaving).
In a parallel environment, each processor has one or more element blocks. Unless the element blocks are all disjoint, some of them share a common set of nodes on the subdomain boundaries. To facilitate setting up interprocessor communications, shared nodes between subdomains on different processors are to be identified and sent to the FEI. Hence, each node in the whole domain is assigned a unique global identifier. The shared node list on each processor contains a subset of the global node list corresponding to the local nodes that are shared with the other processors. The syntax for setting up the shared nodes is

```
feiPtr -> initSharedNodes(nShared, sharedIDs, sharedLengs, sharedProcs);
```

This completes the initialization phase, and a completion signal is sent to the FEI via

```
feiPtr -> initComplete();
```

Next, we begin the load phase. The first entity for loading is the nodal boundary conditions. Here we need to specify the number of boundary equations and the boundary values given by alpha, beta, and gamma. Depending on whether the boundary conditions are Dirichlet, Neumann, or mixed, the three values should be passed into the FEI accordingly.

```
feiPtr -> loadNodeBCs(nBCs, BCEqn, fieldID, alpha, beta, gamma);
```

The element stiffness matrices are to be loaded in the next step. We need to specify the element number $i$, the element block to which element $i$ belongs, the element connectivity information, the element load, and the element matrix format. The element connectivity specifies a set of 8 node global IDs (for hexahedral elements), and the element load is the load or force for each degree of freedom. The element format specifies how the equations are arranged (similar to the interleaving scheme mentioned above). The calling sequence for loading element stiffness matrices is

```
for (i = 0; i < nElems; i++)
    feiPtr -> sumInElem(elemBlkID, elemID, elemConn[i], elemStiff[i],
        elemLoads[i], elemFormat);
```

To complete the assembling of the global stiffness matrix and the corresponding right hand side, a signal is sent to the FEI via

```
feiPtr -> loadComplete();
```


## LINEAR-ALGEBRAIC SYSTEM INTERFACE (IJ)

The IJ interface described in this chapter is the lowest common denominator for specifying linear systems in hypre. This interface provides access to general sparse-matrix solvers in hypre, not to the specialized solvers that require more problem information.

### 5.1 IJ Matrix Interface

As with the other interfaces in hypre, the IJ interface expects to get data in distributed form because this is the only scalable approach for assembling matrices on thousands of processes. Matrices are assumed to be distributed by blocks of rows as follows:

$$
\left[\begin{array}{c}
A_{0} \\
A_{1} \\
\vdots \\
A_{P-1}
\end{array}\right]
$$

In the above example, the matrix is distributed across the $P$ processes, $0,1, \ldots, P-1$ by blocks of rows. Each submatrix $A_{p}$ is "owned" by a single process and its first and last row numbers are given by the global indices ilower and iupper in the Create() call below.

The following example code illustrates the basic usage of the IJ interface for building matrices:

```
MPI_Comm
HYPRE_IJMatrix ij_matrix;
HYPRE_ParCSRMatrix parcsr_matrix;
int ilower, iupper;
int jlower, jupper;
int nrows;
int *ncols;
int *rows;
int *cols;
double *values;
HYPRE_IJMatrixCreate(comm, ilower, iupper, jlower, jupper, &ij_matrix);
HYPRE_IJMatrixSetObjectType(ij_matrix, HYPRE_PARCSR);
HYPRE_IJMatrixInitialize(ij_matrix);
/* set matrix coefficients */
HYPRE_IJMatrixSetValues(ij_matrix, nrows, ncols, rows, cols, values);
/* add-to matrix cofficients, if desired */
```

HYPRE_IJMatrixAddToValues(ij_matrix, nrows, ncols, rows, cols, values);

```
...
```

HYPRE_IJMatrixAssemble(ij_matrix);
HYPRE_IJMatrixGetObject(ij_matrix, (void **) \&parcsr_matrix) ;
The Create() routine creates an empty matrix object that lives on the comm communicator. This is a collective call (i.e., must be called on all processes from a common synchronization point), with each process passing its own row extents, ilower and iupper. The row partitioning must be contiguous, i.e., iupper for process i must equal ilower -1 for process $i+1$. Note that this allows matrices to have 0 - or 1-based indexing. The parameters jlower and jupper define a column partitioning, and should match ilower and iupper when solving square linear systems. See Chapter API for more information.

The SetObjectType() routine sets the underlying matrix object type to HYPRE_PARCSR (this is the only object type currently supported). The Initialize() routine indicates that the matrix coefficients (or values) are ready to be set. This routine may or may not involve the allocation of memory for the coefficient data, depending on the implementation. The optional SetRowSizes() and SetDiagOffdSizes() routines mentioned later in this chapter and in Chapter API, should be called before this step.

The SetValues () routine sets matrix values for some number of rows (nrows) and some number of columns in each row (ncols). The actual row and column numbers of the matrix values to be set are given by rows and cols. The coefficients can be modified with the AddToValues() routine. If AddToValues() is used to add to a value that previously didn't exist, it will set this value. Note that while AddToValues() will add to values on other processors, SetValues () does not set values on other processors. Instead if a user calls SetValues () on processor $i$ to set a matrix coefficient belonging to processor $j$, processor $i$ will erase all previous occurrences of this matrix coefficient, so they will not contribute to this coefficient on processor $j$. The actual coefficient has to be set on processor $j$.

The Assemble() routine is a collective call, and finalizes the matrix assembly, making the matrix "ready to use". The GetObject () routine retrieves the built matrix object so that it can be passed on to hypre solvers that use the ParCSR internal storage format. Note that this is not an expensive routine; the matrix already exists in ParCSR storage format, and the routine simply returns a "handle" or pointer to it. Although we currently only support one underlying data storage format, in the future several different formats may be supported.

One can preset the row sizes of the matrix in order to reduce the execution time for the matrix specification. One can specify the total number of coefficients for each row, the number of coefficients in the row that couple the diagonal unknown to (Diag) unknowns in the same processor domain, and the number of coefficients in the row that couple the diagonal unknown to ( $0 f f d$ ) unknowns in other processor domains:

```
HYPRE_IJMatrixSetRowSizes(ij_matrix, sizes);
HYPRE_IJMatrixSetDiagOffdSizes(matrix, diag_sizes, offdiag_sizes);
```

Once the matrix has been assembled, the sparsity pattern cannot be altered without completely destroying the matrix object and starting from scratch. However, one can modify the matrix values of an already assembled matrix. To do this, first call the Initialize() routine to re-initialize the matrix, then set or add-to values as before, and call the Assemble () routine to re-assemble before using the matrix. Re-initialization and re-assembly are very cheap, essentially a no-op in the current implementation of the code.

### 5.2 IJ Vector Interface

The following example code illustrates the basic usage of the IJ interface for building vectors:

```
MPI_Comm comm;
HYPRE_IJVector ij_vector;
HYPRE_ParVector par_vector;
int jlower, jupper;
int nvalues;
int *indices;
double *values;
HYPRE_IJVectorCreate(comm, jlower, jupper, &ij_vector);
HYPRE_IJVectorSetObjectType(ij_vector, HYPRE_PARCSR);
HYPRE_IJVectorInitialize(ij_vector);
/* set vector values */
HYPRE_IJVectorSetValues(ij_vector, nvalues, indices, values);
...
HYPRE_IJVectorAssemble(ij_vector);
HYPRE_IJVectorGetObject(ij_vector, (void **) &par_vector);
```

The Create() routine creates an empty vector object that lives on the comm communicator. This is a collective call, with each process passing its own index extents, jlower and jupper. The names of these extent parameters begin with a $j$ because we typically think of matrix-vector multiplies as the fundamental operation involving both matrices and vectors. For matrix-vector multiplies, the vector partitioning should match the column partitioning of the matrix (which also uses the $j$ notation). For linear system solves, these extents will typically match the row partitioning of the matrix as well.

The SetObjectType() routine sets the underlying vector storage type to HYPRE_PARCSR (this is the only storage type currently supported). The Initialize() routine indicates that the vector coefficients (or values) are ready to be set. This routine may or may not involve the allocation of memory for the coefficient data, depending on the implementation.

The SetValues () routine sets the vector values for some number (nvalues) of indices. The values can be modified with the AddToValues() routine. Note that while AddToValues() will add to values on other processors, SetValues() does not set values on other processors. Instead if a user calls SetValues() on processor $i$ to set a value belonging to processor $j$, processor $i$ will erase all previous occurrences of this matrix coefficient, so they will not contribute to this value on processor $j$. The actual value has to be set on processor $j$.

The Assemble() routine is a trivial collective call, and finalizes the vector assembly, making the vector "ready to use". The GetObject () routine retrieves the built vector object so that it can be passed on to hypre solvers that use the ParVector internal storage format.

Vector values can be modified in much the same way as with matrices by first re-initializing the vector with the Initialize() routine.

### 5.3 A Scalable Interface

As explained in the previous sections, problem data is passed to the hypre library in its distributed form. However, as is typically the case for a parallel software library, some information regarding the global distribution of the data will be needed for hypre to perform its function. In particular, a solver algorithm requires that a processor obtain "nearby" data from other processors in order to complete the solve. While a processor may easily determine what data it needs from other processors, it may not know which processor owns the data it needs. Therefore, processors must determine their communication partners, or neighbors.
The straightforward approach to determining neighbors involves constructing a global partition of the data. This approach, however, requires $O(P)$ storage and computations and is not scalable for machines with tens of thousands of processors. The assumed partition algorithm was developed to address this problem [BaFY2006]. It is the approach used in hypre.

## SOLVERS AND PRECONDITIONERS

There are several solvers available in hypre via different conceptual interfaces:

|  | System Interfaces |  |  |  |
| :--- | :--- | :--- | :--- | :--- |
| Solvers | Struct | SStruct | FEI | IJ |
| Jacobi | X | X |  |  |
| SMG | X | X |  |  |
| PFMG | X | X |  |  |
| Split |  | X |  |  |
| SysPFMG |  | X |  |  |
| FAC |  | X |  |  |
| Maxwell |  | X |  |  |
| BoomerAMG |  | X | X | X |
| AMS |  | X | X | X |
| ADS |  | X | X | X |
| MLI |  | X | X | X |
| MGR |  |  |  | X |
| FSAI |  |  |  | X |
| ParaSails |  | X | X | X |
| ILU |  |  |  | X |
| Euclid |  | X | X | X |
| PILUT |  | X | X | X |
| PCG | X | X | X | X |
| GMRES | X | X | X | X |
| FlexGMRES | X | X | X | X |
| LGMRES | X | X |  | X |
| BiCGSTAB | X | X | X | X |
| Hybrid | X | X | X | X |
| LOBPCG | X | X |  | X |

Note that there are a few additional solvers and preconditioners not mentioned in the table that can be used only through the FEI interface and are described in Paragraph 6.14. The procedure for setup and use of solvers and preconditioners is largely the same. We will refer to them both as solvers in the sequel except when noted. In normal usage, the preconditioner is chosen and constructed before the solver, and then handed to the solver as part of the solver's setup. In the following, we assume the most common usage pattern in which a single linear system is set up and then solved with a single righthand side. We comment later on considerations for other usage patterns.

## Setup:

1. Pass to the solver the information defining the problem. In the typical user cycle, the user has passed this information into a matrix through one of the conceptual interfaces prior to setting up the solver. In this situation, the problem definition information is then passed to the solver by passing the constructed matrix into the solver.

As described before, the matrix and solver must be compatible, in that the matrix must provide the services needed by the solver. Krylov solvers, for example, need only a matrix-vector multiplication. Most preconditioners, on the other hand, have additional requirements such as access to the matrix coefficients.
2. Create the solver/preconditioner via the Create() routine.
3. Choose parameters for the preconditioner and/or solver. Parameters are chosen through the Set () calls provided by the solver. Throughout hypre, we have made our best effort to give all parameters reasonable defaults if not chosen. However, for some preconditioners/solvers the best choices for parameters depend on the problem to be solved. We give recommendations in the individual sections on how to choose these parameters. Note that in hypre, convergence criteria can be chosen after the preconditioner/solver has been setup. For a complete set of all available parameters see Chapter API.
4. Pass the preconditioner to the solver. For solvers that are not preconditioned, this step is omitted. The preconditioner is passed through the SetPrecond() call.
5. Set up the solver. This is just the Setup () routine. At this point the matrix and right hand side is passed into the solver or preconditioner. Note that the actual right hand side is not used until the actual solve is performed.
At this point, the solver/preconditioner is fully constructed and ready for use.

## Use:

1. Set convergence criteria. Convergence can be controlled by the number of iterations, as well as various tolerances such as relative residual, preconditioned residual, etc. Like all parameters, reasonable defaults are used. Users are free to change these, though care must be taken. For example, if an iterative method is used as a preconditioner for a Krylov method, a constant number of iterations is usually required.
2. Solve the system. This is just the Solve() routine.

## Finalize:

1. Free the solver or preconditioner. This is done using the Destroy() routine.

## Synopsis

In general, a solver (let's call it SOLVER) is set up and run using the following routines, where A is the matrix, b the right hand side and $x$ the solution vector of the linear system to be solved:

```
/* Create Solver */
int HYPRE_SOLVERCreate(MPI_COMM_WORLD, &solver);
/* Set certain parameters if desired */
HYPRE_SOLVERSetTol(solver, 1.e-8);
...
/* Set up Solver */
HYPRE_SOLVERSetup(solver, A, b, x);
/* Solve the system */
HYPRE_SOLVERSolve(solver, A, b, x);
/* Destroy the solver */
HYPRE_SOLVERDestroy(solver);
```

In the following sections, we will give brief descriptions of the available hypre solvers with some suggestions on how to choose the parameters as well as references for users who are interested in a more detailed description and analysis of the solvers. A complete list of all routines that are available can be found in Chapter API.

### 6.1 SMG

SMG is a parallel semicoarsening multigrid solver for the linear systems arising from finite difference, finite volume, or finite element discretizations of the diffusion equation,

$$
\nabla \cdot(D \nabla u)+\sigma u=f
$$

on logically rectangular grids. The code solves both 2D and 3D problems with discretization stencils of up to 9-point in 2D and up to 27-point in 3D. See [Scha1998], [BrFJ2000], [FaJo2000] for details on the algorithm and its parallel implementation/performance.

SMG is a particularly robust method. The algorithm semicoarsens in the z-direction and uses plane smoothing. The xy plane-solves are effected by one V-cycle of the 2D SMG algorithm, which semicoarsens in the $y$-direction and uses line smoothing.

### 6.2 PFMG

PFMG is a parallel semicoarsening multigrid solver similar to SMG. See [AsFa1996], [FaJo2000] for details on the algorithm and its parallel implementation/performance.
The main difference between the two methods is in the smoother: PFMG uses simple pointwise smoothing. As a result, PFMG is not as robust as SMG, but is much more efficient per V-cycle.

### 6.3 SysPFMG

SysPFMG is a parallel semicoarsening multigrid solver for systems of elliptic PDEs. It is a generalization of PFMG, with the interpolation defined only within the same variable. The relaxation is of nodal type- all variables at a given point location are simultaneously solved for in the relaxation.

Although SysPFMG is implemented through the SStruct interface, it can be used only for problems with one grid part. Ideally, the solver should handle any of the seven variable types (cell-, node-, xface-, yface-, zface-, xedge-, yedge-, and zedge-based). However, it has been completed only for cell-based variables.

### 6.4 SplitSolve

SplitSolve is a parallel block Gauss-Seidel solver for semi-structured problems with multiple parts. For problems with only one variable, it can be viewed as a domain-decomposition solver with no grid overlapping.
Consider a multiple part problem given by the linear system $A x=b$. Matrix $A$ can be decomposed into a structured intra-variable block diagonal component $M$ and a component $N$ consisting of the inter-variable blocks and any unstructured connections between the parts. SplitSolve performs the iteration

$$
x_{k+1}=\tilde{M}^{-1}\left(b+N x_{k}\right)
$$

where $\tilde{M}^{-1}$ is a decoupled block-diagonal $\mathrm{V}(1,1)$ cycle, a separate cycle for each part and variable type. There are two V-cycle options, SMG and PFMG.

### 6.5 FAC

FAC is a parallel fast adaptive composite grid solver for finite volume, cell-centred discretizations of smooth diffusion coefficient problems. To be precise, it is a FACx algorithm since the patch solves consist of only relaxation sweeps. For details of the basic overall algorithms, see [McCo1989]. Algorithmic particularities include formation of nonGalerkin coarse-grid operators (i.e., coarse-grid operators underlying refinement patches are automatically generated) and non-stored linear/constant interpolation/restriction operators. Implementation particularities include a processor redistribution of the generated coarse-grid operators so that intra-level communication between adaptive mesh refinement (AMR) levels during the solve phase is kept to a minimum. This redistribution is hidden from the user.
The user input is essentially a linear system describing the composite operator, and the refinement factors between the AMR levels. To form this composite linear system, the AMR grid is described using semi-structured grid parts. Each AMR level grid corresponds to a separate part so that this level grid is simply a collection of boxes, all with the same refinement factor, i.e., it is a struct grid. However, several restrictions are imposed on the patch (box) refinements. First, a refinement box must cover all underlying coarse cells- i.e., refinement of a partial coarse cell is not permitted. Also, the refined/coarse indices must follow a mapping: with $\left[r_{1}, r_{2}, r_{3}\right]$ denoting the refinement factor and $\left[a_{1}, a_{2}, a_{3}\right] \times$ $\left[b_{1}, b_{2}, b_{3}\right]$ denoting the coarse subbox to be refined, the mapping to the refined patch is

$$
\left[r_{1} * a_{1}, r_{2} * a_{2}, r_{3} * a_{3}\right] \times\left[r_{1} * b_{1}+r_{1}-1, r_{2} * b_{2}+r_{2}-1, r_{3} * b_{3}+r_{3}-1\right] .
$$

With the AMR grid constructed under these restrictions, the composite matrix can be formed. Since the AMR levels correspond to semi-structured grid parts, the composite matrix is a semi-structured matrix consisting of structured components within each part, and unstructured components describing the coarse-to-fine/fine-to-coarse connections. The structured and unstructured components can be set using stencils and the HYPRE_SStructGraphAddEntries routine, respectively. The matrix coefficients can be filled after setting these non-zero patterns. Between each pair of successive AMR levels, the coarse matrix underlying the refinement patch must be the identity and the corresponding rows of the rhs must be zero. These can performed using routines HYPRE_SStructFACZeroCFSten (to zero off the stencil values reaching from coarse boxes into refinement boxes), HYPRE_SStructFACZeroFCSten (to zero off the stencil values reaching from refinement boxes into coarse boxes), HYPRE_SStructFACZeroAMRMatrixData (to set
the identity at coarse grid points underlying a refinement patch), and HYPRE_SStructFACZeroAMRVectorData (to zero off a vector at coarse grid points underlying a refinement patch). These routines can simplify the user's matrix setup. For example, consider two successive AMR levels with the coarser level consisting of one box and the finer level consisting of a collection of boxes. Rather than distinguishly setting the stencil values and the identity in the appropriate locations, the user can set the stencil values on the whole coarse grid using the HYPRE_SStructMatrixSetBoxValues routine and then zero off the appropriate values using the above zeroing routines.

The coarse matrix underlying these patches are algebraically generated by operator-collapsing the refinement patch operator and the fine-to-coarse coefficients (this is why stencil values reaching out of a part must be zeroed). This matrix is re-distributed so that each processor has all of its coarse-grid operator.

To solve the coarsest AMR level, a PFMG V cycle is used. Note that a minimum of two AMR levels are needed for this solver.

### 6.6 Maxwell

Maxwell is a parallel solver for edge finite element discretization of the curl-curl formulation of the Maxwell equation

$$
\nabla \times \alpha \nabla \times E+\beta E=f, \beta>0
$$

on semi-structured grids. Details of the algorithm can be found in [JoLe2006]. The solver can be viewed as an operatordependent multiple-coarsening algorithm for the Helmholtz decomposition of the error correction. Input to this solver consist of only the linear system and a gradient operator. In fact, if the orientation of the edge elements conforms to a lexicographical ordering of the nodes of the grid, then the gradient operator can be generated with the routine HYPRE_MaxwellGrad: at grid points $(i, j, k)$ and $(i-1, j, k)$, the produced gradient operator takes values 1 and -1 respectively, which is the correct gradient operator for the appropriate edge orientation. Since the gradient operator is normalized (i.e., $h$ independent) the edge finite element must also be normalized in the discretization.

This solver is currently developed for perfectly conducting boundary condition (Dirichlet). Hence, the rows and columns of the matrix that corresponding to the grid boundary must be set to the identity or zeroed off. This can be achieved with the routines HYPRE_SStructMaxwellPhysBdy and HYPRE_SStructMaxwellEliminateRowsCols. The former identifies the ranks of the rows that are located on the grid boundary, and the latter adjusts the boundary rows and cols. As usual, the rhs of the linear system must also be zeroed off at the boundary rows. This can be done using HYPRE_SStructMaxwellZeroVector.

With the adjusted linear system and a gradient operator, the user can form the Maxwell multigrid solver using several different edge interpolation schemes. For problems with smooth coefficients, the natural Nedelec interpolation operator can be used. This is formed by calling HYPRE_SStructMaxwellSetConstantCoef with the flag $>0$ before setting up the solver, otherwise the default edge interpolation is an operator-collapsing/element-agglomeration scheme. This is suitable for variable coefficients. Also, before setting up the solver, the user must pass the gradient operator, whether user or HYPRE_MaxwellGrad generated, with HYPRE_SStructMaxwellSetGrad. After these preliminary calls, the Maxwell solver can be setup by calling HYPRE_SStructMaxwellSetup.

There are two solver cycling schemes that can be used to solve the linear system. To describe these, one needs to consider the augmented system operator

$$
\mathbf{A}=\left[\begin{array}{ll}
A_{e e} & A_{e n} \\
A_{n e} & A_{n n}
\end{array}\right]
$$

where $A_{e e}$ is the stiffness matrix corresponding to the above curl-curl formulation, $A_{n n}$ is the nodal Poisson operator created by taking the Galerkin product of $A_{e e}$ and the gradient operator, and $A_{n e}$ and $A_{e n}$ are the nodal-edge coupling operators (see [JoLe2006]). The algorithm for this Maxwell solver is based on forming a multigrid hierarchy to this augmented system using the block-diagonal interpolation operator

$$
\mathbf{P}=\left[\begin{array}{ll}
P_{e} & 0 \\
0 & P_{n}
\end{array}\right]
$$

where $P_{e}$ and $P_{n}$ are respectively the edge and nodal interpolation operators determined individually from $A_{e e}$ and $A_{n n}$. Taking a Galerkin product between $\mathbf{A}$ and $\mathbf{P}$ produces the next coarse augmented operator, which also has the nodal-edge coupling operators. Applying this procedure recursively produces nodal-edge coupling operators at all levels. Now, the first solver cycling scheme, HYPRE_SStructMaxwellSolve, keeps these coupling operators on all levels of the V-cycle. The second, cheaper scheme, HYPRE_SStructMaxwellSolve2, keeps the coupling operators only on the finest level, i.e., separate edge and nodal V-cycles that couple only on the finest level.

### 6.7 Hybrid

The hybrid solver is designed to detect whether a multigrid preconditioner is needed when solving a linear system and possibly avoid the expensive setup of a preconditioner if a system can be solved efficiently with a diagonally scaled Krylov solver, e.g. a strongly diagonally dominant system. It first uses a diagonally scaled Krylov solver, which can be chosen by the user (the default is conjugate gradient, but one should use GMRES if the matrix of the linear system to be solved is nonsymmetric). It monitors how fast the Krylov solver converges. If there is not sufficient progress, the algorithm switches to a preconditioned Krylov solver.
If used through the Struct interface, the solver is called StructHybrid and can be used with the preconditioners SMG and PFMG (default). It is called ParCSRHybrid, if used through the IJ interface and is used here with BoomerAMG. The user can determine the average convergence speed by setting a convergence tolerance $0 \leq \theta<1$ via the routine HYPRE_StructHybridSetConvergenceTol or HYPRE_ParCSRHybridSetConvergenceTol. The default setting is 0.9 .

The average convergence factor $\rho_{i}=\left(\frac{\left\|r_{i}\right\|}{\left\|r_{0}\right\|}\right)^{1 / i}$ is monitored within the chosen Krylov solver, where $r_{i}=b-A x_{i}$ is the $i$-th residual. Convergence is considered too slow when

$$
\left(1-\frac{\left|\rho_{i}-\rho_{i-1}\right|}{\max \left(\rho_{i}, \rho_{i-1}\right)}\right) \rho_{i}>\theta .
$$

When this condition is fulfilled the hybrid solver switches from a diagonally scaled Krylov solver to a preconditioned solver.

### 6.8 BoomerAMG

BoomerAMG is a parallel implementation of the algebraic multigrid method [RuSt1987]. It can be used both as a solver or as a preconditioner. The user can choose between various different parallel coarsening techniques, interpolation and relaxation schemes. The default settings for CPUs, HMIS (coarsening 8) combined with a distance-two interpolation (6) truncated to 4 or 5 elements per row, should work fairly well for two- and three-dimensional diffusion problems. Additional reduction in complexity and increased scalability can often be achieved using one or two levels of aggressive coarsening.

### 6.8.1 Parameter Options

Various BoomerAMG functions and options are mentioned below. However, for a complete listing and description of all available functions, see the reference manual.
BoomerAMG's Create function differs from the synopsis in that it has only one parameter HYPRE_BoomerAMGCreate(HYPRE_Solver *solver). It uses the communicator of the matrix A.

### 6.8.2 Coarsening Options

Coarsening can be set by the user using the function HYPRE_BoomerAMGSetCoarsenType. A detailed description of various coarsening techniques can be found in [HeYa2002], [Yang2005].
Various coarsening techniques are available:

- the Cleary-Luby-Jones-Plassman (CLJP) coarsening,
- parallel versions of the classical RS coarsening described in [HeYa2002].
- the Falgout coarsening which is a combination of CLJP and the classical RS coarsening algorithm,
- CGC and CGC-E coarsenings [GrMS2006a], [GrMS2006b],
- PMIS and HMIS coarsening algorithms which lead to coarsenings with lower complexities [DeYH2004] as well as
- aggressive coarsening, which can be applied to any of the coarsening techniques mentioned above a nd thus achieving much lower complexities and lower memory use [Stue 1999].
To use aggressive coarsening users have to set the number of levels to which they want to apply aggressive coarsening (starting with the finest level) via HYPRE_BoomerAMGSetAggNumLevels. Since aggressive coarsening requires long range interpolation, multipass interpolation is always used on levels with aggressive coarsening, unless the user specifies another long-range interpolation suitable for aggressive coarsening via HYPRE_BoomerAMGSetAggInterpType..

Note that the default coarsening for CPUs is HMIS, for GPUs PMIS [DeYH2004].

### 6.8.3 Interpolation Options

Various interpolation techniques can be set using HYPRE_BoomerAMGSetInterpType:

- the "classical" interpolation (0) as defined in [RuSt1987],
- direct interpolation (3) [Stue 1999],
- standard interpolation (8) [Stue1999],
- an extended "classical" interpolation, which is a long range interpolation and is recommended to be used with PMIS and HMIS coarsening for harder problems (6) [DFNY2008],
- distance-two interpolation based on matrix operations (17) [LiSY2021],
- multipass interpolation (4) [Stue1999],
- two-stage interpolation [Yang2010],
- Jacobi interpolation [Stue1999],
- the "classical" interpolation modified for hyperbolic PDEs (2).

Jacobi interpolation is only used to improve certain interpolation operators and can be used with HYPRE_BoomerAMGSetPostInterpType. Since some of the interpolation operators might generate large stencils, it is often possible and recommended to control complexity and truncate the interpolation operators using HYPRE_BoomerAMGSetTruncFactor and/or HYPRE_BoomerAMGSetPMaxElmts, or HYPRE_BoomerAMGSetJacobiTruncTheshold (for Jacobi interpolation only).

Note that the default interpolation is extended+i interpolation [DFNY2008] truncated to 4 elements per row, for CPUs, and a version of this interpolation based on matrix operations for GPUs [LiSY2021].

### 6.8.4 Non-Galerkin Options

In order to reduce communication, there is a non-Galerkin coarse grid sparsification option available [FaSc2014]. This option can be used by itself or with existing strategies to reduce communication such as aggressive coarsening and HMIS coarsening. To use, call HYPRE_BoomerAMGSetNonGalerkTol, which gives BoomerAMG a list of level specific non-Galerkin drop tolerances. It is common to drop more aggressively on coarser levels. A common choice of drop-tolerances is $[0.0,0.01,0.05]$ where the value of 0.0 will skip the non-Galerkin process on the first coarse level (level 1), use a drop-tolerance of 0.01 on the second coarse level (level 2) and then use 0.05 on all subsequent coarse levels. While still experimental, this capability has significantly improved performance on a variety of problems. See the ij driver for an example usage and the reference manual for more details.

### 6.8.5 Smoother Options

A good overview of parallel smoothers and their properties can be found in [BFKY2011]. Various of the described relaxation techniques are available:

- weighted Jacobi relaxation (0),
- a hybrid Gauss-Seidel / Jacobi relaxation scheme (3 4),
- a symmetric hybrid Gauss-Seidel / Jacobi relaxation scheme (6),
- l1-Gauss-Seidel or Jacobi (13 1418 8),
- Chebyshev smoothers (16),
- two-stage Gauss-Seidel smoothers (11 12) [BKRHSMTY2021],
- hybrid block and Schwarz smoothers [Yang2004],
- Incomplete LU factorization, see ILU as Smoother for BoomerAMG.
- Factorized Sparse Approximate Inverse (FSAI), see FSAI as Smoother to BoomerAMG.

Point relaxation schemes can be set using HYPRE_BoomerAMGSetRelaxType or, if one wants to specifically set the up cycle, down cycle or the coarsest grid, with HYPRE_BoomerAMGSetCycleRelaxType. To use the more complicated smoothers, e.g. block, Schwarz, ILU smoothers, it is necessary to use HYPRE_BoomerAMGSetSmoothType and HYPRE_BoomerAMGSetSmoothNumLevels. There are further parameter choices for the individual smoothers, which are described in the reference manual. The default relaxation type is 11-Gauss-Seidel, using a forward solve on the down cycle and a backward solve on the up-cycle, to keep symmetry. Note that if BoomerAMG is used as a preconditioner
for conjugate gradient, it is necessary to use a symmetric smoother. Other symmetric options are weighted Jacobi or hybrid symmetric Gauss-Seidel.

### 6.8.6 AMG for systems of PDEs

If the users wants to solve systems of PDEs and can provide information on which variables belong to which function, BoomerAMG's systems AMG version can also be used. Functions that enable the user to access the systems AMG version are HYPRE_BoomerAMGSetNumFunctions, HYPRE_BoomerAMGSetDofFunc and HYPRE_BoomerAMGSetNodal.

There are basically two approaches to deal with matrices derived from systems of PDEs. The unknown-based approach (which is the default) treats variables corresponding to the same unknown or function separately, i.e., when coarsening or generating interpolation, connections between variables associated with different unknowns are ignored. This can work well for weakly coupled PDEs, but will be problematic for strongly coupled PDEs. For such problems, we recommend to use hypre's multigrid reduction (MGR) solver. The second approach, called the nodal approach, considers all unknowns at a physical grid point together such that coarsening, interpolation and relaxation occur in a point-wise fashion. It is possible and sometimes prefered to combine nodal coarsening with unknown-based interpolation. For this case, HYPRE_BoomerAMGSetNodal should be set > 1. For details see the reference manual.

If the user can provide the near null-space vectors, such as the rigid body modes for linear elasticity problems, an interpolation is available that will incorporate these vectors with HYPRE_BoomerAMGSetInterpVectors and HYPRE_BoomerAMGSetInterpVecVariant. This can lead to improved convergence and scalability [BaKY2010].

### 6.8.7 Special AMG Cycles

The default cycle is a $V(1,1)$-cycle, however it is possible to change the number of sweeps of the up- and down-cycle as well as the coare grid. One can also choose a W-cycle, however for parallel processing this is not recommended, since it is not scalable.

BoomerAMG also provides an additive $\mathrm{V}(1,1)$-cycle as well as a mult-additive $\mathrm{V}(1,1)$-cycle and a simplified versioni [VaYa2014]. The additive variants can only be used with weighted Jacobi or 11-Jacobi smoothing.

### 6.8.8 GPU-supported Options

In general, CUDA unified memory is required for running BoomerAMG solvers on GPUs. However, hypre can also be built without --enable-unified-memory if all the selected parameters have GPU-support. The currently available GPU-supported BoomerAMG options include:

- Coarsening: PMIS (8)
- Interpolation: direct (3), BAMG-direct (15), extended (14), extended+i (6) and extended+e (18)
- Aggressive coarsening
- Second-stage interpolation with aggressive coarsening: extended (5) and extended+e (7)
- Smoother: Jacobi (7), 11-Jacobi (18), hybrid Gauss Seidel/SSOR (3 4 6), two-stage Gauss-Seidel $(11,12)$ [BKRHSMTY2021], and Chebyshev (16)
- Relaxation order can be 0 , lexicographic order, or $C / F$ for (7) and (18)


### 6.8.9 Memory locations and execution policies

Hypre provides two user-level memory locations, HYPRE_MEMORY_HOST and HYPRE_MEMORY_DEVICE, where HYPRE_MEMORY_HOST is always the CPU memory while HYPRE_MEMORY_DEVICE can be mapped to different memory spaces based on the configure options of hypre. When built with --with-cuda, --with-hip, --with-sycl, or --with-device-openmp, HYPRE_MEMORY_DEVICE is the GPU device memory, and when built additionally with --enable-unified-memory, it is the GPU unified memory (UM). For a non-GPU build, HYPRE_MEMORY_DEVICE is also mapped to the CPU memory. The default memory location of hypre's matrix and vector objects is HYPRE_MEMORY_DEVICE, which can be changed at runtime by HYPRE_SetMemoryLocation(. . .).

The execution policies define the platform of running computations based on the memory locations of participating objects. The default policy is HYPRE_EXEC_HOST, i.e., executing on the host if the objects are accessible from the host. It can be adjusted by HYPRE_SetExecutionPolicy (. . .). Clearly, this policy only affects objects in UM, since UM is accessible from both CPUs and GPUs.

A sample code of setting up IJ matrix $A$ and solve $A x=b$ using AMG-preconditioned CG on GPUs is shown below.

```
cudaSetDevice(device_id); /* GPU binding */
HYPRE_Initialize(); /* must be the first HYPRE function call */
..
/* AMG in GPU memory (default) */
HYPRE_SetMemoryLocation(HYPRE_MEMORY_DEVICE);
/* setup AMG on GPUs */
HYPRE_SetExecutionPolicy(HYPRE_EXEC_DEVICE);
/* use hypre's SpGEMM instead of vendor implementation */
HYPRE_SetSpGemmUseVendor(FALSE);
/* use GPU RNG */
HYPRE_SetUseGpuRand(TRUE);
if (useHypreGpuMemPool)
{
    /* use hypre's GPU memory pool */
    HYPRE_SetGPUMemoryPoolSize(bin_growth, min_bin, max_bin, max_bytes);
}
else if (useUmpireGpuMemPool)
{
    /* or use Umpire GPU memory pool */
    HYPRE_SetUmpireUMPoolName("HYPRE_UM_POOL_TEST");
    HYPRE_SetUmpireDevicePoolName("HYPRE_DEVICE_POOL_TEST");
}
/* setup IJ matrix A */
HYPRE_IJMatrixCreate(comm, first_row, last_row, first_col, last_col, &ij_A);
HYPRE_IJMatrixSetObjectType(ij_A, HYPRE_PARCSR);
/* GPU pointers; efficient in large chunks */
HYPRE_IJMatrixAddToValues(ij_A, num_rows, num_cols, rows, cols, data);
HYPRE_IJMatrixAssemble(ij_A);
HYPRE_IJMatrixGetObject(ij_A, (void **) &parcsr_A);
/* setup AMG */
HYPRE_ParCSRPCGCreate(comm, &solver);
HYPRE_BoomerAMGCreate(&precon);
HYPRE_BoomerAMGSetRelaxType(precon, rlx_type); /* 3, 4, 6, 7, 18, 11, 12 %/
HYPRE_BoomerAMGSetRelaxOrder(precon, FALSE); /* must be false */
```

(continues on next page)

```
HYPRE_BoomerAMGSetCoarsenType(precon, coarsen_type); /* 8 */
HYPRE_BoomerAMGSetInterpType(precon, interp_type); /* 3, 15, 6, 14, 18 */
HYPRE_BoomerAMGSetAggNumLevels(precon, agg_num_levels);
HYPRE_BoomerAMGSetAggInterpType(precon, agg_interp_type); /* 5 or 7 %/
HYPRE_BoomerAMGSetKeepTranspose(precon, TRUE); /* keep transpose to avoid SpMTV */
HYPRE_BoomerAMGSetRAP2(precon, FALSE); /* RAP in two multiplications
    (default: FALSE) */
HYPRE_ParCSRPCGSetPrecond(solver, HYPRE_BoomerAMGSolve, HYPRE_BoomerAMGSetup,
            precon);
HYPRE_PCGSetup(solver, parcsr_A, b, x);
/* solve */
HYPRE_PCGSolve(solver, parcsr_A, b, x);
HYPRE_Finalize(); /* must be the last HYPRE function call */
```

HYPRE_Initialize() must be called and precede all the other HYPRE_functions, and HYPRE_Finalize() must be called before exiting.

### 6.8.10 Miscellaneous

For best performance, it might be necessary to set certain parameters, which will affect both coarsening and interpolation. One important parameter is the strong threshold, which can be set using the function HYPRE_BoomerAMGSetStrongThreshold. The default value is 0.25 , which appears to be a good choice for diffusion problems. The choice of the strength threshold is problem dependent. For example, elasticity problems often require a larger strength threshold.

### 6.9 AMS

AMS (the Auxiliary-space Maxwell Solver) is a parallel unstructured Maxwell solver for edge finite element discretizations of the variational problem

$$
\begin{equation*}
\text { Find } \mathbf{u} \in \mathbf{V}_{h}: \quad(\alpha \nabla \times \mathbf{u}, \nabla \times \mathbf{v})+(\beta \mathbf{u}, \mathbf{v})=(\mathbf{f}, \mathbf{v}), \quad \text { for all } \mathbf{v} \in \mathbf{V}_{h} \tag{6.1}
\end{equation*}
$$

Here $\mathbf{V}_{h}$ is the lowest order Nedelec (edge) finite element space, and $\alpha>0$ and $\beta \geq 0$ are scalar, or SPD matrix coefficients. AMS was designed to be scalable on problems with variable coefficients, and allows for $\beta$ to be zero in part or the whole domain. In either case the resulting problem is only semidefinite, and for solvability the right-hand side should be chosen to satisfy compatibility conditions.
AMS is based on the auxiliary space methods for definite Maxwell problems proposed in [HiXu2006]. For more details, see [KoVa2009].

### 6.9.1 Overview

Let $\mathbf{A}$ and $\mathbf{b}$ be the stiffness matrix and the load vector corresponding to (6.1). Then the resulting linear system of interest reads,

$$
\begin{equation*}
\mathbf{A} \mathbf{x}=\mathbf{b} \tag{6.2}
\end{equation*}
$$

The coefficients $\alpha$ and $\beta$ are naturally associated with the "stiffness" and "mass" terms of $\mathbf{A}$. Besides $\mathbf{A}$ and $\mathbf{b}$, AMS requires the following additional user input:

1. The discrete gradient matrix $G$ representing the edges of the mesh in terms of its vertices. $G$ has as many rows as the number of edges in the mesh, with each row having two nonzero entries: +1 and -1 in the columns corresponding to the vertices composing the edge. The sign is determined based on the orientation of the edge. We require that $G$ includes all (interior and boundary) edges and vertices.
2. The representations of the constant vector fields $(1,0,0),(0,1,0)$, and $(0,0,1)$ in the $\mathbf{V}_{h}$ basis, given as three vectors: $G_{x}, G_{y}$, and $G_{z}$. Note that since no boundary conditions are imposed on $G$, the above vectors can be computed as $G_{x}=G x, G_{y}=G y$ and $G_{z}=G z$, where $x, y$, and $z$ are vectors representing the coordinates of the vertices of the mesh.

In addition to the above quantities, AMS can utilize the following (optional) information:

- The Poisson matrices $A_{\alpha}$ and $A_{\beta}$, corresponding to assembling of the forms $(\alpha \nabla u, \nabla v)+(\beta u, v)$ and $(\beta \nabla u, \nabla v)$ using standard linear finite elements on the same mesh.

Internally, AMS proceeds with the construction of the following additional objects:

- $A_{G}$ - a matrix associated with the mass term which is either $G^{T} \mathbf{A} G$ or the Poisson matrix $A_{\beta}$ (if given).
- $\Pi$ - the matrix representation of the interpolation operator from vector linear to edge finite elements.
- $\mathbf{A}_{\boldsymbol{\Pi}}$ - a matrix associated with the stiffness term which is either $\boldsymbol{\Pi}^{T} \mathbf{A} \boldsymbol{\Pi}$ or a block-diagonal matrix with diagonal blocks $A_{\alpha}$ (if given).
- $B_{G}$ and $\mathbf{B}_{\Pi}$ - efficient (AMG) solvers for $A_{G}$ and $\mathbf{A}_{\boldsymbol{\Pi}}$.

The solution procedure then is a three-level method using smoothing in the original edge space and subspace corrections based on $B_{G}$ and $\mathbf{B}_{\boldsymbol{\Pi}}$. We can employ a number of options here utilizing various combinations of the smoother and solvers in additive or multiplicative fashion. If $\beta$ is identically zero one can skip the subspace correction associated with $G$, in which case the solver is a two-level method.

### 6.9.2 Sample Usage

AMS can be used either as a solver or as a preconditioner. Below we list the sequence of hypre calls needed to create and use it as a solver. See example code ex15.c for a complete implementation. We start with the allocation of the HYPRE_Solver object:

```
HYPRE_Solver solver;
HYPRE_AMSCreate(&solver);
```

Next, we set a number of solver parameters. Some of them are optional, while others are necessary in order to perform the solver setup.

AMS offers the option to set the space dimension. By default we consider the dimension to be 3 . The only other option is 2 , and it can be set with the function given below. We note that a 3 D solver will still work for a 2 D problem, but it will be slower and will require more memory than necessary.

```
HYPRE_AMSSetDimension(solver, dim);
```

The user is required to provide the discrete gradient matrix $G$. AMS expects a matrix defined on the whole mesh with no boundary edges/nodes excluded. It is essential to not impose any boundary conditions on $G$. Regardless of which hypre conceptual interface was used to construct $G$, one can obtain a ParCSR version of it. This is the expected format in the following function.

```
HYPRE_AMSSetDiscreteGradient(solver, G);
```

In addition to $G$, we need one additional piece of information in order to construct the solver. The user has the option to choose either the coordinates of the vertices in the mesh or the representations of the constant vector fields in the edge element basis. In both cases three hypre parallel vectors should be provided. For 2D problems, the user can set the third vector to NULL. The corresponding function calls read:

```
HYPRE_AMSSetCoordinateVectors(solver,x,y,z);
```

or
HYPRE_AMSSetEdgeConstantVectors(solver, one_zero_zero, zero_one_zero, zero_zero_one);

The vectors one_zero_zero, zero_one_zero and zero_zero_one above correspond to the constant vector fields $(1,0,0),(0,1,0)$ and $(0,0,1)$.

The remaining solver parameters are optional. For example, the user can choose a different cycle type by calling

```
HYPRE_AMSSetCycleType(solver, cycle_type); /* default value: 1 */
```

The available cycle types in AMS are:

- cycle_type=1: multiplicative solver (01210)
- cycle_type=2: additive solver $(0+1+2)$
- cycle_type=3: multiplicative solver (02120)
- cycle_type=4: additive solver $(010+2)$
- cycle_type=5: multiplicative solver (0102010)
- cycle_type=6: additive solver $(1+020)$
- cycle_type=7: multiplicative solver (0201020)
- cycle_type=8: additive solver $(0(1+2) 0)$
- cycle_type=11: multiplicative solver (013454310)
- cycle_type=12: additive solver $(0+1+3+4+5)$
- cycle_type=13: multiplicative solver (034515430)
- cycle_type=14: additive solver $(01(3+4+5) 10)$

Here we use the following convention for the three subspace correction methods: 0 refers to smoothing, 1 stands for BoomerAMG based on $B_{G}$, and 2 refers to a call to BoomerAMG for $\mathbf{B}_{\boldsymbol{\Pi}}$. The values 3,4 and 5 refer to the scalar subspaces corresponding to the $x, y$ and $z$ components of $\Pi$.

The abbreviation $x y y z$ for $x, y, z \in\{0,1,2,3,4,5\}$ refers to a multiplicative subspace correction based on solvers $x, y$, $y$, and $z$ (in that order). The abbreviation $x+y+z$ stands for an additive subspace correction method based on $x, y$ and $z$ solvers. The additive cycles are meant to be used only when AMS is called as a preconditioner. In our experience the choices cycle_type $=1,5,8,11,13$ often produced fastest solution times, while cycle_type=7 resulted in smallest number of iterations.

Additional solver parameters, such as the maximum number of iterations, the convergence tolerance and the output level, can be set with

```
HYPRE_AMSSetMaxIter(solver, maxit); /* default value: 20 %/
HYPRE_AMSSetTol(solver, tol); /* default value: 1e-6 */
HYPRE_AMSSetPrintLevel(solver, print); /* default value: 1 */
```

More advanced parameters, affecting the smoothing and the internal AMG solvers, can be set with the following three functions:

```
HYPRE_AMSSetSmoothingOptions(solver, 2, 1, 1.0, 1.0);
HYPRE_AMSSetAlphaAMGOptions(solver, 10, 1, 3, 0.25, 0, 0);
HYPRE_AMSSetBetaAMGOptions(solver, 10, 1, 3, 0.25, 0, 0);
```

For (singular) problems where $\beta=0$ in the whole domain, different (in fact simpler) version of the AMS solver is offered. To allow for this simplification, use the following hypre call

```
HYPRE_AMSSetBetaPoissonMatrix(solver, NULL);
```

If $\beta$ is zero only in parts of the domain, the problem is still singular, but the AMS solver will try to detect this and construct a non-singular preconditioner. Though this often works well in practice, AMS also provides a more robust version for solving such singular problems to very low convergence tolerances. This version takes advantage of additional information: the list of nodes which are interior to a zero-conductivity region provided by the function

```
HYPRE_AMSSetInteriorNodes(solver, HYPRE_ParVector interior_nodes);
```

A node is interior, if its entry in the interior_nodes array is 1.0. Based on this array, a restricted discrete gradient operator $G_{0}$ is constructed, and AMS is then defined based on the matrix $\mathbf{A}+\delta G_{0}^{T} G_{0}$ which is non-singular, and a small $\delta>0$ perturbation of $\mathbf{A}$. When iterating with this preconditioner, it is advantageous to project on the compatible subspace $\operatorname{Ker}\left(G_{0}^{T}\right)$. This can be done periodically, or manually through the functions

```
HYPRE_AMSSetProjectionFrequency(solver, int projection_frequency);
HYPRE_AMSProjectOutGradients(solver, HYPRE_ParVector x);
```

Two additional matrices are constructed in the setup of the AMS method-one corresponding to the coefficient $\alpha$ and another corresponding to $\beta$. This may lead to prohibitively high memory requirements, and the next two function calls may help to save some memory. For example, if the Poisson matrix with coefficient $\beta$ (denoted by Abeta) is available then one can avoid one matrix construction by calling

```
HYPRE_AMSSetBetaPoissonMatrix(solver, Abeta);
```

Similarly, if the Poisson matrix with coefficient $\alpha$ is available (denoted by Aalpha) the second matrix construction can also be avoided by calling

```
HYPRE_AMSSetAlphaPoissonMatrix(solver, Aalpha);
```

Note the following regarding these functions:

- Both of them change their input. More specifically, the diagonal entries of the input matrix corresponding to eliminated degrees of freedom (due to essential boundary conditions) are penalized.
- It is assumed that their essential boundary conditions of $\mathbf{A}$, Abeta and Aalpha are on the same part of the boundary.
- HYPRE_AMSSetAlphaPoissonMatrix forces the AMS method to use a simpler, but weaker (in terms of convergence) method. With this option, the multiplicative AMS cycle is not guaranteed to converge with the default parameters. The reason for this is the fact the solver is not variationally obtained from the original matrix (it utilizes the auxiliary Poisson-like matrices Abeta and Aalpha). Therefore, it is recommended in this case to use AMS as preconditioner only.

After the above calls, the solver is ready to be constructed. The user has to provide the stiffness matrix $\mathbf{A}$ (in ParCSR format) and the hypre parallel vectors $\mathbf{b}$ and $\mathbf{x}$. (The vectors are actually not used in the current AMS setup.) The setup call reads,

```
HYPRE_AMSSetup(solver, A, b, x);
```

It is important to note the order of the calling sequence. For example, do not call HYPRE_AMSSetup before calling HYPRE_AMSSetDiscreteGradient and one of the functions HYPRE_AMSSetCoordinateVectors or HYPRE_AMSSetEdgeConstantVectors.

Once the setup has completed, we can solve the linear system by calling

```
HYPRE_AMSSolve(solver, A, b, x);
```

Finally, the solver can be destroyed with

```
HYPRE_AMSDestroy(&solver);
```

More details can be found in the files ams.h and ams.c located in the parcsr_ls directory.

### 6.9.3 High-order Discretizations

In addition to the interface for the lowest-order Nedelec elements described in the previous subsections, AMS also provides support for (arbitrary) high-order Nedelec element discretizations. Since the robustness of AMS depends on the performance of BoomerAMG on the associated (high-order) auxiliary subspace problems, we note that the convergence may not be optimal for large polynomial degrees $k \geq 1$.

In the high-order AMS interface, the user does not need to provide the coordinates of the vertices (or the representations of the constant vector fields in the edge basis), but instead should construct and pass the Nedelec interpolation matrix $\Pi$ which maps (high-order) vector nodal finite elements into the (high-order) Nedelec space. In other words, $\boldsymbol{\Pi}$ is the (parallel) matrix representation of the interpolation mapping from $\mathrm{P}_{k}^{3} / \mathrm{Q}_{k}^{3}$ into $\mathrm{ND}_{k}$, see [HiXu2006], [KoVa2009]. We require this matrix as an input, since in the high-order case its entries very much depend on the particular choice of the basis functions in the edge and nodal spaces, as well as on the geometry of the mesh elements. The columns of $\Pi$ should use a node-based numbering, where the $x / y / z$ components of the first node (vertex or high-order degree of freedom) should be listed first, followed by the $x / y / z$ components of the second node and so on (see the documentation of HYPRE_BoomerAMGSetDofFunc).

Similarly to the Nedelec interpolation, the discrete gradient matrix $G$ should correspond to the mapping $\varphi \in \mathrm{P}_{k}^{3} / \mathrm{Q}_{k}^{3} \mapsto$ $\nabla \varphi \in \mathrm{ND}_{k}$, so even though its values are still independent of the mesh coordinates, they will not be $\pm 1$, but will be determined by the particular form of the high-order basis functions and degrees of freedom.

With these matrices, the high-order setup procedure is simply

```
HYPRE_AMSSetDimension(solver, dim);
HYPRE_AMSSetDiscreteGradient(solver, G);
HYPRE_AMSSetInterpolations(solver, Pi, NULL, NULL, NULL);
```

We remark that the above interface calls can also be used in the lowest-order case (or even other types of discretizations such as those based on the second family of Nedelec elements), but we recommend calling the previously described HYPRE_AMSSetCoordinateVectors instead, since this allows AMS to handle the construction and use of $\Pi$ internally.

Specifying the monolithic $\Pi$ limits the AMS cycle type options to those less than 10. Alternatively one can separately specify the $x, y$ and $z$ components of $\Pi$ :

HYPRE_AMSSetInterpolations(solver, NULL, Pix, Piy, Piz);
which enables the use of AMS cycle types with index greater than 10 . By definition, $\boldsymbol{\Pi}^{x} \varphi=\boldsymbol{\Pi}(\varphi, 0,0)$, and similarly for $\Pi^{y}$ and $\Pi^{z}$. Each of these matrices has the same sparsity pattern as $G$, but their entries depend on the coordinates of the mesh vertices.

Finally, both $\Pi$ and its components can be passed to the solver:

```
HYPRE_AMSSetInterpolations(solver, Pi, Pix, Piy, Piz);
```

which will duplicate some memory, but allows for experimentation with all available AMS cycle types.

### 6.9.4 Non-conforming AMR Grids

AMS could also be applied to problems with adaptive mesh refinement (AMR) posed on non-conforming quadrilat$\mathrm{eral} / \mathrm{hexahedral}$ meshes, see [GrKo2015] for more details.

On non-conforming grids (assuming also arbitrarily high-order elements), each finite element space has two versions: a conforming one, e.g. $\mathrm{Q}_{k}^{c} / \mathrm{ND}_{k}^{c}$, where the hanging degrees of freedom are constrained by the conforming (real) degrees of freedom, and a non-conforming one, e.g. $\mathrm{Q}_{k}^{n c} / \mathrm{ND}_{k}^{n c}$ where the non-conforming degrees of freedom (hanging and real) are unconstrained. These spaces are related with the conforming prolongation and the pure restriction operators $P$ and $R$, as well as the conforming and non-conforming version of the discrete gradient operator as follows:


Since the linear system is posed on $\mathrm{ND}_{k}^{c}$, the user needs to provide the conforming discrete gradient matrix $G_{c}$ to AMS, using HYPRE_AMSSetDiscreteGradient. This matrix is defined by the requirement that the above diagram commutes from $\mathrm{Q}_{k}^{c}$ to $\mathrm{ND}_{k}^{n c}$, corresponding to the definition

$$
G_{c}=R_{\mathrm{ND}} G_{n c} P_{\mathrm{Q}}
$$

i.e. the conforming gradient is computed by starting with a conforming nodal $\mathrm{Q}_{k}$ function, interpolating it in the hanging nodes, computing the gradient locally and representing it in the Nedelec space on each element (the nonconforming discrete gradient $G_{n c}$ in the above formula), and disregarding the values in the hanging $\mathrm{ND}_{k}$ degrees of freedom.

Similar considerations imply that the conforming Nedelec interpolation matrix $\boldsymbol{\Pi}_{c}$ should be defined as

$$
\boldsymbol{\Pi}_{c}=R_{\mathrm{ND}} \boldsymbol{\Pi}_{n c} P_{\mathrm{Q}^{3}},
$$

with $\boldsymbol{\Pi}_{n c}$ computed element-wise as in the previous subsection. Note that in the low-order case, $\boldsymbol{\Pi}_{c}$ can be computed internally in AMS based only $G_{c}$ and the conforming coordinates of the vertices $x_{c} / y_{c} / z_{c}$, see [ GrKo 2015 ].

### 6.10 ADS

The Auxiliary-space Divergence Solver (ADS) is a parallel unstructured solver similar to AMS, but targeting $H$ (div) instead of $H$ (curl) problems. Its usage and options are very similar to those of AMS, and in general the relationship between ADS and AMS is analogous to that between AMS and AMG.

Specifically ADS was designed for the scalable solution of linear systems arising in the finite element discretization of the variational problem

$$
\begin{equation*}
\text { Find } \mathbf{u} \in \mathbf{W}_{h}: \quad(\alpha \nabla \cdot \mathbf{u}, \nabla \cdot \mathbf{v})+(\beta \mathbf{u}, \mathbf{v})=(\mathbf{f}, \mathbf{v}), \quad \text { for all } \mathbf{v} \in \mathbf{W}_{h} \tag{6.3}
\end{equation*}
$$

where $\mathbf{W}_{h}$ is the lowest order Raviart-Thomas (face) finite element space, and $\alpha>0$ and $\beta>0$ are scalar, or SPD matrix variable coefficients. It is based on the auxiliary space methods for $H$ (div) problems proposed in [HiXu2006].

### 6.10.1 Overview

Let $\mathbf{A}$ and $\mathbf{b}$ be the stiffness matrix and the load vector corresponding to (6.3). Then the resulting linear system of interest reads,

$$
\begin{equation*}
\mathbf{A x}=\mathbf{b} \tag{6.4}
\end{equation*}
$$

The coefficients $\alpha$ and $\beta$ are naturally associated with the "stiffness" and "mass" terms of $\mathbf{A}$. Besides $\mathbf{A}$ and $\mathbf{b}$, ADS requires the following additional user input:

1. The discrete curl matrix $C$ representing the faces of the mesh in terms of its edges. $C$ has as many rows as the number of faces in the mesh, with each row having nonzero entries +1 and -1 in the columns corresponding to the edges composing the face. The sign is determined based on the orientation of the edges relative to the face. We require that $C$ includes all (interior and boundary) faces and edges.
2. The discrete gradient matrix $G$ representing the edges of the mesh in terms of its vertices. $G$ has as many rows as the number of edges in the mesh, with each row having two nonzero entries: +1 and -1 in the columns corresponding to the vertices composing the edge. The sign is determined based on the orientation of the edge. We require that $G$ includes all (interior and boundary) edges and vertices.
3. Vectors $x, y$, and $z$ representing the coordinates of the vertices of the mesh.

Internally, ADS proceeds with the construction of the following additional objects:

- $A_{C}$ - the curl-curl matrix $C^{T} \mathbf{A} C$.
- $\Pi$ - the matrix representation of the interpolation operator from vector linear to face finite elements.
- $\mathbf{A}_{\boldsymbol{\Pi}}$ - the vector nodal matrix $\boldsymbol{\Pi}^{T} \mathbf{A} \boldsymbol{\Pi}$.
- $B_{C}$ and $\mathbf{B}_{\Pi}$ - efficient (AMS/AMG) solvers for $A_{C}$ and $\mathbf{A}_{\boldsymbol{\Pi}}$.

The solution procedure then is a three-level method using smoothing in the original face space and subspace corrections based on $B_{C}$ and $\mathbf{B}_{\Pi}$. We can employ a number of options here utilizing various combinations of the smoother and solvers in additive or multiplicative fashion.

### 6.10.2 Sample Usage

ADS can be used either as a solver or as a preconditioner. Below we list the sequence of hypre calls needed to create and use it as a solver. We start with the allocation of the HYPRE_Solver object:

```
HYPRE_Solver solver;
HYPRE_ADSCreate(&solver);
```

Next, we set a number of solver parameters. Some of them are optional, while others are necessary in order to perform the solver setup.

The user is required to provide the discrete curl and gradient matrices $C$ and $G$. ADS expects a matrix defined on the whole mesh with no boundary faces, edges or nodes excluded. It is essential to not impose any boundary conditions on $C$ or $G$. Regardless of which hypre conceptual interface was used to construct the matrices, one can always obtain a ParCSR version of them. This is the expected format in the following functions.

```
HYPRE_ADSSetDiscreteCurl(solver, C);
HYPRE_ADSSetDiscreteGradient(solver, G);
```

Next, ADS requires the coordinates of the vertices in the mesh as three hypre parallel vectors. The corresponding function call reads:

```
HYPRE_ADSSetCoordinateVectors(solver, x, y, z);
```

The remaining solver parameters are optional. For example, the user can choose a different cycle type by calling

```
HYPRE_ADSSetCycleType(solver, cycle_type); /* default value: 1 */
```

The available cycle types in ADS are:

- cycle_type=1: multiplicative solver (01210)
- cycle_type=2: additive solver $(0+1+2)$
- cycle_type=3: multiplicative solver (02120)
- cycle_type=4: additive solver $(010+2)$
- cycle_type=5: multiplicative solver (0102010)
- cycle_type=6: additive solver $(1+020)$
- cycle_type=7: multiplicative solver (0201020)
- cycle_type=8: additive solver $(0(1+2) 0)$
- cycle_type=11: multiplicative solver (013454310)
- cycle_type=12: additive solver $(0+1+3+4+5)$
- cycle_type=13: multiplicative solver (034515430)
- cycle_type=14: additive solver $(01(3+4+5) 10)$

Here we use the following convention for the three subspace correction methods: 0 refers to smoothing, 1 stands for AMS based on $B_{C}$, and 2 refers to a call to BoomerAMG for $\mathbf{B}_{\boldsymbol{\Pi}}$. The values 3,4 and 5 refer to the scalar subspaces corresponding to the $x, y$ and $z$ components of $\Pi$.

The abbreviation $x y y z$ for $x, y, z \in\{0,1,2,3,4,5\}$ refers to a multiplicative subspace correction based on solvers $x, y$, $y$, and $z$ (in that order). The abbreviation $x+y+z$ stands for an additive subspace correction method based on $x, y$ and $z$ solvers. The additive cycles are meant to be used only when ADS is called as a preconditioner. In our experience the
choices cycle_type $=1,5,8,11,13$ often produced fastest solution times, while cycle_type $=7$ resulted in smallest number of iterations.

Additional solver parameters, such as the maximum number of iterations, the convergence tolerance and the output level, can be set with

```
HYPRE_ADSSetMaxIter(solver, maxit); /* default value: 20 */
HYPRE_ADSSetTol(solver, tol); /* default value: 1e-6 */
HYPRE_ADSSetPrintLevel(solver, print); /* default value: 1 */
```

More advanced parameters, affecting the smoothing and the internal AMS and AMG solvers, can be set with the following three functions:

```
HYPRE_ADSSetSmoothingOptions(solver, 2, 1, 1.0, 1.0);
HYPRE_ADSSetAMSOptions(solver, 11, 10, 1, 3, 0.25, 0, 0);
HYPRE_ADSSetAMGOptions(solver, 10, 1, 3, 0.25, 0, 0);
```

We note that the AMS cycle type, which is the second parameter of HYPRE_ADSSetAMSOptions should be greater than 10 , unless the high-order interface of HYPRE_ADSSetInterpolations described in the next subsection is being used.

After the above calls, the solver is ready to be constructed. The user has to provide the stiffness matrix $\mathbf{A}$ (in ParCSR format) and the hypre parallel vectors $\mathbf{b}$ and $\mathbf{x}$. (The vectors are actually not used in the current ADS setup.) The setup call reads,

```
HYPRE_ADSSetup(solver, A, b, x);
```

It is important to note the order of the calling sequence. For example, do not call HYPRE_ADSSetup before calling each of the functions HYPRE_ADSSetDiscreteCurl, HYPRE_ADSSetDiscreteGradient and HYPRE_ADSSetCoordinateVectors.

Once the setup has completed, we can solve the linear system by calling

```
HYPRE_ADSSolve(solver, A, b, x);
```

Finally, the solver can be destroyed with

```
HYPRE_ADSDestroy(&solver);
```

More details can be found in the files ads. h and ads. c located in the parcsr_ls directory.

### 6.10.3 High-order Discretizations

Similarly to AMS, ADS also provides support for (arbitrary) high-order $H$ (div) discretizations. Since the robustness of ADS depends on the performance of AMS and BoomerAMG on the associated (high-order) auxiliary subspace problems, we note that the convergence may not be optimal for large polynomial degrees $k \geq 1$.

In the high-order ADS interface, the user does not need to provide the coordinates of the vertices, but instead should construct and pass the Raviart-Thomas and Nedelec interpolation matrices $\boldsymbol{\Pi}_{R T}$ and $\boldsymbol{\Pi}_{N D}$ which map (high-order) vector nodal finite elements into the (high-order) Raviart-Thomas and Nedelec space. In other words, these are the (parallel) matrix representation of the interpolation mappings from $\mathrm{P}_{k}^{3} / \mathrm{Q}_{k}^{3}$ into $\mathrm{RT}_{k-1}$ and $\mathrm{ND}_{k}$, see [HiXu2006], [KoVa2009]. We require these matrices as inputs, since in the high-order case their entries very much depend on the particular choice of the basis functions in the finite element spaces, as well as on the geometry of the mesh elements. The columns of the $\Pi$ matrices should use a node-based numbering, where the $x / y / z$ components of the first node (vertex or high-order degree of freedom) should be listed first, followed by the $x / y / z$ components of the second node
and so on (see the documentation of HYPRE_BoomerAMGSetDofFunc). Furthermore, each interpolation matrix can be split into $x, y$ and $z$ components by defining $\Pi^{x} \varphi=\boldsymbol{\Pi}(\varphi, 0,0)$, and similarly for $\boldsymbol{\Pi}^{y}$ and $\Pi^{z}$.
The discrete gradient and curl matrices $G$ and $C$ should correspond to the mappings $\varphi \in \mathrm{P}_{k}^{3} / \mathrm{Q}_{k}^{3} \mapsto \nabla \varphi \in \mathrm{ND}_{k}$ and $\mathbf{u} \in \mathrm{ND}_{k} \mapsto \nabla \times \mathbf{u} \in \mathrm{RT}_{k-1}$, so even though their values are still independent of the mesh coordinates, they will not be $\pm 1$, but will be determined by the particular form of the high-order basis functions and degrees of freedom.

With these matrices, the high-order setup procedure is simply

```
HYPRE_ADSSetDiscreteCurl(solver, C);
HYPRE_ADSSetDiscreteGradient(solver, G);
HYPRE_ADSSetInterpolations(solver, RT_Pi, NULL, NULL, NULL,
    ND_Pi, NULL, NULL, NULL);
```

We remark that the above interface calls can also be used in the lowest-order case (or even other types of discretizations), but we recommend calling the previously described HYPRE_ADSSetCoordinateVectors instead, since this allows ADS to handle the construction and use of the interpolations internally.

Specifying the monolithic $\Pi_{R T}$ limits the ADS cycle type options to those less than 10. Alternatively one can separately specify the $x, y$ and $z$ components of $\boldsymbol{\Pi}_{R T}$.

```
HYPRE_ADSSetInterpolations(solver, NULL, RT_Pix, RT_Piy, RT_Piz,
```

ND_Pi, NULL, NULL, NULL);
which enables the use of ADS cycle types with index greater than 10 . The same holds for $\boldsymbol{\Pi}_{N D}$ and its components, e.g. to enable the subspace AMS cycle type greater then 10 we need to call

```
HYPRE_ADSSetInterpolations(solver, NULL, RT_Pix, RT_Piy, RT_Piz,
    NULL, ND_Pix, ND_Piy, ND_Piz);
```

Finally, both $\Pi$ and their components can be passed to the solver:

```
HYPRE_ADSSetInterpolations(solver, RT_Pi, RT_Pix, RT_Piy, RT_Piz
    ND_Pi, ND_Pix, ND_Piy, ND_Piz);
```

which will duplicate some memory, but allows for experimentation with all available ADS and AMS cycle types.

### 6.11 The MLI Package

MLI is an object-oriented module that implements the class of algebraic multigrid algorithms based on Vanek and Brezina's smoothed aggregation method [VaMB1996], [VaBM2001]. There are two main algorithms in this module the original smoothed aggregation algorithm and the modified version that uses the finite element substructure matrices to construct the prolongation operators. As such, the later algorithm can only be used in the finite element context via the finite element interface. In addition, the nodal coordinates obtained via the finite element interface can be used to construct a better prolongation operator than the pure translation modes.

Below is an example on how to set up MLI as a preconditioner for conjugate gradient.

```
HYPRE_LSI_MLICreate(MPI_COMM_WORLD, &pcg_precond);
HYPRE_LSI_MLISetParams(pcg_precond, "MLI strengthThreshold 0.08");
HYPRE_PCGSetPrecond(pcg_solver,
```

(continues on next page)

Note that parameters are set via HYPRE_LSI_MLISetParams. A list of valid parameters that can be set using this routine can be found in the FEI section of the reference manual.

### 6.12 Multigrid Reduction (MGR)

MGR is a parallel multigrid reduction solver and preconditioner designed to take advantage of use-provided information to solve systems of equations with multiple vatiable types. The algorithm is similar to two-stage preconditioner strategies and other reduction techniques like ARMS, but in a standard multigrid framework.
The MGR algorithm accepts information about the variables in block form from the user and uses it to define the appropriate C/F splitting for the multigrid scheme. The linear system solve proceeds with an F-relaxation solve on the F points, folowed by a coarse grid correction. The coarse grid solve is handled by scalar AMG (BoomerAMG). MGR provides users with more control over the coarsening process, and can potentially be a starting point for designing multigrid-based physics-based preconditioners.

The following represents a minimal set of functions, and some optional functions, to call to use the MGR solver. For simplicity, we ignore the function parameters here, and refer the reader to the reference manual for more details on the parameters and their defaults.

- HYPRE_MGRCreate: Create the MGR solver object.
- HYPRE_MGRSetCpointsByBlock: Set up block data with information about coarse indexes for reduction. Here, the user specifies the number of reduction levels, as well as the the coarse nodes for each level of the reduction. These coarse nodes are indexed by their index in the block of unknowns. This is used internally to tag the appropriate indexes of the linear system matrix as coarse nodes.
- (Optional) HYPRE_MGRSetReservedCoarseNodes: Prescribe a subset of nodes to be kept as coarse nodes until the coarsest level. These nodes are transferred onto the coarsest grid of the BoomerAMG coarse grid solver.
- (Optional) HYPRE_MGRSetNonCpointsToFpoints: Set points not prescribed as C points to be fixed as F points for intermediate levels. Setting this to 1 uses the user input to define the C/F splitting. Otherwise, a BoomerAMG
coarsening routine is used to determine the $\mathrm{C} / \mathrm{F}$ splitting for intermediate levels.
- (Optional) HYPRE_MGRSetCoarseSolver: This function sets the BoomerAMG solver to be used for the solve on the coarse grid. The user can define their own BoomerAMG solver with their preferred options and pass this to the MGR solver. Otherwise, an internal BoomerAMG solver is used as the coarse grid solver instead.
- HYPRE_MGRSetup: Setup and MGR solver object.
- HYPRE_MGRSolve: Solve the linear system.
- HYPRE_MGRDestroy: Destroy the MGR solver object

For more details about additional solver options and parameters, please refer to the reference manual. NOTE: The MGR solver is currently only supported by the IJ interface.

### 6.13 FSAI

FSAI is a parallel implementation of the Factorized Sparse Approximate Inverse preconditioner, initially proposed by [KoYe1993]. Given a symmetric positive definite matrix $A$, FSAI computes a triangular matrix $G$ that approximates the inverse of the lower Cholesky factor $(L)$ of $A$. This computation is done by minimizing the Frobenius norm $\|I-G L\|_{F}$ without explicit knowledge of $L$. The resulting preconditioner preserves the positive definiteness of $A$ and is given by $M^{-1}=G^{T} G$.

One of the critical factors determining the quality of sparse approximate inverse preconditioners lies in choosing the sparsity pattern of $G$. While ParaSails employs a priori sparsity patterns, FSAI uses an iterative strategy that generates sparsity patterns on the fly, i.e., while computing their nonzero coefficient values concurrently. At every step of the iterative process, the sparsity pattern of a row of $G$ is augmented with a fixed number of entries, leading to the most significant reduction of the conditioning number of $G A G^{T}$. Such a strategy is also called "adaptive FSAI" or "dynamic FSAI" and it can lead to more robust sparse approximate inverses than ParaSails. For more details on how it works, see [JaFe2015].

### 6.13.1 Parameter Settings

The accuracy and cost of FSAI are determined by three configurations parameters as shown in the table below

| param | type | range | sug. values | default |
| :--- | :--- | :--- | :--- | :--- |
| max_steps | int | $\geq 0$ | $5,10,30$ | 5 |
| max_step_size | int | $\geq 0$ | $1,3,6$ | 3 |
| kap_tolerance | real | $\geq 0$ | $0.0,1 . E-2,1 . E-3$ | 1.E-3 |

The first parameter, max_steps, controls the number of maximum steps used in the iterative algorithm. The second parameter, max_step_size, gives the maximum number of indices added to the sparsity pattern of $G$ at each step. Lastly, the third parameter, kap_tolerance, is a floating-point value used to stop the inclusion of new indices to the sparsity pattern of $G$ when the conditioning number of $G A G^{T}$ stagnates. This can be disabled by setting kap_tolerance = 0 . Naturally, the preconditioner quality increases for denser sparsity patterns of $G$, but so do its setup and solve costs. For a reasonable balance between accuracy and cost, we recommend that max_steps $*$ max_step_size $\leq 30$. The configuration parameters of FSAI can be set via the following calls:

```
HYPRE_FSAISetMaxSteps(HYPRE_Solver solver, HYPRE_Int max_steps);
HYPRE_FSAISetMaxStepSize(HYPRE_Solver solver, HYPRE_Int max_step_size);
HYPRE_FSAISetKapTolerance(HYPRE_Solver solver, HYPRE_Real kap_tolerance);
```


### 6.13.2 FSAI as Smoother to BoomerAMG

As discussed in [PaFa2019], the factorized sparse approximate inverse method can be an effective smoother to AMG for several reasons. Particularly, it leads to a symmetric operator, and thus allows AMG to be used as a preconditioner for the conjugate gradient solver. In hypre, FSAI can be used as a complex smoother to BoomerAMG by calling the functions:

```
HYPRE_BoomerAMGSetSmoothType(HYPRE_Solver solver, 4);
HYPRE_BoomerAMGSetSmoothNumLevels(HYPRE_Solver solver, HYPRE_Int num_levels);
```

where num_levels is the last multigrid level where FSAI is used. The configuration parameters of the FSAI smoother, as described above, can be set via the following calls:

```
HYPRE_BoomerAMGSetFSAIMaxSteps(HYPRE_Solver solver, HYPRE_Int max_steps);
HYPRE_BoomerAMGSetFSAIMaxStepSize(HYPRE_Solver solver, HYPRE_Int max_step_size);
HYPRE_BoomerAMGSetFSAIKapTolerance(HYPRE_Solver solver, HYPRE_Real kap_tolerance);
```


### 6.13.3 Implementation Notes

- When the matrix $A$ is distributed across MPI tasks, FSAI considers only the block diagonal portions of $A$ for computing $G$. The resulting preconditioner is effectively a block-Jacobi sparse approximate inverse in the MPI sense. Although this strategy reduces communication costs, it can degrade convergence performance when several tasks are used, especially when FSAI is employed as a preconditioner to a Krylov solver.
- The CPU version of FSAI supports threading via OpenMP. To enable it, users need to compile hypre with OpenMP support via the configure option --with-openmp. In this case, FSAI relies on an implementation of BLAS/LAPACK that is thread-safe. The one distributed internally with hypre fulfills this criterion, but care must be taken when linking hypre to external BLAS/LAPACK libraries. In HPC platforms, we recommend using vendor implementations of BLAS/LAPACK for better setup performance of FSAI, regardless of whether using OpenMP or not.
- The GPU version of FSAI is under development.


### 6.14 ParaSails

Warning: ParaSails is not actively supported by the hypre development team. We recommend using FSAI for parallel sparse approximate inverse algorithms. This new implementation includes NVIDIA/AMD GPU support through the CUDA/HIP backends.

ParaSails is a parallel implementation of a sparse approximate inverse preconditioner, using a priori sparsity patterns and least-squares (Frobenius norm) minimization. Symmetric positive definite (SPD) problems are handled using a factored SPD sparse approximate inverse. General (nonsymmetric and/or indefinite) problems are handled with an unfactored sparse approximate inverse. It is also possible to precondition nonsymmetric but definite matrices with a factored, SPD preconditioner.

ParaSails uses a priori sparsity patterns that are patterns of powers of sparsified matrices. ParaSails also uses a postfiltering technique to reduce the cost of applying the preconditioner. In advanced usage not described here, the pattern of the preconditioner can also be reused to generate preconditioners for different matrices in a sequence of linear solves.

For more details about the ParaSails algorithm, see [Chow2000].

### 6.14.1 Parameter Settings

The accuracy and cost of ParaSails are parametrized by the real thresh and integer nlevels parameters, $0 \leq$ thresh $\leq 1,0 \leq$ nlevels. Lower values of thresh and higher values of nlevels lead to more accurate, but more expensive preconditioners. More accurate preconditioners are also more expensive per iteration. The default values are thresh $=0.1$ and nlevels $=1$. The parameters are set using HYPRE_ParaSailsSetParams.

Mathematically, given a symmetric matrix $A$, the pattern of the approximate inverse is the pattern of $\tilde{A}^{m}$ where $\tilde{A}$ is a matrix that has been sparsified from $A$. The sparsification is performed by dropping all entries in a symmetrically diagonally scaled $A$ whose values are less than thresh in magnitude. The parameter nlevel is equivalent to $m-1$. Filtering is a post-thresholding procedure. For more details about the algorithm, see [Chow2000].

The storage required for the ParaSails preconditioner depends on the parameters thresh and nlevels. The default parameters often produce a preconditioner that can be stored in less than the space required to store the original matrix. ParaSails does not need a large amount of intermediate storage in order to construct the preconditioner.

ParaSail's Create function differs from the synopsis in the following way:

```
int HYPRE_ParaSailsCreate(MPI_Comm comm, HYPRE_Solver *solver, int symmetry);
```

where comm is the MPI communicator.
The value of symmetry has the following meanings, to indicate the symmetry and definiteness of the problem, and to specify the type of preconditioner to construct:

| value | meaning |
| :--- | :--- |
| 0 | nonsymmetric and/or indefinite problem, and nonsymmetric preconditioner |
| 1 | SPD problem, and SPD (factored) preconditioner |
| 2 | nonsymmetric, definite problem, and SPD (factored) preconditioner |

For more information about the final case, see section Preconditioning Nearly Symmetric Matrices.
Parameters for setting up the preconditioner are specified using

```
int HYPRE_ParaSailsSetParams(HYPRE_Solver solver, double thresh,
    int nlevel, double filter);
```

The parameters are used to specify the sparsity pattern and filtering value (see above), and are described with suggested values as follows:

| param | type | range | sug. values | default | meaning |
| :--- | :--- | :--- | :--- | :--- | :--- |
| nlevel | integer | $\geq 0$ | $0,1,2$ | 1 | $m=1+$ nlevel |
| thresh | real | $\geq 0$ | $0,0.1,0.01$ | 0.1 | thresh $=$ thresh |
|  |  | $<0$ | $-0.75,-0.90$ |  | thresh auto-selected |
|  |  |  |  |  |  |
| filter | real | $\geq 0$ | $0,0.05,0.001$ | 0.05 | filter $=$ filter |
|  |  | $<0$ | -0.90 |  | filter auto-selected |
|  |  |  |  |  |  |

When thresh $<0$, then a threshold is selected such that thresh represents the negative of the fraction of nonzero elements that are dropped. For example, if thresh $=-0.9$ then $\tilde{A}$ will contain approximately ten percent of the nonzeros in $A$.

When filter $<0$, then a filter value is selected such that filter represents the negative of the fraction of nonzero elements that are dropped. For example, if filter $=-0.9$ then approximately 90 percent of the entries in the computed approximate inverse are dropped.

### 6.14.2 Preconditioning Nearly Symmetric Matrices

A nonsymmetric, but definite and nearly symmetric matrix $A$ may be preconditioned with a symmetric preconditioner $M$. Using a symmetric preconditioner has a few advantages, such as guaranteeing positive definiteness of the preconditioner, as well as being less expensive to construct.
The nonsymmetric matrix $A$ must be definite, i.e., $\left(A+A^{T}\right) / 2$ is SPD, and the a priori sparsity pattern to be used must be symmetric. The latter may be guaranteed by 1) constructing the sparsity pattern with a symmetric matrix, or 2) if the matrix is structurally symmetric (has symmetric pattern), then thresholding to construct the pattern is not used (i.e., zero value of the thresh parameter is used).

### 6.15 ILU

ILU is a suite of parallel incomplete LU factorization algorithms featuring dual threshold (ILUT) and level-based (ILUK) variants. The implementation is based on a domain decomposition framework for achieving distributed parallelism. ILU can be used as a standalone iterative solver (this is not recommended), preconditioner for Krylov subspace methods, or smoother for multigrid methods such as BoomerAMG and MGR.

Note: ILU is currently only supported by the IJ interface.

### 6.15.1 Overview

ILU utilizes a domain decomposition framework. A basic block-Jacobi approach involves performing inexact solutions within the local domains owned by the processes, using parallel local ILU factorizations. In a more advanced approach, the unknowns are partitioned into interior and interface points, where the interface points separate the interior points in adjacent domains. In an algebraic context, this is equivalent to dividing the matrix rows into local (processor-owned) and external (off-processor-owned) blocks. The overall parallel ILU strategy is a two-level method that consists of ILU solves within the local blocks and a global solve involving the Schur complement system, which various iterative approaches in this framework can solve.

### 6.15.2 User-level functions

A list of user-level functions for configuring ILU is given below, where each block of functions is marked as Required, Recommended, Optional, or Exclusively required. Note that the last two blocks of function calls are exclusively required, i.e., the first block should be called only when ILU is used as a standalone solver, while the second block should be called only when it is used as a preconditioner to GMRES. In the last case, other Krylov methods can be chosen. We refer the reader to Solvers and Preconditioners for more information.

```
/* (Required) Create ILU solver */
HYPRE_ILUCreate(&ilu_solver);
/* (Recommended) General solver options */
HYPRE_ILUSetType(ilu_solver, ilu_type); /* 0, 1, 10, 11, 20, 21, 30, 31, 40, 41, 50 %/
HYPRE_ILUSetMaxIter(ilu_solver, max_iter);
```

```
HYPRE_ILUSetTol(ilu_solver, tol);
HYPRE_ILUSetLocalReordering(ilu_solver, reordering); /* 0: none, 1: RCM */
HYPRE_ILUSetPrintLevel(ilu_solver, print_level);
/* (Optional) Function calls for ILUK variants */
HYPRE_ILUSetLevelOfFill(ilu_solver, fill);
    (Optional) Function calls for ILUT variants */
HYPRE_ILUSetMaxNnzPerRow(ilu_solver, max_nnz_row);
HYPRE_ILUSetDropThreshold(ilu_solver, threshold);
/* (Optional) Function calls for GMRES-ILU or NSH-ILU */
HYPRE_ILUSetNSHDropThreshold(ilu_solver, threshold);
HYPRE_ILUSetSchurMaxIter(ilu_solver, schur_max_iter);
/* (Optional) Function calls for iterative ILU variants */
HYPRE_ILUSetTriSolve(ilu_solver, 0);
HYPRE_ILUSetLowerJacobiIters(ilu_solver, ljac_iters);
HYPRE_ILUSetUpperJacobiIters(ilu_solver, ujac_iters);
/* (Exclusively required) Function calls for using ILU as standalone solver */
HYPRE_ILUSetup(ilu_solver, parcsr_M, b, x);
HYPRE_ILUSolve(ilu_solver, parcsr_A, b, x);
/* (Exclusively required) Function calls for using ILU as preconditioner to GMRES */
HYPRE_GMRESSetup(gmres_solver, (HYPRE_Matrix)A, (HYPRE_Vector)b, (HYPRE_Vector)x);
HYPRE_GMRESSolve(gmres_solver, (HYPRE_Matrix)A, (HYPRE_Vector)b, (HYPRE_Vector)x);
/* (Required) Free memory */
HYPRE_ILUDestroy(ilu_solver);
```

A short explanation for each of those functions and its parameters is given next.

- HYPRE_ILUCreate Create the hypre_ILU solver object.
- HYPRE_ILUDestroy Destroy the hypre_ILU solver object.
- HYPRE_ILUSetType Set the type of ILU factorization. Options are:
- 0: Block-Jacobi ILUK (BJ-ILUK).
- 1: Block-Jacobi ILUT (BJ-ILUT).
- 10: GMRES with ILUK (GMRES-ILUK).
- 11: GMRES with ILUT (GMRES-ILUT).
- 20: NSH with ILUK (NSH-ILUK).
- 21: NSH with ILUT (NSH-ILUT).
- 30: RAS with ILUK (RAS-ILUK).
- 31: RAS with ILUT (RAS-ILUT).
- 40: ddPQ-GMRES with ILUK (ddPQ-GMRES-ILUK).
- 41: ddPQ-GMRES with ILUT (ddPQ-GMRES-ILUT).
- 50: GMRES with RAP-ILU0 with modified ILU0 (GMRES-RAP-ILU0).
- HYPRE_ILUSetMaxIter Set the maximum number of ILU iterations. We recommend setting this value to one when ILU is used as a preconditioner or smoother.
- HYPRE_ILUSetTol Set the convergence tolerance for ILU. We recommend setting this value to zero when ILU is used as a preconditioner or smoother.
- HYPRE_ILUSetLocalReordering Set the local matrix reordering algorithm.
- 0: No reordering.
- 1: Reverse Cuthill-McKee (RCM).
- HYPRE_ILUSetPrintLevel Set the verbosity level for algorithm statistics.
- 0: No output.
- 1: Print setup info.
- 2: Print solve info.
- 3: Print setup and solve info.
- HYPRE_ILUSetLevel0fFill Set the level of fill used by the level-based ILUK strategy.
- HYPRE_ILUSetMaxNnzPerRow Set the maximum number of nonzero entries per row in the triangular factors for ILUT.
- HYPRE_ILUSetDropThreshold Set the threshold for dropping nonzero entries during the construction of the triangular factors for ILUT.
- HYPRE_ILUSetNSHDropThreshold Set the threshold for dropping nonzero entries during the computation of the approximate inverse matrix via NSH-ILU.
- HYPRE_ILUSetSchurMaxIter Set the maximum number of iterations for solving the Schur complement system (GMRES-ILU or NSH-ILU).
- HYPRE_ILUSetTriSolve Set triangular solve method used in ILU's solve phase. Option zero refers to the iterative approach, which leads to good performance in GPUs, and option one refers to the direct (exact) approach.
- HYPRE_ILUSetLower JacobiIters Set the number of iterations for solving the lower triangular linear system. This option makes sense when enabling the iterative triangular solve approach.
- HYPRE_ILUSetUpperJacobiIters Same as previous function, but for the upper triangular factor.
- HYPRE_ILUSetup Setup a hypre_ILU solver object.
- HYPRE_ILUSolve Solve the linear system with hypre_ILU.
- HYPRE_ILUDestroy Destroy the hypre_ILU solver object.

Note: For more details about ILU options and parameters, including their default values, we refer the reader to hypre's reference manual or section ParCSR Solvers.

### 6.15.3 ILU as Smoother for BoomerAMG

The following functions can be used to configure ILU as a smoother to BoomerAMG:

```
/* (Required) Set ILU as smoother to BoomerAMG */
HYPRE_BoomerAMGSetSmoothType(amg_solver, 5);
HYPRE_BoomerAMGSetSmoothNumLevels(amg_solver, num_levels);
/* (Optional) General ILU configuration parameters */
HYPRE_BoomerAMGSetILUType(amg_solver, ilu_type);
HYPRE_BoomerAMGSetILUMaxIter(amg_solver, ilu_max_iter);
HYPRE_BoomerAMGSetILULocalReordering(amg_solver, ilu_reordering);
/* (Optional) Function calls for ILUK smoother variants */
HYPRE_BoomerAMGSetILULevel(amg_solver, ilu_fill);
/* (Optional) Function calls for ILUT smoother variants */
HYPRE_BoomerAMGSetILUDroptol(amg_solver, ilu_threshold);
HYPRE_BoomerAMGSetILUMaxRowNnz(amg_solver, ilu_max_nnz_row);
/* (Optional) Function calls for iterative ILU smoother variants */
HYPRE_BoomerAMGSetILUTriSolve(amg_solver, 0);
HYPRE_BoomerAMGSetILULowerJacobiIters(amg_solver, ilu_ljac_iters);
HYPRE_BoomerAMGSetILUUpperJacobiIters(amg_solver, ilu_ujac_iters);
```

where:

- HYPRE_BoomerAMGSetSmoothNumLevels Enable smoothing in the first num_levels levels of AMG.
- HYPRE_BoomerAMGSetILUType Set the type of ILU factorization. See HYPRE_ILUSetType.
- HYPRE_BoomerAMGSetILUMaxIter Set the number of ILU smoother sweeps.
- HYPRE_BoomerAMGSetILULocalReordering Set the local matrix reordering algorithm.
- HYPRE_BoomerAMGSetILULevel Set ILUK's fill level.
- HYPRE_BoomerAMGSetILUDroptol Set ILUT's threshold.
- HYPRE_BoomerAMGSetILUMaxRowNnz Set ILUT's maximum number of nonzero entries per row.
- HYPRE_BoomerAMGSetILUTriSolve Set triangular solve method. See HYPRE_ILUSetTriSolve.
- HYPRE_BoomerAMGSetILULowerJacobiIters Set the number of iterations for the L factor.
- HYPRE_BoomerAMGSetILUUpperJacobiIters Same as previous function, but for the U factor.


### 6.15.4 GPU support

The addition of GPU support to ILU is ongoing work. A few algorithm types have already been fully ported to the CUDA and HIP backends, i.e., both their setup (factorization) and solve phases are executed on the device. Below is a detailed list of which phases (setup and solve) of the various ILU algorithms have been ported to GPUs. In the table, UVM-Setup indicates that the setup phase is executed on the CPU (host); at the same time, the triangular factors are stored in a memory space that is accessible from the GPU (device) via unified memory. This feature must be enabled during hypre's configuration.

|  | CUDA (NVIDIA GPUs) | HIP (AMD GPUs) | SYCL (Intel GPUs) |
| :--- | :--- | :--- | :--- |
| BJ-ILU0 | Setup and Solve | Setup and Solve | UVM-Setup and Solve |
| BJ-ILU(K/T) | UVM-Setup and Solve | UVM-Setup and Solve | UVM-Setup and Solve |
| GMRES-ILU0 | Setup and Solve | Setup and Solve | UVM-Setup and Solve |
| GMRES-RAP-ILU0 | UVM-Setup and Solve | UVM-Setup and Solve | UVM-Setup and Solve |
| GMRES-ILU(K/T) | UVM-Setup and Solve | UVM-Setup and Solve | UVM-Setup and Solve |
| ddPQ-GMRES- | UVM-Setup and Solve | UVM-Setup and Solve | UVM-Setup and Solve |
| ILU(K/T) |  |  |  |
| NSH-ILU(K/T) | UVM-Setup and Solve | UVM-Setup and Solve | UVM-Setup and Solve |
| RAS-ILU(K/T) | UVM-Setup and Solve | UVM-Setup and Solve | UVM-Setup and Solve |

Hint: For better setup performance on GPUs, disable local reordering by passing option zero to HYPRE_ILUSetLocalReordering or HYPRE_BoomerAMGSetILULocalReordering. This may degrade convergence of the iterative solver.

Note: hypre must be built with cuSPARSE support when running ILU on NVIDIA GPUs, rocSPARSE when running on AMD GPUs, or oneMKL sparse when running on Intel GPUs.

### 6.16 Euclid

Warning: Euclid is not actively supported by the hypre development team. We recommend using $I L U$ for parallel ILU algorithms. This new ILU implementation includes 64-bit integers support (for linear systems with more than $2,147,483,647$ global unknowns) through both mixedint and bigint builds of hypre and NVIDIA/AMD GPUs support through the CUDA/HIP backends.

The Euclid library is a scalable implementation of the Parallel ILU algorithm that was presented at SC99 [HyPo1999], and published in expanded form in the SIAM Journal on Scientific Computing [HyPo2001]. By scalable we mean that the factorization (setup) and application (triangular solve) timings remain nearly constant when the global problem size is scaled in proportion to the number of processors. As with all ILU preconditioning methods, the number of iterations is expected to increase with global problem size.

Experimental results have shown that PILU preconditioning is in general more effective than Block Jacobi preconditioning for minimizing total solution time. For scaled problems, the relative advantage appears to increase as the number of processors is scaled upwards. Euclid may also be used to good advantage as a smoother within multigrid methods.

### 6.16.1 Overview

Euclid is best thought of as an "extensible ILU preconditioning framework." Extensible means that Euclid can (and eventually will, time and contributing agencies permitting) support many variants of ILU $(k)$ and ILUT preconditioning. (The current release includes Block Jacobi $\operatorname{ILU}(k)$ and Parallel $\operatorname{ILU}(k)$ methods.) Due to this extensibility, and also because Euclid was developed independently of the hypre project, the methods by which one passes runtime parameters to Euclid preconditioners differ in some respects from the hypre norm. While users can directly set options within their code, options can also be passed to Euclid preconditioners via command line switches and/or small text-based configuration files. The latter strategies have the advantage that users will not need to alter their codes as Euclid's capabilities are extended.

The following fragment illustrates the minimum coding required to invoke Euclid preconditioning within hypre application contexts. The next subsection provides examples of the various ways in which Euclid's options can be set. The final subsection lists the options, and provides guidance as to the settings that (in our experience) will likely prove effective for minimizing execution time.

```
#include "HYPRE_parcsr_ls.h"
HYPRE_Solver eu;
HYPRE_Solver pcg_solver;
HYPRE_ParVector b, x;
HYPRE_ParCSRMatrix A;
//Instantiate the preconditioner.
HYPRE_EuclidCreate(comm, &eu);
//Optionally use the following three methods to set runtime options.
// 1. pass options from command line or string array.
HYPRE_EuclidSetParams(eu, argc, argv);
// 2. pass options from a configuration file.
HYPRE_EuclidSetParamsFromFile(eu, "filename");
// 3. pass options using interface functions.
HYPRE_EuclidSetLevel(eu, 3);
...
//Set Euclid as the preconditioning method for some
//other solver, using the function calls HYPRE_EuclidSetup
//and HYPRE_EuclidSolve. We assume that the pcg_solver
//has been properly initialized.
HYPRE_PCGSetPrecond(pcg_solver,
                                    (HYPRE_PtrToSolverFcn) HYPRE_EuclidSolve,
                                    (HYPRE_PtrToSolverFcn) HYPRE_EuclidSetup,
                                    eu) ;
//Solve the system by calling the Setup and Solve methods for,
//in this case, the HYPRE_PCG solver. We assume that A, b, and x
//have been properly initialized.
HYPRE_PCGSetup(pcg_solver, (HYPRE_Matrix)A, (HYPRE_Vector)b, (HYPRE_Vector)x);
HYPRE_PCGSolve(pcg_solver, (HYPRE_Matrix)parcsr_A, (HYPRE_Vector)b, (HYPRE_Vector)x);
//Destroy the Euclid preconditioning object.
HYPRE_EuclidDestroy(eu);
```


### 6.16.2 Setting Options: Examples

For expositional purposes, assume you wish to set the $\operatorname{ILU}(k)$ factorization level to the value $k=3$. There are several methods of accomplishing this. Internal to Euclid, options are stored in a simple database that contains (name, value) pairs. Various of Euclid's internal (private) functions query this database to determine, at runtime, what action the user has requested. If you enter the option -eu_stats 1, a report will be printed when Euclid's destructor is called; this report lists (among other statistics) the options that were in effect during the factorization phase.
Method 1. By default, Euclid always looks for a file titled database in the working directory. If it finds such a file, it opens it and attempts to parse it as a configuration file. Configuration files should be formatted as follows.

```
>cat database
#this is an optional comment
-level 3
```

Any line in a configuration file that contains a "\#" character in the first column is ignored. All other lines should begin with an option name, followed by one or more blanks, followed by the option value. Note that option names always begin with a - character. If you include an option name that is not recognized by Euclid, no harm should ensue.

Method 2. To pass options on the command line, call

```
HYPRE_EuclidSetParams(HYPRE_Solver solver, int argc, char *argv[]);
```

where argc and argv carry the usual connotation: main(int argc, char *argv[]). If your hypre application is called phoo, you can then pass options on the command line per the following example.

```
mpirun -np 2 phoo -level 3
```

Since Euclid looks for the database file when HYPRE_EuclidCreate is called, and parses the command line when HYPRE_EuclidSetParams is called, option values passed on the command line will override any similar settings that may be contained in the database file. Also, if same option name appears more than once on the command line, the final appearance determines the setting.

Some options, such as -bj (see next subsection) are boolean. Euclid always treats these options as the value 1 (true) or 0 (false). When passing boolean options from the command line the value may be committed, in which case it assumed to be 1 . Note, however, that when boolean options are contained in a configuration file, either the 1 or 0 must stated explicitly.

Method 3. There are two ways in which you can read in options from a file whose name is other than database. First, you can call HYPRE_EuclidSetParamsFromFile to specify a configuration filename. Second, if you have passed the command line arguments as described above in Method 2, you can then specify the configuration filename on the command line using the - db_filename filename option, e.g.,

```
mpirun -np 2 phoo -db_filename ../myConfigFile
```

Method 4. One can also set parameters via interface functions, e.g

```
int HYPRE_EuclidSetLevel(HYPRE_Solver solver, int level);
```

For a full set of functions, see the reference manual.

### 6.16.3 Options Summary

- -level $\langle i n t\rangle$ Factorization level for $\operatorname{ILU}(k)$. Default: 1. Guidance: for 2D convection-diffusion and similar problems, fastest solution time is typically obtained with levels 4 through 8. For 3D problems fastest solution time is typically obtained with level 1.
- -bj Use Block Jacobi ILU preconditioning instead of PILU. Default: 0 (false). Guidance: if subdomains contain relatively few nodes (less than 1,000 ), or the problem is not well partitioned, Block Jacobi ILU may give faster solution time than PILU.
- -eu_stats When Euclid's destructor is called a summary of runtime settings and timing information is printed to stdout. Default: 0 (false). The timing marks in the report are the maximum over all processors in the MPI communicator.
- -eu_mem When Euclid's destructor is called a summary of Euclid's memory usage is printed to stdout. Default: 0 (false). The statistics are for the processor whose rank in MPI_COMM_WORLD is 0.
- -printTestData This option is used in our autotest procedures, and should not normally be invoked by users.
- -sparseA $\langle f l o a t\rangle$ Drop-tolerance for $\operatorname{ILU}(k)$ factorization. Default: 0 (no dropping). Entries are treated as zero if their absolute value is less than sparseA * max, where max is the largest absolute value of any entry in the row. Guidance: try this in conjunction with -rowScale. CAUTION: If the coefficient matrix $A$ is symmetric, this setting is likely to cause the filled matrix, $F=L+U-I$, to be non-symmetric. This setting has no effect when ILUT factorization is selected.
- -rowScale Scale values prior to factorization such that the largest value in any row is +1 or -1 . Default: 0 (false). CAUTION: If the coefficient matrix $A$ is symmetric, this setting is likely to cause the filled matrix, $F=L+U-I$, to be non-symmetric. Guidance: if the matrix is poorly scaled, turning on row scaling may help convergence.
- -ilut $\langle$ float $\rangle$ Use ILUT factorization instead of the default, $\operatorname{ILU}(k)$. Here, $\langle$ float $\rangle$ is the drop tolerance, which is relative to the largest absolute value of any entry in the row being factored. CAUTION: If the coefficient matrix $A$ is symmetric, this setting is likely to cause the filled matrix, $F=L+U-I$, to be non-symmetric. NOTE: this option can only be used sequentially!


### 6.17 PILUT: Parallel Incomplete Factorization

Warning: PILUT is not actively supported by the hypre development team. We recommend using $I L U$ for parallel ILU algorithms. This new ILU implementation includes 64-bit integers support (for linear systems with more than $2,147,483,647$ global unknowns) through both mixedint and bigint builds of hypre and NVIDIA/AMD GPUs support through the CUDA/HIP backends.

PILUT is a parallel preconditioner based on Saad's dual-threshold incomplete factorization algorithm. The original version of PILUT was done by Karypis and Kumar [KaKu1998] in terms of the Cray SHMEM library. The code was subsequently modified by the hypre team: SHMEM was replaced by MPI; some algorithmic changes were made; and it was software engineered to be interoperable with several matrix implementations, including hypre's ParCSR format, PETSc's matrices, and ISIS++ RowMatrix. The algorithm produces an approximate factorization $L U$, with the preconditioner $M$ defined by $M=L U$.

Note: PILUT produces a nonsymmetric preconditioner even when the original matrix is symmetric. Thus, it is generally inappropriate for preconditioning symmetric methods such as Conjugate Gradient.

### 6.17.1 Parameters:

- SetMaxNonzerosPerRow( int LFIL ) ; (Default: 20) Set the maximum number of nonzeros to be retained in each row of $L$ and $U$. This parameter can be used to control the amount of memory that $L$ and $U$ occupy. Generally, the larger the value of LFIL, the longer it takes to calculate the preconditioner and to apply the preconditioner and the larger the storage requirements, but this trades off versus a higher quality preconditioner that reduces the number of iterations.
- SetDropTolerance ( double tol) ; (Default: 0.0001) Set the tolerance (relative to the 2-norm of the row) below which entries in $L$ and $U$ are automatically dropped. PILUT first drops entries based on the drop tolerance, and then retains the largest LFIL elements in each row that remain. Smaller values of tol lead to more accurate preconditioners, but can also lead to increases in the time to calculate the preconditioner.


### 6.18 LOBPCG Eigensolver

LOBPCG (Locally Optimal Block Preconditioned Conjugate Gradient) is a simple, yet very efficient, algorithm suggested in [Knya2001], [KLAO2007], [BLOPEWeb] for computing several smallest eigenpairs of the symmetric generalized eigenvalue problem $A x=\lambda B x$ with large, possibly sparse, symmetric matrix $A$ and symmetric positive definite matrix $B$. The matrix $A$ is not assumed to be positive, which also allows one to use LOBPCG to compute the largest eigenpairs of $A x=\lambda B x$ simply by solving $-A x=\mu B x$ for the smallest eigenvalues $\mu=-\lambda$.

LOBPCG simultaneously computes several eigenpairs together, which is controlled by the blockSize parameter, see example ex11.c. The LOBCPG also allows one to impose constraints on the eigenvectors of the form $x^{T} B y_{i}=0$ for a set of vectors $y_{i}$ given to LOBPCG as input parameters. This makes it possible to compute, e.g., 50 eigenpairs by 5 subsequent calls to LOBPCG with the blockSize=10, using deflation. LOBPCG can use preconditioning in two different ways: by running an inner preconditioned PCG linear solver, or by applying the preconditioner directly to the eigenvector residual (option -pcgitr 0). In all other respects, LOBPCG is similar to the PCG linear solver.

The LOBPCG code is available for system interfaces: Struct, SStruct, and IJ. It is also used in the Auxiliary-space Maxwell Eigensolver (AME). The LOBPCG setup is similar to the setup for PCG.

### 6.19 FEI Solvers

Warning: FEI is not actively supported by the hypre development team. For similar functionality, we recommend using Block-Structured Grids with Finite Elements, which allows the representation of block-structured grid problems via hypre's SStruct interface.

After the FEI has been used to assemble the global linear system (as described in Chapter Finite Element Interface), a number of hypre solvers can be called to perform the solution. This is straightforward, if hypre's FEI has been used. If an external FEI is employed, the user needs to link with hypre's implementation of the LinearSystemCore class, as described in Section Using HYPRE in External FEI Implementations.

Solver parameters are specified as an array of strings, and a complete list of the available options can be found in the FEI section of the reference manual. They are passed to the FEI as in the following example:

```
nParams = 5;
paramStrings = new char*[nParams];
for (i = 0; i < nParams; i++) }
    paramStrings[i] = new char[100];
strcpy(paramStrings[0], "solver cg");
strcpy(paramStrings[1], "preconditioner diag");
strcpy(paramStrings[2], "maxiterations 100");
strcpy(paramStrings[3], "tolerance 1.0e-6");
strcpy(paramStrings[4], "outputLevel 1");
feiPtr -> parameters(nParams, paramStrings);
```

To solve the linear system of equations, we call

```
feiPtr -> solve(&status);
```

where the returned value status indicates whether the solve was successful.
Finally, the solution can be retrieved by the following function call:

```
feiPtr -> getBlockNodeSolution(elemBlkID, nNodes, nodeIDList,
    solnOffsets, solnValues);
```

where nodeIDList is a list of nodes in element block elemBlkID, and solnOffsets[i] is the index pointing to the first location where the variables at node $i$ is returned in solnValues.

### 6.19.1 Solvers Available Only through the FEI

While most of the solvers from the previous sections are available through the FEI interface, there are number of additional solvers and preconditioners that are accessible only through the FEI. These solvers are briefly described in this section (see also the reference manual).

## Sequential and Parallel Solvers

hypre currently has many iterative solvers. There is also internally a version of the sequential SuperLU direct solver (developed at U.C. Berkeley) suitable to small problems (may be up to the size of 10000). In the following we list some of these internal solvers.

1. Additional Krylov solvers (FGMRES, TFQMR, symmetric QMR),
2. SuperLU direct solver (sequential),
3. SuperLU direct solver with iterative refinement (sequential),

## Parallel Preconditioners

The performance of the Krylov solvers can be improved by clever selection of preconditioners. Besides those mentioned previously in this chapter, the following preconditioners are available via the LinearSystemCore interface:

1. the modified version of MLI, which requires the finite element substructure matrices to construct the prolongation operators,
2. parallel domain decomposition with inexact local solves (DDIlut),
3. least-squares polynomial preconditioner,
4. $2 \times 2$ block preconditioner, and
5. $2 \times 2$ Uzawa preconditioner.

Some of these preconditioners can be tuned by a number of internal parameters modifiable by users. A description of these parameters is given in the reference manual.

## Matrix Reduction

For some structural mechanics problems with multi-point constraints the discretization matrix is indefinite (eigenvalues lie in both sides of the imaginary axis). Indefinite matrices are much more difficult to solve than definite matrices. Methods have been developed to reduce these indefinite matrices to definite matrices. Two matrix reduction algorithms have been implemented in hypre, as presented in the following subsections.

## Schur Complement Reduction

The incoming linear system of equations is assumed to be in the form:

$$
\left[\begin{array}{cc}
D & B \\
B^{T} & 0
\end{array}\right]\left[\begin{array}{l}
x_{1} \\
x_{2}
\end{array}\right]=\left[\begin{array}{l}
b_{1} \\
b_{2}
\end{array}\right]
$$

where $D$ is a diagonal matrix. After Schur complement reduction is applied, the resulting linear system becomes

$$
-B^{T} D^{-1} B x_{2}=b_{2}-B^{T} D^{-1} b_{1}
$$

## Slide Surface Reduction

With the presence of slide surfaces, the matrix is in the same form as in the case of Schur complement reduction. Here $A$ represents the relationship between the master, slave, and other degrees of freedom. The matrix block $\left[B^{T} 0\right]$ corresponds to the constraint equations. The goal of reduction is to eliminate the constraints. As proposed by Manteuffel, the trick is to re-order the system into a $3 \times 3$ block matrix.

$$
\left[\begin{array}{ccc}
A_{11} & A_{12} & N \\
A_{21} & A_{22} & D \\
N_{T} & D & 0
\end{array}\right]=\left[\begin{array}{cc}
A_{11} & \hat{A}_{12} \\
\hat{A}_{21} & \hat{A}_{22} .
\end{array}\right]
$$

The reduced system has the form :

$$
\left(A_{11}-\hat{A}_{21} \hat{A}_{22}^{-1} \hat{A}_{12}\right) x_{1}=b_{1}-\hat{A}_{21} \hat{A}_{22}^{-1} b_{2}
$$

which is symmetric positive definite (SPD) if the original matrix is PD. In addition, $\hat{A}_{22}^{-1}$ is easy to compute.
There are three slide surface reduction algorithms in hypre. The first follows the matrix formulation in this section. The second is similar except that it replaces the eliminated slave equations with identity rows so that the degree of freedom at each node is preserved. This is essential for certain block algorithms such as the smoothed aggregation multilevel preconditioners. The third is similar to the second except that it is more general and can be applied to problems with intersecting slide surfaces (sequential only for intersecting slide surfaces).

## Other Features

To improve the efficiency of the hypre solvers, a few other features have been incorporated. We list a few of these features below :

1. Preconditioner reuse - For multiple linear solves with matrices that are slightly perturbed from each other, oftentimes the use of the same preconditioners can save preconditioner setup times but suffer little convergence rate degradation.
2. Projection methods - For multiple solves that use the same matrix, previous solution vectors can sometimes be used to give a better initial guess for subsequent solves. Two projection schemes have been implemented in hypre - A-conjugate projection (for SPD matrices) and minimal residual projection (for both SPD and non-SPD matrices).
3. The sparsity pattern of the matrix is in general not destroyed after it has been loaded to an hypre matrix. But if the matrix is not to be reused, an option is provided to clean up this pattern matrix to conserve memory usage.

## GENERAL INFORMATION

### 7.1 Getting the Source Code

The most recent hypre distribution is available at `https://github.com/hypre-space/hypre/tags' _ along with previous distribution versions.

### 7.2 Building the Library

In this and the following several sections, we discuss the steps to install and use hypre. First, we focus on the primary method targeting Unix-like operating systems, such as Linux, AIX, and Mac OS X. Then in CMake instructions, we explain an alternative approach using the CMake build system [CMakeWeb], which is the best approach for building hypre on Windows systems in particular.
After unpacking the hypre tar file, the source code will be in the src sub-directory of a directory named hypreVERSION, where VERSION is the current version number (e.g., hypre-2.29.0).
Move to the src sub-directory to build hypre for the host platform. The simplest method is to configure, compile and install the libraries in ./hypre/lib and ./hypre/include directories, which is accomplished by:

```
./configure
make
```

NOTE: when executing on an IBM platform configure must be executed under the nopoe script (./nopoe ./ configure <option> ...<option>) to force a single task to be run on the log-in node.
There are many options to configure and make to customize such things as installation directories, compilers used, compile and load flags, etc.

Executing configure results in the creation of platform specific files that are used when building the library. The information may include such things as the system type being used for building and executing, compilers being used, libraries being searched, option flags being set, etc. When all of the searching is done two files are left in the src directory; config.status contains information to recreate the current configuration and config.log contains compiler messages which may help in debugging configure errors.
Upon successful completion of configure the file config/Makefile.config is created from its template config/ Makefile.config.in and hypre is ready to be built.
Executing make, with or without targets being specified, in the src directory initiates compiling of all of the source code and building of the hypre library. If any errors occur while compiling, the user can edit the file config/Makefile. config directly then run make again; without having to re-run configure.

When building hypre without the install target, the libraries and include files will be copied into the default directories, src/hypre/lib and src/hypre/include, respectively.

When building hypre using the install target, the libraries and include files will be copied into the directories that the user specified in the options to configure, e.g. --prefix=/usr/apps. If none were specified the default directories, src/hypre/lib and src/hypre/include, are used.

### 7.2.1 Configure Options

There are many options to configure to allow the user to override and refine the defaults for any system. The best way to find out what options are available is to display the help package, by executing ./configure --help, which also includes the usage information. The user can mix and match the configure options and variable settings to meet their needs.

Some commonly used options include:

```
--enable-debug
--enable-shared
--with-print-errors
--with-openmp
--enable-bigint
--enable-mixedint
```

```
Sets compiler flags to generate information
```

Sets compiler flags to generate information
needed for debugging.
needed for debugging.
Build shared libraries.
Build shared libraries.
NOTE: in order to use the resulting shared
NOTE: in order to use the resulting shared
libraries the user MUST have the path to
libraries the user MUST have the path to
the libraries defined in the environment
the libraries defined in the environment
variable LD_LIBRARY_PATH.
variable LD_LIBRARY_PATH.
Print HYPRE errors
Print HYPRE errors
Use OpenMP. This may affect which compiler is
Use OpenMP. This may affect which compiler is
chosen.
chosen.
Use long long int for HYPRE_Int (default is NO).
Use long long int for HYPRE_Int (default is NO).
NOTE: This option is not available for Nvidia
NOTE: This option is not available for Nvidia
and AMD GPUs.
and AMD GPUs.
Use long long int for HYPRE_BigInt and int for
Use long long int for HYPRE_BigInt and int for
HYPRE_Int.
HYPRE_Int.
NOTE: This option disables Euclid, ParaSails,
NOTE: This option disables Euclid, ParaSails,
PILUT and CGC coarsening.

```
    PILUT and CGC coarsening.
```

The user can mix and match the configure options and variable settings to meet their needs. It should be noted that hypre can be configured with external BLAS and LAPACK libraries, which can be combined with any other option. This is done as follows (currently, both libraries must be configured as external together):

```
./configure --with-blas-lib="blas-lib-name" \
    --with-blas-lib-dirs="path-to-blas-lib" \
    --with-lapack-lib="lapack-lib-name" \
    --with-lapack-lib-dirs="path-to-lapack-lib"
```

The output from configure is several pages long. It reports the system type being used for building and executing, compilers being used, libraries being searched, option flags being set, etc.

### 7.2.2 Make Targets

The make step in building hypre is where the compiling, loading and creation of libraries occurs. Make has several options that are called targets. These include:

```
help prints the details of each target
all default target in all directories
    compile the entire library
    does NOT rebuild documentation
```

```
clean deletes all files from the current directory that are
    created by building the library
distclean deletes all files from the current directory that are created
    by configuring or building the library
install compile the source code, build the library and copy executables,
        libraries, etc to the appropriate directories for user access
uninstall deletes all files that the install target created
tags runs etags to create a tags table
    file is named TAGS and is saved in the top-level directory
test depends on the all target to be completed
    removes existing temporary installation directories
    creates temporary installation directories
    copies all libHYPRE* and *.h files to the temporary locations
    builds the test drivers; linking to the temporary locations to
    simulate how application codes will link to HYPRE
```


### 7.2.3 GPU build

Hypre can support NVIDIA GPUs with CUDA and OpenMP ( $\geq 4.5$ ). The related configure options are

| - -with-cuda | Use CUDA. Require cuda-9.0 or higher (default is <br> NO). |
| :--- | :--- |
| --with-device-openmp | Use OpenMP 4.5 Device Directives. This may affect <br> which compiler is chosen. |

The related environment variables

```
HYPRE_CUDA_SM (default 70)
CUDA_HOME the CUDA home directory
```

need to be set properly, which can be also set by

```
--with-gpu-arch=ARG
    (e.g., --with-gpu-arch='60 70')
--with-cuda-home=DIR
```

When configured with --with-cuda or --with-device-openmp, the memory allocated on the GPUs, by default, is the GPU device memory, which is not accessible from the CPUs. Hypre's structured solvers can run with device memory, whereas only selected unstructured solvers can run with device memory. See GPU-supported Options for details. Some solver options for BoomerAMG require unified (CUDA managed) memory. To use these options add the following configure option:
--enable-unified-memory Use unified memory for allocating the memory (default is NO).

Hypre's Struct solvers can also choose RAJA and Kokkos as the backend. The configure options are

```
--with-raja Use RAJA. Require RAJA package to be compiled
    properly (default is NO).
Use Kokkos. Require kokkos package to be compiled
properly(default is NO).
```

To run on the GPUs with RAJA and Kokkos, the options --with-cuda and --with-device-openmp are also needed, and the RAJA and Kokkos libraries should be built with CUDA or OpenMP 4.5 correspondingly.
The other NVIDIA GPU related options include:

- --enable-gpu-profiling Use NVTX on CUDA, rocTX on HIP (default is NO)
- --enable-cusparse Use cuSPARSE for GPU sparse kernels (default is YES)
- --enable-cublas Use cuBLAS for GPU dense kernels (default is YES)
- --enable-curand Use random numbers generators on GPUs (default is YES)

Allocations and deallocations of GPU memory are expensive. Memory pooling is a common approach to reduce such overhead and improve performance. hypre provides caching allocators for GPU device memory and unified memory, enabled by
--enable-device-memory-pool Enable the caching GPU memory allocator in hypre (default is NO)
hypre also supports Umpire [Umpire]. To enable Umpire pool, include the following options:

```
--with-umpire Use Umpire Allocator for device and unified memory
                    (default is NO)
--with-umpire-include=/path-of-umpire-install/include
--with-umpire-lib-dirs=/path-of-umpire-install/lib
--with-umpire-libs=umpire
```

For running on AMD GPUs, configure with

```
--with-hip Use HIP for AMD GPUs. (default is NO)
--with-gpu-arch=ARG Use appropriate AMD GPU architecture
```

The other AMD GPU related options include:

- --enable-gpu-profiling Use NVTX on CUDA, rocTX on HIP (default is NO)
- --enable-rocsparse Use rocSPARSE (default is YES)
- --enable-rocblas Use rocBLAS (default is NO)
- --enable-rocrand Use rocRAND (default is YES)

All the options supported by CUDA are also supported with HIP. Note that the "-enable-bigint" option is not supported with CUDA or HIP.

For running on Intel GPUs, configure with

```
--with-sycl Use SYCL for Intel GPUs. (default is NO).
--with-sycl-target=ARG User specifies sycl targets for AOT compilation in
    ARG, where ARG is a comma-separated list (enclosed
    in quotes), e.g. "spir64_gen".
```

--with-sycl-target-backend=ARG
User specifies additional options for the sycl
target backend for AOT compilation in ARG, where ARG
contains the desired options (enclosed in
double+single quotes), e.g.
--with-sycl-target-backend="'-device
12.1.0,12.4.0'".

Intel oneMKL functionality is also used by default (and required for certain hypre solvers):
--enable-onemklsparse Use oneMKL sparse (default is YES).
--enable-onemklblas Use oneMKL blas (default is YES).
--enable-onemklrand Use oneMKL rand (default is YES).

The SYCL backend now supports all GPU-enabled hypre functionality currently supported by CUDA/HIP except for FSAI (work in progress). The --enable-bigint option is supported with SYCL (not supported for CUDA/HIP).

### 7.2.4 Testing the Library

The examples subdirectory contains several codes that can be used to test the newly created hypre library. To create the executable versions, move into the examples subdirectory, enter make then execute the codes as described in the initial comments section of each source code.

### 7.3 CMake-based Build Instructions

This section describes hypre's CMake build system, which is particularly useful for building the code on Windows machines. CMake-based installation provides a platform-independent build system. CMake can generate Unix and Linux Makefiles, as well as Visual Studio and (Apple) XCode project files from the same configuration file. In addition, CMake also provides a GUI front end and which allows an interactive build and installation process. For more detailed information on using CMake, see CMake's User Interaction Guide.

Note: Not all options are currently supported when using CMake. This is an on-going effort to support all hypre configure options.

Here are the basic steps to configure, make, and install hypre using CMake:

1. Ensure that CMake version 3.13 .0 or later is installed on the system.
2. After unpacking the hypre tar file or cloning, move to the src sub-directory.
3. To build the library, run CMake on the top-level hypre source directory to generate files appropriate for the native build system. To prevent writing over the Makefiles in hypre's configure/make system above, only out-of-source builds are allowed with CMake, that is, it is required to use a separate build directory.
The directory src/cmbuild is provided in the release for convenience, but alternative build directories may be created by the user. To configure with the default options:

- Unix: From the src/cmbuild directory, type cmake ...
- Windows Visual Studio: Set the source and build directories to src and src/cmbuild, then click on Configure following by Generate.

4. To build the library, compile with the native build system:

- Unix: From the src/cmbuild directory, type make or make -j 4 (for a faster parallel build with 4 threads).
- Windows Visual Studio: Open the 'hypre' VS solution file generated by CMake and build the $A L L \_B U I L D$ target.

5. To install hypre to the installation directory specified in the configuration:

- Unix: From the src/cmbuild directory, type make install.
- Windows Visual Studio: Open the hypre VS solution file generated by CMake and build the INSTALL target.
- Note: The default installation location is set to src/hypre. Use the HYPRE_INSTALL_PREFIX option to change this location if desired.


### 7.3.1 Changing Default CMake Configuration Options

Various configuration options can be set from within CMake (see CMake options). One option is to specify these options in the command-line CMake invocation, e.g., to enabling building of the examples:

```
cmake -DHYPRE_BUILD_EXAMPLES=ON ..
```

Another option is to use the CMake GUI (ccmake or cmake-gui) to change the default options as appropriate, then reconfigure / generate:

- Unix: From the src/cmbuild directory, type ccmake ...
- Change options to desired settings:
* To set a variable, move the cursor to the variable and press enter.
* If it is a boolean (ON/OFF) it will toggle the value.
* If it is string or file, it will allow editing of the string.
- Then configure (c key).
- Repeat until all values are set as desired and then generate (g key).
- Windows Visual Studio: Change options, then click on Configure then Generate.

Then the re-build and re-install with the updated configuration options.

### 7.3.2 CMake Configure Options

There are many options to allow the user to override and refine the defaults for any system. The best way to find out what options are available is to use cmake, cmake-gui, or inspect using Windows Visual Studio.

Some commonly used options (default value) include:

```
HYPRE_INSTALL_PREFIX (src/hypre) Installation location.
HYPRE_BUILD_EXAMPLES (OFF) Compile test cases for examples of using the library.
HYPRE_BUILD_TYPE (Release) Sets compiler flags to generate information.
    needed for debugging.
HYPRE_ENABLE_SHARED (OFF) Build shared libraries.
HYPRE_PRINT_ERRORS (OFF) Print HYPRE errors.
HYPRE_WITH_OPENMP (OFF) Use OpenMP.
```

(continues on next page)

```
HYPRE_ENABLE_BIGINT (OFF) Use long long int for HYPRE_Int.
HYPRE_ENABLE_MIXEDINT (OFF) Use long long int for HYPRE_BigInt and int for
HYPRE_Int.
```


## GPU CMake Build Options

Some of the commonly used options for GPU CMake builds of hypre are listed below.

- CUDA support for NVIDIA GPUs relevant options:

```
HYPRE_WITH_CUDA (OFF) Use CUDA v9.0 or higher.
HYPRE_CUDA_SM (70) Target CUDA architecture.
```

When configured with CUDA, the memory allocated on the GPUs, by default, is the GPU device memory, which is not accessible from the CPUs. Hypre's structured solvers can run with device memory, whereas only selected unstructured solvers can run with device memory. See GPU-supported Options for details. Some solver options for BoomerAMG require unified (CUDA managed) memory. To use these options turn the following option on:

```
HYPRE_ENABLE_UNIFIED_MEMORY (OFF) Use unified memory for allocating the memory.
```

The other NVIDIA GPU related options include:

```
HYPRE_ENABLE_GPU_PROFILING (OFF) Use NVTX.
HYPRE_ENABLE_CUSPARSE (ON) Use cuSPARSE for GPU sparse kernels.
HYPRE_ENABLE_CUBLAS (OFF) Use cuBLAS for GPU dense kernels.
HYPRE_ENABLE_CURAND (ON) Use random numbers generators on GPUs.
```

Allocations and deallocations of GPU memory are expensive. Memory pooling is a common approach to reduce such overhead and improve performance. hypre provides caching allocators for GPU device memory and unified memory, enabled by

```
HYPRE_ENABLE_DEVICE_POOL (OFF) Enable the caching GPU memory allocator in hypre
```

hypre also supports Umpire [Umpire]. To enable Umpire pool, include the following options:

```
HYPRE_WITH_UMPIRE (OFF) Use Umpire Allocator for device and unified memory.
TPL_UMPIRE_LIBRARIES List of absolute paths to Umpire link libraries.
TPL_UMPIRE_INCLUDE_DIRS List of absolute paths to Umpire include directories.
```

SYCL support for Intel GPUs relevant options:

```
HYPRE_WITH_SYCL (OFF)
HYPRE_SYCL_TARGET
HYPRE_SYCL_TARGET_BACKEND
๑12.4.0'.
```

Enable SYCL support.
Target SYCL architecture, e.g. 'spir64_gen'.
Additional SYCL backend options, e.g. '-device 12.1.0,

### 7.3.3 Testing the Library with CMake Build Process

The examples subdirectory contains several codes that can be used to test the newly created hypre library. The CMake option HYPRE_BUILD_EXAMPLES should be enabled so ensure the executables in the examples subdirectory are built.

### 7.4 Linking to the Library

An application code linking with hypre must be compiled with -I\$PREFIX/include and linked with -L\$PREFIX/lib -lHYPRE, where \$PREFIX is the directory where hypre is installed, default is hypre, or as defined by the configure option --prefix=PREFIX. As noted above, if hypre was built as a shared library the user MUST have its location defined in the environment variable LD_LIBRARY_PATH.

As an example of linking with hypre, a user may refer to the Makefile in the examples sub-directory. It is designed to build codes similar to user applications that link with and call hypre. All include and linking flags are defined in the Makefile.config file by configure.

### 7.5 Error Flags

Every hypre function returns an integer, which is used to indicate errors during execution. Note that the error flag returned by a given function reflects the errors from all previous calls to hypre functions. In particular, a value of zero means that all hypre functions up to (and including) the current one have completed successfully. This new error flag system is being implemented throughout the library, but currently there are still functions that do not support it. The error flag value is a combination of one or a few of the following error codes:

1. HYPRE_ERROR_GENERIC - describes a generic error
2. HYPRE_ERROR_MEMORY - hypre was unable to allocate memory
3. HYPRE_ERROR_ARG - error in one of the arguments of a hypre function
4. HYPRE_ERROR_CONV - a hypre solver did not converge as expected

One can use the HYPRE_CheckError function to determine exactly which errors have occurred:

```
/* call some HYPRE functions */
int hypre_ierr;
hypre_ierr = HYPRE_Function();
/* check if the previously called hypre functions returned error(s) */
if (hypre_ierr)
    /* check if the error with code HYPRE_ERROR_CODE has occurred */
    if (HYPRE_CheckError(hypre_ierr,HYPRE_ERROR_CODE))
```

The corresponding FORTRAN code is

```
! header file with hypre error codes
include 'HYPRE_error_f.h'
! call some HYPRE functions
integer hypre_ierr
call HYPRE_Function(hypre_ierr)
! check if the previously called hypre functions returned error(s)
```

(continues on next page)

```
if (hypre_ierr .ne. 0) then
    ! check if the error with code HYPRE_ERROR_CODE has occurred
    call HYPRE_CheckError(hypre_ierr, HYPRE_ERROR_CODE, check)
    if (check .ne. 0) then
```

The global error flag can also be obtained directly, between calls to other hypre functions, by calling HYPRE_GetError (). If an argument error (HYPRE_ERROR_ARG) has occurred, the argument index (counting from 1) can be obtained from HYPRE_GetErrorArg(). To get a character string with a description of all errors in a given error flag, use

HYPRE_DescribeError(int hypre_ierr, char *descr);
The global error flag can be cleared manually by calling HYPRE_ClearAllErrors(), which will essentially ignore all previous hypre errors. To only clear a specific error code, the user can call HYPRE_ClearError (HYPRE_ERROR_CODE). Finally, if hypre was configured with --with-print-errors, additional error information will be printed to the standard error during execution.

### 7.6 Bug Reporting and General Support

Simply create an issue at `https://github.com/hypre-space/hypre/issues _ to report bugs, request features, or ask general usage questions.

Users should include as much relevant information as possible in their issue report, including at a minimum, the hypre version number being used. For compile and runtime problems, please also include the machine type, operating system, MPI implementation, compiler, and any error messages produced.

### 7.7 Using HYPRE in External FEl Implementations

Warning: FEI is not actively supported by the hypre development team. For similar functionality, we recommend using Block-Structured Grids with Finite Elements, which allows the representation of block-structured grid problems via hypre's SStruct interface.

To set up hypre for use in external, e.g. Sandia's, FEI implementations one needs to follow the following steps:

1. obtain the hypre and Sandia's FEI source codes,
2. compile Sandia's FEI (fei-2.5.0) to create the fei_base library.
3. compile hypre

- unpack the archive and go into the src directory
- do a configure with the --with-fei-inc-dir option set to the FEI include directory plus other compile options
- compile with make install to create the HYPRE_LSI library in hypre/lib.

4. call the FEI functions in your application code (as shown in Chapters Finite Element Interface and Solvers and Preconditioners)

- include cfei-hypre.h in your file
- include FEI_Implementation.h in your file


## 5. Modify your Makefile

- include hypre's include and lib directories in the search paths.
- Link with -lfei_base -lHYPRE_LSI. Note that the order in which the libraries are listed may be important.

Building an application executable often requires linking with many different software packages, and many software packages use some LAPACK and/or BLAS functions. In order to alleviate the problem of multiply defined functions at link time, it is recommended that all software libraries are stripped of all LAPACK and BLAS function definitions. These LAPACK and BLAS functions should then be resolved at link time by linking with the system LAPACK and BLAS libraries (e.g. dxml on DEC cluster). Both hypre and SuperLU were built with this in mind. However, some other software library files needed may have the BLAS functions defined in them. To avoid the problem of multiply defined functions, it is recommended that the offending library files be stripped of the BLAS functions.

### 7.8 Calling HYPRE from Other Languages

The hypre library currently supports two languages: C (native) and Fortran (in version 2.10 .1 and earlier, additional language interfaces were also provided through a tool called Babel). The Fortran interface is manually supported to mirror the "native" C interface used throughout most of this manual. We describe this interface next.
Typically, the Fortran subroutine name is the same as the C name, unless it is longer than 31 characters. In these situations, the name is condensed to 31 characters, usually by simple truncation. For now, users should look at the Fortran test drivers (*.f codes) in the test directory for the correct condensed names. In the future, this aspect of the interface conversion will be made consistent and straightforward.

The Fortran subroutine argument list is always the same as the corresponding C routine, except that the error return code ierr is always last. Conversion from C parameter types to Fortran argument type is summarized in following table:

| C parameter | Fortran argument |
| :--- | :--- |
| int i | integer i |
| double d | double precision d |
| int *array | integer array(*) |
| double *array | double precision array(*) |
| char *string | character string(*) |
| HYPRE_Type object | integer*8 object |
| HYPRE_Type *object | integer*8 object |

Array arguments in hypre are always of type (int *) or (double *), and the corresponding Fortran types are simply integer or double precision arrays. Note that the Fortran arrays may be indexed in any manner. For example, an integer array of length N may be declared in fortran as either of the following:

```
integer array(N)
integer array(0:N-1)
```

hypre objects can usually be declared as in the table because integer*8 usually corresponds to the length of a pointer. However, there may be some machines where this is not the case. On such machines, the Fortran type for a hypre object should be an integer of the appropriate length.
This simple example illustrates the above information:
C prototype:

```
int HYPRE_IJMatrixSetValues(HYPRE_IJMatrix matrix,
    int nrows, int *ncols,
    const int *rows, const int *cols,
    const double *values);
```

The corresponding Fortran code for calling this routine is as follows:

```
integer*8 matrix
integer nrows, ncols(MAX_NCOLS)
integer rows(MAX_ROWS), cols(MAX_COLS)
double precision values(MAX_COLS)
integer ierr
call HYPRE_IJMatrixSetValues(matrix, nrows, ncols, rows, cols, values, ierr)
```


### 8.1 Struct System Interface

## group StructSystemInterface

A structured-grid conceptual interface.
This interface represents a structured-grid conceptual view of a linear system.

## Struct Grids

## typedef struct hypre_StructGrid_struct *HYPRE_StructGrid

A grid object is constructed out of several "boxes", defined on a global abstract index space.
HYPRE_Int HYPRE_StructGridCreate(MPI_Comm comm, HYPRE_Int ndim, HYPRE_StructGrid *grid)
Create an ndim-dimensional grid object.
HYPRE_Int HYPRE_StructGridDestroy (HYPRE_StructGrid grid)
Destroy a grid object.
An object should be explicitly destroyed using this destructor when the user's code no longer needs direct access to it. Once destroyed, the object must not be referenced again. Note that the object may not be deallocated at the completion of this call, since there may be internal package references to the object. The object will then be destroyed when all internal reference counts go to zero.
HYPRE_Int HYPRE_StructGridSetExtents(HYPRE_StructGrid grid, HYPRE_Int *ilower, HYPRE_Int *iupper)
Set the extents for a box on the grid.
HYPRE_Int HYPRE_StructGridAssemble(HYPRE_StructGrid grid)
Finalize the construction of the grid before using.
HYPRE_Int HYPRE_StructGridSetPeriodic (HYPRE_StructGrid grid, HYPRE_Int *periodic)
Set the periodicity for the grid.
The argument periodic is an ndim-dimensional integer array that contains the periodicity for each dimension. A zero value for a dimension means non-periodic, while a nonzero value means periodic and contains the actual period. For example, periodicity in the first and third dimensions for a $10 \times 11 \times 12$ grid is indicated by the array $[10,0,12]$.

NOTE: Some of the solvers in hypre have power-of-two restrictions on the size of the periodic dimensions.

HYPRE_Int HYPRE_StructGridSetNumGhost (HYPRE_StructGrid grid, HYPRE_Int *num_ghost)
Set the ghost layer in the grid object.

## Struct Stencils

typedef struct hypre_StructStencil_struct *HYPRE_StructStencil
The stencil object.
HYPRE_Int HYPRE_StructStencilCreate(HYPRE_Int ndim, HYPRE_Int size, HYPRE_StructStencil *stencil)
Create a stencil object for the specified number of spatial dimensions and stencil entries.
HYPRE_Int HYPRE_StructStencilDestroy (HYPRE_StructStencil stencil)
Destroy a stencil object.
HYPRE_Int HYPRE_StructStencilSetElement(HYPRE_StructStencil stencil, HYPRE_Int entry, HYPRE_Int *offset)

Set a stencil entry.
NOTE: The name of this routine will eventually be changed to HYPRE_StructStencilSetEntry.

## Struct Matrices

typedef struct hypre_StructMatrix_struct *HYPRE_StructMatrix
The matrix object.
HYPRE_Int HYPRE_StructMatrixCreate(MPI_Comm comm, HYPRE_StructGrid grid, HYPRE_StructStencil stencil, HYPRE_StructMatrix *matrix)

Create a matrix object.
HYPRE_Int HYPRE_StructMatrixDestroy (HYPRE_StructMatrix matrix)
Destroy a matrix object.
HYPRE_Int HYPRE_StructMatrixInitialize(HYPRE_StructMatrix matrix)
Prepare a matrix object for setting coefficient values.
HYPRE_Int HYPRE_StructMatrixSetValues(HYPRE_StructMatrix matrix, HYPRE_Int *index, HYPRE_Int nentries, HYPRE_Int *entries, HYPRE_Complex *values)

Set matrix coefficients index by index.
The values array is of length nentries.
NOTE: For better efficiency, use HYPRE_StructMatrixSetBoxValues to set coefficients a box at a time.
HYPRE_Int HYPRE_StructMatrixAddToValues (HYPRE_StructMatrix matrix, HYPRE_Int *index, HYPRE_Int nentries, HYPRE_Int *entries, HYPRE_Complex *values)

Add to matrix coefficients index by index.
The values array is of length nentries.
NOTE: For better efficiency, use HYPRE_StructMatrixAddToBoxValues to set coefficients a box at a time.

HYPRE_Int HYPRE_StructMatrixSetConstantValues (HYPRE_StructMatrix matrix, HYPRE_Int nentries, HYPRE_Int *entries, HYPRE_Complex *values)

Set matrix coefficients which are constant over the grid.
The values array is of length nentries.
HYPRE_Int HYPRE_StructMatrixAddToConstantValues(HYPRE_StructMatrix matrix, HYPRE_Int nentries, HYPRE_Int *entries, HYPRE_Complex *values)
Add to matrix coefficients which are constant over the grid.
The values array is of length nentries.
HYPRE_Int HYPRE_StructMatrixSetBoxValues(HYPRE_StructMatrix matrix, HYPRE_Int *ilower,
HYPRE_Int *iupper, HYPRE_Int nentries, HYPRE_Int *entries, HYPRE_Complex *values)
Set matrix coefficients a box at a time.
The data in values is ordered as follows:

```
m = 0;
for (k = ilower[2]; k <= iupper[2]; k++)
    for (j = ilower[1]; j <= iupper[1]; j++)
        for (i = ilower[0]; i <= iupper[0]; i++)
            for (entry = 0; entry < nentries; entry++)
            {
            values[m] = ...;
            m++;
            }
```

HYPRE_Int HYPRE_StructMatrixAddToBoxValues(HYPRE_StructMatrix matrix, HYPRE_Int *ilower, HYPRE_Int *iupper, HYPRE_Int nentries, HYPRE_Int *entries, HYPRE_Complex *values)
Add to matrix coefficients a box at a time.
The data in values is ordered as in HYPRE_StructMatrixSetBoxValues.
HYPRE_Int HYPRE_StructMatrixSetBoxValues2 (HYPRE_StructMatrix matrix, HYPRE_Int *ilower, HYPRE_Int *iupper, HYPRE_Int nentries, HYPRE_Int *entries, HYPRE_Int *vilower, HYPRE_Int *viupper, HYPRE_Complex *values)
Set matrix coefficients a box at a time.
The values array is logically box shaped with value-box extents vilower and viupper that must contain the set-box extents ilower and iupper. The data in the values array is ordered as in HYPRE_StructMatrixSetBoxValues, but based on the value-box extents.

HYPRE_Int HYPRE_StructMatrixAddToBoxValues2 (HYPRE_StructMatrix matrix, HYPRE_Int *ilower, HYPRE_Int *iupper, HYPRE_Int nentries, HYPRE_Int *entries, HYPRE_Int *vilower, HYPRE_Int *viupper, HYPRE_Complex *values)
Add to matrix coefficients a box at a time.
The data in values is ordered as in HYPRE_StructMatrixSetBoxValues 2.

HYPRE_Int HYPRE_StructMatrixAssemble(HYPRE_StructMatrix matrix)
Finalize the construction of the matrix before using.
HYPRE_Int HYPRE_StructMatrixGetValues(HYPRE_StructMatrix matrix, HYPRE_Int *index, HYPRE_Int nentries, HYPRE_Int *entries, HYPRE_Complex *values)
Get matrix coefficients index by index.
The values array is of length nentries.
NOTE: For better efficiency, use HYPRE_StructMatrixGetBoxValues to get coefficients a box at a time.
HYPRE_Int HYPRE_StructMatrixGetBoxValues(HYPRE_StructMatrix matrix, HYPRE_Int *ilower, HYPRE_Int *iupper, HYPRE_Int nentries, HYPRE_Int *entries, HYPRE_Complex *values)
Get matrix coefficients a box at a time.
The data in values is ordered as in HYPRE_StructMatrixSetBoxValues.
HYPRE_Int HYPRE_StructMatrixGetBoxValues2 (HYPRE_StructMatrix matrix, HYPRE_Int *ilower, HYPRE_Int *iupper, HYPRE_Int nentries, HYPRE_Int *entries, HYPRE_Int *vilower, HYPRE_Int *viupper, HYPRE_Complex *values)
Get matrix coefficients a box at a time.
The data in values is ordered as in HYPRE_StructMatrixSetBoxValues 2 .
HYPRE_Int HYPRE_StructMatrixSetSymmetric (HYPRE_StructMatrix matrix, HYPRE_Int symmetric)
Define symmetry properties of the matrix.
By default, matrices are assumed to be nonsymmetric. Significant storage savings can be made if the matrix is symmetric.

HYPRE_Int HYPRE_StructMatrixSetConstantEntries(HYPRE_StructMatrix matrix, HYPRE_Int nentries, HYPRE_Int *entries)
Specify which stencil entries are constant over the grid.
Declaring entries to be "constant over the grid" yields significant memory savings because the value for each declared entry will only be stored once. However, not all solvers are able to utilize this feature.

Presently supported:

- no entries constant (this function need not be called)
- all entries constant
- all but the diagonal entry constant

HYPRE_Int HYPRE_StructMatrixSetNumGhost (HYPRE_StructMatrix matrix, HYPRE_Int *num_ghost)
Set the ghost layer in the matrix.
HYPRE_Int HYPRE_StructMatrixPrint (const char *filename, HYPRE_StructMatrix matrix, HYPRE_Int all)

Print the matrix to file.
This is mainly for debugging purposes.
HYPRE_Int HYPRE_StructMatrixRead(MPI_Comm comm, const char *filename, HYPRE_Int *num_ghost, HYPRE_StructMatrix *matrix)
Read the matrix from file.
This is mainly for debugging purposes.

HYPRE_Int HYPRE_StructMatrixMatvec (HYPRE_Complex alpha, HYPRE_StructMatrix A, HYPRE_StructVector x, HYPRE_Complex beta, HYPRE_StructVector y)
Matvec operator.
This operation is $y=\alpha A x+\beta y$. Note that you can do a simple matrix-vector multiply by setting $\alpha=1$ and $\beta=0$.

## Struct Vectors

HYPRE_Int HYPRE_StructVectorCreate(MPI_Comm comm, HYPRE_StructGrid grid, HYPRE_StructVector *vector)
The vector object.
Create a vector object.
HYPRE_Int HYPRE_StructVectorDestroy (HYPRE_StructVector vector)
Destroy a vector object.
HYPRE_Int HYPRE_StructVectorInitialize(HYPRE_StructVector vector)
Prepare a vector object for setting coefficient values.
HYPRE_Int HYPRE_StructVectorSetValues(HYPRE_StructVector vector, HYPRE_Int *index, HYPRE_Complex value)
Set vector coefficients index by index.
NOTE: For better efficiency, use HYPRE_StructVectorSetBoxValues to set coefficients a box at a time.
HYPRE_Int HYPRE_StructVectorAddToValues(HYPRE_StructVector vector, HYPRE_Int *index, HYPRE_Complex value)
Add to vector coefficients index by index.
NOTE: For better efficiency, use HYPRE_StructVectorAddToBoxValues to set coefficients a box at a time.
HYPRE_Int HYPRE_StructVectorSetBoxValues(HYPRE_StructVector vector, HYPRE_Int *ilower, HYPRE_Int *iupper, HYPRE_Complex *values)
Set vector coefficients a box at a time.
The data in values is ordered as follows:

```
m = 0;
for (k = ilower[2]; k <= iupper[2]; k++)
    for (j = ilower[1]; j <= iupper[1]; j++)
        for (i = ilower[0]; i <= iupper[0]; i++)
        {
            values[m] = ...;
            m++;
        }
```

HYPRE_Int HYPRE_StructVectorAddToBoxValues(HYPRE_StructVector vector, HYPRE_Int *ilower, HYPRE_Int *iupper, HYPRE_Complex *values)
Add to vector coefficients a box at a time.
The data in values is ordered as in HYPRE_StructVectorSetBoxValues.

HYPRE_Int HYPRE_StructVectorSetBoxValues2 (HYPRE_StructVector vector, HYPRE_Int *ilower, HYPRE_Int *iupper, HYPRE_Int *vilower, HYPRE_Int *viupper, HYPRE_Complex *values)
Set vector coefficients a box at a time.
The values array is logically box shaped with value-box extents vilower and viupper that must contain the set-box extents ilower and iupper. The data in the values array is ordered as in HYPRE_StructVectorSetBoxValues, but based on the value-box extents.

HYPRE_Int HYPRE_StructVectorAddToBoxValues2 (HYPRE_StructVector vector, HYPRE_Int *ilower, HYPRE_Int *iupper, HYPRE_Int *vilower, HYPRE_Int *viupper, HYPRE_Complex *values)
Add to vector coefficients a box at a time.
The data in values is ordered as in HYPRE_StructVectorSetBoxValues 2.
HYPRE_Int HYPRE_StructVectorAssemble(HYPRE_StructVector vector)
Finalize the construction of the vector before using.
HYPRE_Int HYPRE_StructVectorGetValues(HYPRE_StructVector vector, HYPRE_Int *index, HYPRE_Complex *value)

Get vector coefficients index by index.
NOTE: For better efficiency, use HYPRE_StructVectorGetBoxValues to get coefficients a box at a time.
HYPRE_Int HYPRE_StructVectorGetBoxValues(HYPRE_StructVector vector, HYPRE_Int *ilower, HYPRE_Int *iupper, HYPRE_Complex *values)

Get vector coefficients a box at a time.
The data in values is ordered as in HYPRE_StructVectorSetBoxValues.
HYPRE_Int HYPRE_StructVectorGetBoxValues2 (HYPRE_StructVector vector, HYPRE_Int *ilower, HYPRE_Int *iupper, HYPRE_Int *vilower, HYPRE_Int *viupper, HYPRE_Complex *values)

Get vector coefficients a box at a time.
The data in values is ordered as in HYPRE_StructVectorSetBoxValues 2.
HYPRE_Int HYPRE_StructVectorPrint (const char *filename, HYPRE_StructVector vector, HYPRE_Int all)

Print the vector to file.
This is mainly for debugging purposes.
HYPRE_Int HYPRE_StructVectorRead(MPI_Comm comm, const char *filename, HYPRE_Int *num_ghost, HYPRE_StructVector *vector)
Read the vector from file.
This is mainly for debugging purposes.

### 8.2 SStruct System Interface

## group SStructSystemInterface

A semi-structured-grid conceptual interface.
This interface represents a semi-structured-grid conceptual view of a linear system.

## SStruct Grids

typedef struct hypre_SStructGrid_struct *HYPRE_SStructGrid
A grid object is constructed out of several structured "parts" and an optional unstructured "part".
Each structured part has its own abstract index space.
typedef HYPRE_Int HYPRE_SStructVariable
An enumerated type that supports cell centered, node centered, face centered, and edge centered variables.
Face centered variables are split into $x$-face, $y$-face, and $z$-face variables, and edge centered variables are split into x-edge, y-edge, and z-edge variables. The edge centered variable types are only used in 3D. In 2 D , edge centered variables are handled by the face centered types.
Variables are referenced relative to an abstract (cell centered) index in the following way:

- cell centered variables are aligned with the index;
- node centered variables are aligned with the cell corner at relative index ( $1 / 2,1 / 2,1 / 2$ );
- x-face, $y$-face, and $z$-face centered variables are aligned with the faces at relative indexes $(1 / 2,0,0)$, ( $0,1 / 2,0$ ), and ( $0,0,1 / 2$ ), respectively;
- x-edge, y-edge, and $z$-edge centered variables are aligned with the edges at relative indexes $(0,1 / 2$, $1 / 2),(1 / 2,0,1 / 2)$, and ( $1 / 2,1 / 2,0$ ), respectively.

The supported identifiers are:

- HYPRE_SSTRUCT_VARIABLE_CELL
- HYPRE_SSTRUCT_VARIABLE_NODE
- HYPRE_SSTRUCT_VARIABLE_XFACE
- HYPRE_SSTRUCT_VARIABLE_YFACE
- HYPRE_SSTRUCT_VARIABLE_ZFACE
- HYPRE_SSTRUCT_VARIABLE_XEDGE
- HYPRE_SSTRUCT_VARIABLE_YEDGE
- HYPRE_SSTRUCT_VARIABLE_ZEDGE

NOTE: Although variables are referenced relative to a unique abstract cell-centered index, some variables are associated with multiple grid cells. For example, node centered variables in 3D are associated with 8 cells (away from boundaries). Although grid cells are distributed uniquely to different processes, variables may be owned by multiple processes because they may be associated with multiple cells.

HYPRE_Int HYPRE_SStructGridCreate(MPI_Comm comm, HYPRE_Int ndim, HYPRE_Int nparts, HYPRE_SStructGrid *grid)

Create an ndim-dimensional grid object with nparts structured parts.

HYPRE_Int HYPRE_SStructGridDestroy(HYPRE_SStructGrid grid)
Destroy a grid object.
An object should be explicitly destroyed using this destructor when the user's code no longer needs direct access to it. Once destroyed, the object must not be referenced again. Note that the object may not be deallocated at the completion of this call, since there may be internal package references to the object. The object will then be destroyed when all internal reference counts go to zero.

HYPRE_Int HYPRE_SStructGridSetExtents(HYPRE_SStructGrid grid, HYPRE_Int part, HYPRE_Int *ilower, HYPRE_Int *iupper)
Set the extents for a box on a structured part of the grid.
HYPRE_Int HYPRE_SStructGridSetVariables(HYPRE_SStructGrid grid, HYPRE_Int part, HYPRE_Int nvars, HYPRE_SStructVariable *vartypes)
Describe the variables that live on a structured part of the grid.
HYPRE_Int HYPRE_SStructGridAddVariables(HYPRE_SStructGrid grid, HYPRE_Int part, HYPRE_Int
*index, HYPRE_Int nvars, HYPRE_SStructVariable
*vartypes)
Describe additional variables that live at a particular index.
These variables are appended to the array of variables set in HYPRE_SStructGridSetVariables, and are referenced as such.

NOTE: This routine is not yet supported.
HYPRE_Int HYPRE_SStructGridSetFEMOrdering(HYPRE_SStructGrid grid, HYPRE_Int part, HYPRE_Int *ordering)

Set the ordering of variables in a finite element problem.
This overrides the default ordering described below.
Array ordering is composed of blocks of size $(1+n d i m)$. Each block indicates a specific variable in the element and the ordering of the blocks defines the ordering of the variables. A block contains a variable number followed by an offset direction relative to the element's center. For example, a block containing ( 2 , $1,-1,0)$ means variable 2 on the edge located in the $(1,-1,0)$ direction from the center of the element. Note that here variable 2 must be of type ZEDGE for this to make sense. The ordering array must account for all variables in the element. This routine can only be called after HYPRE_SStructGridSetVariables.

The default ordering for element variables (var, $i, j, k$ ) varies fastest in index $i$, followed by $j$, then $k$, then var. For example, if var 0, var 1, and var 2 are declared to be XFACE, YFACE, and NODE variables, respectively, then the default ordering (in 2D) would first list the two XFACE variables, then the two YFACE variables, then the four NODE variables as follows:
$(0,-1,0),(0,1,0),(1,0,-1),(1,0,1),(2,-1,-1),(2,1,-1),(2,-1,1),(2,1,1)$
HYPRE_Int HYPRE_SStructGridSetNeighborPart (HYPRE_SStructGrid grid, HYPRE_Int part, HYPRE_Int *ilower, HYPRE_Int *iupper, HYPRE_Int nbor_part, HYPRE_Int *nbor_ilower, HYPRE_Int *nbor_iupper, HYPRE_Int *index_map, HYPRE_Int *index_dir)

Describe how regions just outside of a part relate to other parts.
This is done a box at a time.
Parts part and nbor_part must be different, except in the case where only cell-centered data is used.
Indexes should increase from ilower to iupper. It is not necessary that indexes increase from nbor_ilower to nbor_iupper.

The index_map describes the mapping of indexes 0,1 , and 2 on part part to the corresponding indexes on part nbor_part. For example, triple ( $1,2,0$ ) means that indexes 0,1 , and 2 on part part map to indexes 1 , 2 , and 0 on part nbor_part, respectively.

The index_dir describes the direction of the mapping in index_map. For example, triple $(1,1,-1)$ means that for indexes 0 and 1, increasing values map to increasing values on nbor_part, while for index 2, decreasing values map to increasing values.

NOTE: All parts related to each other via this routine must have an identical list of variables and variable types. For example, if part 0 has only two variables on it, a cell centered variable and a node centered variable, and we declare part 1 to be a neighbor of part 0 , then part 1 must also have only two variables on it, and they must be of type cell and node. In addition, variables associated with FACEs or EDGEs must be grouped together and listed in X, Y, Z order. This is to enable the code to correctly associate variables on one part with variables on its neighbor part when a coordinate transformation is specified. For example, an XFACE variable on one part may correspond to a YFACE variable on a neighbor part under a particular tranformation, and the code determines this association by assuming that the variable lists are as noted here.

## HYPRE_Int HYPRE_SStructGridSetSharedPart (HYPRE_SStructGrid grid, HYPRE_Int part, HYPRE_Int *ilower, HYPRE_Int *iupper, HYPRE_Int *offset, HYPRE_Int shared_part, HYPRE_Int *shared_ilower, HYPRE_Int *shared_iupper, HYPRE_Int *shared_offset, HYPRE_Int *index_map, HYPRE_Int *index_dir)

Describe how regions inside a part are shared with regions in other parts.
Parts part and shared_part must be different.
Indexes should increase from ilower to iupper. It is not necessary that indexes increase from shared_ilower to shared_iupper. This is to maintain consistency with the SetNeighborPart function, which is also able to describe shared regions but in a more limited fashion.

The offset is a triple (in 3D) used to indicate the dimensionality of the shared set of data and its position with respect to the box extents ilower and iupper on part part. The dimensionality is given by the number of 0 's in the triple, and the position is given by plus or minus 1 's. For example: $(0,0,0)$ indicates sharing of all data in the given box; $(1,0,0)$ indicates sharing of data on the faces in the $(1,0,0)$ direction; $(1,0$, $-1)$ indicates sharing of data on the edges in the $(1,0,-1)$ direction; and $(1,-1,1)$ indicates sharing of data on the nodes in the $(1,-1,1)$ direction. To ensure the dimensionality, it is required that for every nonzero entry, the corresponding extents of the box are the same. For example, if offset is $(0,1,0)$, then $(2,1,3)$ and $(10,1,15)$ are valid box extents, whereas $(2,1,3)$ and $(10,7,15)$ are invalid (because 1 and 7 are not the same).

The shared_offset is used in the same way as offset, but with respect to the box extents shared_ilower and shared_iupper on part shared_part.

The index_map describes the mapping of indexes 0,1 , and 2 on part part to the corresponding indexes on part shared_part. For example, triple ( $1,2,0$ ) means that indexes 0,1 , and 2 on part part map to indexes 1,2 , and 0 on part shared_part, respectively.

The index_dir describes the direction of the mapping in index_map. For example, triple $(1,1,-1)$ means that for indexes 0 and 1 , increasing values map to increasing values on shared_part, while for index 2 , decreasing values map to increasing values.

NOTE: All parts related to each other via this routine must have an identical list of variables and variable types. For example, if part 0 has only two variables on it, a cell centered variable and a node centered variable, and we declare part 1 to have shared regions with part 0 , then part 1 must also have only two variables on it, and they must be of type cell and node. In addition, variables associated with FACEs or EDGEs must be grouped together and listed in X, Y, Z order. This is to enable the code to correctly associate variables on one part with variables on a shared part when a coordinate transformation is specified. For example, an XFACE variable on one part may correspond to a YFACE variable on a shared part under a
particular tranformation, and the code determines this association by assuming that the variable lists are as noted here

HYPRE_Int HYPRE_SStructGridAddUnstructuredPart (HYPRE_SStructGrid grid, HYPRE_Int ilower, HYPRE_Int iupper)

Add an unstructured part to the grid.
The variables in the unstructured part of the grid are referenced by a global rank between 0 and the total number of unstructured variables minus one. Each process owns some unique consecutive range of variables, defined by ilower and iupper.
NOTE: This is just a placeholder. This part of the interface is not finished.
HYPRE_Int HYPRE_SStructGridAssemble(HYPRE_SStructGrid grid)
Finalize the construction of the grid before using.
HYPRE_Int HYPRE_SStructGridSetPeriodic(HYPRE_SStructGrid grid, HYPRE_Int part, HYPRE_Int *periodic)
Set the periodicity on a particular part.
The argument periodic is an ndim-dimensional integer array that contains the periodicity for each dimension. A zero value for a dimension means non-periodic, while a nonzero value means periodic and contains the actual period. For example, periodicity in the first and third dimensions for a $10 \times 11 \times 12$ part is indicated by the array $[10,0,12]$.

NOTE: Some of the solvers in hypre have power-of-two restrictions on the size of the periodic dimensions.
HYPRE_Int HYPRE_SStructGridSetNumGhost (HYPRE_SStructGrid grid, HYPRE_Int *num_ghost)
Setting ghost in the sgrids.

HYPRE_SSTRUCT_VARIABLE_UNDEFINED

HYPRE_SSTRUCT_VARIABLE_CELL

HYPRE_SSTRUCT_VARIABLE_NODE

HYPRE_SSTRUCT_VARIABLE_XFACE

HYPRE_SSTRUCT_VARIABLE_YFACE

HYPRE_SSTRUCT_VARIABLE_ZFACE

HYPRE_SSTRUCT_VARIABLE_XEDGE

HYPRE_SSTRUCT_VARIABLE_YEDGE

HYPRE_SSTRUCT_VARIABLE_ZEDGE

## SStruct Stencils

typedef struct hypre_SStructStencil_struct *HYPRE_SStructStencil
The stencil object.
HYPRE_Int HYPRE_SStructStencilCreate(HYPRE_Int ndim, HYPRE_Int size, HYPRE_SStructStencil *stencil)

Create a stencil object for the specified number of spatial dimensions and stencil entries.
HYPRE_Int HYPRE_SStructStencilDestroy (HYPRE_SStructStencil stencil)
Destroy a stencil object.
HYPRE_Int HYPRE_SStructStencilSetEntry(HYPRE_SStructStencil stencil, HYPRE_Int entry, HYPRE_Int *offset, HYPRE_Int var)
Set a stencil entry.

## SStruct Graphs

typedef struct hypre_SStructGraph_struct *HYPRE_SStructGraph
The graph object is used to describe the nonzero structure of a matrix.
HYPRE_Int HYPRE_SStructGraphCreate(MPI_Comm comm, HYPRE_SStructGrid grid, HYPRE_SStructGraph *graph)

Create a graph object.
HYPRE_Int HYPRE_SStructGraphDestroy (HYPRE_SStructGraph graph)
Destroy a graph object.
HYPRE_Int HYPRE_SStructGraphSetDomainGrid(HYPRE_SStructGraph graph, HYPRE_SStructGrid domain_grid)
Set the domain grid.
HYPRE_Int HYPRE_SStructGraphSetStencil (HYPRE_SStructGraph graph, HYPRE_Int part, HYPRE_Int var, HYPRE_SStructStencil stencil)
Set the stencil for a variable on a structured part of the grid.
HYPRE_Int HYPRE_SStructGraphSetFEM (HYPRE_SStructGraph graph, HYPRE_Int part)
Indicate that an FEM approach will be used to set matrix values on this part.
HYPRE_Int HYPRE_SStructGraphSetFEMSparsity(HYPRE_SStructGraph graph, HYPRE_Int part,
HYPRE_Int nsparse, HYPRE_Int *sparsity)
Set the finite element stiffness matrix sparsity.
This overrides the default full sparsity pattern described below.
Array sparsity contains nsparse row/column tuples (I,J) that indicate the nonzeroes of the local stiffness matrix. The layout of the values passed into the routine HYPRE_SStructMatrixAddFEMValues is determined here.

The default sparsity is full (each variable is coupled to all others), and the values passed into the routine HYPRE_SStructMatrixAddFEMValues are assumed to be by rows (that is, column indices vary fastest).

HYPRE_Int HYPRE_SStructGraphAddEntries(HYPRE_SStructGraph graph, HYPRE_Int part, HYPRE_Int *index, HYPRE_Int var, HYPRE_Int to_part, HYPRE_Int *to_index, HYPRE_Int to_var)

Add a non-stencil graph entry at a particular index.
This graph entry is appended to the existing graph entries, and is referenced as such.
NOTE: Users are required to set graph entries on all processes that own the associated variables. This means that some data will be multiply defined.

## HYPRE_Int HYPRE_SStructGraphAssemble(HYPRE_SStructGraph graph)

Finalize the construction of the graph before using.

## HYPRE_Int HYPRE_SStructGraphSetObjectType(HYPRE_SStructGraph graph, HYPRE_Int type)

Set the storage type of the associated matrix object.
It is used before AddEntries and Assemble to compute the right ranks in the graph.
NOTE: This routine is only necessary for implementation reasons, and will eventually be removed.

## See also:

HYPRE_SStructMatrixSetObjectType

## SStruct Matrices

typedef struct hypre_SStructMatrix_struct *HYPRE_SStructMatrix
The matrix object.
HYPRE_Int HYPRE_SStructMatrixCreate(MPI_Comm comm, HYPRE_SStructGraph graph, HYPRE_SStructMatrix *matrix)

Create a matrix object.
HYPRE_Int HYPRE_SStructMatrixDestroy (HYPRE_SStructMatrix matrix)
Destroy a matrix object.
HYPRE_Int HYPRE_SStructMatrixInitialize(HYPRE_SStructMatrix matrix)
Prepare a matrix object for setting coefficient values.
HYPRE_Int HYPRE_SStructMatrixSetValues(HYPRE_SStructMatrix matrix, HYPRE_Int part, HYPRE_Int *index, HYPRE_Int var, HYPRE_Int nentries, HYPRE_Int *entries, HYPRE_Complex *values)

Set matrix coefficients index by index.
The values array is of length nentries.
NOTE: For better efficiency, use HYPRE_SStructMatrixSetBoxValues to set coefficients a box at a time.
NOTE: Users are required to set values on all processes that own the associated variables. This means that some data will be multiply defined.

NOTE: The entries in this routine must all be of the same type: either stencil or non-stencil, but not both. Also, if they are stencil entries, they must all represent couplings to the same variable type (there are no such restrictions for non-stencil entries).

HYPRE_Int HYPRE_SStructMatrixAddToValues (HYPRE_SStructMatrix matrix, HYPRE_Int part, HYPRE_Int *index, HYPRE_Int var, HYPRE_Int nentries, HYPRE_Int *entries, HYPRE_Complex *values)

Add to matrix coefficients index by index.
The values array is of length nentries.
NOTE: For better efficiency, use HYPRE_SStructMatrixAddToBoxValues to set coefficients a box at a time.
NOTE: Users are required to set values on all processes that own the associated variables. This means that some data will be multiply defined.
NOTE: The entries in this routine must all be of the same type: either stencil or non-stencil, but not both. Also, if they are stencil entries, they must all represent couplings to the same variable type.
HYPRE_Int HYPRE_SStructMatrixAddFEMValues (HYPRE_SStructMatrix matrix, HYPRE_Int part,
HYPRE_Int *index, HYPRE_Complex *values)
Add finite element stiffness matrix coefficients index by index.
The layout of the data in values is determined by the routines HYPRE_SStructGridSetFEMOrdering and HYPRE_SStructGraphSetFEMSparsity.
NOTE: For better efficiency, use HYPRE_SStructMatrixAddFEMBoxValues to set coefficients a box at a time.

HYPRE_Int HYPRE_SStructMatrixGetValues(HYPRE_SStructMatrix matrix, HYPRE_Int part, HYPRE_Int *index, HYPRE_Int var, HYPRE_Int nentries, HYPRE_Int *entries, HYPRE_Complex *values)
Get matrix coefficients index by index.
The values array is of length nentries.
NOTE: For better efficiency, use HYPRE_SStructMatrixGetBoxValues to get coefficients a box at a time.
NOTE: Users may get values on any process that owns the associated variables.
NOTE: The entries in this routine must all be of the same type: either stencil or non-stencil, but not both. Also, if they are stencil entries, they must all represent couplings to the same variable type (there are no such restrictions for non-stencil entries).
HYPRE_Int HYPRE_SStructMatrixGetFEMValues(HYPRE_SStructMatrix matrix, HYPRE_Int part, HYPRE_Int *index, HYPRE_Complex *values)
Get finite element stiffness matrix coefficients index by index.
The layout of the data in values is determined by the routines HYPRE_SStructGridSetFEMOrdering and HYPRE_SStructGraphSetFEMSparsity.

HYPRE_Int HYPRE_SStructMatrixSetBoxValues(HYPRE_SStructMatrix matrix, HYPRE_Int part, HYPRE_Int *ilower, HYPRE_Int *iupper, HYPRE_Int var, HYPRE_Int nentries, HYPRE_Int *entries, HYPRE_Complex *values)
Set matrix coefficients a box at a time.
The data in values is ordered as follows:

```
m = 0;
for (k = ilower[2]; k <= iupper[2]; k++)
    for (j = ilower[1]; j <= iupper[1]; j++)
        for (i = ilower[0]; i <= iupper[0]; i++)
```

(continues on next page)

```
for (entry = 0; entry < nentries; entry++)
{
    values[m] = ...;
    m++;
}
```

NOTE: Users are required to set values on all processes that own the associated variables. This means that some data will be multiply defined.
NOTE: The entries in this routine must all be of the same type: either stencil or non-stencil, but not both. Also, if they are stencil entries, they must all represent couplings to the same variable type (there are no such restrictions for non-stencil entries).

HYPRE_Int HYPRE_SStructMatrixAddToBoxValues(HYPRE_SStructMatrix matrix, HYPRE_Int part, HYPRE_Int *ilower, HYPRE_Int *iupper, HYPRE_Int var, HYPRE_Int nentries, HYPRE_Int *entries, HYPRE_Complex *values)
Add to matrix coefficients a box at a time.
The data in values is ordered as in HYPRE_SStructMatrixSetBoxValues.
NOTE: Users are required to set values on all processes that own the associated variables. This means that some data will be multiply defined.

NOTE: The entries in this routine must all be of stencil type. Also, they must all represent couplings to the same variable type.

HYPRE_Int HYPRE_SStructMatrixSetBoxValues2 (HYPRE_SStructMatrix matrix, HYPRE_Int part,
HYPRE_Int *ilower, HYPRE_Int *iupper, HYPRE_Int var, HYPRE_Int nentries, HYPRE_Int *entries, HYPRE_Int *vilower, HYPRE_Int *viupper, HYPRE_Complex *values)
Set matrix coefficients a box at a time.
The values array is logically box shaped with value-box extents vilower and viupper that must contain the set-box extents ilower and iupper. The data in the values array is ordered as in HYPRE_SStructMatrixSetBoxValues, but based on the value-box extents.
HYPRE_Int HYPRE_SStructMatrixAddToBoxValues2 (HYPRE_SStructMatrix matrix, HYPRE_Int part, HYPRE_Int *ilower, HYPRE_Int *iupper, HYPRE_Int var, HYPRE_Int nentries, HYPRE_Int *entries, HYPRE_Int *vilower, HYPRE_Int *viupper, HYPRE_Complex *values)
Add to matrix coefficients a box at a time.
The data in values is ordered as in HYPRE_SStructMatrixSetBoxValues 2.
HYPRE_Int HYPRE_SStructMatrixAddFEMBoxValues(HYPRE_SStructMatrix matrix, HYPRE_Int part, HYPRE_Int *ilower, HYPRE_Int *iupper, HYPRE_Complex *values)

Add finite element stiffness matrix coefficients a box at a time.
The data in values is organized as an array of element matrices ordered as in HYPRE_SStructMatrixSetBoxValues. The layout of the data entries of each element matrix is determined by the routines HYPRE_SStructGridSetFEMOrdering and HYPRE_SStructGraphSetFEMSparsity.

HYPRE_Int HYPRE_SStructMatrixAssemble(HYPRE_SStructMatrix matrix)
Finalize the construction of the matrix before using.
HYPRE_Int HYPRE_SStructMatrixGetBoxValues(HYPRE_SStructMatrix matrix, HYPRE_Int part, HYPRE_Int *ilower, HYPRE_Int *iupper, HYPRE_Int var, HYPRE_Int nentries, HYPRE_Int *entries, HYPRE_Complex *values)
Get matrix coefficients a box at a time
The data in values is ordered as in HYPRE_SStructMatrixSetBoxValues.
NOTE: Users may get values on any process that owns the associated variables.
NOTE: The entries in this routine must all be of stencil type. Also, they must all represent couplings to the same variable type.

HYPRE_Int HYPRE_SStructMatrixGetBoxValues2 (HYPRE_SStructMatrix matrix, HYPRE_Int part, HYPRE_Int *ilower, HYPRE_Int *iupper, HYPRE_Int var, HYPRE_Int nentries, HYPRE_Int *entries, HYPRE_Int * vilower, HYPRE_Int *viupper, HYPRE_Complex *values)
Get matrix coefficients a box at a time.
The data in values is ordered as in HYPRE_SStructMatrixSetBoxValues 2.
HYPRE_Int HYPRE_SStructMatrixGetFEMBoxValues (HYPRE_SStructMatrix matrix, HYPRE_Int part, HYPRE_Int *ilower, HYPRE_Int *iupper, HYPRE_Complex *values)
Does this even make sense to implement?
HYPRE_Int HYPRE_SStructMatrixSetSymmetric(HYPRE_SStructMatrix matrix, HYPRE_Int part, HYPRE_Int var, HYPRE_Int to_var, HYPRE_Int symmetric)
Define symmetry properties for the stencil entries in the matrix.
The boolean argument symmetric is applied to stencil entries on part part that couple variable var to variable to_var. A value of -1 may be used for part, var, or to_var to specify "all". For example, if part and to_var are set to -1 , then the boolean is applied to stencil entries on all parts that couple variable var to all other variables.

By default, matrices are assumed to be nonsymmetric. Significant storage savings can be made if the matrix is symmetric.

## HYPRE_Int HYPRE_SStructMatrixSetNSSymmetric(HYPRE_SStructMatrix matrix, HYPRE_Int symmetric)

Define symmetry properties for all non-stencil matrix entries.
HYPRE_Int HYPRE_SStructMatrixSetObjectType (HYPRE_SStructMatrix matrix, HYPRE_Int type)
Set the storage type of the matrix object to be constructed.
Currently, type can be either HYPRE_SSTRUCT (the default), HYPRE_STRUCT, or HYPRE_PARCSR.

## See also:

HYPRE_SStructMatrixGetObject

HYPRE_Int HYPRE_SStructMatrixGetObject (HYPRE_SStructMatrix matrix, void **object)
Get a reference to the constructed matrix object.

## See also:

HYPRE_SStructMatrixSetObjectType
HYPRE_Int HYPRE_SStructMatrixPrint (const char *filename, HYPRE_SStructMatrix matrix, HYPRE_Int all)

Print the matrix to file.
This is mainly for debugging purposes.
HYPRE_Int HYPRE_SStructMatrixRead(MPI_Comm comm, const char *filename, HYPRE_SStructMatrix *matrix_ptr)
Read the matrix from file.
This is mainly for debugging purposes.

## SStruct Vectors

typedef struct hypre_SStructVector_struct *HYPRE_SStructVector
The vector object.

## HYPRE_Int HYPRE_SStructVectorCreate(MPI_Comm comm, HYPRE_SStructGrid grid,

 HYPRE_SStructVector * vector)Create a vector object.
HYPRE_Int HYPRE_SStructVectorDestroy (HYPRE_SStructVector vector)
Destroy a vector object.
HYPRE_Int HYPRE_SStructVectorInitialize(HYPRE_SStructVector vector)
Prepare a vector object for setting coefficient values.
HYPRE_Int HYPRE_SStructVectorSetValues(HYPRE_SStructVector vector, HYPRE_Int part, HYPRE_Int *index, HYPRE_Int var, HYPRE_Complex *value)

Set vector coefficients index by index.
NOTE: For better efficiency, use HYPRE_SStructVectorSetBoxValues to set coefficients a box at a time.
NOTE: Users are required to set values on all processes that own the associated variables. This means that some data will be multiply defined.

HYPRE_Int HYPRE_SStructVectorAddToValues (HYPRE_SStructVector vector, HYPRE_Int part, HYPRE_Int *index, HYPRE_Int var, HYPRE_Complex *value)

Add to vector coefficients index by index.
NOTE: For better efficiency, use HYPRE_SStructVectorAddToBoxValues to set coefficients a box at a time.
NOTE: Users are required to set values on all processes that own the associated variables. This means that some data will be multiply defined.

HYPRE_Int HYPRE_SStructVectorAddFEMValues (HYPRE_SStructVector vector, HYPRE_Int part, HYPRE_Int *index, HYPRE_Complex *values)
Add finite element vector coefficients index by index.
The layout of the data in values is determined by the routine HYPRE_SStructGridSetFEMOrdering.
NOTE: For better efficiency, use HYPRE_SStructVectorAddFEMBoxValues to set coefficients a box at a time.

HYPRE_Int HYPRE_SStructVectorGetValues(HYPRE_SStructVector vector, HYPRE_Int part, HYPRE_Int *index, HYPRE_Int var, HYPRE_Complex *value)
Get vector coefficients index by index.
Users must first call the routine HYPRE_SStructVectorGather to ensure that data owned by multiple processes is correct.

NOTE: For better efficiency, use HYPRE_SStructVectorGetBoxValues to get coefficients a box at a time.
NOTE: Users may only get values on processes that own the associated variables.
HYPRE_Int HYPRE_SStructVectorGetFEMValues (HYPRE_SStructVector vector, HYPRE_Int part, HYPRE_Int *index, HYPRE_Complex *values)
Get finite element vector coefficients index by index.
The layout of the data in values is determined by the routine HYPRE_SStructGridSetFEMOrdering. Users must first call the routine HYPRE_SStructVectorGather to ensure that data owned by multiple processes is correct.
HYPRE_Int HYPRE_SStructVectorSetBoxValues (HYPRE_SStructVector vector, HYPRE_Int part,
HYPRE_Int *ilower, HYPRE_Int *iupper, HYPRE_Int var, HYPRE_Complex *values)
Set vector coefficients a box at a time.
The data in values is ordered as follows:

```
m = 0;
for (k = ilower[2]; k <= iupper[2]; k++)
    for (j = ilower[1]; j <= iupper[1]; j++)
        for (i = ilower[0]; i <= iupper[0]; i++)
        {
            values[m] = ...;
            m++;
        }
```

NOTE: Users are required to set values on all processes that own the associated variables. This means that some data will be multiply defined.
HYPRE_Int HYPRE_SStructVectorAddToBoxValues (HYPRE_SStructVector vector, HYPRE_Int part, HYPRE_Int *ilower, HYPRE_Int *iupper, HYPRE_Int var, HYPRE_Complex *values)

Add to vector coefficients a box at a time.
The data in values is ordered as in HYPRE_SStructVectorSetBoxValues.
NOTE: Users are required to set values on all processes that own the associated variables. This means that some data will be multiply defined.

HYPRE_Int HYPRE_SStructVectorSetBoxValues2 (HYPRE_SStructVector vector, HYPRE_Int part, HYPRE_Int *ilower, HYPRE_Int *iupper, HYPRE_Int var, HYPRE_Int *vilower, HYPRE_Int *viupper, HYPRE_Complex *values)
Set vector coefficients a box at a time.
The values array is logically box shaped with value-box extents vilower and viupper that must contain the set-box extents ilower and iupper . The data in the values array is ordered as in HYPRE_SStructVectorSetBoxValues, but based on the value-box extents.

HYPRE_Int HYPRE_SStructVectorAddToBoxValues2 (HYPRE_SStructVector vector, HYPRE_Int part, HYPRE_Int *ilower, HYPRE_Int *iupper, HYPRE_Int var, HYPRE_Int *vilower, HYPRE_Int *viupper, HYPRE_Complex *values)
Add to vector coefficients a box at a time.
The data in values is ordered as in HYPRE_SStructVectorSetBoxValues2.
HYPRE_Int HYPRE_SStructVectorAddFEMBoxValues(HYPRE_SStructVector vector, HYPRE_Int part, HYPRE_Int *ilower, HYPRE_Int *iupper, HYPRE_Complex *values)
Add finite element vector coefficients a box at a time.
The data in values is organized as an array of element vectors ordered as in HYPRE_SStructVectorSetBoxValues. The layout of the data entries of each element vector is determined by the routine HYPRE_SStructGridSetFEMOrdering.
HYPRE_Int HYPRE_SStructVectorAssemble (HYPRE_SStructVector vector)
Finalize the construction of the vector before using.
HYPRE_Int HYPRE_SStructVectorGetBoxValues(HYPRE_SStructVector vector, HYPRE_Int part, HYPRE_Int *ilower, HYPRE_Int *iupper, HYPRE_Int var, HYPRE_Complex *values)
Get vector coefficients a box at a time.
The data in values is ordered as in HYPRE_SStructVectorSetBoxValues. Users must first call the routine HYPRE_SStructVectorGather to ensure that data owned by multiple processes is correct.

NOTE: Users may only get values on processes that own the associated variables.
HYPRE_Int HYPRE_SStructVectorGetBoxValues2 (HYPRE_SStructVector vector, HYPRE_Int part, HYPRE_Int *ilower, HYPRE_Int *iupper, HYPRE_Int var, HYPRE_Int *vilower, HYPRE_Int *viupper, HYPRE_Complex *values)
Get vector coefficients a box at a time.
The data in values is ordered as in HYPRE_SStructVectorSetBoxValues2.
HYPRE_Int HYPRE_SStructVectorGetFEMBoxValues(HYPRE_SStructVector vector, HYPRE_Int part, HYPRE_Int *ilower, HYPRE_Int *iupper, HYPRE_Complex *values)
Does this even make sense to implement?
HYPRE_Int HYPRE_SStructVectorGather(HYPRE_SStructVector vector)
Gather vector data so that efficient GetValues can be done.
This routine must be called prior to calling GetValues to ensure that correct and consistent values are returned, especially for non cell-centered data that is shared between more than one processor.

HYPRE_Int HYPRE_SStructVectorSetObjectType (HYPRE_SStructVector vector, HYPRE_Int type)
Set the storage type of the vector object to be constructed.
Currently, type can be either HYPRE_SSTRUCT (the default), HYPRE_STRUCT, or HYPRE_PARCSR.

## See also:

HYPRE_SStructVectorGetObject
HYPRE_Int HYPRE_SStructVectorGetObject (HYPRE_SStructVector vector, void **object)
Get a reference to the constructed vector object.

## See also:

HYPRE_SStructVectorSetObjectType
HYPRE_Int HYPRE_SStructVectorPrint (const char *filename, HYPRE_SStructVector vector, HYPRE_Int all)

Print the vector to file.
This is mainly for debugging purposes.
HYPRE_Int HYPRE_SStructVectorRead(MPI_Comm comm, const char *filename, HYPRE_SStructVector *vector_ptr)
Read the vector from file.
This is mainly for debugging purposes.

### 8.3 IJ System Interface

## group IJSystemInterface

A linear-algebraic conceptual interface.
This interface represents a linear-algebraic conceptual view of a linear system. The ' I ' and ' J ' in the name are meant to be mnemonic for the traditional matrix notation $A(I, J)$.

## IJ Matrices

typedef struct hypre_IJMatrix_struct *HYPRE_IJMatrix
The matrix object.
HYPRE_Int HYPRE_IJMatrixCreate(MPI_Comm comm, HYPRE_BigInt ilower, HYPRE_BigInt iupper, HYPRE_BigInt jlower, HYPRE_BigInt jupper, HYPRE_IJMatrix *matrix)
Create a matrix object.
Each process owns some unique consecutive range of rows, indicated by the global row indices ilower and iupper. The row data is required to be such that the value of ilower on any process $p$ be exactly one more than the value of iupper on process $p-1$. Note that the first row of the global matrix may start with any integer value. In particular, one may use zero- or one-based indexing.

For square matrices, jlower and jupper typically should match ilower and iupper, respectively. For rectangular matrices, jlower and jupper should define a partitioning of the columns. This partitioning must be used for any vector $v$ that will be used in matrix-vector products with the rectangular matrix. The matrix data structure may use jlower and jupper to store the diagonal blocks (rectangular in general) of the matrix separately from the rest of the matrix.

Collective.

## HYPRE_Int HYPRE_IJMatrixDestroy (HYPRE_IJMatrix matrix)

Destroy a matrix object.
An object should be explicitly destroyed using this destructor when the user's code no longer needs direct access to it. Once destroyed, the object must not be referenced again. Note that the object may not be deallocated at the completion of this call, since there may be internal package references to the object. The object will then be destroyed when all internal reference counts go to zero.

## HYPRE_Int HYPRE_IJMatrixInitialize(HYPRE_IJMatrix matrix)

Prepare a matrix object for setting coefficient values.
This routine will also re-initialize an already assembled matrix, allowing users to modify coefficient values.

## HYPRE_Int HYPRE_IJMatrixInitialize_v2(HYPRE_IJMatrix matrix, HYPRE_MemoryLocation memory_location)

Prepare a matrix object for setting coefficient values.
This routine will also re-initialize an already assembled matrix, allowing users to modify coefficient values. This routine also specifies the memory location, i.e. host or device.
HYPRE_Int HYPRE_IJMatrixSetValues (HYPRE_IJMatrix matrix, HYPRE_Int nrows, HYPRE_Int *ncols, const HYPRE_BigInt *rows, const HYPRE_BigInt * cols, const HYPRE_Complex *values)
Sets values for nrows rows or partial rows of the matrix.
The arrays ncols and rows are of dimension nrows and contain the number of columns in each row and the row indices, respectively. The array cols contains the column indices for each of the rows, and is ordered by rows. The data in the values array corresponds directly to the column entries in cols. Erases any previous values at the specified locations and replaces them with new ones, or, if there was no value there before, inserts a new one if set locally. Note that it is not possible to set values on other processors. If one tries to set a value from proc i on proc j , proc i will erase all previous occurrences of this value in its stack (including values generated with AddToValues), and treat it like a zero value. The actual value needs to be set on proc j .

Note that a threaded version (threaded over the number of rows) will be called if HYPRE_IJMatrixSetOMPFlag is set to a value $!=0$. This requires that rows $[i]!=r o w s[j]$ for $\mathrm{i}!=\mathrm{j}$ and is only efficient if a large number of rows is set in one call to HYPRE_IJMatrixSetValues.
Not collective.
HYPRE_Int HYPRE_IJMatrixSetConstantValues (HYPRE_IJMatrix matrix, HYPRE_Complex value)
Sets all matrix coefficients of an already assembled matrix to value.
HYPRE_Int HYPRE_IJMatrixAddToValues (HYPRE_IJMatrix matrix, HYPRE_Int nrows, HYPRE_Int *ncols, const HYPRE_BigInt *rows, const HYPRE_BigInt *cols, const HYPRE_Complex *values)
Adds to values for nrows rows or partial rows of the matrix.
Usage details are analogous to HYPRE_IJMatrixSetValues. Adds to any previous values at the specified locations, or, if there was no value there before, inserts a new one. AddToValues can be used to add to values on other processors.

Note that a threaded version (threaded over the number of rows) will be called if HYPRE_IJMatrixSetOMPFlag is set to a value $!=0$. This requires that rows $[i]!=$ rows $[j]$ for $\mathrm{i}!=\mathrm{j}$ and is only efficient if a large number of rows is added in one call to HYPRE_IJMatrixAddToValues.

Not collective.
HYPRE_Int HYPRE_IJMatrixSetValues2 (HYPRE_IJMatrix matrix, HYPRE_Int nrows, HYPRE_Int
*ncols, const HYPRE_BigInt *rows, const HYPRE_Int
*row_indexes, const HYPRE_BigInt *cols, const
HYPRE_Complex *values)
Sets values for nrows rows or partial rows of the matrix.
Same as IJMatrixSetValues, but with an additional row_indexes array that provides indexes into the cols and values arrays. Because of this, there can be gaps between the row data in these latter two arrays.
HYPRE_Int HYPRE_IJMatrixAddToValues2 (HYPRE_IJMatrix matrix, HYPRE_Int nrows, HYPRE_Int
*ncols, const HYPRE_BigInt *rows, const HYPRE_Int
*row_indexes, const HYPRE_BigInt * cols, const
HYPRE_Complex *values)
Adds to values for nrows rows or partial rows of the matrix.
Same as IJMatrixAddToValues, but with an additional row_indexes array that provides indexes into the cols and values arrays. Because of this, there can be gaps between the row data in these latter two arrays.

HYPRE_Int HYPRE_IJMatrixAssemble(HYPRE_IJMatrix matrix)
Finalize the construction of the matrix before using.
HYPRE_Int HYPRE_IJMatrixGetRowCounts(HYPRE_IJMatrix matrix, HYPRE_Int nrows, HYPRE_BigInt *rows, HYPRE_Int *ncols)

Gets number of nonzeros elements for nrows rows specified in rows and returns them in ncols, which needs to be allocated by the user.
HYPRE_Int HYPRE_IJMatrixGetValues(HYPRE_IJMatrix matrix, HYPRE_Int nrows, HYPRE_Int *ncols, HYPRE_BigInt *rows, HYPRE_BigInt *cols, HYPRE_Complex *values)
Gets values for nrows rows or partial rows of the matrix.
Usage details are mostly analogous to HYPRE_IJMatrixSetValues. Note that if nrows is negative, the routine will return the column_indices and matrix coefficients of the (-nrows) rows contained in rows.
HYPRE_Int HYPRE_IJMatrixSetObjectType (HYPRE_IJMatrix matrix, HYPRE_Int type)
Set the storage type of the matrix object to be constructed.
Currently, type can only be HYPRE_PARCSR.
Not collective, but must be the same on all processes.

## See also:

HYPRE_IJMatrixGetObject
HYPRE_Int HYPRE_IJMatrixGetObjectType (HYPRE_IJMatrix matrix, HYPRE_Int *type)
Get the storage type of the constructed matrix object.
HYPRE_Int HYPRE_IJMatrixGetLocalRange (HYPRE_IJMatrix matrix, HYPRE_BigInt *ilower, HYPRE_BigInt *iupper, HYPRE_BigInt *jlower, HYPRE_BigInt *jupper)
Gets range of rows owned by this processor and range of column partitioning for this processor.

HYPRE_Int HYPRE_IJMatrixGetObject(HYPRE_IJMatrix matrix, void **object)
Get a reference to the constructed matrix object.

## See also:

HYPRE_IJMatrixSetObjectType
HYPRE_Int HYPRE_IJMatrixSetRowSizes (HYPRE_IJMatrix matrix, const HYPRE_Int *sizes)
(Optional) Set the max number of nonzeros to expect in each row.
The array sizes contains estimated sizes for each row on this process. This call can significantly improve the efficiency of matrix construction, and should always be utilized if possible.

Not collective.
HYPRE_Int HYPRE_IJMatrixSetDiagOffdSizes (HYPRE_IJMatrix matrix, const HYPRE_Int *diag_sizes, const HYPRE_Int *offdiag_sizes)
(Optional) Sets the exact number of nonzeros in each row of the diagonal and off-diagonal blocks.
The diagonal block is the submatrix whose column numbers correspond to rows owned by this process, and the off-diagonal block is everything else. The arrays diag_sizes and offdiag_sizes contain estimated sizes for each row of the diagonal and off-diagonal blocks, respectively. This routine can significantly improve the efficiency of matrix construction, and should always be utilized if possible.
Not collective.
HYPRE_Int HYPRE_IJMatrixSetMaxOffProcElmts(HYPRE_IJMatrix matrix, HYPRE_Int max_off_proc_elmts)
(Optional) Sets the maximum number of elements that are expected to be set (or added) on other processors from this processor This routine can significantly improve the efficiency of matrix construction, and should always be utilized if possible.

Not collective.
HYPRE_Int HYPRE_IJMatrixSetPrintLevel (HYPRE_IJMatrix matrix, HYPRE_Int print_level)
(Optional) Sets the print level, if the user wants to print error messages.
The default is 0 , i.e. no error messages are printed.
HYPRE_Int HYPRE_IJMatrixSetOMPFlag (HYPRE_IJMatrix matrix, HYPRE_Int omp_flag)
(Optional) if set, will use a threaded version of HYPRE_IJMatrixSetValues and HYPRE_IJMatrixAddToValues.

This is only useful if a large number of rows is set or added to at once.
NOTE that the values in the rows array of HYPRE_IJMatrixSetValues or HYPRE_IJMatrixAddToValues must be different from each other !!!

This option is VERY inefficient if only a small number of rows is set or added at once and/or if reallocation of storage is required and/or if values are added to off processor values.

HYPRE_Int HYPRE_IJMatrixRead(const char *filename, MPI_Comm comm, HYPRE_Int type, HYPRE_IJMatrix *matrix)

Read the matrix from file.
This is mainly for debugging purposes.

HYPRE_Int HYPRE_IJMatrixReadMM (const char *filename, MPI_Comm comm, HYPRE_Int type, HYPRE_IJMatrix *matrix)
Read the matrix from MM file.
This is mainly for debugging purposes.
HYPRE_Int HYPRE_IJMatrixPrint (HYPRE_IJMatrix matrix, const char *filename)
Print the matrix to file.
This is mainly for debugging purposes.
HYPRE_Int HYPRE_IJMatrixPrintBinary (HYPRE_IJMatrix matrix, const char *filename)
Print the matrix to file in binary format.
This is mainly for debugging purposes.
HYPRE_Int HYPRE_IJMatrixReadBinary (const char *filename, MPI_Comm comm, HYPRE_Int type, HYPRE_IJMatrix *matrix_ptr)

Read the matrix from file stored in binary format.
This is mainly for debugging purposes.

## IJ Vectors

typedef struct hypre_IJVector_struct *HYPRE_IJVector
The vector object.
HYPRE_Int HYPRE_IJVectorCreate(MPI_Comm comm, HYPRE_BigInt jlower, HYPRE_BigInt jupper, HYPRE_IJVector ${ }^{*}$ vector)

Create a vector object.
Each process owns some unique consecutive range of vector unknowns, indicated by the global indices jlower and jupper. The data is required to be such that the value of jlower on any process $p$ be exactly one more than the value of jupper on process $p-1$. Note that the first index of the global vector may start with any integer value. In particular, one may use zero- or one-based indexing.
Collective.
HYPRE_Int HYPRE_IJVectorDestroy (HYPRE_IJVector vector)
Destroy a vector object.
An object should be explicitly destroyed using this destructor when the user's code no longer needs direct access to it. Once destroyed, the object must not be referenced again. Note that the object may not be deallocated at the completion of this call, since there may be internal package references to the object. The object will then be destroyed when all internal reference counts go to zero.

## HYPRE_Int HYPRE_IJVectorInitialize(HYPRE_IJVector vector)

Prepare a vector object for setting coefficient values.
This routine will also re-initialize an already assembled vector, allowing users to modify coefficient values.
HYPRE_Int HYPRE_IJVectorInitialize_v2(HYPRE_IJVector vector, HYPRE_MemoryLocation memory_location)
Prepare a vector object for setting coefficient values.
This routine will also re-initialize an already assembled vector, allowing users to modify coefficient values. This routine also specifies the memory location, i.e. host or device.

HYPRE_Int HYPRE_IJVectorSetMaxOffProcElmts(HYPRE_IJVector vector, HYPRE_Int max_off_proc_elmts)
(Optional) Sets the maximum number of elements that are expected to be set (or added) on other processors from this processor This routine can significantly improve the efficiency of matrix construction, and should always be utilized if possible.

Not collective.
HYPRE_Int HYPRE_IJVectorSetNumComponents (HYPRE_IJVector vector, HYPRE_Int num_components) (Optional) Sets the number of components (vectors) of a multivector.

A vector is assumed to have a single component when this function is not called. This function must be called prior to HYPRE_IJVectorInitialize.

HYPRE_Int HYPRE_IJVectorSetComponent (HYPRE_IJVector vector, HYPRE_Int component) (Optional) Sets the component identifier of a vector with multiple components (multivector).

This can be used for Set/AddTo/Get purposes.
HYPRE_Int HYPRE_IJVectorSetValues(HYPRE_IJVector vector, HYPRE_Int nvalues, const HYPRE_BigInt *indices, const HYPRE_Complex *values)
Sets values in vector.
The arrays values and indices are of dimension nvalues and contain the vector values to be set and the corresponding global vector indices, respectively. Erases any previous values at the specified locations and replaces them with new ones. Note that it is not possible to set values on other processors. If one tries to set a value from proc i on proc j , proc i will erase all previous occurrences of this value in its stack (including values generated with AddToValues), and treat it like a zero value. The actual value needs to be set on proc j.

Not collective.
HYPRE_Int HYPRE_IJVectorAddToValues (HYPRE_IJVector vector, HYPRE_Int nvalues, const
HYPRE_BigInt *indices, const HYPRE_Complex *values)
Adds to values in vector.
Usage details are analogous to HYPRE_IJVectorSetValues. Adds to any previous values at the specified locations, or, if there was no value there before, inserts a new one. AddToValues can be used to add to values on other processors.

Not collective.
HYPRE_Int HYPRE_IJVectorAssemble(HYPRE_IJVector vector)
Finalize the construction of the vector before using.
HYPRE_Int HYPRE_IJVectorUpdateValues(HYPRE_IJVector vector, HYPRE_Int nvalues, const HYPRE_BigInt *indices, const HYPRE_Complex *values, HYPRE_Int action)
Update vectors by setting (action 1) or adding to (action 0 ) values in 'vector'.
Note that this function cannot update values owned by other processes and does not allow repeated index values in 'indices'.

Not collective.
HYPRE_Int HYPRE_IJVectorGetValues (HYPRE_IJVector vector, HYPRE_Int nvalues, const
HYPRE_BigInt *indices, HYPRE_Complex *values)
Gets values in vector.
Usage details are analogous to HYPRE_IJVectorSetValues.

Not collective.
HYPRE_Int HYPRE_IJVectorSetObjectType (HYPRE_IJVector vector, HYPRE_Int type)
Set the storage type of the vector object to be constructed.
Currently, type can only be HYPRE_PARCSR.
Not collective, but must be the same on all processes.

## See also:

HYPRE_IJVectorGetObject
HYPRE_Int HYPRE_IJVectorGetObjectType (HYPRE_IJVector vector, HYPRE_Int *type)
Get the storage type of the constructed vector object.
HYPRE_Int HYPRE_IJVectorGetLocalRange (HYPRE_IJVector vector, HYPRE_BigInt *jlower, HYPRE_BigInt *jupper)
Returns range of the part of the vector owned by this processor.
HYPRE_Int HYPRE_IJVectorGetObject(HYPRE_IJVector vector, void **object)
Get a reference to the constructed vector object.

See also:
HYPRE_IJVectorSetObjectType
HYPRE_Int HYPRE_IJVectorSetPrintLevel (HYPRE_IJVector vector, HYPRE_Int print_level)
(Optional) Sets the print level, if the user wants to print error messages.
The default is 0 , i.e. no error messages are printed.
HYPRE_Int HYPRE_IJVectorRead(const char *filename, MPI_Comm comm, HYPRE_Int type, HYPRE_IJVector *vector)
Read the vector from file.
This is mainly for debugging purposes.
HYPRE_Int HYPRE_IJVectorReadBinary (const char *filename, MPI_Comm comm, HYPRE_Int type, HYPRE_IJVector $*$ vector)
Read the vector from binary file.
This is mainly for debugging purposes.
HYPRE_Int HYPRE_IJVectorPrint (HYPRE_IJVector vector, const char *filename)
Print the vector to file.
This is mainly for debugging purposes.
HYPRE_Int HYPRE_IJVectorPrintBinary (HYPRE_IJVector vector, const char *filename)
Print the vector to binary file.
This is mainly for debugging purposes.
HYPRE_Int HYPRE_IJVectorInnerProd(HYPRE_IJVector x, HYPRE_IJVector y, HYPRE_Real *prod)
Computes the inner product between two vectors.

### 8.4 Struct Solvers

## group StructSolvers

Linear solvers for structured grids.
These solvers use matrix/vector storage schemes that are tailored to structured grid problems.

## Struct Solvers

typedef struct hypre_StructSolver_struct *HYPRE_StructSolver
The solver object.
typedef HYPRE_Int (*HYPRE_PtrToStructSolverFcn)(HYPRE_StructSolver, HYPRE_StructMatrix, HYPRE_StructVector, HYPRE_StructVector)
typedef HYPRE_Int (*HYPRE_PtrToModifyPCFcn)(HYPRE_Solver, HYPRE_Int, HYPRE_Real)

## HYPRE_MODIFYPC

## Struct Jacobi Solver

HYPRE_Int HYPRE_StructJacobiCreate(MPI_Comm comm, HYPRE_StructSolver *solver)
Create a solver object.
HYPRE_Int HYPRE_StructJacobiDestroy (HYPRE_StructSolver solver)
Destroy a solver object.
An object should be explicitly destroyed using this destructor when the user's code no longer needs direct access to it. Once destroyed, the object must not be referenced again. Note that the object may not be deallocated at the completion of this call, since there may be internal package references to the object. The object will then be destroyed when all internal reference counts go to zero.

HYPRE_Int HYPRE_StructJacobiSetup(HYPRE_StructSolver solver, HYPRE_StructMatrix A, HYPRE_StructVector b, HYPRE_StructVector x)

Prepare to solve the system.
The coefficient data in $b$ and $x$ is ignored here, but information about the layout of the data may be used.
HYPRE_Int HYPRE_StructJacobiSolve(HYPRE_StructSolver solver, HYPRE_StructMatrix A, HYPRE_StructVector b, HYPRE_StructVector x)
Solve the system.
HYPRE_Int HYPRE_StructJacobiSetTol (HYPRE_StructSolver solver, HYPRE_Real tol) (Optional) Set the convergence tolerance.

HYPRE_Int HYPRE_StructJacobiGetTol (HYPRE_StructSolver solver, HYPRE_Real *tol)
HYPRE_Int HYPRE_StructJacobiSetMaxIter (HYPRE_StructSolver solver, HYPRE_Int max_iter)
(Optional) Set maximum number of iterations.
HYPRE_Int HYPRE_StructJacobiGetMaxIter(HYPRE_StructSolver solver, HYPRE_Int *max_iter)

HYPRE_Int HYPRE_StructJacobiSetZeroGuess (HYPRE_StructSolver solver)
(Optional) Use a zero initial guess.
This allows the solver to cut corners in the case where a zero initial guess is needed (e.g., for preconditioning) to reduce compuational cost.

HYPRE_Int HYPRE_StructJacobiGetZeroGuess (HYPRE_StructSolver solver, HYPRE_Int *zeroguess)
HYPRE_Int HYPRE_StructJacobiSetNonZeroGuess (HYPRE_StructSolver solver)
(Optional) Use a nonzero initial guess.
This is the default behavior, but this routine allows the user to switch back after using SetZeroGuess.
HYPRE_Int HYPRE_StructJacobiGetNumIterations (HYPRE_StructSolver solver, HYPRE_Int *num_iterations)
Return the number of iterations taken.
HYPRE_Int HYPRE_StructJacobiGetFinalRelativeResidualNorm (HYPRE_StructSolver solver,
HYPRE_Real *norm)
Return the norm of the final relative residual.

## Struct PFMG Solver

PFMG is a semicoarsening multigrid solver that uses pointwise relaxation.
For periodic problems, users should try to set the grid size in periodic dimensions to be as close to a power-of-two as possible. That is, if the grid size in a periodic dimension is given by $N=2^{m} * M$ where $M$ is not a power-of-two, then $M$ should be as small as possible. Large values of $M$ will generally result in slower convergence rates.

HYPRE_Int HYPRE_StructPFMGCreate(MPI_Comm comm, HYPRE_StructSolver *solver)
Create a solver object.
HYPRE_Int HYPRE_StructPFMGDestroy (HYPRE_StructSolver solver)
Destroy a solver object.
HYPRE_Int HYPRE_StructPFMGSetup (HYPRE_StructSolver solver, HYPRE_StructMatrix A, HYPRE_StructVector b, HYPRE_StructVector x)
Prepare to solve the system.
The coefficient data in $b$ and $x$ is ignored here, but information about the layout of the data may be used.
HYPRE_Int HYPRE_StructPFMGSolve(HYPRE_StructSolver solver, HYPRE_StructMatrix A, HYPRE_StructVector b, HYPRE_StructVector x)
Solve the system.
HYPRE_Int HYPRE_StructPFMGSetTol(HYPRE_StructSolver solver, HYPRE_Real tol)
(Optional) Set the convergence tolerance.
HYPRE_Int HYPRE_StructPFMGGetTol (HYPRE_StructSolver solver, HYPRE_Real *tol)
HYPRE_Int HYPRE_StructPFMGSetMaxIter (HYPRE_StructSolver solver, HYPRE_Int max_iter)
(Optional) Set maximum number of iterations.
HYPRE_Int HYPRE_StructPFMGGetMaxIter (HYPRE_StructSolver solver, HYPRE_Int *max_iter)
HYPRE_Int HYPRE_StructPFMGSetMaxLevels(HYPRE_StructSolver solver, HYPRE_Int max_levels)
(Optional) Set maximum number of multigrid grid levels.

HYPRE_Int HYPRE_StructPFMGGetMaxLevels(HYPRE_StructSolver solver, HYPRE_Int *max_levels)
HYPRE_Int HYPRE_StructPFMGSetRelChange (HYPRE_StructSolver solver, HYPRE_Int rel_change)
(Optional) Additionally require that the relative difference in successive iterates be small.
HYPRE_Int HYPRE_StructPFMGGetRelChange (HYPRE_StructSolver solver, HYPRE_Int *rel_change)
HYPRE_Int HYPRE_StructPFMGSetZeroGuess (HYPRE_StructSolver solver)
(Optional) Use a zero initial guess.
This allows the solver to cut corners in the case where a zero initial guess is needed (e.g., for preconditioning) to reduce compuational cost.

HYPRE_Int HYPRE_StructPFMGGetZeroGuess(HYPRE_StructSolver solver, HYPRE_Int *zeroguess)
HYPRE_Int HYPRE_StructPFMGSetNonZeroGuess(HYPRE_StructSolver solver)
(Optional) Use a nonzero initial guess.
This is the default behavior, but this routine allows the user to switch back after using SetZeroGuess.
HYPRE_Int HYPRE_StructPFMGSetRelaxType (HYPRE_StructSolver solver, HYPRE_Int relax_type) (Optional) Set relaxation type.
Current relaxation methods set by relax_type are:

- 0 : Jacobi
- 1: Weighted Jacobi (default)
- 2 : Red/Black Gauss-Seidel (symmetric: RB pre-relaxation, BR post-relaxation)
- 3 : Red/Black Gauss-Seidel (nonsymmetric: RB pre- and post-relaxation)

HYPRE_Int HYPRE_StructPFMGGetRelaxType (HYPRE_StructSolver solver, HYPRE_Int *relax_type)
HYPRE_Int HYPRE_StructPFMGSetJacobiWeight (HYPRE_StructSolver solver, HYPRE_Real weight)
HYPRE_Int HYPRE_StructPFMGGetJacobiWeight (HYPRE_StructSolver solver, HYPRE_Real *weight)
HYPRE_Int HYPRE_StructPFMGSetRAPType(HYPRE_StructSolver solver, HYPRE_Int rap_type)
(Optional) Set type of coarse-grid operator to use.
Current operators set by rap_type are:

- 0 : Galerkin (default)
- 1 : non-Galerkin 5-pt or 7-pt stencils

Both operators are constructed algebraically. The non-Galerkin option maintains a 5-pt stencil in 2D and a 7-pt stencil in 3D on all grid levels. The stencil coefficients are computed by averaging techniques.

HYPRE_Int HYPRE_StructPFMGGetRAPType(HYPRE_StructSolver solver, HYPRE_Int *rap_type)
HYPRE_Int HYPRE_StructPFMGSetNumPreRelax (HYPRE_StructSolver solver, HYPRE_Int num_pre_relax)
(Optional) Set number of relaxation sweeps before coarse-grid correction.
HYPRE_Int HYPRE_StructPFMGGetNumPreRelax (HYPRE_StructSolver solver, HYPRE_Int *num_pre_relax)

HYPRE_Int HYPRE_StructPFMGSetNumPostRelax (HYPRE_StructSolver solver, HYPRE_Int num_post_relax)
(Optional) Set number of relaxation sweeps after coarse-grid correction.
HYPRE_Int HYPRE_StructPFMGGetNumPostRelax (HYPRE_StructSolver solver, HYPRE_Int *num_post_relax)

HYPRE_Int HYPRE_StructPFMGSetSkipRelax (HYPRE_StructSolver solver, HYPRE_Int skip_relax) (Optional) Skip relaxation on certain grids for isotropic problems. This can greatly improve efficiency by eliminating unnecessary relaxations when the underlying problem is isotropic.
HYPRE_Int HYPRE_StructPFMGGetSkipRelax (HYPRE_StructSolver solver, HYPRE_Int *skip_relax)
HYPRE_Int HYPRE_StructPFMGSetDxyz (HYPRE_StructSolver solver, HYPRE_Real *dxyz)
HYPRE_Int HYPRE_StructPFMGSetLogging (HYPRE_StructSolver solver, HYPRE_Int logging)
(Optional) Set the amount of logging to do.
HYPRE_Int HYPRE_StructPFMGGetLogging (HYPRE_StructSolver solver, HYPRE_Int *logging)
HYPRE_Int HYPRE_StructPFMGSetPrintLevel (HYPRE_StructSolver solver, HYPRE_Int print_level) (Optional) Set the amount of printing to do to the screen.

HYPRE_Int HYPRE_StructPFMGGetPrintLevel (HYPRE_StructSolver solver, HYPRE_Int *print_level)
HYPRE_Int HYPRE_StructPFMGGetNumIterations(HYPRE_StructSolver solver, HYPRE_Int
*num_iterations)
Return the number of iterations taken.
HYPRE_Int HYPRE_StructPFMGGetFinalRelativeResidualNorm(HYPRE_StructSolver solver, HYPRE_Real *norm)
Return the norm of the final relative residual.

## Struct SMG Solver

SMG is a semicoarsening multigrid solver that uses plane smoothing (in 3D).
The plane smoother calls a 2D SMG algorithm with line smoothing, and the line smoother is cyclic reduction (1D SMG). For periodic problems, the grid size in periodic dimensions currently must be a power-of-two.

HYPRE_Int HYPRE_StructSMGCreate(MPI_Comm comm, HYPRE_StructSolver *solver)
Create a solver object.
HYPRE_Int HYPRE_StructSMGDestroy (HYPRE_StructSolver solver)
Destroy a solver object.
HYPRE_Int HYPRE_StructSMGSetup(HYPRE_StructSolver solver, HYPRE_StructMatrix A, HYPRE_StructVector b, HYPRE_StructVector x)

Prepare to solve the system.
The coefficient data in $b$ and $x$ is ignored here, but information about the layout of the data may be used.
HYPRE_Int HYPRE_StructSMGSolve(HYPRE_StructSolver solver, HYPRE_StructMatrix A, HYPRE_StructVector b, HYPRE_StructVector x)
Solve the system.

HYPRE_Int HYPRE_StructSMGSetMemoryUse(HYPRE_StructSolver solver, HYPRE_Int memory_use)
HYPRE_Int HYPRE_StructSMGGetMemoryUse(HYPRE_StructSolver solver, HYPRE_Int *memory_use)
HYPRE_Int HYPRE_StructSMGSetTol (HYPRE_StructSolver solver, HYPRE_Real tol)
(Optional) Set the convergence tolerance.
HYPRE_Int HYPRE_StructSMGGetTol (HYPRE_StructSolver solver, HYPRE_Real *tol)
HYPRE_Int HYPRE_StructSMGSetMaxIter (HYPRE_StructSolver solver, HYPRE_Int max_iter) (Optional) Set maximum number of iterations.

HYPRE_Int HYPRE_StructSMGGetMaxIter (HYPRE_StructSolver solver, HYPRE_Int *max_iter)
HYPRE_Int HYPRE_StructSMGSetRelChange (HYPRE_StructSolver solver, HYPRE_Int rel_change) (Optional) Additionally require that the relative difference in successive iterates be small.

HYPRE_Int HYPRE_StructSMGGetRelChange (HYPRE_StructSolver solver, HYPRE_Int *rel_change)
HYPRE_Int HYPRE_StructSMGSetZeroGuess (HYPRE_StructSolver solver)
(Optional) Use a zero initial guess.
This allows the solver to cut corners in the case where a zero initial guess is needed (e.g., for preconditioning) to reduce compuational cost.

HYPRE_Int HYPRE_StructSMGGetZeroGuess(HYPRE_StructSolver solver, HYPRE_Int *zeroguess)
HYPRE_Int HYPRE_StructSMGSetNonZeroGuess(HYPRE_StructSolver solver)
(Optional) Use a nonzero initial guess.
This is the default behavior, but this routine allows the user to switch back after using SetZeroGuess.
HYPRE_Int HYPRE_StructSMGSetNumPreRelax (HYPRE_StructSolver solver, HYPRE_Int num_pre_relax) (Optional) Set number of relaxation sweeps before coarse-grid correction.

HYPRE_Int HYPRE_StructSMGGetNumPreRelax(HYPRE_StructSolver solver, HYPRE_Int
*num_pre_relax)
HYPRE_Int HYPRE_StructSMGSetNumPostRelax (HYPRE_StructSolver solver, HYPRE_Int num_post_relax)
(Optional) Set number of relaxation sweeps after coarse-grid correction.
HYPRE_Int HYPRE_StructSMGGetNumPostRelax (HYPRE_StructSolver solver, HYPRE_Int *num_post_relax)

HYPRE_Int HYPRE_StructSMGSetLogging (HYPRE_StructSolver solver, HYPRE_Int logging) (Optional) Set the amount of logging to do.

HYPRE_Int HYPRE_StructSMGGetLogging(HYPRE_StructSolver solver, HYPRE_Int *logging)
HYPRE_Int HYPRE_StructSMGSetPrintLevel (HYPRE_StructSolver solver, HYPRE_Int print_level) (Optional) Set the amount of printing to do to the screen.

HYPRE_Int HYPRE_StructSMGGetPrintLevel(HYPRE_StructSolver solver, HYPRE_Int *print_level)
HYPRE_Int HYPRE_StructSMGGetNumIterations(HYPRE_StructSolver solver, HYPRE_Int *num_iterations)
Return the number of iterations taken.

HYPRE_Int HYPRE_StructSMGGetFinalRelativeResidualNorm(HYPRE_StructSolver solver, HYPRE_Real *norm)
Return the norm of the final relative residual.

## Struct CycRed Solver

CycRed is a cyclic reduction solver that simultaneously solves a collection of 1D tridiagonal systems embedded in a d-dimensional grid.

## HYPRE_Int HYPRE_StructCycRedCreate(MPI_Comm comm, HYPRE_StructSolver *solver)

Create a solver object.
HYPRE_Int HYPRE_StructCycRedDestroy (HYPRE_StructSolver solver)
Destroy a solver object.
HYPRE_Int HYPRE_StructCycRedSetup(HYPRE_StructSolver solver, HYPRE_StructMatrix A, HYPRE_StructVector b, HYPRE_StructVector x)
Prepare to solve the system.
The coefficient data in $b$ and $x$ is ignored here, but information about the layout of the data may be used.
HYPRE_Int HYPRE_StructCycRedSolve(HYPRE_StructSolver solver, HYPRE_StructMatrix A, HYPRE_StructVector b, HYPRE_StructVector x)
Solve the system.
HYPRE_Int HYPRE_StructCycRedSetTDim(HYPRE_StructSolver solver, HYPRE_Int tdim)
(Optional) Set the dimension number for the embedded 1D tridiagonal systems.
The default is $t d i m=0$.
HYPRE_Int HYPRE_StructCycRedSetBase(HYPRE_StructSolver solver, HYPRE_Int ndim, HYPRE_Int
*base_index, HYPRE_Int *base_stride)
(Optional) Set the base index and stride for the embedded 1D systems.
The stride must be equal one in the dimension corresponding to the 1 D systems (see HYPRE_StructCycRedSetTDim).

## Struct PCG Solver

These routines should be used in conjunction with the generic interface in Krylov Solvers.
HYPRE_Int HYPRE_StructPCGCreate(MPI_Comm comm, HYPRE_StructSolver *solver)
Create a solver object.
HYPRE_Int HYPRE_StructPCGDestroy (HYPRE_StructSolver solver)
Destroy a solver object.
HYPRE_Int HYPRE_StructPCGSetup(HYPRE_StructSolver solver, HYPRE_StructMatrix A, HYPRE_StructVector b, HYPRE_StructVector x)

HYPRE_Int HYPRE_StructPCGSolve(HYPRE_StructSolver solver, HYPRE_StructMatrix A, HYPRE_StructVector b, HYPRE_StructVector x)

HYPRE_Int HYPRE_StructPCGSetTol (HYPRE_StructSolver solver, HYPRE_Real tol)
HYPRE_Int HYPRE_StructPCGSetAbsoluteTol(HYPRE_StructSolver solver, HYPRE_Real tol)

HYPRE_Int HYPRE_StructPCGSetMaxIter (HYPRE_StructSolver solver, HYPRE_Int max_iter)
HYPRE_Int HYPRE_StructPCGSetTwoNorm (HYPRE_StructSolver solver, HYPRE_Int two_norm)
HYPRE_Int HYPRE_StructPCGSetRelChange (HYPRE_StructSolver solver, HYPRE_Int rel_change)
HYPRE_Int HYPRE_StructPCGSetPrecond(HYPRE_StructSolver solver, HYPRE_PtrToStructSolverFcn precond, HYPRE_PtrToStructSolverFcn precond_setup, HYPRE_StructSolver precond_solver)

HYPRE_Int HYPRE_StructPCGSetLogging(HYPRE_StructSolver solver, HYPRE_Int logging)
HYPRE_Int HYPRE_StructPCGSetPrintLevel (HYPRE_StructSolver solver, HYPRE_Int level)
HYPRE_Int HYPRE_StructPCGGetNumIterations(HYPRE_StructSolver solver, HYPRE_Int *num_iterations)

HYPRE_Int HYPRE_StructPCGGetFinalRelativeResidualNorm(HYPRE_StructSolver solver, HYPRE_Real *norm)

HYPRE_Int HYPRE_StructPCGGetResidual (HYPRE_StructSolver solver, void **residual)
HYPRE_Int HYPRE_StructDiagScaleSetup(HYPRE_StructSolver solver, HYPRE_StructMatrix A, HYPRE_StructVector y, HYPRE_StructVector x)
Setup routine for diagonal preconditioning.
HYPRE_Int HYPRE_StructDiagScale(HYPRE_StructSolver solver, HYPRE_StructMatrix HA, HYPRE_StructVector Hy, HYPRE_StructVector Hx)
Solve routine for diagonal preconditioning.

## Struct GMRES Solver

These routines should be used in conjunction with the generic interface in Krylov Solvers.
HYPRE_Int HYPRE_StructGMRESCreate(MPI_Comm comm, HYPRE_StructSolver *solver)
Create a solver object.
HYPRE_Int HYPRE_StructGMRESDestroy (HYPRE_StructSolver solver)
Destroy a solver object.
HYPRE_Int HYPRE_StructGMRESSetup(HYPRE_StructSolver solver, HYPRE_StructMatrix A, HYPRE_StructVector b, HYPRE_StructVector x)

HYPRE_Int HYPRE_StructGMRESSolve(HYPRE_StructSolver solver, HYPRE_StructMatrix A,
HYPRE_StructVector b, HYPRE_StructVector x)
HYPRE_Int HYPRE_StructGMRESSetTol (HYPRE_StructSolver solver, HYPRE_Real tol)
HYPRE_Int HYPRE_StructGMRESSetAbsoluteTol (HYPRE_StructSolver solver, HYPRE_Real tol)
HYPRE_Int HYPRE_StructGMRESSetMaxIter (HYPRE_StructSolver solver, HYPRE_Int max_iter)
HYPRE_Int HYPRE_StructGMRESSetKDim(HYPRE_StructSolver solver, HYPRE_Int k_dim)
HYPRE_Int HYPRE_StructGMRESSetPrecond(HYPRE_StructSolver solver, HYPRE_PtrToStructSolverFcn precond, HYPRE_PtrToStructSolverFcn precond_setup, HYPRE_StructSolver precond_solver)

HYPRE_Int HYPRE_StructGMRESSetLogging (HYPRE_StructSolver solver, HYPRE_Int logging) HYPRE_Int HYPRE_StructGMRESSetPrintLevel (HYPRE_StructSolver solver, HYPRE_Int level)

HYPRE_Int HYPRE_StructGMRESGetNumIterations(HYPRE_StructSolver solver, HYPRE_Int *num_iterations)

HYPRE_Int HYPRE_StructGMRESGetFinalRelativeResidualNorm(HYPRE_StructSolver solver, HYPRE_Real *norm)

HYPRE_Int HYPRE_StructGMRESGetResidual (HYPRE_StructSolver solver, void **residual)

## Struct FlexGMRES Solver

These routines should be used in conjunction with the generic interface in Krylov Solvers.
HYPRE_Int HYPRE_StructFlexGMRESCreate(MPI_Comm comm, HYPRE_StructSolver *solver) Create a solver object.

HYPRE_Int HYPRE_StructFlexGMRESDestroy (HYPRE_StructSolver solver)
Destroy a solver object.
HYPRE_Int HYPRE_StructFlexGMRESSetup(HYPRE_StructSolver solver, HYPRE_StructMatrix A, HYPRE_StructVector b, HYPRE_StructVector x)

HYPRE_Int HYPRE_StructFlexGMRESSolve(HYPRE_StructSolver solver, HYPRE_StructMatrix A, HYPRE_StructVector b, HYPRE_StructVector x)

HYPRE_Int HYPRE_StructFlexGMRESSetTol (HYPRE_StructSolver solver, HYPRE_Real tol)
HYPRE_Int HYPRE_StructFlexGMRESSetAbsoluteTol (HYPRE_StructSolver solver, HYPRE_Real tol)
HYPRE_Int HYPRE_StructFlexGMRESSetMaxIter (HYPRE_StructSolver solver, HYPRE_Int max_iter)
HYPRE_Int HYPRE_StructFlexGMRESSetKDim(HYPRE_StructSolver solver, HYPRE_Int k_dim)
HYPRE_Int HYPRE_StructFlexGMRESSetPrecond(HYPRE_StructSolver solver,
HYPRE_PtrToStructSolverFcn precond,
HYPRE_PtrToStructSolverFcn precond_setup,
HYPRE_StructSolver precond_solver)
HYPRE_Int HYPRE_StructFlexGMRESSetLogging(HYPRE_StructSolver solver, HYPRE_Int logging)
HYPRE_Int HYPRE_StructFlexGMRESSetPrintLevel (HYPRE_StructSolver solver, HYPRE_Int level)
HYPRE_Int HYPRE_StructFlexGMRESGetNumIterations(HYPRE_StructSolver solver, HYPRE_Int *num_iterations)

HYPRE_Int HYPRE_StructFlexGMRESGetFinalRelativeResidualNorm(HYPRE_StructSolver solver, HYPRE_Real *norm)

HYPRE_Int HYPRE_StructFlexGMRESGetResidual (HYPRE_StructSolver solver, void **residual)
HYPRE_Int HYPRE_StructFlexGMRESSetModifyPC(HYPRE_StructSolver solver,
HYPRE_PtrToModifyPCFcn modify_pc)

## Struct LGMRES Solver

These routines should be used in conjunction with the generic interface in Krylov Solvers.
HYPRE_Int HYPRE_StructLGMRESCreate(MPI_Comm comm, HYPRE_StructSolver *solver)
Create a solver object.
HYPRE_Int HYPRE_StructLGMRESDestroy (HYPRE_StructSolver solver)
Destroy a solver object.
HYPRE_Int HYPRE_StructLGMRESSetup(HYPRE_StructSolver solver, HYPRE_StructMatrix A, HYPRE_StructVector b, HYPRE_StructVector x)

HYPRE_Int HYPRE_StructLGMRESSolve(HYPRE_StructSolver solver, HYPRE_StructMatrix A, HYPRE_StructVector b , HYPRE_StructVector x )

HYPRE_Int HYPRE_StructLGMRESSetTol (HYPRE_StructSolver solver, HYPRE_Real tol)
HYPRE_Int HYPRE_StructLGMRESSetAbsoluteTol (HYPRE_StructSolver solver, HYPRE_Real tol)
HYPRE_Int HYPRE_StructLGMRESSetMaxIter(HYPRE_StructSolver solver, HYPRE_Int max_iter)
HYPRE_Int HYPRE_StructLGMRESSetKDim(HYPRE_StructSolver solver, HYPRE_Int k_dim)
HYPRE_Int HYPRE_StructLGMRESSetAugDim (HYPRE_StructSolver solver, HYPRE_Int aug_dim)
HYPRE_Int HYPRE_StructLGMRESSetPrecond(HYPRE_StructSolver solver, HYPRE_PtrToStructSolverFcn precond, HYPRE_PtrToStructSolverFcn precond_setup, HYPRE_StructSolver precond_solver)

HYPRE_Int HYPRE_StructLGMRESSetLogging(HYPRE_StructSolver solver, HYPRE_Int logging)
HYPRE_Int HYPRE_StructLGMRESSetPrintLevel (HYPRE_StructSolver solver, HYPRE_Int level)
HYPRE_Int HYPRE_StructLGMRESGetNumIterations(HYPRE_StructSolver solver, HYPRE_Int *num_iterations)

HYPRE_Int HYPRE_StructLGMRESGetFinalRelativeResidualNorm(HYPRE_StructSolver solver, HYPRE_Real *norm)

HYPRE_Int HYPRE_StructLGMRESGetResidual (HYPRE_StructSolver solver, void **residual)

## Struct BiCGSTAB Solver

These routines should be used in conjunction with the generic interface in Krylov Solvers.
HYPRE_Int HYPRE_StructBiCGSTABCreate(MPI_Comm comm, HYPRE_StructSolver *solver)
Create a solver object.
HYPRE_Int HYPRE_StructBiCGSTABDestroy (HYPRE_StructSolver solver)
Destroy a solver object.
HYPRE_Int HYPRE_StructBiCGSTABSetup (HYPRE_StructSolver solver, HYPRE_StructMatrix A, HYPRE_StructVector b , HYPRE_StructVector x )

HYPRE_Int HYPRE_StructBiCGSTABSolve(HYPRE_StructSolver solver, HYPRE_StructMatrix A, HYPRE_StructVector b, HYPRE_StructVector x)

HYPRE_Int HYPRE_StructBiCGSTABSetTol (HYPRE_StructSolver solver, HYPRE_Real tol)
HYPRE_Int HYPRE_StructBiCGSTABSetAbsoluteTol (HYPRE_StructSolver solver, HYPRE_Real tol)
HYPRE_Int HYPRE_StructBiCGSTABSetMaxIter (HYPRE_StructSolver solver, HYPRE_Int max_iter)
HYPRE_Int HYPRE_StructBiCGSTABSetPrecond(HYPRE_StructSolver solver, HYPRE_PtrToStructSolverFcn precond, HYPRE_PtrToStructSolverFcn precond_setup, HYPRE_StructSolver precond_solver)

HYPRE_Int HYPRE_StructBiCGSTABSetLogging (HYPRE_StructSolver solver, HYPRE_Int logging)
HYPRE_Int HYPRE_StructBiCGSTABSetPrintLevel (HYPRE_StructSolver solver, HYPRE_Int level)
HYPRE_Int HYPRE_StructBiCGSTABGetNumIterations(HYPRE_StructSolver solver, HYPRE_Int *num_iterations)

HYPRE_Int HYPRE_StructBiCGSTABGetFinalRelativeResidualNorm(HYPRE_StructSolver solver, HYPRE_Real *norm)

HYPRE_Int HYPRE_StructBiCGSTABGetResidual (HYPRE_StructSolver solver, void **residual)

## Struct Hybrid Solver

HYPRE_Int HYPRE_StructHybridCreate(MPI_Comm comm, HYPRE_StructSolver *solver)
Create a solver object.
HYPRE_Int HYPRE_StructHybridDestroy (HYPRE_StructSolver solver)
Destroy a solver object.
HYPRE_Int HYPRE_StructHybridSetup(HYPRE_StructSolver solver, HYPRE_StructMatrix A, HYPRE_StructVector b , HYPRE_StructVector x )

Prepare to solve the system.
The coefficient data in $b$ and $x$ is ignored here, but information about the layout of the data may be used.
HYPRE_Int HYPRE_StructHybridSolve(HYPRE_StructSolver solver, HYPRE_StructMatrix A, HYPRE_StructVector b, HYPRE_StructVector x)
Solve the system.
HYPRE_Int HYPRE_StructHybridSetTol (HYPRE_StructSolver solver, HYPRE_Real tol)
(Optional) Set the convergence tolerance.
HYPRE_Int HYPRE_StructHybridSetConvergenceTol (HYPRE_StructSolver solver, HYPRE_Real cf_tol)
(Optional) Set an accepted convergence tolerance for diagonal scaling (DS).
The solver will switch preconditioners if the convergence of DS is slower than cf_tol.
HYPRE_Int HYPRE_StructHybridSetDSCGMaxIter (HYPRE_StructSolver solver, HYPRE_Int ds_max_its)
(Optional) Set maximum number of iterations for diagonal scaling (DS).
The solver will switch preconditioners if DS reaches $d s \_m a x \_i t s$.
HYPRE_Int HYPRE_StructHybridSetPCGMaxIter (HYPRE_StructSolver solver, HYPRE_Int pre_max_its)
(Optional) Set maximum number of iterations for general preconditioner (PRE).
The solver will stop if PRE reaches pre_max_its.

HYPRE_Int HYPRE_StructHybridSetTwoNorm(HYPRE_StructSolver solver, HYPRE_Int two_norm) (Optional) Use the two-norm in stopping criteria.

HYPRE_Int HYPRE_StructHybridSetStopCrit(HYPRE_StructSolver solver, HYPRE_Int stop_crit)
HYPRE_Int HYPRE_StructHybridSetRelChange (HYPRE_StructSolver solver, HYPRE_Int rel_change) (Optional) Additionally require that the relative difference in successive iterates be small.

HYPRE_Int HYPRE_StructHybridSetSolverType(HYPRE_StructSolver solver, HYPRE_Int solver_type) (Optional) Set the type of Krylov solver to use.

Current krylov methods set by solver_type are:

- 0 : PCG (default)
- 1: GMRES
- 2 : BiCGSTAB

HYPRE_Int HYPRE_StructHybridSetRecomputeResidual (HYPRE_StructSolver solver, HYPRE_Int recompute_residual)
(Optional) Set recompute residual (don't rely on 3-term recurrence).
HYPRE_Int HYPRE_StructHybridGetRecomputeResidual (HYPRE_StructSolver solver, HYPRE_Int *recompute_residual)
(Optional) Get recompute residual option.
HYPRE_Int HYPRE_StructHybridSetRecomputeResidualP(HYPRE_StructSolver solver, HYPRE_Int recompute_residual_p)
(Optional) Set recompute residual period (don't rely on 3-term recurrence).
Recomputes residual after every specified number of iterations.
HYPRE_Int HYPRE_StructHybridGetRecomputeResidualP (HYPRE_StructSolver solver, HYPRE_Int *recompute_residual_p)
(Optional) Get recompute residual period option.
HYPRE_Int HYPRE_StructHybridSetKDim (HYPRE_StructSolver solver, HYPRE_Int k_dim) (Optional) Set the maximum size of the Krylov space when using GMRES.

HYPRE_Int HYPRE_StructHybridSetPrecond(HYPRE_StructSolver solver, HYPRE_PtrToStructSolverFcn precond, HYPRE_PtrToStructSolverFcn precond_setup, HYPRE_StructSolver precond_solver)
(Optional) Set the preconditioner to use.
HYPRE_Int HYPRE_StructHybridSetLogging (HYPRE_StructSolver solver, HYPRE_Int logging) (Optional) Set the amount of logging to do.
HYPRE_Int HYPRE_StructHybridSetPrintLevel (HYPRE_StructSolver solver, HYPRE_Int print_level) (Optional) Set the amount of printing to do to the screen.
HYPRE_Int HYPRE_StructHybridGetNumIterations(HYPRE_StructSolver solver, HYPRE_Int *num_its)
Return the number of iterations taken.

HYPRE_Int HYPRE_StructHybridGetDSCGNumIterations(HYPRE_StructSolver solver, HYPRE_Int *ds_num_its)
Return the number of diagonal scaling iterations taken.
HYPRE_Int HYPRE_StructHybridGetPCGNumIterations(HYPRE_StructSolver solver, HYPRE_Int *pre_num_its)
Return the number of general preconditioning iterations taken.
HYPRE_Int HYPRE_StructHybridGetFinalRelativeResidualNorm(HYPRE_StructSolver solver, HYPRE_Real *norm)

Return the norm of the final relative residual.
HYPRE_Int HYPRE_StructHybridSetPCGAbsoluteTolFactor (HYPRE_StructSolver solver, HYPRE_Real pcg_atolf)

## Struct LOBPCG Eigensolver

These routines should be used in conjunction with the generic interface in Eigensolvers.
HYPRE_Int HYPRE_StructSetupInterpreter (mv_InterfaceInterpreter *i)
Load interface interpreter.
Vector part loaded with hypre_StructKrylov functions and multivector part loaded with mv_TempMultiVector functions.
HYPRE_Int HYPRE_StructSetupMatvec (HYPRE_MatvecFunctions *mv)
Load Matvec interpreter with hypre_StructKrylov functions.

## Functions

HYPRE_Int HYPRE_StructSparseMSGCreate(MPI_Comm comm, HYPRE_StructSolver *solver)
HYPRE_Int HYPRE_StructSparseMSGDestroy (HYPRE_StructSolver solver)
HYPRE_Int HYPRE_StructSparseMSGSetup (HYPRE_StructSolver solver, HYPRE_StructMatrix A, HYPRE_StructVector b, HYPRE_StructVector x)

HYPRE_Int HYPRE_StructSparseMSGSolve(HYPRE_StructSolver solver, HYPRE_StructMatrix A, HYPRE_StructVector b, HYPRE_StructVector x)

HYPRE_Int HYPRE_StructSparseMSGSetTol (HYPRE_StructSolver solver, HYPRE_Real tol)
HYPRE_Int HYPRE_StructSparseMSGSetMaxIter (HYPRE_StructSolver solver, HYPRE_Int max_iter)
HYPRE_Int HYPRE_StructSparseMSGSetJump (HYPRE_StructSolver solver, HYPRE_Int jump)
HYPRE_Int HYPRE_StructSparseMSGSetRelChange (HYPRE_StructSolver solver, HYPRE_Int rel_change)

HYPRE_Int HYPRE_StructSparseMSGSetZeroGuess (HYPRE_StructSolver solver)
HYPRE_Int HYPRE_StructSparseMSGSetNonZeroGuess(HYPRE_StructSolver solver)
HYPRE_Int HYPRE_StructSparseMSGSetRelaxType (HYPRE_StructSolver solver, HYPRE_Int relax_type)

HYPRE_Int HYPRE_StructSparseMSGSetJacobiWeight (HYPRE_StructSolver solver, HYPRE_Real weight)

HYPRE_Int HYPRE_StructSparseMSGSetNumPreRelax (HYPRE_StructSolver solver, HYPRE_Int num_pre_relax)

HYPRE_Int HYPRE_StructSparseMSGSetNumPostRelax (HYPRE_StructSolver solver, HYPRE_Int num_post_relax)

HYPRE_Int HYPRE_StructSparseMSGSetNumFineRelax (HYPRE_StructSolver solver, HYPRE_Int num_fine_relax)

HYPRE_Int HYPRE_StructSparseMSGSetLogging(HYPRE_StructSolver solver, HYPRE_Int logging)
HYPRE_Int HYPRE_StructSparseMSGSetPrintLevel (HYPRE_StructSolver solver, HYPRE_Int print_level)

HYPRE_Int HYPRE_StructSparseMSGGetNumIterations(HYPRE_StructSolver solver, HYPRE_Int *num_iterations)

HYPRE_Int HYPRE_StructSparseMSGGetFinalRelativeResidualNorm(HYPRE_StructSolver solver, HYPRE_Real *norm)

### 8.5 SStruct Solvers

group SStructSolvers
Linear solvers for semi-structured grids.
These solvers use matrix/vector storage schemes that are taylored to semi-structured grid problems.

## SStruct Solvers

typedef struct hypre_SStructSolver_struct *HYPRE_SStructSolver
The solver object.
typedef HYPRE_Int (*HYPRE_PtrToSStructSolverFcn)(HYPRE_SStructSolver, HYPRE_SStructMatrix, HYPRE_SStructVector, HYPRE_SStructVector)
typedef HYPRE_Int (*HYPRE_PtrToModifyPCFcn)(HYPRE_Solver, HYPRE_Int, HYPRE_Real)

HYPRE_MODIFYPC

## SStruct SysPFMG Solver

SysPFMG is a semicoarsening multigrid solver similar to PFMG, but for systems of PDEs.
For periodic problems, users should try to set the grid size in periodic dimensions to be as close to a power-of-two as possible (for more details, see Struct PFMG Solver).

HYPRE_Int HYPRE_SStructSysPFMGCreate(MPI_Comm comm, HYPRE_SStructSolver *solver)
Create a solver object.
HYPRE_Int HYPRE_SStructSysPFMGDestroy (HYPRE_SStructSolver solver)
Destroy a solver object.
An object should be explicitly destroyed using this destructor when the user's code no longer needs direct access to it. Once destroyed, the object must not be referenced again. Note that the object may not be deallocated at the completion of this call, since there may be internal package references to the object. The object will then be destroyed when all internal reference counts go to zero.

## HYPRE_Int HYPRE_SStructSysPFMGSetup (HYPRE_SStructSolver solver, HYPRE_SStructMatrix A, HYPRE_SStructVector $\mathbf{b}$, HYPRE_SStructVector x)

Prepare to solve the system.
The coefficient data in $b$ and $x$ is ignored here, but information about the layout of the data may be used.

## HYPRE_Int HYPRE_SStructSysPFMGSolve(HYPRE_SStructSolver solver, HYPRE_SStructMatrix A,

 HYPRE_SStructVector b, HYPRE_SStructVector x)Solve the system.
HYPRE_Int HYPRE_SStructSysPFMGSetTol (HYPRE_SStructSolver solver, HYPRE_Real tol)
(Optional) Set the convergence tolerance.
HYPRE_Int HYPRE_SStructSysPFMGSetMaxIter (HYPRE_SStructSolver solver, HYPRE_Int max_iter)
(Optional) Set maximum number of iterations.
HYPRE_Int HYPRE_SStructSysPFMGSetRelChange (HYPRE_SStructSolver solver, HYPRE_Int rel_change)
(Optional) Additionally require that the relative difference in successive iterates be small.
HYPRE_Int HYPRE_SStructSysPFMGSetZeroGuess (HYPRE_SStructSolver solver)
(Optional) Use a zero initial guess.
This allows the solver to cut corners in the case where a zero initial guess is needed (e.g., for preconditioning) to reduce compuational cost.

HYPRE_Int HYPRE_SStructSysPFMGSetNonZeroGuess (HYPRE_SStructSolver solver)
(Optional) Use a nonzero initial guess.
This is the default behavior, but this routine allows the user to switch back after using SetZeroGuess.
HYPRE_Int HYPRE_SStructSysPFMGSetRelaxType (HYPRE_SStructSolver solver, HYPRE_Int relax_type)
(Optional) Set relaxation type.
Current relaxation methods set by relax_type are:

- 0 : Jacobi
- 1 : Weighted Jacobi (default)
- 2 : Red/Black Gauss-Seidel (symmetric: RB pre-relaxation, BR post-relaxation)

HYPRE_Int HYPRE_SStructSysPFMGSetJacobiWeight (HYPRE_SStructSolver solver, HYPRE_Real weight)
(Optional) Set Jacobi Weight.
HYPRE_Int HYPRE_SStructSysPFMGSetNumPreRelax(HYPRE_SStructSolver solver, HYPRE_Int num_pre_relax)
(Optional) Set number of relaxation sweeps before coarse-grid correction.
HYPRE_Int HYPRE_SStructSysPFMGSetNumPostRelax (HYPRE_SStructSolver solver, HYPRE_Int num_post_relax)
(Optional) Set number of relaxation sweeps after coarse-grid correction.
HYPRE_Int HYPRE_SStructSysPFMGSetSkipRelax (HYPRE_SStructSolver solver, HYPRE_Int skip_relax) (Optional) Skip relaxation on certain grids for isotropic problems.

This can greatly improve efficiency by eliminating unnecessary relaxations when the underlying problem is isotropic.

HYPRE_Int HYPRE_SStructSysPFMGSetDxyz (HYPRE_SStructSolver solver, HYPRE_Real *dxyz)
HYPRE_Int HYPRE_SStructSysPFMGSetLogging(HYPRE_SStructSolver solver, HYPRE_Int logging) (Optional) Set the amount of logging to do.
HYPRE_Int HYPRE_SStructSysPFMGSetPrintLevel (HYPRE_SStructSolver solver, HYPRE_Int print_level)
(Optional) Set the amount of printing to do to the screen.
HYPRE_Int HYPRE_SStructSysPFMGGetNumIterations(HYPRE_SStructSolver solver, HYPRE_Int *num_iterations)
Return the number of iterations taken.
HYPRE_Int HYPRE_SStructSysPFMGGetFinalRelativeResidualNorm(HYPRE_SStructSolver solver, HYPRE_Real *norm)
Return the norm of the final relative residual.

## SStruct Split Solver

HYPRE_Int HYPRE_SStructSplitCreate(MPI_Comm comm, HYPRE_SStructSolver *solver)
Create a solver object.
HYPRE_Int HYPRE_SStructSplitDestroy (HYPRE_SStructSolver solver)
Destroy a solver object.
An object should be explicitly destroyed using this destructor when the user's code no longer needs direct access to it. Once destroyed, the object must not be referenced again. Note that the object may not be deallocated at the completion of this call, since there may be internal package references to the object. The object will then be destroyed when all internal reference counts go to zero.

HYPRE_Int HYPRE_SStructSplitSetup (HYPRE_SStructSolver solver, HYPRE_SStructMatrix A, HYPRE_SStructVector $\mathbf{b}$, HYPRE_SStructVector x)
Prepare to solve the system.
The coefficient data in $b$ and $x$ is ignored here, but information about the layout of the data may be used.

HYPRE_Int HYPRE_SStructSplitSolve(HYPRE_SStructSolver solver, HYPRE_SStructMatrix A, HYPRE_SStructVector $\mathbf{b}$, HYPRE_SStructVector x )
Solve the system.
HYPRE_Int HYPRE_SStructSplitSetTol(HYPRE_SStructSolver solver, HYPRE_Real tol)
(Optional) Set the convergence tolerance.
HYPRE_Int HYPRE_SStructSplitSetMaxIter (HYPRE_SStructSolver solver, HYPRE_Int max_iter)
(Optional) Set maximum number of iterations.
HYPRE_Int HYPRE_SStructSplitSetZeroGuess (HYPRE_SStructSolver solver)
(Optional) Use a zero initial guess.
This allows the solver to cut corners in the case where a zero initial guess is needed (e.g., for preconditioning) to reduce compuational cost.
HYPRE_Int HYPRE_SStructSplitSetNonZeroGuess (HYPRE_SStructSolver solver)
(Optional) Use a nonzero initial guess.
This is the default behavior, but this routine allows the user to switch back after using SetZeroGuess.
HYPRE_Int HYPRE_SStructSplitSetStructSolver (HYPRE_SStructSolver solver, HYPRE_Int ssolver)
(Optional) Set up the type of diagonal struct solver.
Either ssolver is set to HYPRE_SMG or HYPRE_PFMG.
HYPRE_Int HYPRE_SStructSplitGetNumIterations(HYPRE_SStructSolver solver, HYPRE_Int *num_iterations)
Return the number of iterations taken.
HYPRE_Int HYPRE_SStructSplitGetFinalRelativeResidualNorm(HYPRE_SStructSolver solver, HYPRE_Real *norm)
Return the norm of the final relative residual.

## HYPRE_PFMG

HYPRE_SMG

HYPRE_Jacobi

## SStruct FAC Solver

HYPRE_Int HYPRE_SStructFACCreate(MPI_Comm comm, HYPRE_SStructSolver *solver)
Create a solver object.
HYPRE_Int HYPRE_SStructFACDestroy2 (HYPRE_SStructSolver solver)
Destroy a solver object.
An object should be explicitly destroyed using this destructor when the user's code no longer needs direct access to it. Once destroyed, the object must not be referenced again. Note that the object may not be deallocated at the completion of this call, since there may be internal package references to the object. The object will then be destroyed when all internal reference counts go to zero.

HYPRE_Int HYPRE_SStructFACAMR_RAP(HYPRE_SStructMatrix A, HYPRE_Int
(*rfactors)[HYPRE_MAXDIM], HYPRE_SStructMatrix *fac_A)
Re-distribute the composite matrix so that the amr hierachy is approximately nested.
Coarse underlying operators are also formed.
HYPRE_Int HYPRE_SStructFACSetup2 (HYPRE_SStructSolver solver, HYPRE_SStructMatrix A, HYPRE_SStructVector b, HYPRE_SStructVector x)
Set up the FAC solver structure .
HYPRE_Int HYPRE_SStructFACSolve3 (HYPRE_SStructSolver solver, HYPRE_SStructMatrix A, HYPRE_SStructVector b, HYPRE_SStructVector x)

Solve the system.
HYPRE_Int HYPRE_SStructFACSetPLevels(HYPRE_SStructSolver solver, HYPRE_Int nparts, HYPRE_Int *plevels)
Set up amr structure.
HYPRE_Int HYPRE_SStructFACSetPRefinements(HYPRE_SStructSolver solver, HYPRE_Int nparts, HYPRE_Int (*rfactors)[HYPRE_MAXDIM])
Set up amr refinement factors.
HYPRE_Int HYPRE_SStructFACZeroCFSten(HYPRE_SStructMatrix A, HYPRE_SStructGrid grid, HYPRE_Int part, HYPRE_Int rfactors[HYPRE_MAXDIM]) (Optional, but user must make sure that they do this function otherwise.) Zero off the coarse level stencils reaching into a fine level grid.

HYPRE_Int HYPRE_SStructFACZeroFCSten(HYPRE_SStructMatrix A, HYPRE_SStructGrid grid, HYPRE_Int part)
(Optional, but user must make sure that they do this function otherwise.) Zero off the fine level stencils reaching into a coarse level grid.

HYPRE_Int HYPRE_SStructFACZeroAMRMatrixData(HYPRE_SStructMatrix A, HYPRE_Int part_crse, HYPRE_Int rfactors[HYPRE_MAXDIM])
(Optional, but user must make sure that they do this function otherwise.) Places the identity in the coarse grid matrix underlying the fine patches.

Required between each pair of amr levels.
HYPRE_Int HYPRE_SStructFACZeroAMRVectorData(HYPRE_SStructVector b, HYPRE_Int *plevels, HYPRE_Int (*rfactors)[HYPRE_MAXDIM]) (Optional, but user must make sure that they do this function otherwise.) Places zeros in the coarse grid vector underlying the fine patches.

Required between each pair of amr levels.
HYPRE_Int HYPRE_SStructFACSetMaxLevels(HYPRE_SStructSolver solver, HYPRE_Int max_levels) (Optional) Set maximum number of FAC levels.
HYPRE_Int HYPRE_SStructFACSetTol (HYPRE_SStructSolver solver, HYPRE_Real tol) (Optional) Set the convergence tolerance.

HYPRE_Int HYPRE_SStructFACSetMaxIter (HYPRE_SStructSolver solver, HYPRE_Int max_iter) (Optional) Set maximum number of iterations.

HYPRE_Int HYPRE_SStructFACSetRelChange(HYPRE_SStructSolver solver, HYPRE_Int rel_change) (Optional) Additionally require that the relative difference in successive iterates be small.

HYPRE_Int HYPRE_SStructFACSetZeroGuess (HYPRE_SStructSolver solver)
(Optional) Use a zero initial guess.
This allows the solver to cut corners in the case where a zero initial guess is needed (e.g., for preconditioning) to reduce compuational cost.

HYPRE_Int HYPRE_SStructFACSetNonZeroGuess(HYPRE_SStructSolver solver)
(Optional) Use a nonzero initial guess.
This is the default behavior, but this routine allows the user to switch back after using SetZeroGuess.
HYPRE_Int HYPRE_SStructFACSetRelaxType (HYPRE_SStructSolver solver, HYPRE_Int relax_type) (Optional) Set relaxation type.

See HYPRE_SStructSysPFMGSetRelaxType for appropriate values of relax_type.
HYPRE_Int HYPRE_SStructFACSetJacobiWeight (HYPRE_SStructSolver solver, HYPRE_Real weight)
(Optional) Set Jacobi weight if weighted Jacobi is used.
HYPRE_Int HYPRE_SStructFACSetNumPreRelax (HYPRE_SStructSolver solver, HYPRE_Int num_pre_relax)
(Optional) Set number of relaxation sweeps before coarse-grid correction.
HYPRE_Int HYPRE_SStructFACSetNumPostRelax(HYPRE_SStructSolver solver, HYPRE_Int num_post_relax)
(Optional) Set number of relaxation sweeps after coarse-grid correction.
HYPRE_Int HYPRE_SStructFACSetCoarseSolverType(HYPRE_SStructSolver solver, HYPRE_Int csolver_type)
(Optional) Set coarsest solver type.
Current solver types set by csolver_type are:

- 1 : SysPFMG-PCG (default)
- 2 : SysPFMG

HYPRE_Int HYPRE_SStructFACSetLogging (HYPRE_SStructSolver solver, HYPRE_Int logging) (Optional) Set the amount of logging to do.

HYPRE_Int HYPRE_SStructFACGetNumIterations(HYPRE_SStructSolver solver, HYPRE_Int *num_iterations)
Return the number of iterations taken.
HYPRE_Int HYPRE_SStructFACGetFinalRelativeResidualNorm (HYPRE_SStructSolver solver, HYPRE_Real *norm)

Return the norm of the final relative residual.

## SStruct Maxwell Solver

HYPRE_Int HYPRE_SStructMaxwellCreate(MPI_Comm comm, HYPRE_SStructSolver *solver)
Create a solver object.
HYPRE_Int HYPRE_SStructMaxwellDestroy (HYPRE_SStructSolver solver)
Destroy a solver object.
An object should be explicitly destroyed using this destructor when the user's code no longer needs direct access to it. Once destroyed, the object must not be referenced again. Note that the object may not be deallocated at the completion of this call, since there may be internal package references to the object. The object will then be destroyed when all internal reference counts go to zero.

HYPRE_Int HYPRE_SStructMaxwellSetup (HYPRE_SStructSolver solver, HYPRE_SStructMatrix A, HYPRE_SStructVector b, HYPRE_SStructVector x)
Prepare to solve the system.
The coefficient data in $b$ and $x$ is ignored here, but information about the layout of the data may be used.
HYPRE_Int HYPRE_SStructMaxwellSolve(HYPRE_SStructSolver solver, HYPRE_SStructMatrix A, HYPRE_SStructVector b, HYPRE_SStructVector x)
Solve the system.
Full coupling of the augmented system used throughout the multigrid hierarchy.
HYPRE_Int HYPRE_SStructMaxwellSolve2 (HYPRE_SStructSolver solver, HYPRE_SStructMatrix A, HYPRE_SStructVector $\mathbf{b}$, HYPRE_SStructVector x)

Solve the system.
Full coupling of the augmented system used only on the finest level, i.e., the node and edge multigrid cycles are coupled only on the finest level.
HYPRE_Int HYPRE_SStructMaxwellSetGrad(HYPRE_SStructSolver solver, HYPRE_ParCSRMatrix T)
Sets the gradient operator in the Maxwell solver.
HYPRE_Int HYPRE_SStructMaxwellSetRfactors(HYPRE_SStructSolver solver, HYPRE_Int rfactors[HYPRE_MAXDIM])

Sets the coarsening factor.
HYPRE_Int HYPRE_SStructMaxwellPhysBdy (HYPRE_SStructGrid *grid_l, HYPRE_Int num_levels, HYPRE_Int rfactors[HYPRE_MAXDIM], HYPRE_Int ***BdryRanks_ptr, HYPRE_Int **BdryRanksCnt_ptr)
Finds the physical boundary row ranks on all levels.
HYPRE_Int HYPRE_SStructMaxwellEliminateRowsCols(HYPRE_ParCSRMatrix parA, HYPRE_Int nrows, HYPRE_Int *rows)
Eliminates the rows and cols corresponding to the physical boundary in a parcsr matrix.
HYPRE_Int HYPRE_SStructMaxwellZeroVector (HYPRE_ParVector b, HYPRE_Int *rows, HYPRE_Int nrows)
Zeros the rows corresponding to the physical boundary in a par vector.
HYPRE_Int HYPRE_SStructMaxwellSetSetConstantCoef (HYPRE_SStructSolver solver, HYPRE_Int flag)
(Optional) Set the constant coefficient flag- Nedelec interpolation used.

HYPRE_Int HYPRE_SStructMaxwellGrad(HYPRE_SStructGrid grid, HYPRE_ParCSRMatrix *T)
(Optional) Creates a gradient matrix from the grid.
This presupposes a particular orientation of the edge elements.
HYPRE_Int HYPRE_SStructMaxwellSetTol(HYPRE_SStructSolver solver, HYPRE_Real tol) (Optional) Set the convergence tolerance.

HYPRE_Int HYPRE_SStructMaxwellSetMaxIter (HYPRE_SStructSolver solver, HYPRE_Int max_iter)
(Optional) Set maximum number of iterations.
HYPRE_Int HYPRE_SStructMaxwellSetRelChange (HYPRE_SStructSolver solver, HYPRE_Int rel_change)
(Optional) Additionally require that the relative difference in successive iterates be small.
HYPRE_Int HYPRE_SStructMaxwellSetNumPreRelax (HYPRE_SStructSolver solver, HYPRE_Int num_pre_relax)
(Optional) Set number of relaxation sweeps before coarse-grid correction.
HYPRE_Int HYPRE_SStructMaxwellSetNumPostRelax (HYPRE_SStructSolver solver, HYPRE_Int num_post_relax)
(Optional) Set number of relaxation sweeps after coarse-grid correction.
HYPRE_Int HYPRE_SStructMaxwellSetLogging (HYPRE_SStructSolver solver, HYPRE_Int logging) (Optional) Set the amount of logging to do.
HYPRE_Int HYPRE_SStructMaxwellGetNumIterations(HYPRE_SStructSolver solver, HYPRE_Int *num_iterations)
Return the number of iterations taken.
HYPRE_Int HYPRE_SStructMaxwellGetFinalRelativeResidualNorm(HYPRE_SStructSolver solver, HYPRE_Real *norm)
Return the norm of the final relative residual.

## SStruct PCG Solver

These routines should be used in conjunction with the generic interface in Krylov Solvers.
HYPRE_Int HYPRE_SStructPCGCreate(MPI_Comm comm, HYPRE_SStructSolver *solver)
Create a solver object.
HYPRE_Int HYPRE_SStructPCGDestroy (HYPRE_SStructSolver solver)
Destroy a solver object.
An object should be explicitly destroyed using this destructor when the user's code no longer needs direct access to it. Once destroyed, the object must not be referenced again. Note that the object may not be deallocated at the completion of this call, since there may be internal package references to the object. The object will then be destroyed when all internal reference counts go to zero.

```
HYPRE_Int HYPRE_SStructPCGSetup (HYPRE_SStructSolver solver, HYPRE_SStructMatrix A, HYPRE_SStructVector b, HYPRE_SStructVector x)
HYPRE_Int HYPRE_SStructPCGSolve(HYPRE_SStructSolver solver, HYPRE_SStructMatrix A, HYPRE_SStructVector b, HYPRE_SStructVector x)
```

HYPRE_Int HYPRE_SStructPCGSetTol (HYPRE_SStructSolver solver, HYPRE_Real tol)

HYPRE_Int HYPRE_SStructPCGSetAbsoluteTol (HYPRE_SStructSolver solver, HYPRE_Real tol)
HYPRE_Int HYPRE_SStructPCGSetMaxIter (HYPRE_SStructSolver solver, HYPRE_Int max_iter)
HYPRE_Int HYPRE_SStructPCGSetTwoNorm (HYPRE_SStructSolver solver, HYPRE_Int two_norm)
HYPRE_Int HYPRE_SStructPCGSetRelChange (HYPRE_SStructSolver solver, HYPRE_Int rel_change)
HYPRE_Int HYPRE_SStructPCGSetPrecond(HYPRE_SStructSolver solver, HYPRE_PtrToSStructSolverFcn precond, HYPRE_PtrToSStructSolverFcn precond_setup, void *precond_solver)

HYPRE_Int HYPRE_SStructPCGSetLogging (HYPRE_SStructSolver solver, HYPRE_Int logging)
HYPRE_Int HYPRE_SStructPCGSetPrintLevel (HYPRE_SStructSolver solver, HYPRE_Int level)
HYPRE_Int HYPRE_SStructPCGGetNumIterations (HYPRE_SStructSolver solver, HYPRE_Int *num_iterations)

HYPRE_Int HYPRE_SStructPCGGetFinalRelativeResidualNorm (HYPRE_SStructSolver solver, HYPRE_Real *norm)

HYPRE_Int HYPRE_SStructPCGGetResidual (HYPRE_SStructSolver solver, void **residual)
HYPRE_Int HYPRE_SStructDiagScaleSetup(HYPRE_SStructSolver solver, HYPRE_SStructMatrix A, HYPRE_SStructVector y, HYPRE_SStructVector x)

Setup routine for diagonal preconditioning.
HYPRE_Int HYPRE_SStructDiagScale(HYPRE_SStructSolver solver, HYPRE_SStructMatrix A, HYPRE_SStructVector y, HYPRE_SStructVector x)
Solve routine for diagonal preconditioning.

## SStruct GMRES Solver

These routines should be used in conjunction with the generic interface in Krylov Solvers.
HYPRE_Int HYPRE_SStructGMRESCreate(MPI_Comm comm, HYPRE_SStructSolver *solver) Create a solver object.
HYPRE_Int HYPRE_SStructGMRESDestroy (HYPRE_SStructSolver solver)
Destroy a solver object.
An object should be explicitly destroyed using this destructor when the user's code no longer needs direct access to it. Once destroyed, the object must not be referenced again. Note that the object may not be deallocated at the completion of this call, since there may be internal package references to the object. The object will then be destroyed when all internal reference counts go to zero.

HYPRE_Int HYPRE_SStructGMRESSetup (HYPRE_SStructSolver solver, HYPRE_SStructMatrix A, HYPRE_SStructVector b, HYPRE_SStructVector x)

HYPRE_Int HYPRE_SStructGMRESSolve(HYPRE_SStructSolver solver, HYPRE_SStructMatrix A, HYPRE_SStructVector $\mathbf{b}$, HYPRE_SStructVector x )

HYPRE_Int HYPRE_SStructGMRESSetTol (HYPRE_SStructSolver solver, HYPRE_Real tol)
HYPRE_Int HYPRE_SStructGMRESSetAbsoluteTol (HYPRE_SStructSolver solver, HYPRE_Real tol)

```
HYPRE_Int HYPRE_SStructGMRESSetMinIter(HYPRE_SStructSolver solver, HYPRE_Int min_iter)
HYPRE_Int HYPRE_SStructGMRESSetMaxIter(HYPRE_SStructSolver solver, HYPRE_Int max_iter)
HYPRE_Int HYPRE_SStructGMRESSetKDim (HYPRE_SStructSolver solver, HYPRE_Int k_dim)
HYPRE_Int HYPRE_SStructGMRESSetStopCrit(HYPRE_SStructSolver solver, HYPRE_Int stop_crit)
HYPRE_Int HYPRE_SStructGMRESSetPrecond(HYPRE_SStructSolver solver,
HYPRE_PtrToSStructSolverFcn precond,
HYPRE_PtrToSStructSolverFcn precond_setup, void
*precond_solver)
HYPRE_Int HYPRE_SStructGMRESSetLogging (HYPRE_SStructSolver solver, HYPRE_Int logging)
HYPRE_Int HYPRE_SStructGMRESSetPrintLevel (HYPRE_SStructSolver solver, HYPRE_Int print_level)
HYPRE_Int HYPRE_SStructGMRESGetNumIterations(HYPRE_SStructSolver solver, HYPRE_Int
*num_iterations)
HYPRE_Int HYPRE_SStructGMRESGetFinalRelativeResidualNorm(HYPRE_SStructSolver solver, HYPRE_Real *norm)
HYPRE_Int HYPRE_SStructGMRESGetResidual (HYPRE_SStructSolver solver, void **residual)
```


## SStruct FlexGMRES Solver

These routines should be used in conjunction with the generic interface in Krylov Solvers.
HYPRE_Int HYPRE_SStructFlexGMRESCreate(MPI_Comm comm, HYPRE_SStructSolver *solver)
Create a solver object.
HYPRE_Int HYPRE_SStructFlexGMRESDestroy (HYPRE_SStructSolver solver)
Destroy a solver object.
An object should be explicitly destroyed using this destructor when the user's code no longer needs direct access to it. Once destroyed, the object must not be referenced again. Note that the object may not be deallocated at the completion of this call, since there may be internal package references to the object. The object will then be destroyed when all internal reference counts go to zero.
HYPRE_Int HYPRE_SStructFlexGMRESSetup (HYPRE_SStructSolver solver, HYPRE_SStructMatrix A, HYPRE_SStructVector $\mathbf{b}$, HYPRE_SStructVector $\mathbf{x}$ )

HYPRE_Int HYPRE_SStructFlexGMRESSolve(HYPRE_SStructSolver solver, HYPRE_SStructMatrix A, HYPRE_SStructVector b, HYPRE_SStructVector x)

HYPRE_Int HYPRE_SStructFlexGMRESSetTol (HYPRE_SStructSolver solver, HYPRE_Real tol)
HYPRE_Int HYPRE_SStructFlexGMRESSetAbsoluteTol (HYPRE_SStructSolver solver, HYPRE_Real tol)
HYPRE_Int HYPRE_SStructFlexGMRESSetMinIter (HYPRE_SStructSolver solver, HYPRE_Int min_iter)
HYPRE_Int HYPRE_SStructFlexGMRESSetMaxIter (HYPRE_SStructSolver solver, HYPRE_Int max_iter)
HYPRE_Int HYPRE_SStructFlexGMRESSetKDim(HYPRE_SStructSolver solver, HYPRE_Int k_dim)
$\begin{aligned} \text { HYPRE_Int HYPRE_SStructFlexGMRESSetPrecond } & \text { HYPRE_SStructSolver solver, } \\ & \text { HYPRE_PtrToSStructSolverFcn precond, } \\ & \text { HYPRE_PtrToSStructSolverFcn precond_setup, void } \\ & \text { *precond_Solver) }\end{aligned}$
HYPRE_Int HYPRE_SStructFlexGMRESSetLogging(HYPRE_SStructSolver solver, HYPRE_Int logging)
HYPRE_Int HYPRE_SStructFlexGMRESSetPrintLevel (HYPRE_SStructSolver solver, HYPRE_Int print_level)

HYPRE_Int HYPRE_SStructFlexGMRESGetNumIterations(HYPRE_SStructSolver solver, HYPRE_Int *num_iterations)

HYPRE_Int HYPRE_SStructFlexGMRESGetFinalRelativeResidualNorm(HYPRE_SStructSolver solver, HYPRE_Real *norm)

HYPRE_Int HYPRE_SStructFlexGMRESGetResidual (HYPRE_SStructSolver solver, void **residual)
HYPRE_Int HYPRE_SStructFlexGMRESSetModifyPC(HYPRE_SStructSolver solver, HYPRE_PtrToModifyPCFcn modify_pc)

## SStruct LGMRES Solver

These routines should be used in conjunction with the generic interface in Krylov Solvers.
HYPRE_Int HYPRE_SStructLGMRESCreate(MPI_Comm comm, HYPRE_SStructSolver *solver)
Create a solver object.
HYPRE_Int HYPRE_SStructLGMRESDestroy (HYPRE_SStructSolver solver)
Destroy a solver object.
An object should be explicitly destroyed using this destructor when the user's code no longer needs direct access to it. Once destroyed, the object must not be referenced again. Note that the object may not be deallocated at the completion of this call, since there may be internal package references to the object. The object will then be destroyed when all internal reference counts go to zero.
HYPRE_Int HYPRE_SStructLGMRESSetup (HYPRE_SStructSolver solver, HYPRE_SStructMatrix A, HYPRE_SStructVector b , HYPRE_SStructVector x )

HYPRE_Int HYPRE_SStructLGMRESSolve(HYPRE_SStructSolver solver, HYPRE_SStructMatrix A, HYPRE_SStructVector b, HYPRE_SStructVector x)

HYPRE_Int HYPRE_SStructLGMRESSetTol (HYPRE_SStructSolver solver, HYPRE_Real tol)
HYPRE_Int HYPRE_SStructLGMRESSetAbsoluteTol (HYPRE_SStructSolver solver, HYPRE_Real tol)
HYPRE_Int HYPRE_SStructLGMRESSetMinIter(HYPRE_SStructSolver solver, HYPRE_Int min_iter)
HYPRE_Int HYPRE_SStructLGMRESSetMaxIter (HYPRE_SStructSolver solver, HYPRE_Int max_iter)
HYPRE_Int HYPRE_SStructLGMRESSetKDim(HYPRE_SStructSolver solver, HYPRE_Int k_dim)
HYPRE_Int HYPRE_SStructLGMRESSetAugDim(HYPRE_SStructSolver solver, HYPRE_Int aug_dim)

HYPRE_Int HYPRE_SStructLGMRESSetPrecond(HYPRE_SStructSolver solver, HYPRE_PtrToSStructSolverFcn precond, HYPRE_PtrToSStructSolverFcn precond_setup, void *precond_solver)

HYPRE_Int HYPRE_SStructLGMRESSetLogging(HYPRE_SStructSolver solver, HYPRE_Int logging)
HYPRE_Int HYPRE_SStructLGMRESSetPrintLevel (HYPRE_SStructSolver solver, HYPRE_Int print_level)

HYPRE_Int HYPRE_SStructLGMRESGetNumIterations(HYPRE_SStructSolver solver, HYPRE_Int *num_iterations)

HYPRE_Int HYPRE_SStructLGMRESGetFinalRelativeResidualNorm(HYPRE_SStructSolver solver, HYPRE_Real *norm)

HYPRE_Int HYPRE_SStructLGMRESGetResidual (HYPRE_SStructSolver solver, void **residual)

## SStruct BiCGSTAB Solver

These routines should be used in conjunction with the generic interface in Krylov Solvers.
HYPRE_Int HYPRE_SStructBiCGSTABCreate(MPI_Comm comm, HYPRE_SStructSolver *solver)
Create a solver object.
HYPRE_Int HYPRE_SStructBiCGSTABDestroy (HYPRE_SStructSolver solver)
Destroy a solver object.
An object should be explicitly destroyed using this destructor when the user's code no longer needs direct access to it. Once destroyed, the object must not be referenced again. Note that the object may not be deallocated at the completion of this call, since there may be internal package references to the object. The object will then be destroyed when all internal reference counts go to zero.

HYPRE_Int HYPRE_SStructBiCGSTABSetup(HYPRE_SStructSolver solver, HYPRE_SStructMatrix A, HYPRE_SStructVector b, HYPRE_SStructVector x)

HYPRE_Int HYPRE_SStructBiCGSTABSolve(HYPRE_SStructSolver solver, HYPRE_SStructMatrix A, HYPRE_SStructVector b , HYPRE_SStructVector x )

HYPRE_Int HYPRE_SStructBiCGSTABSetTol (HYPRE_SStructSolver solver, HYPRE_Real tol)
HYPRE_Int HYPRE_SStructBiCGSTABSetAbsoluteTol (HYPRE_SStructSolver solver, HYPRE_Real tol)
HYPRE_Int HYPRE_SStructBiCGSTABSetMinIter (HYPRE_SStructSolver solver, HYPRE_Int min_iter)
HYPRE_Int HYPRE_SStructBiCGSTABSetMaxIter (HYPRE_SStructSolver solver, HYPRE_Int max_iter)
HYPRE_Int HYPRE_SStructBiCGSTABSetStopCrit (HYPRE_SStructSolver solver, HYPRE_Int stop_crit)
HYPRE_Int HYPRE_SStructBiCGSTABSetPrecond(HYPRE_SStructSolver solver,
HYPRE_PtrToSStructSolverFcn precond, HYPRE_PtrToSStructSolverFcn precond_setup, void *precond_solver)

HYPRE_Int HYPRE_SStructBiCGSTABSetLogging (HYPRE_SStructSolver solver, HYPRE_Int logging)
HYPRE_Int HYPRE_SStructBiCGSTABSetPrintLevel (HYPRE_SStructSolver solver, HYPRE_Int level)

HYPRE_Int HYPRE_SStructBiCGSTABGetNumIterations(HYPRE_SStructSolver solver, HYPRE_Int *num_iterations)

HYPRE_Int HYPRE_SStructBiCGSTABGetFinalRelativeResidualNorm(HYPRE_SStructSolver solver, HYPRE_Real *norm)

HYPRE_Int HYPRE_SStructBiCGSTABGetResidual (HYPRE_SStructSolver solver, void **residual)

## SStruct LOBPCG Eigensolver

These routines should be used in conjunction with the generic interface in Eigensolvers.
HYPRE_Int HYPRE_SStructSetupInterpreter (mv_InterfaceInterpreter *i)
Load interface interpreter.
Vector part loaded with hypre_SStructKrylov functions and multivector part loaded with mv_TempMultiVector functions.

HYPRE_Int HYPRE_SStructSetupMatvec (HYPRE_MatvecFunctions *mv)
Load Matvec interpreter with hypre_SStructKrylov functions.

### 8.6 ParCSR Solvers

## group ParCSRSolvers

Linear solvers for sparse matrix systems.
These solvers use matrix/vector storage schemes that are taylored for general sparse matrix systems.

## ParCSR Solvers

typedef HYPRE_Int (*HYPRE_PtrToParSolverFcn)(HYPRE_Solver, HYPRE_ParCSRMatrix, HYPRE_ParVector, HYPRE_ParVector)

The solver object.
typedef HYPRE_Int (*HYPRE_PtrToModifyPCFcn)(HYPRE_Solver, HYPRE_Int, HYPRE_Real)

HYPRE_MODIFYPC

## ParCSR BoomerAMG Solver and Preconditioner

Parallel unstructured algebraic multigrid solver and preconditioner
HYPRE_Int HYPRE_BoomerAMGCreate(HYPRE_Solver *solver)
Create a solver object.
HYPRE_Int HYPRE_BoomerAMGDestroy (HYPRE_Solver solver)
Destroy a solver object.

HYPRE_Int HYPRE_BoomerAMGSetup(HYPRE_Solver solver, HYPRE_ParCSRMatrix A, HYPRE_ParVector b, HYPRE_ParVector $x$ )
Set up the BoomerAMG solver or preconditioner.
If used as a preconditioner, this function should be passed to the iterative solver SetPrecond function.

## Parameters

- solver - [IN] object to be set up.
- A - [IN] ParCSR matrix used to construct the solver/preconditioner.
- b-Ignored by this function.
- $\mathbf{x}$ - Ignored by this function.

HYPRE_Int HYPRE_BoomerAMGSolve(HYPRE_Solver solver, HYPRE_ParCSRMatrix A, HYPRE_ParVector b, HYPRE_ParVector x )
Solve the system or apply AMG as a preconditioner.
If used as a preconditioner, this function should be passed to the iterative solver SetPrecond function.

## Parameters

- solver - [IN] solver or preconditioner object to be applied.
- A - [IN] ParCSR matrix, matrix of the linear system to be solved
- $\mathbf{b}$ - [IN] right hand side of the linear system to be solved
- $\mathbf{x}$ - [OUT] approximated solution of the linear system to be solved

HYPRE_Int HYPRE_BoomerAMGSolveT(HYPRE_Solver solver, HYPRE_ParCSRMatrix A, HYPRE_ParVector b, HYPRE_ParVector x)
Solve the transpose system $A^{T} x=b$ or apply AMG as a preconditioner to the transpose system .
Note that this function should only be used when preconditioning CGNR with BoomerAMG. It can only be used with Jacobi smoothing (relax_type 0 or 7) and without CF smoothing, i.e relax_order needs to be set to 0 . If used as a preconditioner, this function should be passed to the iterative solver SetPrecond function.

## Parameters

- solver - [IN] solver or preconditioner object to be applied.
- A - [IN] ParCSR matrix
- $\mathbf{b}-[\mathrm{IN}]$ right hand side of the linear system to be solved
- $\mathbf{x}-$ [OUT] approximated solution of the linear system to be solved

HYPRE_Int HYPRE_BoomerAMGSet01dDefault (HYPRE_Solver solver)
Recovers old default for coarsening and interpolation, i.e Falgout coarsening and untruncated modified classical interpolation.

This option might be preferred for 2 dimensional problems.
HYPRE_Int HYPRE_BoomerAMGGetResidual (HYPRE_Solver solver, HYPRE_ParVector *residual)
Returns the residual.
HYPRE_Int HYPRE_BoomerAMGGetNumIterations(HYPRE_Solver solver, HYPRE_Int *num_iterations)
Returns the number of iterations taken.
HYPRE_Int HYPRE_BoomerAMGGetCumNnzAP (HYPRE_Solver solver, HYPRE_Real *cum_nnz_AP)
Returns cumulative num of nonzeros for A and P operators.

HYPRE_Int HYPRE_BoomerAMGSetCumNnzAP(HYPRE_Solver solver, HYPRE_Real cum_nnz_AP)
Activates cumulative num of nonzeros for A and P operators.
Needs to be set to a positive number for activation.
HYPRE_Int HYPRE_BoomerAMGGetFinalRelativeResidualNorm(HYPRE_Solver solver, HYPRE_Real *rel_resid_norm)
Returns the norm of the final relative residual.
HYPRE_Int HYPRE_BoomerAMGSetNumFunctions (HYPRE_Solver solver, HYPRE_Int num_functions)
(Optional) Sets the size of the system of PDEs, if using the systems version.
The default is 1 , i.e. a scalar system.
HYPRE_Int HYPRE_BoomerAMGSetDoffunc (HYPRE_Solver solver, HYPRE_Int *dof_func)
(Optional) Sets the mapping that assigns the function to each variable, if using the systems version.
If no assignment is made and the number of functions is $k>1$, the mapping generated is $(0,1, \ldots, k-$ $1,0,1, \ldots, k-1, \ldots)$.
HYPRE_Int HYPRE_BoomerAMGSetConvergeType (HYPRE_Solver solver, HYPRE_Int type)
(Optional) Set the type convergence checking 0: (default) norm(r)/norm(b), or norm(r) when $\mathrm{b}==01$ : nomr(r) / norm(r_0)
HYPRE_Int HYPRE_BoomerAMGSetTol(HYPRE_Solver solver, HYPRE_Real tol)
(Optional) Set the convergence tolerance, if BoomerAMG is used as a solver.
If it is used as a preconditioner, it should be set to 0 . The default is $1 . e-6$.
HYPRE_Int HYPRE_BoomerAMGSetMaxIter(HYPRE_Solver solver, HYPRE_Int max_iter)
(Optional) Sets maximum number of iterations, if BoomerAMG is used as a solver.
If it is used as a preconditioner, it should be set to 1 . The default is 20 .
HYPRE_Int HYPRE_BoomerAMGSetMinIter(HYPRE_Solver solver, HYPRE_Int min_iter) (Optional)
HYPRE_Int HYPRE_BoomerAMGSetMaxCoarseSize(HYPRE_Solver solver, HYPRE_Int max_coarse_size) (Optional) Sets maximum size of coarsest grid.
The default is 9 .
HYPRE_Int HYPRE_BoomerAMGSetMinCoarseSize(HYPRE_Solver solver, HYPRE_Int min_coarse_size) (Optional) Sets minimum size of coarsest grid.
The default is 1 .
HYPRE_Int HYPRE_BoomerAMGSetMaxLevels(HYPRE_Solver solver, HYPRE_Int max_levels)
(Optional) Sets maximum number of multigrid levels.
The default is 25 .
HYPRE_Int HYPRE_BoomerAMGSetCoarsenCutFactor(HYPRE_Solver solver, HYPRE_Int coarsen_cut_factor)
(Optional) Sets cut factor for choosing isolated points during coarsening according to the rows' density.
The default is 0 . If nnzrow > coarsen_cut_factor*avg_nnzrow, where avg_nnzrow is the average number of nonzeros per row of the global matrix, holds for a given row, it is set as fine, and interpolation weights are not computed.

HYPRE_Int HYPRE_BoomerAMGSetStrongThreshold(HYPRE_Solver solver, HYPRE_Real strong_threshold)
(Optional) Sets AMG strength threshold.
The default is 0.25 . For 2D Laplace operators, 0.25 is a good value, for 3D Laplace operators, 0.5 or 0.6 is a better value. For elasticity problems, a large strength threshold, such as 0.9 , is often better.
HYPRE_Int HYPRE_BoomerAMGSetStrongThresholdR(HYPRE_Solver solver, HYPRE_Real strong_threshold)
(Optional) The strong threshold for R is strong connections used in building an approximate ideal restriction. Default value is 0.25 .

HYPRE_Int HYPRE_BoomerAMGSetFilterThresholdR(HYPRE_Solver solver, HYPRE_Real filter_threshold)
(Optional) The filter threshold for R is used to eliminate small entries of the approximate ideal restriction after building it.
Default value is 0.0 , which disables filtering.
HYPRE_Int HYPRE_BoomerAMGSetSCommPkgSwitch(HYPRE_Solver solver, HYPRE_Real S_commpkg_switch)
(Optional) Deprecated.
This routine now has no effect.
HYPRE_Int HYPRE_BoomerAMGSetMaxRowSum(HYPRE_Solver solver, HYPRE_Real max_row_sum)
(Optional) Sets a parameter to modify the definition of strength for diagonal dominant portions of the matrix.
The default is 0.9 . If max_row_sum is 1 , no checking for diagonally dominant rows is performed.
HYPRE_Int HYPRE_BoomerAMGSetCoarsenType (HYPRE_Solver solver, HYPRE_Int coarsen_type)
(Optional) Defines which parallel coarsening algorithm is used.
There are the following options for coarsen_type:

- 0: CLJP-coarsening (a parallel coarsening algorithm using independent sets.
- 1 : classical Ruge-Stueben coarsening on each processor, no boundary treatment (not recommended!)
- 3 : classical Ruge-Stueben coarsening on each processor, followed by a third pass, which adds coarse points on the boundaries
- 6 : Falgout coarsening (uses 1 first, followed by CLJP using the interior coarse points generated by 1 as its first independent set)
- 7 : CLJP-coarsening (using a fixed random vector, for debugging purposes only)
- 8 : PMIS-coarsening (a parallel coarsening algorithm using independent sets, generating lower complexities than CLJP, might also lead to slower convergence)
- 9 : PMIS-coarsening (using a fixed random vector, for debugging purposes only)
- 10 : HMIS-coarsening (uses one pass Ruge-Stueben on each processor independently, followed by PMIS using the interior C-points generated as its first independent set)
- 11 : one-pass Ruge-Stueben coarsening on each processor, no boundary treatment (not recommended!)
- 21 : CGC coarsening by M. Griebel, B. Metsch and A. Schweitzer
- 22 : CGC-E coarsening by M. Griebel, B. Metsch and A.Schweitzer

The default is 10 .
HYPRE_Int HYPRE_BoomerAMGSetNonGalerkinTol(HYPRE_Solver solver, HYPRE_Real nongalerkin_tol)
(Optional) Defines the non-Galerkin drop-tolerance for sparsifying coarse grid operators and thus reducing communication.

Value specified here is set on all levels. This routine should be used before HYPRE_BoomerAMGSetLevelNonGalerkinTol, which then can be used to change individual levels if desired

HYPRE_Int HYPRE_BoomerAMGSetLevelNonGalerkinTol (HYPRE_Solver solver, HYPRE_Real nongalerkin_tol, HYPRE_Int level)
(Optional) Defines the level specific non-Galerkin drop-tolerances for sparsifying coarse grid operators and thus reducing communication.

A drop-tolerance of 0.0 means to skip doing non-Galerkin on that level. The maximum drop tolerance for a level is 1.0 , although much smaller values such as 0.03 or 0.01 are recommended.

Note that if the user wants to set a specific tolerance on all levels, HYPRE_BooemrAMGSetNonGalerkinTol should be used. Individual levels can then be changed using this routine.
In general, it is safer to drop more aggressively on coarser levels. For instance, one could use 0.0 on the finest level, 0.01 on the second level and then using 0.05 on all remaining levels. The best way to achieve this is to set 0.05 on all levels with HYPRE_BoomerAMGSetNonGalerkinTol and then change the tolerance on level 0 to 0.0 and the tolerance on level 1 to 0.01 with HYPRE_BoomerAMGSetLevelNonGalerkinTol. Like many AMG parameters, these drop tolerances can be tuned. It is also common to delay the start of the non-Galerkin process further to a later level than level 1.

## Parameters

- solver - [IN] solver or preconditioner object to be applied.
- nongalerkin_tol - [IN] level specific drop tolerance
- level - [IN] level on which drop tolerance is used

HYPRE_Int HYPRE_BoomerAMGSetNonGalerkTol (HYPRE_Solver solver, HYPRE_Int nongalerk_num_tol, HYPRE_Real *nongalerk_tol)
(Optional) Defines the non-Galerkin drop-tolerance (old version)
HYPRE_Int HYPRE_BoomerAMGSetMeasureType(HYPRE_Solver solver, HYPRE_Int measure_type)
(Optional) Defines whether local or global measures are used.
HYPRE_Int HYPRE_BoomerAMGSetAggNumLevels(HYPRE_Solver solver, HYPRE_Int agg_num_levels)
(Optional) Defines the number of levels of aggressive coarsening.
The default is 0 , i.e. no aggressive coarsening.
HYPRE_Int HYPRE_BoomerAMGSetNumPaths(HYPRE_Solver solver, HYPRE_Int num_paths)
(Optional) Defines the degree of aggressive coarsening.
The default is 1 . Larger numbers lead to less aggressive coarsening.
HYPRE_Int HYPRE_BoomerAMGSetCGCIts(HYPRE_Solver solver, HYPRE_Int its)
(optional) Defines the number of pathes for CGC-coarsening.
HYPRE_Int HYPRE_BoomerAMGSetNodal(HYPRE_Solver solver, HYPRE_Int nodal)
(Optional) Sets whether to use the nodal systems coarsening.

Should be used for linear systems generated from systems of PDEs. The default is 0 (unknown-based coarsening, only coarsens within same function). For the remaining options a nodal matrix is generated by applying a norm to the nodal blocks and applying the coarsening algorithm to this matrix.

- 1 : Frobenius norm
- 2 : sum of absolute values of elements in each block
- 3 : largest element in each block (not absolute value)
- 4 : row-sum norm
- 6 : sum of all values in each block

HYPRE_Int HYPRE_BoomerAMGSetNodalDiag(HYPRE_Solver solver, HYPRE_Int nodal_diag)
(Optional) Sets whether to give special treatment to diagonal elements in the nodal systems version.
The default is 0 . If set to 1 , the diagonal entry is set to the negative sum of all off diagonal entries. If set to 2 , the signs of all diagonal entries are inverted.

HYPRE_Int HYPRE_BoomerAMGSetKeepSameSign(HYPRE_Solver solver, HYPRE_Int keep_same_sign)
HYPRE_Int HYPRE_BoomerAMGSetInterpType(HYPRE_Solver solver, HYPRE_Int interp_type)
(Optional) Defines which parallel interpolation operator is used.
There are the following options for interp_type:

- 0 : classical modified interpolation
- 1 : LS interpolation (for use with GSMG)
- 2 : classical modified interpolation for hyperbolic PDEs
- 3 : direct interpolation (with separation of weights) (also for GPU use)
- 4 : multipass interpolation
- 5 : multipass interpolation (with separation of weights)
- 6 : extended+i interpolation (also for GPU use)
- 7 : extended+i (if no common C neighbor) interpolation
- 8 : standard interpolation
- 9: standard interpolation (with separation of weights)
- 10 : classical block interpolation (for use with nodal systems version only)
- 11 : classical block interpolation (for use with nodal systems version only) with diagonalized diagonal blocks
- 12 : FF interpolation
- 13 : FF1 interpolation
- 14 : extended interpolation (also for GPU use)
- 15 : interpolation with adaptive weights (GPU use only)
- 16 : extended interpolation in matrix-matrix form
- 17 : extended+i interpolation in matrix-matrix form
- 18 : extended+e interpolation in matrix-matrix form

The default is ext+i interpolation (interp_type 6) trunctated to at most 4 elements per row. (see HYPRE_BoomerAMGSetPMaxElmts).

HYPRE_Int HYPRE_BoomerAMGSetTruncFactor(HYPRE_Solver solver, HYPRE_Real trunc_factor)
(Optional) Defines a truncation factor for the interpolation.
The default is 0 .
HYPRE_Int HYPRE_BoomerAMGSetPMaxElmts (HYPRE_Solver solver, HYPRE_Int P_max_elmts)
(Optional) Defines the maximal number of elements per row for the interpolation.
The default is 4 . To turn off truncation, it needs to be set to 0 .
HYPRE_Int HYPRE_BoomerAMGSetSepWeight (HYPRE_Solver solver, HYPRE_Int sep_weight)
(Optional) Defines whether separation of weights is used when defining strength for standard interpolation or multipass interpolation.

Default: 0, i.e. no separation of weights used.
HYPRE_Int HYPRE_BoomerAMGSetAggInterpType(HYPRE_Solver solver, HYPRE_Int agg_interp_type) (Optional) Defines the interpolation used on levels of aggressive coarsening The default is 4, i.e. multipass interpolation. The following options exist:

- 1: 2-stage extended+i interpolation
- 2: 2-stage standard interpolation
- 3: 2-stage extended interpolation
- 4 : multipass interpolation
- 5: 2-stage extended interpolation in matrix-matrix form
- 6:2-stage extended+i interpolation in matrix-matrix form
- 7: 2-stage extended+e interpolation in matrix-matrix form

HYPRE_Int HYPRE_BoomerAMGSetAggTruncFactor(HYPRE_Solver solver, HYPRE_Real agg_trunc_factor)
(Optional) Defines the truncation factor for the interpolation used for aggressive coarsening. The default is 0 .

HYPRE_Int HYPRE_BoomerAMGSetAggP12TruncFactor (HYPRE_Solver solver, HYPRE_Real agg_P12_trunc_factor)
(Optional) Defines the truncation factor for the matrices P1 and P2 which are used to build 2-stage interpolation.
The default is 0 .
HYPRE_Int HYPRE_BoomerAMGSetAggPMaxElmts (HYPRE_Solver solver, HYPRE_Int agg_P_max_elmts)
(Optional) Defines the maximal number of elements per row for the interpolation used for aggressive coarsening.
The default is 0 .
HYPRE_Int HYPRE_BoomerAMGSetAggP12MaxElmts (HYPRE_Solver solver, HYPRE_Int agg_P12_max_elmts)
(Optional) Defines the maximal number of elements per row for the matrices P1 and P2 which are used to build 2-stage interpolation.

The default is 0 .
HYPRE_Int HYPRE_BoomerAMGSetInterpVectors(HYPRE_Solver solver, HYPRE_Int num_vectors, HYPRE_ParVector *interp_vectors)
(Optional) Allows the user to incorporate additional vectors into the interpolation for systems AMG, e.g. rigid body modes for linear elasticity problems. This can only be used in context with nodal coarsening and still requires the user to choose an interpolation.

HYPRE_Int HYPRE_BoomerAMGSetInterpVecVariant (HYPRE_Solver solver, HYPRE_Int var) (Optional) Defines the interpolation variant used for HYPRE_BoomerAMGSetInterpVectors:

- 1 : GM approach 1
- 2 : GM approach 2 (to be preferred over 1)
- 3 : LN approach

HYPRE_Int HYPRE_BoomerAMGSetInterpVecQMax (HYPRE_Solver solver, HYPRE_Int q_max)
(Optional) Defines the maximal elements per row for Q , the additional columns added to the original interpolation matrix P , to reduce complexity.

The default is no truncation.
HYPRE_Int HYPRE_BoomerAMGSetInterpVecAbsQTrunc (HYPRE_Solver solver, HYPRE_Real q_trunc)
(Optional) Defines a truncation factor for Q , the additional columns added to the original interpolation matrix P, to reduce complexity.

The default is no truncation.
HYPRE_Int HYPRE_BoomerAMGSetGSMG(HYPRE_Solver solver, HYPRE_Int gsmg)
(Optional) Specifies the use of GSMG - geometrically smooth coarsening and interpolation.
Currently any nonzero value for gsmg will lead to the use of GSMG. The default is 0 , i.e. (GSMG is not used)

HYPRE_Int HYPRE_BoomerAMGSetNumSamples (HYPRE_Solver solver, HYPRE_Int num_samples)
(Optional) Defines the number of sample vectors used in GSMG or LS interpolation.
HYPRE_Int HYPRE_BoomerAMGSetCycleType(HYPRE_Solver solver, HYPRE_Int cycle_type)
(Optional) Defines the type of cycle.
For a V-cycle, set cycle_type to 1 , for a W-cycle set cycle_type to 2 . The default is 1 .
HYPRE_Int HYPRE_BoomerAMGSetFCycle(HYPRE_Solver solver, HYPRE_Int fcycle)
(Optional) Specifies the use of Full multigrid cycle.
The default is 0 .
HYPRE_Int HYPRE_BoomerAMGSetAdditive(HYPRE_Solver solver, HYPRE_Int addlvl)
(Optional) Defines use of an additive $\mathrm{V}(1,1)$-cycle using the classical additive method starting at level 'addlvl'.

The multiplicative approach is used on levels $0, \ldots$ 'addlvl +1 '. 'addlvl' needs to be $>-1$ for this to have an effect. Can only be used with weighted Jacobi and 11-Jacobi(default).

Can only be used when AMG is used as a preconditioner !!!

HYPRE_Int HYPRE_BoomerAMGSetMultAdditive(HYPRE_Solver solver, HYPRE_Int addlvl)
(Optional) Defines use of an additive $\mathrm{V}(1,1)$-cycle using the mult-additive method starting at level 'addlvl'.
The multiplicative approach is used on levels $0, \ldots$ 'addlvl +1 '. 'addlvl' needs to be $>-1$ for this to have an effect. Can only be used with weighted Jacobi and 11-Jacobi(default).

Can only be used when AMG is used as a preconditioner !!!
HYPRE_Int HYPRE_BoomerAMGSetSimple(HYPRE_Solver solver, HYPRE_Int addlvl)
(Optional) Defines use of an additive $\mathrm{V}(1,1)$-cycle using the simplified mult-additive method starting at level 'addlvl'.

The multiplicative approach is used on levels $0, \ldots$ 'addlvl +1 '. 'addlvl' needs to be $>-1$ for this to have an effect. Can only be used with weighted Jacobi and 11-Jacobi(default).

Can only be used when AMG is used as a preconditioner !!!
HYPRE_Int HYPRE_BoomerAMGSetAddLastLvl (HYPRE_Solver solver, HYPRE_Int add_last_lvl)
(Optional) Defines last level where additive, mult-additive or simple cycle is used.
The multiplicative approach is used on levels > add_last_lvl.
Can only be used when AMG is used as a preconditioner !!!
HYPRE_Int HYPRE_BoomerAMGSetMultAddTruncFactor (HYPRE_Solver solver, HYPRE_Real add_trunc_factor)
(Optional) Defines the truncation factor for the smoothed interpolation used for mult-additive or simple method.

The default is 0 .
HYPRE_Int HYPRE_BoomerAMGSetMultAddPMaxElmts(HYPRE_Solver solver, HYPRE_Int add_P_max_elmts)
(Optional) Defines the maximal number of elements per row for the smoothed interpolation used for multadditive or simple method.

The default is 0 .
HYPRE_Int HYPRE_BoomerAMGSetAddRelaxType(HYPRE_Solver solver, HYPRE_Int add_rlx_type)
(Optional) Defines the relaxation type used in the (mult)additive cycle portion (also affects simple method.) The default is 18 (L1-Jacobi).

Currently the only other option allowed is 0 (Jacobi) which should be used in combination with HYPRE_BoomerAMGSetAddRelaxWt.

HYPRE_Int HYPRE_BoomerAMGSetAddRelaxWt (HYPRE_Solver solver, HYPRE_Real add_rlx_wt) (Optional) Defines the relaxation weight used for Jacobi within the (mult)additive or simple cycle portion. The default is 1 . The weight only affects the Jacobi method, and has no effect on L1-Jacobi

HYPRE_Int HYPRE_BoomerAMGSetSeqThreshold(HYPRE_Solver solver, HYPRE_Int seq_threshold) (Optional) Sets maximal size for agglomeration or redundant coarse grid solve.

When the system is smaller than this threshold, sequential AMG is used on process 0 or on all remaining active processes (if redundant =1).

HYPRE_Int HYPRE_BoomerAMGSetRedundant (HYPRE_Solver solver, HYPRE_Int redundant)
(Optional) operates switch for redundancy.
Needs to be used with HYPRE_BoomerAMGSetSeqThreshold. Default is 0, i.e. no redundancy.

HYPRE_Int HYPRE_BoomerAMGSetNumGridSweeps(HYPRE_Solver solver, HYPRE_Int
*num_grid_sweeps)
(Optional) Defines the number of sweeps for the fine and coarse grid, the up and down cycle.
Note: This routine will be phased out!!!! Use HYPRE_BoomerAMGSetNumSweeps or HYPRE_BoomerAMGSetCycleNumSweeps instead.

HYPRE_Int HYPRE_BoomerAMGSetNumSweeps (HYPRE_Solver solver, HYPRE_Int num_sweeps)
(Optional) Sets the number of sweeps.
On the finest level, the up and the down cycle the number of sweeps are set to num_sweeps and on the coarsest level to 1 . The default is 1 .

HYPRE_Int HYPRE_BoomerAMGSetCycleNumSweeps(HYPRE_Solver solver, HYPRE_Int num_sweeps, HYPRE_Int k)
(Optional) Sets the number of sweeps at a specified cycle.
There are the following options for $k$ :

- 1 : the down cycle
- 2 : the up cycle
- 3 : the coarsest level

HYPRE_Int HYPRE_BoomerAMGSetGridRelaxType(HYPRE_Solver solver, HYPRE_Int *grid_relax_type)
(Optional) Defines which smoother is used on the fine and coarse grid, the up and down cycle.
Note: This routine will be phased out!!!! Use HYPRE_BoomerAMGSetRelaxType or HYPRE_BoomerAMGSetCycleRelaxType instead.

HYPRE_Int HYPRE_BoomerAMGSetRelaxType(HYPRE_Solver solver, HYPRE_Int relax_type)
(Optional) Defines the smoother to be used.
It uses the given smoother on the fine grid, the up and the down cycle and sets the solver on the coarsest level to Gaussian elimination (9). The default is $\ell_{1}$-Gauss-Seidel, forward solve (13) on the down cycle and backward solve (14) on the up cycle.

There are the following options for relax_type:

- 0 : Jacobi
- 1: Gauss-Seidel, sequential (very slow!)
- 2 : Gauss-Seidel, interior points in parallel, boundary sequential (slow!)
- 3 : hybrid Gauss-Seidel or SOR, forward solve
- 4 : hybrid Gauss-Seidel or SOR, backward solve
- 5 : hybrid chaotic Gauss-Seidel (works only with OpenMP)
- 6 : hybrid symmetric Gauss-Seidel or SSOR
- 7 : Jacobi (uses Matvec)
- $8: \ell_{1}$-scaled hybrid symmetric Gauss-Seidel
- 9 : Gaussian elimination (only on coarsest level)
- 10 : On-processor direct forward solve for matrices with triangular structure
- 11: Two Stage approximation to GS. Uses the strict lower part of the diagonal matrix
- 12 : Two Stage approximation to GS. Uses the strict lower part of the diagonal matrix and a second iteration for additional error approximation
- $13: \ell_{1}$ Gauss-Seidel, forward solve
- 14 : $\ell_{1}$ Gauss-Seidel, backward solve
- 15 : CG (warning - not a fixed smoother - may require FGMRES)
- 16 : Chebyshev
- 17 : FCF-Jacobi
- $18: \ell_{1}$-scaled jacobi
- 19 : Gaussian elimination (old version)
- 21 : The same as 8 except forcing serialization on CPU (\#OMP-thread $=1$ )
- 29 : Direct solve: use Gaussian elimination \& BLAS (with pivoting) (old version)
- 30 : Kaczmarz
- 88: The same methods as 8 with a convergent 11 -term
- 89: Symmetric 11-hybrid Gauss-Seidel (i.e., 13 followed by 14)
- 98 : LU with pivoting
- 99 : LU with pivoting -199: Matvec with the inverse

HYPRE_Int HYPRE_BoomerAMGSetCycleRelaxType(HYPRE_Solver solver, HYPRE_Int relax_type, HYPRE_Int k)
(Optional) Defines the smoother at a given cycle.
For options of relax_type see description of HYPRE_BoomerAMGSetRelaxType. In addition, the following options for relax_type are available when choosing the coarsest level solver $(\mathrm{k}=3)$ :

For coarsest level systems formed via a sub-communicator defined with active ranks:

- 9 : hypre's internal Gaussian elimination (host only).
- 99 : LU factorization with pivoting.
- 199 : explicit (dense) inverse.

For coarsest level systems formed via hypre_DataExchangeList:

- 19 : hypre's internal Gaussian elimination (host only).
- 98 : LU factorization with pivoting.
- 198 : explicit (dense) inverse.

Options for $k$ are

- 1 : the down cycle
- 2 : the up cycle
- 3 : the coarsest level

HYPRE_Int HYPRE_BoomerAMGSetRelaxOrder (HYPRE_Solver solver, HYPRE_Int relax_order)
(Optional) Defines in which order the points are relaxed.
There are the following options for relax_order:

- 0 : the points are relaxed in natural or lexicographic order on each processor
- 1: CF-relaxation is used, i.e on the fine grid and the down cycle the coarse points are relaxed first, followed by the fine points; on the up cycle the F-points are relaxed first, followed by the C-points. On the coarsest level, if an iterative scheme is used, the points are relaxed in lexicographic order.
The default is 0 .
HYPRE_Int HYPRE_BoomerAMGSetGridRelaxPoints(HYPRE_Solver solver, HYPRE_Int **grid_relax_points)
(Optional) Defines in which order the points are relaxed.
See also HYPRE_BoomerAMGSetRelaxOrder.
HYPRE_Int HYPRE_BoomerAMGSetRelaxWeight (HYPRE_Solver solver, HYPRE_Real *relax_weight)
(Optional) Defines the relaxation weight for smoothed Jacobi and hybrid SOR.
Note: This routine will be phased out!!!! Use HYPRE_BoomerAMGSetRelaxWt or HYPRE_BoomerAMGSetLevelRelaxWt instead.
HYPRE_Int HYPRE_BoomerAMGSetRelaxWt (HYPRE_Solver solver, HYPRE_Real relax_weight)
(Optional) Defines the relaxation weight for smoothed Jacobi and hybrid SOR on all levels.
Values for relax_weight are
- >0 : this assigns the given relaxation weight on all levels
- = 0 : the weight is determined on each level with the estimate $\frac{3}{4\left\|D^{-1 / 2} A D^{-1 / 2}\right\|}$, where $D$ is the diagonal of $A$ (this should only be used with Jacobi)
- = -k : the relaxation weight is determined with at most k CG steps on each level (this should only be used for symmetric positive definite problems)
The default is 1 .
HYPRE_Int HYPRE_BoomerAMGSetLevelRelaxWt (HYPRE_Solver solver, HYPRE_Real relax_weight, HYPRE_Int level)
(Optional) Defines the relaxation weight for smoothed Jacobi and hybrid SOR on the user defined level.
Note that the finest level is denoted 0 , the next coarser level 1 , etc. For nonpositive relax_weight, the parameter is determined on the given level as described for HYPRE_BoomerAMGSetRelaxWt. The default is 1 .

HYPRE_Int HYPRE_BoomerAMGSetOmega(HYPRE_Solver solver, HYPRE_Real *omega)
(Optional) Defines the outer relaxation weight for hybrid SOR.
Note: This routine will be phased out!!!! Use HYPRE_BoomerAMGSetOuterWt or HYPRE_BoomerAMGSetLevelOuterWt instead.

HYPRE_Int HYPRE_BoomerAMGSetOuterWt (HYPRE_Solver solver, HYPRE_Real omega)
(Optional) Defines the outer relaxation weight for hybrid SOR and SSOR on all levels.
Values for omega are

- > 0 : this assigns the same outer relaxation weight omega on each level
- = -k : an outer relaxation weight is determined with at most k CG steps on each level (this only makes sense for symmetric positive definite problems and smoothers such as SSOR)
The default is 1 .
HYPRE_Int HYPRE_BoomerAMGSetLevelOuterWt (HYPRE_Solver solver, HYPRE_Real omega, HYPRE_Int level)
(Optional) Defines the outer relaxation weight for hybrid SOR or SSOR on the user defined level.
Note that the finest level is denoted 0 , the next coarser level 1, etc. For nonpositive omega, the parameter is determined on the given level as described for HYPRE_BoomerAMGSetOuterWt. The default is 1.
HYPRE_Int HYPRE_BoomerAMGSetChebyOrder (HYPRE_Solver solver, HYPRE_Int order)
(Optional) Defines the Order for Chebyshev smoother.
The default is 2 (valid options are 1-4).
HYPRE_Int HYPRE_BoomerAMGSetChebyFraction(HYPRE_Solver solver, HYPRE_Real ratio)
(Optional) Fraction of the spectrum to use for the Chebyshev smoother.
The default is .3 (i.e., damp on upper $30 \%$ of the spectrum).
HYPRE_Int HYPRE_BoomerAMGSetChebyScale(HYPRE_Solver solver, HYPRE_Int scale)
(Optional) Defines whether matrix should be scaled.
The default is 1 (i.e., scaled).
HYPRE_Int HYPRE_BoomerAMGSetChebyVariant (HYPRE_Solver solver, HYPRE_Int variant)
(Optional) Defines which polynomial variant should be used.
The default is 0 (i.e., scaled).
HYPRE_Int HYPRE_BoomerAMGSetChebyEigEst (HYPRE_Solver solver, HYPRE_Int eig_est)
(Optional) Defines how to estimate eigenvalues.
The default is 10 (i.e., 10 CG iterations are used to find extreme eigenvalues.) If eig_est=0, the largest eigenvalue is estimated using Gershgorin, the smallest is set to 0 . If eig_est is a positive number n , n iterations of CG are used to determine the smallest and largest eigenvalue.

HYPRE_Int HYPRE_BoomerAMGSetSmoothType(HYPRE_Solver solver, HYPRE_Int smooth_type)
(Optional) Enables the use of more complex smoothers.
The following options exist for smooth_type:

- 4 : FSAI (routines needed to set: HYPRE_BoomerAMGSetFSAIMaxSteps, HYPRE_BoomerAMGSetFSAIMaxStepSize, HYPRE_BoomerAMGSetFSAIEigMaxIters, HYPRE_BoomerAMGSetFSAIKapTolerance)
- 5 : ParILUK (routines needed to set: HYPRE_ILUSetLevelOfFill, HYPRE_ILUSetType)
- 6 : Schwarz (routines needed to set: HYPRE_BoomerAMGSetDomainType, HYPRE_BoomerAMGSetOverlap, HYPRE_BoomerAMGSetVariant, HYPRE_BoomerAMGSetSchwarzR1xWeight)
- 7 : Pilut (routines needed to set: HYPRE_BoomerAMGSetDropTol, HYPRE_BoomerAMGSetMaxNzPerRow)
- 8 : ParaSails (routines needed to set: HYPRE_BoomerAMGSetSym, HYPRE_BoomerAMGSetLevel, HYPRE_BoomerAMGSetFilter, HYPRE_BoomerAMGSetThreshold)
- 9 : Euclid (routines needed to set: HYPRE_BoomerAMGSetEuclidFile)

The default is 6 . Also, if no smoother parameters are set via the routines mentioned in the table above, default values are used.

HYPRE_Int HYPRE_BoomerAMGSetSmoothNumLevels(HYPRE_Solver solver, HYPRE_Int smooth_num_levels)
(Optional) Sets the number of levels for more complex smoothers.
The smoothers, as defined by HYPRE_BoomerAMGSetSmoothType, will be used on level 0 (the finest level) through level smooth_num_levels-1. The default is 0 , i.e. no complex smoothers are used.

HYPRE_Int HYPRE_BoomerAMGSetSmoothNumSweeps(HYPRE_Solver solver, HYPRE_Int smooth_num_sweeps)
(Optional) Sets the number of sweeps for more complex smoothers.
The default is 1 .
HYPRE_Int HYPRE_BoomerAMGSetVariant (HYPRE_Solver solver, HYPRE_Int variant)
(Optional) Defines which variant of the Schwarz method is used.
The following options exist for variant:

- 0 : hybrid multiplicative Schwarz method (no overlap across processor boundaries)
- 1 : hybrid additive Schwarz method (no overlap across processor boundaries)
- 2 : additive Schwarz method
- 3 : hybrid multiplicative Schwarz method (with overlap across processor boundaries)

The default is 0 .
HYPRE_Int HYPRE_BoomerAMGSetOverlap(HYPRE_Solver solver, HYPRE_Int overlap)
(Optional) Defines the overlap for the Schwarz method.
The following options exist for overlap:

- 0 : no overlap
- 1 : minimal overlap (default)
- 2 : overlap generated by including all neighbors of domain boundaries

HYPRE_Int HYPRE_BoomerAMGSetDomainType(HYPRE_Solver solver, HYPRE_Int domain_type)
(Optional) Defines the type of domain used for the Schwarz method.
The following options exist for domain_type:

- 0 : each point is a domain
- 1 : each node is a domain (only of interest in "systems" AMG)
- 2 : each domain is generated by agglomeration (default)

HYPRE_Int HYPRE_BoomerAMGSetSchwarzRlxWeight (HYPRE_Solver solver, HYPRE_Real schwarz_rlx_weight)
(Optional) Defines a smoothing parameter for the additive Schwarz method.

HYPRE_Int HYPRE_BoomerAMGSetSchwarzUseNonSymm(HYPRE_Solver solver, HYPRE_Int use_nonsymm)
(Optional) Indicates that the aggregates may not be SPD for the Schwarz method.
The following options exist for use_nonsymm:

- 0 : assume SPD (default)
- 1: assume non-symmetric

HYPRE_Int HYPRE_BoomerAMGSetSym(HYPRE_Solver solver, HYPRE_Int sym)
(Optional) Defines symmetry for ParaSAILS.
For further explanation see description of ParaSAILS.
HYPRE_Int HYPRE_BoomerAMGSetLevel (HYPRE_Solver solver, HYPRE_Int level)
(Optional) Defines number of levels for ParaSAILS.
For further explanation see description of ParaSAILS.
HYPRE_Int HYPRE_BoomerAMGSetThreshold(HYPRE_Solver solver, HYPRE_Real threshold)
(Optional) Defines threshold for ParaSAILS.
For further explanation see description of ParaSAILS.
HYPRE_Int HYPRE_BoomerAMGSetFilter (HYPRE_Solver solver, HYPRE_Real filter)
(Optional) Defines filter for ParaSAILS.
For further explanation see description of ParaSAILS.
HYPRE_Int HYPRE_BoomerAMGSetDropTol (HYPRE_Solver solver, HYPRE_Real drop_tol)
(Optional) Defines drop tolerance for PILUT.
For further explanation see description of PILUT.
HYPRE_Int HYPRE_BoomerAMGSetMaxNzPerRow(HYPRE_Solver solver, HYPRE_Int max_nz_per_row)
(Optional) Defines maximal number of nonzeros for PILUT.
For further explanation see description of PILUT.
HYPRE_Int HYPRE_BoomerAMGSetEuclidFile(HYPRE_Solver solver, char *euclidfile)
(Optional) Defines name of an input file for Euclid parameters.
For further explanation see description of Euclid.
HYPRE_Int HYPRE_BoomerAMGSetEuLevel (HYPRE_Solver solver, HYPRE_Int eu_level)
(Optional) Defines number of levels for $\operatorname{ILU}(\mathrm{k})$ in Euclid.
For further explanation see description of Euclid.
HYPRE_Int HYPRE_BoomerAMGSetEuSparseA(HYPRE_Solver solver, HYPRE_Real eu_sparse_A)
(Optional) Defines filter for ILU(k) for Euclid.
For further explanation see description of Euclid.
HYPRE_Int HYPRE_BoomerAMGSetEuBJ (HYPRE_Solver solver, HYPRE_Int eu_bj)
(Optional) Defines use of block jacobi ILUT for Euclid.
For further explanation see description of Euclid.

HYPRE_Int HYPRE_BoomerAMGSetILUType(HYPRE_Solver solver, HYPRE_Int ilu_type) Defines type of ILU smoother to use For further explanation see description of ILU.

HYPRE_Int HYPRE_BoomerAMGSetILULevel (HYPRE_Solver solver, HYPRE_Int ilu_lfil) Defines level $k$ for ILU(k) smoother For further explanation see description of ILU.

HYPRE_Int HYPRE_BoomerAMGSetILUMaxRowNnz (HYPRE_Solver solver, HYPRE_Int ilu_max_row_nnz) Defines max row nonzeros for ILUT smoother For further explanation see description of ILU.

HYPRE_Int HYPRE_BoomerAMGSetILUMaxIter(HYPRE_Solver solver, HYPRE_Int ilu_max_iter) Defines number of iterations for ILU smoother on each level For further explanation see description of ILU.

HYPRE_Int HYPRE_BoomerAMGSetILUDroptol (HYPRE_Solver solver, HYPRE_Real ilu_droptol) Defines drop tolorance for iLUT smoother For further explanation see description of ILU.

HYPRE_Int HYPRE_BoomerAMGSetILUTriSolve(HYPRE_Solver solver, HYPRE_Int ilu_tri_solve) (Optional) Defines triangular solver for ILU(k,T) smoother: 0-iterative, 1-direct (default) For further explanation see description of ILU.

HYPRE_Int HYPRE_BoomerAMGSetILULowerJacobiIters(HYPRE_Solver solver, HYPRE_Int ilu_lower_jacobi_iters)
(Optional) Defines number of lower Jacobi iterations for ILU(k,T) smoother triangular solve. For further explanation see description of ILU.
HYPRE_Int HYPRE_BoomerAMGSetILUUpperJacobiIters(HYPRE_Solver solver, HYPRE_Int ilu_upper_jacobi_iters)
(Optional) Defines number of upper Jacobi iterations for ILU(k,T) smoother triangular solve. For further explanation see description of ILU.

HYPRE_Int HYPRE_BoomerAMGSetILULocalReordering(HYPRE_Solver solver, HYPRE_Int ilu_reordering_type)
(Optional) Set Local Reordering paramter ( $1==\mathrm{RCM}, 0==$ None) For further explanation see description of ILU.

HYPRE_Int HYPRE_BoomerAMGSetILUIterSetupType(HYPRE_Solver solver, HYPRE_Int ilu_iter_setup_type)
(Optional) Set iterative ILU's algorithm type.
For further explanation see HYPRE_ILUSetIterativeSetupType.
HYPRE_Int HYPRE_BoomerAMGSetILUIterSetupOption(HYPRE_Solver solver, HYPRE_Int ilu_iter_setup_option)
(Optional) Set iterative ILU's option.
For further explanation see HYPRE_ILUSetIterativeSetupOption.
HYPRE_Int HYPRE_BoomerAMGSetILUIterSetupMaxIter(HYPRE_Solver solver, HYPRE_Int ilu_iter_setup_max_iter) (Optional) Set iterative ILU's max. number of iterations. For further explanation see HYPRE_ILUSetIterativeSetupMaxIter.
HYPRE_Int HYPRE_BoomerAMGSetILUIterSetupTolerance(HYPRE_Solver solver, HYPRE_Real ilu_iter_setup_tolerance)
(Optional) Set iterative ILU's tolerance.
For further explanation see HYPRE_ILUSetIterativeSetupTolerance.

HYPRE_Int HYPRE_BoomerAMGSetFSAIAlgoType(HYPRE_Solver solver, HYPRE_Int algo_type) (Optional) Defines the algorithm type for setting up FSAI For further explanation see HYPRE_FSAISetAlgoType.

HYPRE_Int HYPRE_BoomerAMGSetFSAILocalSolveType(HYPRE_Solver solver, HYPRE_Int local_solve_type)
(Optional) Sets the solver type for solving local linear systems in FSAI.
For further explanation see HYPRE_FSAISetLocalSolveType.
HYPRE_Int HYPRE_BoomerAMGSetFSAIMaxSteps (HYPRE_Solver solver, HYPRE_Int max_steps) (Optional) Defines maximum number of steps for FSAI.
For further explanation see HYPRE_FSAISetMaxSteps.
HYPRE_Int HYPRE_BoomerAMGSetFSAIMaxStepSize(HYPRE_Solver solver, HYPRE_Int max_step_size)
(Optional) Defines maximum step size for FSAI.
For further explanation see HYPRE_FSAISetMaxStepSize.
HYPRE_Int HYPRE_BoomerAMGSetFSAIMaxNnzRow (HYPRE_Solver solver, HYPRE_Int max_nnz_row)
(Optional) Defines maximum number of nonzero entries per row for FSAI.
For further explanation see HYPRE_FSAISetMaxNnzRow.
HYPRE_Int HYPRE_BoomerAMGSetFSAINumLevels(HYPRE_Solver solver, HYPRE_Int num_levels)
(Optional) Defines number of levels for computing the candidate pattern for FSAI For further explanation see HYPRE_FSAISetNumLevels.

HYPRE_Int HYPRE_BoomerAMGSetFSAIThreshold(HYPRE_Solver solver, HYPRE_Real threshold) (Optional) Defines the threshold for computing the candidate pattern for FSAI For further explanation see HYPRE_FSAISetThreshold.

HYPRE_Int HYPRE_BoomerAMGSetFSAIEigMaxIters(HYPRE_Solver solver, HYPRE_Int eig_max_iters) (Optional) Defines maximum number of iterations for estimating the largest eigenvalue of the FSAI preconditioned matrix $\left(\mathrm{G}^{\wedge} \mathrm{T} * \mathrm{G} * \mathrm{~A}\right)$.

For further explanation see HYPRE_FSAISetEigMaxIters.
HYPRE_Int HYPRE_BoomerAMGSetFSAIKapTolerance(HYPRE_Solver solver, HYPRE_Real kap_tolerance)
(Optional) Defines the kaporin dropping tolerance.
For further explanation see HYPRE_FSAISetKapTolerance.
HYPRE_Int HYPRE_BoomerAMGSetRestriction(HYPRE_Solver solver, HYPRE_Int restr_par)
(Optional) Defines which parallel restriction operator is used.
There are the following options for restr_type:

- $0: P^{T}$ - Transpose of the interpolation operator
- 1 : AIR-1 - Approximate Ideal Restriction (distance 1)
- 2 : AIR-2 - Approximate Ideal Restriction (distance 2)

The default is 0 .

HYPRE_Int HYPRE_BoomerAMGSetIsTriangular(HYPRE_Solver solver, HYPRE_Int is_triangular)
(Optional) Assumes the matrix is triangular in some ordering to speed up the setup time of approximate ideal restriction.

The default is 0 .
HYPRE_Int HYPRE_BoomerAMGSetGMRESSwitchR (HYPRE_Solver solver, HYPRE_Int gmres_switch) (Optional) Set local problem size at which GMRES is used over a direct solve in approximating ideal restriction.

The default is 0 .
HYPRE_Int HYPRE_BoomerAMGSetADropTol (HYPRE_Solver solver, HYPRE_Real A_drop_tol) (Optional) Defines the drop tolerance for the A-matrices from the 2nd level of AMG. The default is 0 .

HYPRE_Int HYPRE_BoomerAMGSetADropType(HYPRE_Solver solver, HYPRE_Int A_drop_type)
(Optional) Drop the entries that are not on the diagonal and smaller than its row norm: type 1: 1-norm, 2:
2-norm, -1: infinity norm
HYPRE_Int HYPRE_BoomerAMGSetPrintFileName(HYPRE_Solver solver, const char *print_file_name)
(Optional) Name of file to which BoomerAMG will print; cf HYPRE_BoomerAMGSetPrintLevel.
(Presently this is ignored).
HYPRE_Int HYPRE_BoomerAMGSetPrintLevel (HYPRE_Solver solver, HYPRE_Int print_level)
(Optional) Requests automatic printing of setup and solve information.

- 0 : no printout (default)
- 1 : print setup information
- 2 : print solve information
- 3 : print both setup and solve information

Note, that if one desires to print information and uses BoomerAMG as a preconditioner, suggested print_level is 1 to avoid excessive output, and use print_level of solver for solve phase information.

HYPRE_Int HYPRE_BoomerAMGSetLogging (HYPRE_Solver solver, HYPRE_Int logging)
(Optional) Requests additional computations for diagnostic and similar data to be logged by the user.
Default to 0 to do nothing. The latest residual will be available if logging $>1$.
HYPRE_Int HYPRE_BoomerAMGSetDebugFlag(HYPRE_Solver solver, HYPRE_Int debug_flag) (Optional)

HYPRE_Int HYPRE_BoomerAMGInitGridRelaxation(HYPRE_Int **num_grid_sweeps_ptr, HYPRE_Int **grid_relax_type_ptr, HYPRE_Int ***grid_relax_points_ptr, HYPRE_Int coarsen_type, HYPRE_Real **relax_weights_ptr, HYPRE_Int max_levels)
(Optional) This routine will be eliminated in the future.
HYPRE_Int HYPRE_BoomerAMGSetRAP2 (HYPRE_Solver solver, HYPRE_Int rap2)
(Optional) If rap2 not equal 0 , the triple matrix product RAP is replaced by two matrix products.
(Required for triple matrix product generation on GPUs)

HYPRE_Int HYPRE_BoomerAMGSetModuleRAP2 (HYPRE_Solver solver, HYPRE_Int mod_rap2)
(Optional) If mod_rap2 not equal 0, the triple matrix product RAP is replaced by two matrix products with modularized kernels (Required for triple matrix product generation on GPUs)

HYPRE_Int HYPRE_BoomerAMGSetKeepTranspose(HYPRE_Solver solver, HYPRE_Int keepTranspose)
(Optional) If set to 1 , the local interpolation transposes will be saved to use more efficient matvecs instead of matvecTs (Recommended for efficient use on GPUs)

HYPRE_Int HYPRE_BoomerAMGSetPlotGrids(HYPRE_Solver solver, HYPRE_Int plotgrids) HYPRE_BoomerAMGSetPlotGrids.
HYPRE_Int HYPRE_BoomerAMGSetPlotFileName(HYPRE_Solver solver, const char *plotfilename)
HYPRE_BoomerAMGSetPlotFilename.
HYPRE_Int HYPRE_BoomerAMGSetCoordDim(HYPRE_Solver solver, HYPRE_Int coorddim)
HYPRE_BoomerAMGSetCoordDim.
HYPRE_Int HYPRE_BoomerAMGSetCoordinates (HYPRE_Solver solver, float * coordinates)
HYPRE_BoomerAMGSetCoordinates.
HYPRE_Int HYPRE_BoomerAMGGetGridHierarchy (HYPRE_Solver solver, HYPRE_Int *cgrid)
(Optional) Get the coarse grid hierarchy.
Assumes input/ output array is preallocated to the size of the local matrix. On return, cgrid[i] returns the last grid level containing node $i$.

## Parameters

- solver - [IN] solver or preconditioner
- cgrid - [IN/ OUT] preallocated array. On return, contains grid hierarchy info.

HYPRE_Int HYPRE_BoomerAMGSetCPoints(HYPRE_Solver solver, HYPRE_Int cpt_coarse_level, HYPRE_Int num_cpt_coarse, HYPRE_BigInt *cpt_coarse_index)
(Optional) Fix C points to be kept till a specified coarse level.

## Parameters

- solver - [IN] solver or preconditioner
- cpt_coarse_level - [IN] coarse level up to which to keep C points
- num_cpt_coarse - [IN] number of $C$ points to be kept
- cpt_coarse_index - [IN] indexes of C points to be kept

HYPRE_Int HYPRE_BoomerAMGSetCpointsToKeep(HYPRE_Solver solver, HYPRE_Int cpt_coarse_level, HYPRE_Int num_cpt_coarse, HYPRE_BigInt *cpt_coarse_index)
(Optional) Deprecated function.
Use HYPRE_BoomerAMGSetCPoints instead.
HYPRE_Int HYPRE_BoomerAMGSetFPoints(HYPRE_Solver solver, HYPRE_Int num_fpt, HYPRE_BigInt *fpt_index)
(Optional) Set fine points in the first level.
Parameters

- solver - [IN] solver or preconditioner
- num_fpt - [IN] number of fine points
- fpt_index - [IN] global indices of fine points

HYPRE_Int HYPRE_BoomerAMGSetIsolatedFPoints(HYPRE_Solver solver, HYPRE_Int num_isolated_fpt, HYPRE_BigInt *isolated_fpt_index)
(Optional) Set isolated fine points in the first level.
Interpolation weights are not computed for these points.

## Parameters

- solver - [IN] solver or preconditioner
- num_isolated_fpt - [IN] number of isolated fine points
- isolated_fpt_index - [IN] global indices of isolated fine points

HYPRE_Int HYPRE_BoomerAMGSetSabs (HYPRE_Solver solver, HYPRE_Int Sabs)
(Optional) if Sabs equals 1, the strength of connection test is based on the absolute value of the matrix coefficients

## ParCSR BoomerAMGDD Solver and Preconditioner

Communication reducing solver and preconditioner built on top of algebraic multigrid
HYPRE_Int HYPRE_BoomerAMGDDCreate(HYPRE_Solver *solver)
Create a solver object.
HYPRE_Int HYPRE_BoomerAMGDDDestroy (HYPRE_Solver solver)
Destroy a solver object.
HYPRE_Int HYPRE_BoomerAMGDDSetup(HYPRE_Solver solver, HYPRE_ParCSRMatrix A, HYPRE_ParVector b, HYPRE_ParVector x)
Set up the BoomerAMGDD solver or preconditioner.
If used as a preconditioner, this function should be passed to the iterative solver SetPrecond function.

## Parameters

- solver - [IN] object to be set up.
- A $-[$ IN $]$ ParCSR matrix used to construct the solver/preconditioner.
- $\mathbf{b}$ - Ignored by this function.
- $\mathbf{x}$ - Ignored by this function.

HYPRE_Int HYPRE_BoomerAMGDDSolve(HYPRE_Solver solver, HYPRE_ParCSRMatrix A, HYPRE_ParVector b, HYPRE_ParVector x)
Solve the system or apply AMG-DD as a preconditioner.
If used as a preconditioner, this function should be passed to the iterative solver SetPrecond function.

## Parameters

- solver - [IN] solver or preconditioner object to be applied.
- A - [IN] ParCSR matrix, matrix of the linear system to be solved
- $\mathbf{b}-[\mathrm{IN}]$ right hand side of the linear system to be solved
- $\mathbf{x}-$ [OUT] approximated solution of the linear system to be solved

HYPRE_Int HYPRE_BoomerAMGDDSetFACNumRelax (HYPRE_Solver solver, HYPRE_Int amgdd_fac_num_relax)
(Optional) Set the number of pre- and post-relaxations per level for AMG-DD inner FAC cycles.
Default is 1 .
HYPRE_Int HYPRE_BoomerAMGDDSetFACNumCycles(HYPRE_Solver solver, HYPRE_Int amgdd_fac_num_cycles)
(Optional) Set the number of inner FAC cycles per AMG-DD iteration.
Default is 2 .
HYPRE_Int HYPRE_BoomerAMGDDSetFACCycleType(HYPRE_Solver solver, HYPRE_Int amgdd_fac_cycle_type)
(Optional) Set the cycle type for the AMG-DD inner FAC cycles.
1 (default) $=$ V-cycle, $2=\mathrm{W}$-cycle, $3=$ F-cycle
HYPRE_Int HYPRE_BoomerAMGDDSetFACRelaxType (HYPRE_Solver solver, HYPRE_Int amgdd_fac_relax_type)
(Optional) Set the relaxation type for the AMG-DD inner FAC cycles.
0 = Jacobi, 1 = Gauss-Seidel, 2 = ordered Gauss-Seidel, 3 (default) = C/F L1-scaled Jacobi
HYPRE_Int HYPRE_BoomerAMGDDSetFACRelaxWeight(HYPRE_Solver solver, HYPRE_Real amgdd_fac_relax_weight)
(Optional) Set the relaxation weight for the AMG-DD inner FAC cycles.
Default is 1.0 .
HYPRE_Int HYPRE_BoomerAMGDDSetStartLevel (HYPRE_Solver solver, HYPRE_Int start_level) (Optional) Set the AMG-DD start level.

Default is 0 .
HYPRE_Int HYPRE_BoomerAMGDDSetPadding(HYPRE_Solver solver, HYPRE_Int padding) (Optional) Set the AMG-DD padding.

Default is 1 .
HYPRE_Int HYPRE_BoomerAMGDDSetNumGhostLayers(HYPRE_Solver solver, HYPRE_Int num_ghost_layers)
(Optional) Set the AMG-DD number of ghost layers.
Default is 1 .
HYPRE_Int HYPRE_BoomerAMGDDSetUserFACRelaxation(HYPRE_Solver solver, HYPRE_Int (*userFACRelaxation)(void *amgdd_vdata, HYPRE_Int level, HYPRE_Int cycle_param))
(Optional) Pass a custom user-defined function as a relaxation method for the AMG-DD FAC cycles.
Function should have the following form, where amgdd_solver is of type hypre_ParAMGDDData* and level is the level on which to relax: HYPRE_Int userFACRelaxation(HYPRE_Solver amgdd_solver, HYPRE_Int level)

HYPRE_Int HYPRE_BoomerAMGDDGetAMG(HYPRE_Solver solver, HYPRE_Solver *amg_solver)
(Optional) Get the underlying AMG hierarchy as a HYPRE_Solver object.

HYPRE_Int HYPRE_BoomerAMGDDGetFinalRelativeResidualNorm(HYPRE_Solver solver, HYPRE_Real *rel_resid_norm)
Returns the norm of the final relative residual.
HYPRE_Int HYPRE_BoomerAMGDDGetNumIterations(HYPRE_Solver solver, HYPRE_Int
*num_iterations)
Returns the number of iterations taken.

## ParCSR FSAI Solver and Preconditioner

An adaptive factorized sparse approximate inverse solver/preconditioner/smoother that computes a sparse approximation $G$ to the inverse of the lower cholesky factor of $A$ such that $M^{\wedge}\{-1\}$ ไapprox $G^{\wedge} T * G$.
HYPRE_Int HYPRE_FSAICreate(HYPRE_Solver *solver)
Create a solver object.
HYPRE_Int HYPRE_FSAIDestroy (HYPRE_Solver solver)
Destroy a solver object.
HYPRE_Int HYPRE_FSAISetup(HYPRE_Solver solver, HYPRE_ParCSRMatrix A, HYPRE_ParVector b, HYPRE_ParVector x )
Set up the FSAI solver or preconditioner.
If used as a preconditioner, this function should be passed to the iterative solver SetPrecond function.

## Parameters

- solver - [IN] object to be set up.
- A - [IN] ParCSR matrix used to construct the solver/preconditioner.
- $\mathbf{b}$ - Ignored by this function.
- $\mathbf{x}$ - Ignored by this function.

HYPRE_Int HYPRE_FSAISolve(HYPRE_Solver solver, HYPRE_ParCSRMatrix A, HYPRE_ParVector b, HYPRE_ParVector x)

Solve the system or apply FSAI as a preconditioner.
If used as a preconditioner, this function should be passed to the iterative solver SetPrecond function.

## Parameters

- solver - [IN] solver or preconditioner object to be applied.
- A - [IN] ParCSR matrix, matrix of the linear system to be solved
- $\mathbf{b}-[\mathrm{IN}]$ right hand side of the linear system to be solved
- $\mathbf{x}$ - [OUT] approximated solution of the linear system to be solved

HYPRE_Int HYPRE_FSAISetAlgoType (HYPRE_Solver solver, HYPRE_Int algo_type)
(Optional) Sets the algorithm type used to compute the lower triangular factor G

```
- 1: Adaptive (can use OpenMP with static scheduling)
- 2: Adaptive OpenMP with dynamic scheduling
- 3: Static - power pattern
```

HYPRE_Int HYPRE_FSAISetLocalSolveType (HYPRE_Solver solver, HYPRE_Int local_solve_type)
(Optional) Sets the solver type for solving local linear systems in FSAI.
This option makes sense only for GPU runs.

```
- Q: Gauss-Jordan solver
- 1: Vendor solver (cuSOLVER/rocSOLVER)
- 2: MAGMA solver
```

HYPRE_Int HYPRE_FSAISetMaxSteps (HYPRE_Solver solver, HYPRE_Int max_steps)
(Optional) Sets the maximum number of steps for computing the sparsity pattern of $G$. This input parameter makes sense when using adaptive FSAI, i.e., algorithm type 1 or 2.
HYPRE_Int HYPRE_FSAISetMaxStepSize(HYPRE_Solver solver, HYPRE_Int max_step_size)
(Optional) Sets the maximum step size for computing the sparsity pattern of $G$.
This input parameter makes sense when using adaptive FSAI, i.e., algorithm type 1 or 2.
HYPRE_Int HYPRE_FSAISetMaxNnzRow (HYPRE_Solver solver, HYPRE_Int max_nnz_row)
(Optional) Sets the maximum number of off-diagonal entries per row of $G$.
This input parameter makes sense when using static FSAI, i.e., algorithm type 3.
HYPRE_Int HYPRE_FSAISetNumLevels (HYPRE_Solver solver, HYPRE_Int num_levels)
(Optional) Sets the number of levels for computing the candidate pattern of $G$.
This input parameter must be a positive integer and it makes sense when using static FSAI, i.e., algorithm type 3.

HYPRE_Int HYPRE_FSAISetThreshold(HYPRE_Solver solver, HYPRE_Real threshold)
(Optional) Sets the threshold for computing the candidate pattern of G This input parameter makes sense when using static FSAI, i.e., algorithm type 3.

HYPRE_Int HYPRE_FSAISetKapTolerance(HYPRE_Solver solver, HYPRE_Real kap_tolerance)
(Optional) Sets the kaporin gradient reduction factor for computing the sparsity pattern of G.
This input parameter makes sense when using adaptive FSAI, i.e., algorithm types 1 or 2.
HYPRE_Int HYPRE_FSAISetOmega(HYPRE_Solver solver, HYPRE_Real omega)
(Optional) Sets the relaxation factor for FSAI.
This input parameter makes sense to all algorithm types for setting up FSAI.
HYPRE_Int HYPRE_FSAISetMaxIterations(HYPRE_Solver solver, HYPRE_Int max_iterations)
(Optional) Sets the maximum number of iterations (sweeps) for FSAI.
This input parameter makes sense to all algorithm types for setting up FSAI.
HYPRE_Int HYPRE_FSAISetEigMaxIters(HYPRE_Solver solver, HYPRE_Int eig_max_iters)
(Optional) Set number of iterations for computing maximum eigenvalue of the preconditioned operator.
This input parameter makes sense to all algorithm types for setting up FSAI.
HYPRE_Int HYPRE_FSAISetTolerance(HYPRE_Solver solver, HYPRE_Real tolerance)
(Optional) Set the convergence tolerance, if FSAI is used as a solver.
This input parameter makes sense to all algorithm types for setting up FSAI. When using FSAI as a preconditioner, set the tolerance to 0.0 . The default is $10^{-6}$.

HYPRE_Int HYPRE_FSAISetPrintLevel (HYPRE_Solver solver, HYPRE_Int print_level)
(Optional) Requests automatic printing of setup information.

- 0 : no printout (default)
- 1 : print setup information

HYPRE_Int HYPRE_FSAISetZeroGuess(HYPRE_Solver solver, HYPRE_Int zero_guess)
(Optional) Use a zero initial guess.
This allows the solver to cut corners in the case where a zero initial guess is needed (e.g., for preconditioning) to reduce compuational cost.

## ParCSR ParaSails Preconditioner

Parallel sparse approximate inverse preconditioner for the ParCSR matrix format.
HYPRE_Int HYPRE_ParaSailsCreate(MPI_Comm comm, HYPRE_Solver *solver)
Create a ParaSails preconditioner.
HYPRE_Int HYPRE_ParaSailsDestroy (HYPRE_Solver solver)
Destroy a ParaSails preconditioner.
HYPRE_Int HYPRE_ParaSailsSetup(HYPRE_Solver solver, HYPRE_ParCSRMatrix A, HYPRE_ParVector b, HYPRE_ParVector x)

Set up the ParaSails preconditioner.
This function should be passed to the iterative solver SetPrecond function.

## Parameters

- solver - [IN] Preconditioner object to set up.
- A - [IN] ParCSR matrix used to construct the preconditioner.
- $\mathbf{b}$ - Ignored by this function.
- $\mathbf{x}$ - Ignored by this function.

HYPRE_Int HYPRE_ParaSailsSolve(HYPRE_Solver solver, HYPRE_ParCSRMatrix A, HYPRE_ParVector b, HYPRE_ParVector x )
Apply the ParaSails preconditioner.
This function should be passed to the iterative solver SetPrecond function.

## Parameters

- solver - [IN] Preconditioner object to apply.
- A - Ignored by this function.
- $\mathbf{b}-[\mathrm{IN}]$ Vector to precondition.
- $\mathbf{x}$ - [OUT] Preconditioned vector.

HYPRE_Int HYPRE_ParaSailsSetParams(HYPRE_Solver solver, HYPRE_Real thresh, HYPRE_Int nlevels)

Set the threshold and levels parameter for the ParaSails preconditioner.

The accuracy and cost of ParaSails are parameterized by these two parameters. Lower values of the threshold parameter and higher values of levels parameter lead to more accurate, but more expensive preconditioners.

## Parameters

- solver - [IN] Preconditioner object for which to set parameters.
- thresh $-[\mathrm{IN}]$ Value of threshold parameter, $0 \leq$ thresh $\leq 1$. The default value is 0.1 .
- nlevels - [IN] Value of levels parameter, $0 \leq$ nlevels. The default value is 1.


## HYPRE_Int HYPRE_ParaSailsSetFilter(HYPRE_Solver solver, HYPRE_Real filter)

Set the filter parameter for the ParaSails preconditioner.

## Parameters

- solver - [IN] Preconditioner object for which to set filter parameter.
- filter - [IN] Value of filter parameter. The filter parameter is used to drop small nonzeros in the preconditioner, to reduce the cost of applying the preconditioner. Values from 0.05 to 0.1 are recommended. The default value is 0.1 .

HYPRE_Int HYPRE_ParaSailsSetSym(HYPRE_Solver solver, HYPRE_Int sym)
Set the symmetry parameter for the ParaSails preconditioner.
Values for sym

- 0 : nonsymmetric and/or indefinite problem, and nonsymmetric preconditioner
- 1 : SPD problem, and SPD (factored) preconditioner
- 2 : nonsymmetric, definite problem, and SPD (factored) preconditioner


## Parameters

- solver - [IN] Preconditioner object for which to set symmetry parameter.
- sym - [IN] Symmetry parameter.


## HYPRE_Int HYPRE_ParaSailsSetLoadbal (HYPRE_Solver solver, HYPRE_Real loadbal)

Set the load balance parameter for the ParaSails preconditioner.

## Parameters

- solver - [IN] Preconditioner object for which to set the load balance parameter.
- loadbal - [IN] Value of the load balance parameter, $0 \leq$ loadbal $\leq 1$. A zero value indicates that no load balance is attempted; a value of unity indicates that perfect load balance will be attempted. The recommended value is 0.9 to balance the overhead of data exchanges for load balancing. No load balancing is needed if the preconditioner is very sparse and fast to construct. The default value when this parameter is not set is 0 .
HYPRE_Int HYPRE_ParaSailsSetReuse(HYPRE_Solver solver, HYPRE_Int reuse)
Set the pattern reuse parameter for the ParaSails preconditioner.


## Parameters

- solver - [IN] Preconditioner object for which to set the pattern reuse parameter.
- reuse - [IN] Value of the pattern reuse parameter. A nonzero value indicates that the pattern of the preconditioner should be reused for subsequent constructions of the preconditioner. A zero value indicates that the preconditioner should be constructed from scratch. The default value when this parameter is not set is 0 .

HYPRE_Int HYPRE_ParaSailsSetLogging(HYPRE_Solver solver, HYPRE_Int logging)
Set the logging parameter for the ParaSails preconditioner.

## Parameters

- solver - [IN] Preconditioner object for which to set the logging parameter.
- logging - [IN] Value of the logging parameter. A nonzero value sends statistics of the setup procedure to stdout. The default value when this parameter is not set is 0 .
HYPRE_Int HYPRE_ParaSailsBuildIJMatrix (HYPRE_Solver solver, HYPRE_IJMatrix *pij_A)
Build IJ Matrix of the sparse approximate inverse (factor).
This function explicitly creates the IJ Matrix corresponding to the sparse approximate inverse or the inverse factor. Example: HYPRE_IJMatrix ij_A; HYPRE_ParaSailsBuildIJMatrix(solver, \&ij_A);


## Parameters

- solver - [IN] Preconditioner object.
- pij_A - [OUT] Pointer to the IJ Matrix.

HYPRE_Int HYPRE_ParCSRParaSailsCreate(MPI_Comm comm, HYPRE_Solver *solver)
HYPRE_Int HYPRE_ParCSRParaSailsDestroy (HYPRE_Solver solver)
HYPRE_Int HYPRE_ParCSRParaSailsSetup(HYPRE_Solver solver, HYPRE_ParCSRMatrix A, HYPRE_ParVector b, HYPRE_ParVector x)

HYPRE_Int HYPRE_ParCSRParaSailsSolve(HYPRE_Solver solver, HYPRE_ParCSRMatrix A, HYPRE_ParVector b, HYPRE_ParVector x)

HYPRE_Int HYPRE_ParCSRParaSailsSetParams(HYPRE_Solver solver, HYPRE_Real thresh, HYPRE_Int nlevels)

HYPRE_Int HYPRE_ParCSRParaSailsSetFilter(HYPRE_Solver solver, HYPRE_Real filter)
HYPRE_Int HYPRE_ParCSRParaSailsSetSym(HYPRE_Solver solver, HYPRE_Int sym)
HYPRE_Int HYPRE_ParCSRParaSailsSetLoadbal (HYPRE_Solver solver, HYPRE_Real loadbal)
HYPRE_Int HYPRE_ParCSRParaSailsSetReuse(HYPRE_Solver solver, HYPRE_Int reuse)
HYPRE_Int HYPRE_ParCSRParaSailsSetLogging(HYPRE_Solver solver, HYPRE_Int logging)

## ParCSR Euclid Preconditioner

MPI Parallel ILU preconditioner
Options summary:

| Option | Default | Synopsis |
| :--- | :--- | :--- |
| -level | 1 | ILU(k) factorization level |
| -bj | 0 (false) | Use Block Jacobi ILU instead of PILU |
| -eu_stats | 0 (false) | Print internal timing and statistics |
| -eu_mem | 0 (false) | Print internal memory usage |

HYPRE_Int HYPRE_EuclidCreate(MPI_Comm comm, HYPRE_Solver *solver)
Create a Euclid object.
HYPRE_Int HYPRE_EuclidDestroy (HYPRE_Solver solver)
Destroy a Euclid object.
HYPRE_Int HYPRE_EuclidSetup(HYPRE_Solver solver, HYPRE_ParCSRMatrix A, HYPRE_ParVector b, HYPRE_ParVector x)

Set up the Euclid preconditioner.
This function should be passed to the iterative solver SetPrecond function.

## Parameters

- solver - [IN] Preconditioner object to set up.
- A - [IN] ParCSR matrix used to construct the preconditioner.
- $\mathbf{b}$ - Ignored by this function.
- $\mathbf{x}$ - Ignored by this function.

HYPRE_Int HYPRE_EuclidSolve(HYPRE_Solver solver, HYPRE_ParCSRMatrix A, HYPRE_ParVector b, HYPRE_ParVector x)
Apply the Euclid preconditioner.
This function should be passed to the iterative solver SetPrecond function.

## Parameters

- solver - [IN] Preconditioner object to apply.
- A - Ignored by this function.
- $\mathbf{b}-[\mathrm{IN}]$ Vector to precondition.
- $\mathbf{x}$ - [OUT] Preconditioned vector.

HYPRE_Int HYPRE_EuclidSetParams (HYPRE_Solver solver, HYPRE_Int argc, char *argv[])
Insert (name, value) pairs in Euclid's options database by passing Euclid the command line (or an array of strings).

All Euclid options (e.g, level, drop-tolerance) are stored in this database. If a (name, value) pair already exists, this call updates the value. See also: HYPRE_EuclidSetParamsFromFile.

## Parameters

- argc - [IN] Length of argv array
- argv - [IN] Array of strings

HYPRE_Int HYPRE_EuclidSetParamsFromFile(HYPRE_Solver solver, char *filename)
Insert (name, value) pairs in Euclid's options database.
Each line of the file should either begin with a " \#", indicating a comment line, or contain a (name value) pair, e.g:

```
>cat optionsFile
\#sample runtime parameter file
-blockJacobi 3
-matFile /home/hysom/myfile.euclid
-doSomething true
-xx_coeff -1.0
```

See also: HYPRE_EuclidSetParams.

## Parameters

filename[IN] - Pathname/filename to read
HYPRE_Int HYPRE_EuclidSetLevel(HYPRE_Solver solver, HYPRE_Int level)
Set level k for ILU(k) factorization, default: 1.
HYPRE_Int HYPRE_EuclidSetBJ (HYPRE_Solver solver, HYPRE_Int bj)
Use block Jacobi ILU preconditioning instead of PILU.
HYPRE_Int HYPRE_EuclidSetStats(HYPRE_Solver solver, HYPRE_Int eu_stats)
If eu_stats not equal 0 , a summary of runtime settings and timing information is printed to stdout.
HYPRE_Int HYPRE_EuclidSetMem(HYPRE_Solver solver, HYPRE_Int eu_mem)
If eu_mem not equal 0 , a summary of Euclid's memory usage is printed to stdout.
HYPRE_Int HYPRE_EuclidSetSparseA(HYPRE_Solver solver, HYPRE_Real sparse_A)
Defines a drop tolerance for ILU(k).
Default: 0 Use with HYPRE_EuclidSetRowScale. Note that this can destroy symmetry in a matrix.
HYPRE_Int HYPRE_EuclidSetRowScale(HYPRE_Solver solver, HYPRE_Int row_scale)
If row_scale not equal 0 , values are scaled prior to factorization so that largest value in any row is +1 or -1 .
Note that this can destroy symmetry in a matrix.
HYPRE_Int HYPRE_EuclidSetILUT (HYPRE_Solver solver, HYPRE_Real drop_tol)
uses ILUT and defines a drop tolerance relative to the largest absolute value of any entry in the row being factored.

## ParCSR Pilut Preconditioner

HYPRE_Int HYPRE_ParCSRPilutCreate(MPI_Comm comm, HYPRE_Solver *solver)
Create a preconditioner object.
HYPRE_Int HYPRE_ParCSRPilutDestroy (HYPRE_Solver solver)
Destroy a preconditioner object.
HYPRE_Int HYPRE_ParCSRPilutSetup(HYPRE_Solver solver, HYPRE_ParCSRMatrix A, HYPRE_ParVector b, HYPRE_ParVector x)

HYPRE_Int HYPRE_ParCSRPilutSolve(HYPRE_Solver solver, HYPRE_ParCSRMatrix A, HYPRE_ParVector b, HYPRE_ParVector x)
Precondition the system.
HYPRE_Int HYPRE_ParCSRPilutSetMaxIter (HYPRE_Solver solver, HYPRE_Int max_iter)
(Optional) Set maximum number of iterations.
HYPRE_Int HYPRE_ParCSRPilutSetDropTolerance(HYPRE_Solver solver, HYPRE_Real tol) (Optional)

HYPRE_Int HYPRE_ParCSRPilutSetFactorRowSize(HYPRE_Solver solver, HYPRE_Int size) (Optional)

HYPRE_Int HYPRE_ParCSRPilutSetLogging(HYPRE_Solver solver, HYPRE_Int logging)

## ParCSR AMS Solver and Preconditioner

Parallel auxiliary space Maxwell solver and preconditioner
HYPRE_Int HYPRE_AMSCreate(HYPRE_Solver *solver)
Create an AMS solver object.
HYPRE_Int HYPRE_AMSDestroy (HYPRE_Solver solver)
Destroy an AMS solver object.
HYPRE_Int HYPRE_AMSSetup(HYPRE_Solver solver, HYPRE_ParCSRMatrix A, HYPRE_ParVector b, HYPRE_ParVector x)

Set up the AMS solver or preconditioner.
If used as a preconditioner, this function should be passed to the iterative solver SetPrecond function.

## Parameters

- solver - [IN] object to be set up.
- A - [IN] ParCSR matrix used to construct the solver/preconditioner.
- $\mathbf{b}$ - Ignored by this function.
- $\mathbf{x}$ - Ignored by this function.

HYPRE_Int HYPRE_AMSSolve(HYPRE_Solver solver, HYPRE_ParCSRMatrix A, HYPRE_ParVector b, HYPRE_ParVector x)

Solve the system or apply AMS as a preconditioner.
If used as a preconditioner, this function should be passed to the iterative solver SetPrecond function.

## Parameters

- solver - [IN] solver or preconditioner object to be applied.
- A - [IN] ParCSR matrix, matrix of the linear system to be solved
- $\mathbf{b}-[\mathrm{IN}]$ right hand side of the linear system to be solved
- $\mathbf{x}-$ [OUT] approximated solution of the linear system to be solved

HYPRE_Int HYPRE_AMSSetDimension(HYPRE_Solver solver, HYPRE_Int dim)
(Optional) Sets the problem dimension (2 or 3).
The default is 3 .
HYPRE_Int HYPRE_AMSSetDiscreteGradient (HYPRE_Solver solver, HYPRE_ParCSRMatrix G)
Sets the discrete gradient matrix $G$.
This function should be called before HYPRE_AMSSetup()!
HYPRE_Int HYPRE_AMSSetCoordinateVectors(HYPRE_Solver solver, HYPRE_ParVector x, HYPRE_ParVector y, HYPRE_ParVector z)
Sets the $x, y$ and $z$ coordinates of the vertices in the mesh.
Either HYPRE_AMSSetCoordinateVectors() or HYPRE_AMSSetEdgeConstantVectors() should be called before HYPRE_AMSSetup()!

HYPRE_Int HYPRE_AMSSetEdgeConstantVectors(HYPRE_Solver solver, HYPRE_ParVector Gx,
HYPRE_ParVector Gy, HYPRE_ParVector Gz)

Sets the vectors $G x, G y$ and $G z$ which give the representations of the constant vector fields $(1,0,0),(0,1,0)$ and $(0,0,1)$ in the edge element basis.

Either HYPRE_AMSSetCoordinateVectors() or HYPRE_AMSSetEdgeConstantVectors() should be called before HYPRE_AMSSetup()!
HYPRE_Int HYPRE_AMSSetInterpolations(HYPRE_Solver solver, HYPRE_ParCSRMatrix Pi, HYPRE_ParCSRMatrix Pix, HYPRE_ParCSRMatrix Piy, HYPRE_ParCSRMatrix Piz)
(Optional) Set the (components of) the Nedelec interpolation matrix $\Pi=\left[\Pi^{x}, \Pi^{y}, \Pi^{z}\right]$.
This function is generally intended to be used only for high-order Nedelec discretizations (in the lowest order case, $\Pi$ is constructed internally in AMS from the discreet gradient matrix and the coordinates of the vertices), though it can also be used in the lowest-order case or for other types of discretizations (e.g. ones based on the second family of Nedelec elements).

By definition, $\Pi$ is the matrix representation of the linear operator that interpolates (high-order) vector nodal finite elements into the (high-order) Nedelec space. The component matrices are defined as $\Pi^{x} \varphi=$ $\Pi(\varphi, 0,0)$ and similarly for $\Pi^{y}$ and $\Pi^{z}$. Note that all these operators depend on the choice of the basis and degrees of freedom in the high-order spaces.

The column numbering of Pi should be node-based, i.e. the $x / y / z$ components of the first node (vertex or high-order dof) should be listed first, followed by the $x / y / z$ components of the second node and so on (see the documentation of HYPRE_BoomerAMGSetDofFunc).

If used, this function should be called before $\operatorname{HYPRE} A M S S e t u p()$ and there is no need to provide the vertex coordinates. Furthermore, only one of the sets $\{\Pi\}$ and $\left\{\Pi^{x}, \Pi^{y}, \Pi^{z}\right\}$ needs to be specified (though it is OK to provide both). If Pix is NULL, then scalar ח-based AMS cycles, i.e. those with cycle_type $>10$, will be unavailable. Similarly, AMS cycles based on monolithic $\Pi$ (cycle_type $<10$ ) require that Pi is not NULL.

HYPRE_Int HYPRE_AMSSetAlphaPoissonMatrix(HYPRE_Solver solver, HYPRE_ParCSRMatrix A_alpha)
(Optional) Sets the matrix $A_{\alpha}$ corresponding to the Poisson problem with coefficient $\alpha$ (the curl-curl term coefficient in the Maxwell problem).
If this function is called, the coarse space solver on the range of $\Pi^{T}$ is a block-diagonal version of $A_{\Pi}$. If this function is not called, the coarse space solver on the range of $\Pi^{T}$ is constructed as $\Pi^{T} A \Pi$ in HYPRE_AMSSetup(). See the user's manual for more details.

HYPRE_Int HYPRE_AMSSetBetaPoissonMatrix (HYPRE_Solver solver, HYPRE_ParCSRMatrix A_beta)
(Optional) Sets the matrix $A_{\beta}$ corresponding to the Poisson problem with coefficient $\beta$ (the mass term coefficient in the Maxwell problem).

If not given, the Poisson matrix will be computed in HYPRE_AMSSetup(). If the given matrix is NULL, we assume that $\beta$ is identically 0 and use two-level (instead of three-level) methods. See the user's manual for more details.

HYPRE_Int HYPRE_AMSSetInteriorNodes(HYPRE_Solver solver, HYPRE_ParVector interior_nodes)
(Optional) Set the list of nodes which are interior to a zero-conductivity region.
This way, a more robust solver is constructed, that can be iterated to lower tolerance levels. A node is interior if its entry in the array is 1.0. This function should be called before HYPRE_AMSSetup()!
HYPRE_Int HYPRE_AMSSetProjectionFrequency(HYPRE_Solver solver, HYPRE_Int
projection_frequency)
(Optional) Set the frequency at which a projection onto the compatible subspace for problems with zeroconductivity regions is performed.

The default value is 5 .
HYPRE_Int HYPRE_AMSSetMaxIter(HYPRE_Solver solver, HYPRE_Int maxit)
(Optional) Sets maximum number of iterations, if AMS is used as a solver.
To use AMS as a preconditioner, set the maximum number of iterations to 1 . The default is 20 .
HYPRE_Int HYPRE_AMSSetTol (HYPRE_Solver solver, HYPRE_Real tol)
(Optional) Set the convergence tolerance, if AMS is used as a solver.
When using AMS as a preconditioner, set the tolerance to 0.0 . The default is $10^{-6}$.
HYPRE_Int HYPRE_AMSSetCycleType(HYPRE_Solver solver, HYPRE_Int cycle_type)
(Optional) Choose which three-level solver to use.
Possible values are:

- 1: 3-level multiplicative solver (01210)
- 2 : 3-level additive solver $(0+1+2)$
- 3 : 3-level multiplicative solver (02120)
- 4 : 3-level additive solver $(010+2)$
- 5 : 3-level multiplicative solver (0102010)
- 6:3-level additive solver ( $1+020$ )
- 7 : 3-level multiplicative solver (0201020)
- 8 : 3-level additive solver $(0(1+2) 0)$
- 11 : 5-level multiplicative solver (013454310)
- 12 : 5-level additive solver $(0+1+3+4+5)$
- 13 : 5-level multiplicative solver (034515430)
- 14 : 5-level additive solver $(01(3+4+5) 10)$

The default is 1 . See the user's manual for more details.
HYPRE_Int HYPRE_AMSSetPrintLevel (HYPRE_Solver solver, HYPRE_Int print_level)
(Optional) Control how much information is printed during the solution iterations.
The default is 1 (print residual norm at each step).
HYPRE_Int HYPRE_AMSSetSmoothingOptions(HYPRE_Solver solver, HYPRE_Int relax_type, HYPRE_Int relax_times, HYPRE_Real relax_weight, HYPRE_Real omega)
(Optional) Sets relaxation parameters for $A$.
The defaults are $2,1,1.0,1.0$.
The available options for relax_type are:

- $1: \ell_{1}$-scaled Jacobi
- $2: \ell_{1}$-scaled block symmetric Gauss-Seidel/SSOR
- 3 : Kaczmarz
- 4 : truncated version of $\ell_{1}$-scaled block symmetric Gauss-Seidel/SSOR
- 16 : Chebyshev

HYPRE_Int HYPRE_AMSSetAlphaAMGOptions(HYPRE_Solver solver, HYPRE_Int alpha_coarsen_type, HYPRE_Int alpha_agg_levels, HYPRE_Int alpha_relax_type, HYPRE_Real alpha_strength_threshold, HYPRE_Int alpha_interp_type, HYPRE_Int alpha_Pmax)
(Optional) Sets AMG parameters for $B_{\Pi}$.
The defaults are $10,1,3,0.25,0,0$. See the user's manual for more details.
HYPRE_Int HYPRE_AMSSetAlphaAMGCoarseRelaxType(HYPRE_Solver solver, HYPRE_Int alpha_coarse_relax_type)
(Optional) Sets the coarsest level relaxation in the AMG solver for $B_{\Pi}$.
The default is 8 (11-GS). Use 9, 19, 29 or 99 for a direct solver.
HYPRE_Int HYPRE_AMSSetBetaAMGOptions(HYPRE_Solver solver, HYPRE_Int beta_coarsen_type, HYPRE_Int beta_agg_levels, HYPRE_Int beta_relax_type, HYPRE_Real beta_strength_threshold, HYPRE_Int beta_interp_type, HYPRE_Int beta_Pmax)
(Optional) Sets AMG parameters for $B_{G}$.
The defaults are $10,1,3,0.25,0,0$. See the user's manual for more details.
HYPRE_Int HYPRE_AMSSetBetaAMGCoarseRelaxType(HYPRE_Solver solver, HYPRE_Int beta_coarse_relax_type)
(Optional) Sets the coarsest level relaxation in the AMG solver for $B_{G}$.
The default is 8 (11-GS). Use 9, 19, 29 or 99 for a direct solver.
HYPRE_Int HYPRE_AMSGetNumIterations(HYPRE_Solver solver, HYPRE_Int *num_iterations)
Returns the number of iterations taken.
HYPRE_Int HYPRE_AMSGetFinalRelativeResidualNorm(HYPRE_Solver solver, HYPRE_Real
*rel_resid_norm)
Returns the norm of the final relative residual.
HYPRE_Int HYPRE_AMSProjectOutGradients(HYPRE_Solver solver, HYPRE_ParVector x)
For problems with zero-conductivity regions, project the vector onto the compatible subspace: $x=(I-$ $\left.G_{0}\left(G_{0}^{t} G_{0}\right)^{-1} G_{0}^{T}\right) x$, where $G_{0}$ is the discrete gradient restricted to the interior nodes of the regions with zero conductivity.
This ensures that x is orthogonal to the gradients in the range of $G_{0}$.
This function is typically called after the solution iteration is complete, in order to facilitate the visualization of the computed field. Without it the values in the zero-conductivity regions contain kernel components.
HYPRE_Int HYPRE_AMSConstructDiscreteGradient(HYPRE_ParCSRMatrix A, HYPRE_ParVector x_coord, HYPRE_BigInt *edge_vertex, HYPRE_Int edge_orientation, HYPRE_ParCSRMatrix *G)
Construct and return the lowest-order discrete gradient matrix G using some edge and vertex information. We assume that edge_vertex lists the edge vertices consecutively, and that the orientation of all edges is consistent.
If edge_orientation $=1$, the edges are already oriented.
If edge_orientation $=2$, the orientation of edge i depends only on the sign of edge_vertex[2*i+1] edge_vertex[2*i].

## ParCSR ADS Solver and Preconditioner

Parallel auxiliary space divergence solver and preconditioner
HYPRE_Int HYPRE_ADSCreate(HYPRE_Solver *solver)
Create an ADS solver object.
HYPRE_Int HYPRE_ADSDestroy (HYPRE_Solver solver)
Destroy an ADS solver object.
HYPRE_Int HYPRE_ADSSetup(HYPRE_Solver solver, HYPRE_ParCSRMatrix A, HYPRE_ParVector b, HYPRE_ParVector x)
Set up the ADS solver or preconditioner.
If used as a preconditioner, this function should be passed to the iterative solver SetPrecond function.

## Parameters

- solver - [IN] object to be set up.
- A - [IN] ParCSR matrix used to construct the solver/preconditioner.
- $\mathbf{b}$ - Ignored by this function.
- $\mathbf{x}$ - Ignored by this function.

HYPRE_Int HYPRE_ADSSolve(HYPRE_Solver solver, HYPRE_ParCSRMatrix A, HYPRE_ParVector b, HYPRE_ParVector x )

Solve the system or apply ADS as a preconditioner.
If used as a preconditioner, this function should be passed to the iterative solver SetPrecond function.

## Parameters

- solver - [IN] solver or preconditioner object to be applied.
- A - [IN] ParCSR matrix, matrix of the linear system to be solved
- $\mathbf{b}-[\mathrm{IN}]$ right hand side of the linear system to be solved
- $\mathbf{x}-$ [OUT] approximated solution of the linear system to be solved

HYPRE_Int HYPRE_ADSSetDiscreteCurl (HYPRE_Solver solver, HYPRE_ParCSRMatrix C)
Sets the discrete curl matrix $C$.
This function should be called before HYPRE_ADSSetup()!
HYPRE_Int HYPRE_ADSSetDiscreteGradient (HYPRE_Solver solver, HYPRE_ParCSRMatrix G)
Sets the discrete gradient matrix $G$.
This function should be called before HYPRE_ADSSetup()!
HYPRE_Int HYPRE_ADSSetCoordinateVectors(HYPRE_Solver solver, HYPRE_ParVector x, HYPRE_ParVector y, HYPRE_ParVector z)
Sets the $x, y$ and $z$ coordinates of the vertices in the mesh.
This function should be called before HYPRE_ADSSetup()!
HYPRE_Int HYPRE_ADSSetInterpolations(HYPRE_Solver solver, HYPRE_ParCSRMatrix RT_Pi, HYPRE_ParCSRMatrix RT_Pix, HYPRE_ParCSRMatrix RT_Piy, HYPRE_ParCSRMatrix RT_Piz, HYPRE_ParCSRMatrix ND_Pi, HYPRE_ParCSRMatrix ND_Pix, HYPRE_ParCSRMatrix ND_Piy, HYPRE_ParCSRMatrix ND_Piz)
(Optional) Set the (components of) the Raviart-Thomas $\left(\Pi_{R T}\right)$ and the Nedelec ( $\Pi_{N D}$ ) interpolation matrices.

This function is generally intended to be used only for high-order $H($ div $)$ discretizations (in the lowest order case, these matrices are constructed internally in ADS from the discreet gradient and curl matrices and the coordinates of the vertices), though it can also be used in the lowest-order case or for other types of discretizations.

By definition, $R T \_P i$ and $N D \_P i$ are the matrix representations of the linear operators $\Pi_{R T}$ and $\Pi_{N D}$ that interpolate (high-order) vector nodal finite elements into the (high-order) Raviart-Thomas and Nedelec spaces. The component matrices are defined in both cases as $\Pi^{x} \varphi=\Pi(\varphi, 0,0)$ and similarly for $\Pi^{y}$ and $\Pi^{z}$. Note that all these operators depend on the choice of the basis and degrees of freedom in the high-order spaces.

The column numbering of $R T \_P i$ and $N D \_P i$ should be node-based, i.e. the $x / y / z$ components of the first node (vertex or high-order dof) should be listed first, followed by the $x / y / z$ components of the second node and so on (see the documentation of HYPRE_BoomerAMGSetDofFunc).
If used, this function should be called before hypre_ADSSetup() and there is no need to provide the vertex coordinates. Furthermore, only one of the sets $\left\{\Pi_{R T}\right\}$ and $\left\{\Pi_{R T}^{x}, \Pi_{R T}^{y}, \Pi_{R T}^{z}\right\}$ needs to be specified (though it is OK to provide both). If RT_Pix is NULL, then scalar ח-based ADS cycles, i.e. those with cycle_type $>10$, will be unavailable. Similarly, ADS cycles based on monolithic $\Pi$ (cycle_type $<10$ ) require that $R T_{-} P i$ is not NULL. The same restrictions hold for the sets $\left\{\Pi_{N D}\right\}$ and $\left\{\Pi_{N D}^{x}, \Pi_{N D}^{y}, \Pi_{N D}^{z}\right\}$ $\& \# 8212$; only one of them needs to be specified, and the availability of each enables different AMS cycle type options.

HYPRE_Int HYPRE_ADSSetMaxIter(HYPRE_Solver solver, HYPRE_Int maxit)
(Optional) Sets maximum number of iterations, if ADS is used as a solver.
To use ADS as a preconditioner, set the maximum number of iterations to 1 . The default is 20 .
HYPRE_Int HYPRE_ADSSetTol (HYPRE_Solver solver, HYPRE_Real tol)
(Optional) Set the convergence tolerance, if ADS is used as a solver.
When using ADS as a preconditioner, set the tolerance to 0.0 . The default is $10^{-6}$.
HYPRE_Int HYPRE_ADSSetCycleType(HYPRE_Solver solver, HYPRE_Int cycle_type)
(Optional) Choose which auxiliary-space solver to use.
Possible values are:

- 1:3-level multiplicative solver (01210)
- 2 : 3-level additive solver $(0+1+2)$
- 3: 3-level multiplicative solver (02120)
- 4 : 3-level additive solver $(010+2)$
- 5 : 3-level multiplicative solver (0102010)
- 6:3-level additive solver ( $1+020$ )
- 7 : 3-level multiplicative solver (0201020)
- 8 : 3-level additive solver $(0(1+2) 0)$
- 11 : 5-level multiplicative solver (013454310)
- 12 : 5 -level additive solver $(0+1+3+4+5)$
- 13: 5-level multiplicative solver (034515430)
- 14 : 5-level additive solver $(01(3+4+5) 10)$

The default is 1 . See the user's manual for more details.
HYPRE_Int HYPRE_ADSSetPrintLevel (HYPRE_Solver solver, HYPRE_Int print_level)
(Optional) Control how much information is printed during the solution iterations.
The default is 1 (print residual norm at each step).
HYPRE_Int HYPRE_ADSSetSmoothingOptions(HYPRE_Solver solver, HYPRE_Int relax_type, HYPRE_Int relax_times, HYPRE_Real relax_weight, HYPRE_Real omega)
(Optional) Sets relaxation parameters for $A$.
The defaults are $2,1,1.0,1.0$.
The available options for relax_type are:

- $1: \ell_{1}$-scaled Jacobi
- 2 : $\ell_{1}$-scaled block symmetric Gauss-Seidel/SSOR
- 3 : Kaczmarz
- 4 : truncated version of $\ell_{1}$-scaled block symmetric Gauss-Seidel/SSOR
- 16 : Chebyshev

HYPRE_Int HYPRE_ADSSetChebySmoothingOptions (HYPRE_Solver solver, HYPRE_Int cheby_order, HYPRE_Real cheby_fraction)
(Optional) Sets parameters for Chebyshev relaxation.
The defaults are 2, 0.3 .
HYPRE_Int HYPRE_ADSSetAMSOptions(HYPRE_Solver solver, HYPRE_Int cycle_type, HYPRE_Int coarsen_type, HYPRE_Int agg_levels, HYPRE_Int relax_type, HYPRE_Real strength_threshold, HYPRE_Int interp_type, HYPRE_Int Pmax)
(Optional) Sets AMS parameters for $B_{C}$.
The defaults are $11,10,1,3,0.25,0,0$. Note that cycle_type should be greater than 10 , unless the high-order interface of HYPRE_ADSSetInterpolations is being used! See the user's manual for more details.

HYPRE_Int HYPRE_ADSSetAMGOptions(HYPRE_Solver solver, HYPRE_Int coarsen_type, HYPRE_Int agg_levels, HYPRE_Int relax_type, HYPRE_Real strength_threshold, HYPRE_Int interp_type, HYPRE_Int Pmax) (Optional) Sets AMG parameters for $B_{\Pi}$. The defaults are $10,1,3,0.25,0,0$. See the user's manual for more details.

HYPRE_Int HYPRE_ADSGetNumIterations(HYPRE_Solver solver, HYPRE_Int *num_iterations) Returns the number of iterations taken.

HYPRE_Int HYPRE_ADSGetFinalRelativeResidualNorm(HYPRE_Solver solver, HYPRE_Real *rel_resid_norm)

Returns the norm of the final relative residual.

## ParCSR PCG Solver

These routines should be used in conjunction with the generic interface in Krylov Solvers.
HYPRE_Int HYPRE_ParCSRPCGCreate(MPI_Comm comm, HYPRE_Solver *solver) Create a solver object.

HYPRE_Int HYPRE_ParCSRPCGDestroy (HYPRE_Solver solver)
Destroy a solver object.
HYPRE_Int HYPRE_ParCSRPCGSetup(HYPRE_Solver solver, HYPRE_ParCSRMatrix A, HYPRE_ParVector b, HYPRE_ParVector x )

HYPRE_Int HYPRE_ParCSRPCGSolve(HYPRE_Solver solver, HYPRE_ParCSRMatrix A, HYPRE_ParVector b, HYPRE_ParVector x)

HYPRE_Int HYPRE_ParCSRPCGSetTol(HYPRE_Solver solver, HYPRE_Real tol)
HYPRE_Int HYPRE_ParCSRPCGSetAbsoluteTol (HYPRE_Solver solver, HYPRE_Real tol)
HYPRE_Int HYPRE_ParCSRPCGSetMaxIter(HYPRE_Solver solver, HYPRE_Int max_iter)
HYPRE_Int HYPRE_ParCSRPCGSetStopCrit(HYPRE_Solver solver, HYPRE_Int stop_crit)
HYPRE_Int HYPRE_ParCSRPCGSetTwoNorm(HYPRE_Solver solver, HYPRE_Int two_norm)
HYPRE_Int HYPRE_ParCSRPCGSetRelChange(HYPRE_Solver solver, HYPRE_Int rel_change)
HYPRE_Int HYPRE_ParCSRPCGSetPrecond(HYPRE_Solver solver, HYPRE_PtrToParSolverFcn precond, HYPRE_PtrToParSolverFcn precond_setup, HYPRE_Solver precond_solver)

HYPRE_Int HYPRE_ParCSRPCGSetPreconditioner(HYPRE_Solver solver, HYPRE_Solver precond)
HYPRE_Int HYPRE_ParCSRPCGGetPrecond(HYPRE_Solver solver, HYPRE_Solver *precond_data)
HYPRE_Int HYPRE_ParCSRPCGSetLogging(HYPRE_Solver solver, HYPRE_Int logging)
HYPRE_Int HYPRE_ParCSRPCGSetPrintLevel (HYPRE_Solver solver, HYPRE_Int print_level)
HYPRE_Int HYPRE_ParCSRPCGGetNumIterations(HYPRE_Solver solver, HYPRE_Int *num_iterations)
HYPRE_Int HYPRE_ParCSRPCGGetFinalRelativeResidualNorm(HYPRE_Solver solver, HYPRE_Real *norm)

HYPRE_Int HYPRE_ParCSRPCGGetResidual (HYPRE_Solver solver, HYPRE_ParVector *residual)
Returns the residual.
HYPRE_Int HYPRE_ParCSRDiagScaleSetup(HYPRE_Solver solver, HYPRE_ParCSRMatrix A, HYPRE_ParVector y, HYPRE_ParVector x)
Setup routine for diagonal preconditioning.
HYPRE_Int HYPRE_ParCSRDiagScale(HYPRE_Solver solver, HYPRE_ParCSRMatrix HA, HYPRE_ParVector Hy, HYPRE_ParVector Hx)
Solve routine for diagonal preconditioning.
HYPRE_Int HYPRE_ParCSROnProcTriSetup(HYPRE_Solver solver, HYPRE_ParCSRMatrix HA, HYPRE_ParVector Hy, HYPRE_ParVector Hx)
Setup routine for on-processor triangular solve as preconditioning.

HYPRE_Int HYPRE_ParCSROnProcTriSolve(HYPRE_Solver solver, HYPRE_ParCSRMatrix HA, HYPRE_ParVector Hy, HYPRE_ParVector Hx)
Solve routine for on-processor triangular solve as preconditioning.

## ParCSR GMRES Solver

These routines should be used in conjunction with the generic interface in Krylov Solvers.
HYPRE_Int HYPRE_ParCSRGMRESCreate(MPI_Comm comm, HYPRE_Solver *solver) Create a solver object.
HYPRE_Int HYPRE_ParCSRGMRESDestroy (HYPRE_Solver solver)
Destroy a solver object.
HYPRE_Int HYPRE_ParCSRGMRESSetup(HYPRE_Solver solver, HYPRE_ParCSRMatrix A, HYPRE_ParVector b, HYPRE_ParVector x)

HYPRE_Int HYPRE_ParCSRGMRESSolve(HYPRE_Solver solver, HYPRE_ParCSRMatrix A, HYPRE_ParVector b, HYPRE_ParVector x)

HYPRE_Int HYPRE_ParCSRGMRESSetKDim(HYPRE_Solver solver, HYPRE_Int k_dim)
HYPRE_Int HYPRE_ParCSRGMRESSetTol (HYPRE_Solver solver, HYPRE_Real tol)
HYPRE_Int HYPRE_ParCSRGMRESSetAbsoluteTol (HYPRE_Solver solver, HYPRE_Real a_tol)
HYPRE_Int HYPRE_ParCSRGMRESSetMinIter(HYPRE_Solver solver, HYPRE_Int min_iter)
HYPRE_Int HYPRE_ParCSRGMRESSetMaxIter(HYPRE_Solver solver, HYPRE_Int max_iter)
HYPRE_Int HYPRE_ParCSRGMRESSetStopCrit (HYPRE_Solver solver, HYPRE_Int stop_crit)
HYPRE_Int HYPRE_ParCSRGMRESSetPrecond(HYPRE_Solver solver, HYPRE_PtrToParSolverFcn precond, HYPRE_PtrToParSolverFcn precond_setup, HYPRE_Solver precond_solver)

HYPRE_Int HYPRE_ParCSRGMRESGetPrecond(HYPRE_Solver solver, HYPRE_Solver *precond_data)
HYPRE_Int HYPRE_ParCSRGMRESSetLogging(HYPRE_Solver solver, HYPRE_Int logging)
HYPRE_Int HYPRE_ParCSRGMRESSetPrintLevel (HYPRE_Solver solver, HYPRE_Int print_level)
HYPRE_Int HYPRE_ParCSRGMRESGetNumIterations(HYPRE_Solver solver, HYPRE_Int *num_iterations)

HYPRE_Int HYPRE_ParCSRGMRESGetFinalRelativeResidualNorm(HYPRE_Solver solver, HYPRE_Real *norm)

HYPRE_Int HYPRE_ParCSRGMRESGetResidual (HYPRE_Solver solver, HYPRE_ParVector *residual)
Returns the residual.
HYPRE_Int HYPRE_ParCSRCOGMRESCreate(MPI_Comm comm, HYPRE_Solver *solver) Create a solver object.
HYPRE_Int HYPRE_ParCSRCOGMRESDestroy (HYPRE_Solver solver)
Destroy a solver object.

[^0]HYPRE_Int HYPRE_ParCSRFlexGMRESSetTol(HYPRE_Solver solver, HYPRE_Real tol)
HYPRE_Int HYPRE_ParCSRFlexGMRESSetAbsoluteTol(HYPRE_Solver solver, HYPRE_Real a_tol)
HYPRE_Int HYPRE_ParCSRFlexGMRESSetMinIter(HYPRE_Solver solver, HYPRE_Int min_iter)
$\begin{array}{r}\text { HYPRE_Int HYPRE_ParCSRFlexGMRESSetMaxIter(HYPRE_Solver solver, HYPRE_Int max_iter) } \\ \text { HYPRE_Int HYPRE_ParCSRFlexGMRESSetPrecond(HYPRE_Solver solver, HYPRE_PtrToParSolverFcn } \\ \text { precond, HYPRE_PtrToParSolverFcn precond_setup, } \\ \text { HYPRE_Solver precond_solver) }\end{array}$
$\begin{array}{r}\text { HYPRE_Int HYPRE_ParCSRFlexGMRESGetPrecond(HYPRE_Solver solver, HYPRE_Solver *precond_data) }\end{array}$
$\begin{array}{r}\text { HYPRE_Int HYPRE_ParCSRFlexGMRESSetLogging(HYPRE_Solver solver, HYPRE_Int logging) }\end{array}$
HYPRE_Int HYPRE_ParCSRFlexGMRESSetPrintLevel(HYPRE_Solver solver, HYPRE_Int print_level)
HYPRE_Int HYPRE_ParCSRFlexGMRESGetNumIterations(HYPRE_Solver solver, HYPRE_Int
*num_iterations)

## ParCSR LGMRES Solver

These routines should be used in conjunction with the generic interface in Krylov Solvers.
HYPRE_Int HYPRE_ParCSRLGMRESCreate(MPI_Comm comm, HYPRE_Solver *solver)
Create a solver object.
HYPRE_Int HYPRE_ParCSRLGMRESDestroy(HYPRE_Solver solver)
Destroy a solver object.
HYPRE_Int HYPRE_ParCSRLGMRESSetup(HYPRE_Solver solver, HYPRE_ParCSRMatrix A, HYPRE_ParVector b, HYPRE_ParVector x)

HYPRE_Int HYPRE_ParCSRLGMRESSolve(HYPRE_Solver solver, HYPRE_ParCSRMatrix A, HYPRE_ParVector b, HYPRE_ParVector x)

HYPRE_Int HYPRE_ParCSRLGMRESSetKDim(HYPRE_Solver solver, HYPRE_Int k_dim)
HYPRE_Int HYPRE_ParCSRLGMRESSetAugDim(HYPRE_Solver solver, HYPRE_Int aug_dim)
HYPRE_Int HYPRE_ParCSRLGMRESSetTol (HYPRE_Solver solver, HYPRE_Real tol)
HYPRE_Int HYPRE_ParCSRLGMRESSetAbsoluteTol (HYPRE_Solver solver, HYPRE_Real a_tol)
HYPRE_Int HYPRE_ParCSRLGMRESSetMinIter (HYPRE_Solver solver, HYPRE_Int min_iter)
HYPRE_Int HYPRE_ParCSRLGMRESSetMaxIter (HYPRE_Solver solver, HYPRE_Int max_iter)

# HYPRE_Int HYPRE_ParCSRLGMRESSetPrecond(HYPRE_Solver solver, HYPRE_PtrToParSolverFcn precond, HYPRE_PtrToParSolverFcn precond_setup, HYPRE_Solver precond_solver) <br> HYPRE_Int HYPRE_ParCSRLGMRESGetPrecond(HYPRE_Solver solver, HYPRE_Solver *precond_data) <br> HYPRE_Int HYPRE_ParCSRLGMRESSetLogging(HYPRE_Solver solver, HYPRE_Int logging) <br> HYPRE_Int HYPRE_ParCSRLGMRESSetPrintLevel (HYPRE_Solver solver, HYPRE_Int print_level) <br> HYPRE_Int HYPRE_ParCSRLGMRESGetNumIterations(HYPRE_Solver solver, HYPRE_Int *num_iterations) <br> HYPRE_Int HYPRE_ParCSRLGMRESGetFinalRelativeResidualNorm(HYPRE_Solver solver, HYPRE_Real *norm) <br> HYPRE_Int HYPRE_ParCSRLGMRESGetResidual(HYPRE_Solver solver, HYPRE_ParVector *residual) <br> <br> ParCSR BiCGSTAB Solver 

 <br> <br> ParCSR BiCGSTAB Solver}

These routines should be used in conjunction with the generic interface in Krylov Solvers.
HYPRE_Int HYPRE_ParCSRBiCGSTABCreate(MPI_Comm comm, HYPRE_Solver *solver)
Create a solver object.
HYPRE_Int HYPRE_ParCSRBiCGSTABDestroy (HYPRE_Solver solver)
Destroy a solver object.
HYPRE_Int HYPRE_ParCSRBiCGSTABSetup(HYPRE_Solver solver, HYPRE_ParCSRMatrix A, HYPRE_ParVector b, HYPRE_ParVector x)

HYPRE_Int HYPRE_ParCSRBiCGSTABSolve(HYPRE_Solver solver, HYPRE_ParCSRMatrix A, HYPRE_ParVector b, HYPRE_ParVector x)

HYPRE_Int HYPRE_ParCSRBiCGSTABSetTol (HYPRE_Solver solver, HYPRE_Real tol)
HYPRE_Int HYPRE_ParCSRBiCGSTABSetAbsoluteTol (HYPRE_Solver solver, HYPRE_Real a_tol)
HYPRE_Int HYPRE_ParCSRBiCGSTABSetMinIter (HYPRE_Solver solver, HYPRE_Int min_iter)
HYPRE_Int HYPRE_ParCSRBiCGSTABSetMaxIter (HYPRE_Solver solver, HYPRE_Int max_iter)
HYPRE_Int HYPRE_ParCSRBiCGSTABSetStopCrit(HYPRE_Solver solver, HYPRE_Int stop_crit)
HYPRE_Int HYPRE_ParCSRBiCGSTABSetPrecond(HYPRE_Solver solver, HYPRE_PtrToParSolverFcn precond, HYPRE_PtrToParSolverFcn precond_setup, HYPRE_Solver precond_solver)

HYPRE_Int HYPRE_ParCSRBiCGSTABGetPrecond(HYPRE_Solver solver, HYPRE_Solver *precond_data)
HYPRE_Int HYPRE_ParCSRBiCGSTABSetLogging(HYPRE_Solver solver, HYPRE_Int logging)
HYPRE_Int HYPRE_ParCSRBiCGSTABSetPrintLevel (HYPRE_Solver solver, HYPRE_Int print_level)
HYPRE_Int HYPRE_ParCSRBiCGSTABGetNumIterations(HYPRE_Solver solver, HYPRE_Int
*num_iterations)

HYPRE_Int HYPRE_ParCSRBiCGSTABGetFinalRelativeResidualNorm(HYPRE_Solver solver, HYPRE_Real *norm)

HYPRE_Int HYPRE_ParCSRBiCGSTABGetResidual (HYPRE_Solver solver, HYPRE_ParVector *residual)

## ParCSR Hybrid Solver

HYPRE_Int HYPRE_ParCSRHybridCreate(HYPRE_Solver *solver)
Create solver object.
HYPRE_Int HYPRE_ParCSRHybridDestroy (HYPRE_Solver solver)
Destroy solver object.
HYPRE_Int HYPRE_ParCSRHybridSetup(HYPRE_Solver solver, HYPRE_ParCSRMatrix A, HYPRE_ParVector b, HYPRE_ParVector x)

Setup the hybrid solver.

## Parameters

- solver - [IN] object to be set up.
- A - [IN] ParCSR matrix used to construct the solver/preconditioner.
- $\mathbf{b}$ - Ignored by this function.
- $\mathbf{x}$ - Ignored by this function.

HYPRE_Int HYPRE_ParCSRHybridSolve(HYPRE_Solver solver, HYPRE_ParCSRMatrix A, HYPRE_ParVector b, HYPRE_ParVector x)
Solve linear system.

## Parameters

- solver - [IN] solver or preconditioner object to be applied.
- A - [IN] ParCSR matrix, matrix of the linear system to be solved
- $\mathbf{b}-[\mathrm{IN}]$ right hand side of the linear system to be solved
- $\mathbf{x}$ - [OUT] approximated solution of the linear system to be solved

HYPRE_Int HYPRE_ParCSRHybridSetTol(HYPRE_Solver solver, HYPRE_Real tol)
Set the convergence tolerance for the Krylov solver.
The default is 1.e-6.
HYPRE_Int HYPRE_ParCSRHybridSetAbsoluteTol (HYPRE_Solver solver, HYPRE_Real tol)
Set the absolute convergence tolerance for the Krylov solver.
The default is 0 .
HYPRE_Int HYPRE_ParCSRHybridSetConvergenceTol (HYPRE_Solver solver, HYPRE_Real cf_tol)
Set the desired convergence factor.
HYPRE_Int HYPRE_ParCSRHybridSetDSCGMaxIter (HYPRE_Solver solver, HYPRE_Int dscg_max_its)
Set the maximal number of iterations for the diagonally preconditioned solver.
HYPRE_Int HYPRE_ParCSRHybridSetPCGMaxIter(HYPRE_Solver solver, HYPRE_Int pcg_max_its)
Set the maximal number of iterations for the AMG preconditioned solver.

HYPRE_Int HYPRE_ParCSRHybridSetSetupType(HYPRE_Solver solver, HYPRE_Int setup_type)
HYPRE_Int HYPRE_ParCSRHybridSetSolverType(HYPRE_Solver solver, HYPRE_Int solver_type)
Set the desired solver type.
There are the following options:

- 1: PCG (default)
- 2 : GMRES
- 3 : BiCGSTAB

HYPRE_Int HYPRE_ParCSRHybridSetRecomputeResidual (HYPRE_Solver solver, HYPRE_Int recompute_residual)
(Optional) Set recompute residual (don't rely on 3-term recurrence).
HYPRE_Int HYPRE_ParCSRHybridGetRecomputeResidual (HYPRE_Solver solver, HYPRE_Int *recompute_residual)
(Optional) Get recompute residual option.
HYPRE_Int HYPRE_ParCSRHybridSetRecomputeResidualP(HYPRE_Solver solver, HYPRE_Int recompute_residual_p)
(Optional) Set recompute residual period (don't rely on 3-term recurrence).
Recomputes residual after every specified number of iterations.
HYPRE_Int HYPRE_ParCSRHybridGetRecomputeResidualP(HYPRE_Solver solver, HYPRE_Int *recompute_residual_p)
(Optional) Get recompute residual period option.
HYPRE_Int HYPRE_ParCSRHybridSetKDim(HYPRE_Solver solver, HYPRE_Int k_dim)
Set the Krylov dimension for restarted GMRES.
The default is 5 .
HYPRE_Int HYPRE_ParCSRHybridSetTwoNorm(HYPRE_Solver solver, HYPRE_Int two_norm) Set the type of norm for PCG.

HYPRE_Int HYPRE_ParCSRHybridSetStopCrit(HYPRE_Solver solver, HYPRE_Int stop_crit) RE-VISIT.

HYPRE_Int HYPRE_ParCSRHybridSetRelChange(HYPRE_Solver solver, HYPRE_Int rel_change)
HYPRE_Int HYPRE_ParCSRHybridSetPrecond(HYPRE_Solver solver, HYPRE_PtrToParSolverFcn precond, HYPRE_PtrToParSolverFcn precond_setup, HYPRE_Solver precond_solver) Set preconditioner if wanting to use one that is not set up by the hybrid solver.
HYPRE_Int HYPRE_ParCSRHybridSetLogging (HYPRE_Solver solver, HYPRE_Int logging)
Set logging parameter (default: 0, no logging).
HYPRE_Int HYPRE_ParCSRHybridSetPrintLevel (HYPRE_Solver solver, HYPRE_Int print_level)
Set print level (default: 0 , no printing) 2 will print residual norms per iteration 10 will print AMG setup information if AMG is used 12 both Setup information and iterations.

HYPRE_Int HYPRE_ParCSRHybridSetStrongThreshold(HYPRE_Solver solver, HYPRE_Real strong_threshold)
(Optional) Sets AMG strength threshold.
The default is 0.25 . For elasticity problems, a larger strength threshold, such as 0.7 or 0.8 , is often better.

HYPRE_Int HYPRE_ParCSRHybridSetMaxRowSum(HYPRE_Solver solver, HYPRE_Real max_row_sum) (Optional) Sets a parameter to modify the definition of strength for diagonal dominant portions of the matrix.

The default is 0.9 . If max_row_sum is 1 , no checking for diagonally dominant rows is performed.
HYPRE_Int HYPRE_ParCSRHybridSetTruncFactor(HYPRE_Solver solver, HYPRE_Real trunc_factor) (Optional) Defines a truncation factor for the interpolation.

The default is 0 .
HYPRE_Int HYPRE_ParCSRHybridSetPMaxElmts (HYPRE_Solver solver, HYPRE_Int P_max_elmts)
(Optional) Defines the maximal number of elements per row for the interpolation.
The default is 0 .
HYPRE_Int HYPRE_ParCSRHybridSetMaxLevels(HYPRE_Solver solver, HYPRE_Int max_levels)
(Optional) Defines the maximal number of levels used for AMG.
The default is 25 .
HYPRE_Int HYPRE_ParCSRHybridSetMeasureType(HYPRE_Solver solver, HYPRE_Int measure_type)
(Optional) Defines whether local or global measures are used.
HYPRE_Int HYPRE_ParCSRHybridSetCoarsenType(HYPRE_Solver solver, HYPRE_Int coarsen_type)
(Optional) Defines which parallel coarsening algorithm is used.
There are the following options for coarsen_type:

- 0 : CLJP-coarsening (a parallel coarsening algorithm using independent sets).
- 1 : classical Ruge-Stueben coarsening on each processor, no boundary treatment
- 3 : classical Ruge-Stueben coarsening on each processor, followed by a third pass, which adds coarse points on the boundaries
- 6 : Falgout coarsening (uses 1 first, followed by CLJP using the interior coarse points generated by 1 as its first independent set)
- 7 : CLJP-coarsening (using a fixed random vector, for debugging purposes only)
- 8: PMIS-coarsening (a parallel coarsening algorithm using independent sets with lower complexities than CLJP, might also lead to slower convergence)
- 9 : PMIS-coarsening (using a fixed random vector, for debugging purposes only)
- 10 : HMIS-coarsening (uses one pass Ruge-Stueben on each processor independently, followed by PMIS using the interior C-points as its first independent set)
- 11 : one-pass Ruge-Stueben coarsening on each processor, no boundary treatment

The default is 10 .
HYPRE_Int HYPRE_ParCSRHybridSetInterpType(HYPRE_Solver solver, HYPRE_Int interp_type)
(Optional) Specifies which interpolation operator is used The default is ext+i interpolation truncated to at most 4 elements per row.

HYPRE_Int HYPRE_ParCSRHybridSetCycleType(HYPRE_Solver solver, HYPRE_Int cycle_type) (Optional) Defines the type of cycle.
For a V-cycle, set cycle_type to 1 , for a W-cycle set cycle_type to 2 . The default is 1 .

HYPRE_Int HYPRE_ParCSRHybridSetGridRelaxType(HYPRE_Solver solver, HYPRE_Int *grid_relax_type)

HYPRE_Int HYPRE_ParCSRHybridSetGridRelaxPoints(HYPRE_Solver solver, HYPRE_Int **grid_relax_points)

HYPRE_Int HYPRE_ParCSRHybridSetNumSweeps (HYPRE_Solver solver, HYPRE_Int num_sweeps)
(Optional) Sets the number of sweeps.
On the finest level, the up and the down cycle the number of sweeps are set to num_sweeps and on the coarsest level to 1 . The default is 1 .
HYPRE_Int HYPRE_ParCSRHybridSetCycleNumSweeps (HYPRE_Solver solver, HYPRE_Int num_sweeps, HYPRE_Int k)
(Optional) Sets the number of sweeps at a specified cycle.
There are the following options for $k$ :

- 1 : the down cycle
- 2 : the up cycle
- 3 : the coarsest level

HYPRE_Int HYPRE_ParCSRHybridSetRelaxType(HYPRE_Solver solver, HYPRE_Int relax_type)
(Optional) Defines the smoother to be used.
It uses the given smoother on the fine grid, the up and the down cycle and sets the solver on the coarsest level to Gaussian elimination (9). The default is 11 -Gauss-Seidel, forward solve on the down cycle (13) and backward solve on the up cycle (14).
There are the following options for relax_type:

- 0 : Jacobi
- 1 : Gauss-Seidel, sequential (very slow!)
- 2 : Gauss-Seidel, interior points in parallel, boundary sequential (slow!)
- 3 : hybrid Gauss-Seidel or SOR, forward solve
- 4 : hybrid Gauss-Seidel or SOR, backward solve
- 6 : hybrid symmetric Gauss-Seidel or SSOR
- 8 : hybrid symmetric 11 -Gauss-Seidel or SSOR
- 13 : 11-Gauss-Seidel, forward solve
- 14 : 11-Gauss-Seidel, backward solve
- 18 : 11-Jacobi
- 9 : Gaussian elimination (only on coarsest level)

HYPRE_Int HYPRE_ParCSRHybridSetCycleRelaxType(HYPRE_Solver solver, HYPRE_Int relax_type, HYPRE_Int k)
(Optional) Defines the smoother at a given cycle.
For options of relax_type see description of HYPRE_BoomerAMGSetRelaxType). Options for k are

- 1: the down cycle
- 2 : the up cycle
- 3 : the coarsest level

HYPRE_Int HYPRE_ParCSRHybridSetRelax0rder (HYPRE_Solver solver, HYPRE_Int relax_order)
(Optional) Defines in which order the points are relaxed.
There are the following options for relax_order:

- 0 : the points are relaxed in natural or lexicographic order on each processor
- 1: CF-relaxation is used, i.e on the fine grid and the down cycle the coarse points are relaxed first, followed by the fine points; on the up cycle the F-points are relaxed first, followed by the C-points. On the coarsest level, if an iterative scheme is used, the points are relaxed in lexicographic order.

The default is 0 (CF-relaxation).
HYPRE_Int HYPRE_ParCSRHybridSetRelaxWt (HYPRE_Solver solver, HYPRE_Real relax_wt)
(Optional) Defines the relaxation weight for smoothed Jacobi and hybrid SOR on all levels.
Values for relax_wt are

- >0 : this assigns the given relaxation weight on all levels
- $=0$ : the weight is determined on each level with the estimate $\frac{3}{4\left\|D^{-1 / 2} A D^{-1 / 2}\right\|}$, where $D$ is the diagonal of $A$ (this should only be used with Jacobi)
- = -k : the relaxation weight is determined with at most k CG steps on each level (this should only be used for symmetric positive definite problems)

The default is 1 .
HYPRE_Int HYPRE_ParCSRHybridSetLevelRelaxWt (HYPRE_Solver solver, HYPRE_Real relax_wt, HYPRE_Int level)
(Optional) Defines the relaxation weight for smoothed Jacobi and hybrid SOR on the user defined level.
Note that the finest level is denoted 0, the next coarser level 1, etc. For nonpositive relax_weight, the parameter is determined on the given level as described for HYPRE_BoomerAMGSetRelaxWt. The default is 1 .

HYPRE_Int HYPRE_ParCSRHybridSetOuterWt (HYPRE_Solver solver, HYPRE_Real outer_wt)
(Optional) Defines the outer relaxation weight for hybrid SOR and SSOR on all levels.
Values for outer_wt are

- > 0 : this assigns the same outer relaxation weight omega on each level
- = -k : an outer relaxation weight is determined with at most k CG steps on each level (this only makes sense for symmetric positive definite problems and smoothers such as SSOR)

The default is 1 .
HYPRE_Int HYPRE_ParCSRHybridSetLevelOuterWt (HYPRE_Solver solver, HYPRE_Real outer_wt, HYPRE_Int level)
(Optional) Defines the outer relaxation weight for hybrid SOR or SSOR on the user defined level.
Note that the finest level is denoted 0 , the next coarser level 1 , etc. For nonpositive omega, the parameter is determined on the given level as described for HYPRE_BoomerAMGSetOuterWt. The default is 1 .

HYPRE_Int HYPRE_ParCSRHybridSetMaxCoarseSize(HYPRE_Solver solver, HYPRE_Int max_coarse_size)
(Optional) Defines the maximal coarse grid size.
The default is 9 .
HYPRE_Int HYPRE_ParCSRHybridSetMinCoarseSize(HYPRE_Solver solver, HYPRE_Int min_coarse_size)
(Optional) Defines the minimal coarse grid size. The default is 0 .

HYPRE_Int HYPRE_ParCSRHybridSetSeqThreshold(HYPRE_Solver solver, HYPRE_Int seq_threshold) (Optional) enables redundant coarse grid size.
If the system size becomes smaller than seq_threshold, sequential AMG is used on all remaining processors. The default is 0 .

HYPRE_Int HYPRE_ParCSRHybridSetRelaxWeight (HYPRE_Solver solver, HYPRE_Real *relax_weight)
HYPRE_Int HYPRE_ParCSRHybridSetOmega(HYPRE_Solver solver, HYPRE_Real *omega)
HYPRE_Int HYPRE_ParCSRHybridSetAggNumLevels(HYPRE_Solver solver, HYPRE_Int agg_num_levels)
(Optional) Defines the number of levels of aggressive coarsening, starting with the finest level. The default is 0 , i.e. no aggressive coarsening.

HYPRE_Int HYPRE_ParCSRHybridSetAggInterpType(HYPRE_Solver solver, HYPRE_Int agg_interp_type)
(Optional) Defines the interpolation used on levels of aggressive coarsening The default is 4, i.e. multipass interpolation. The following options exist:

- 1:2-stage extended+i interpolation
- 2:2-stage standard interpolation
- 3:2-stage extended interpolation
- 4 : multipass interpolation
- 5: 2-stage extended interpolation in matrix-matrix form
- 6:2-stage extended+i interpolation in matrix-matrix form
- 7: 2-stage extended+e interpolation in matrix-matrix form

HYPRE_Int HYPRE_ParCSRHybridSetNumPaths(HYPRE_Solver solver, HYPRE_Int num_paths)
(Optional) Defines the degree of aggressive coarsening.
The default is 1 , which leads to the most aggressive coarsening. Setting num_paths to 2 will increase complexity somewhat, but can lead to better convergence.
HYPRE_Int HYPRE_ParCSRHybridSetNumFunctions (HYPRE_Solver solver, HYPRE_Int num_functions) (Optional) Sets the size of the system of PDEs, if using the systems version.
The default is 1 .

HYPRE_Int HYPRE_ParCSRHybridSetDofFunc (HYPRE_Solver solver, HYPRE_Int *dof_func)
(Optional) Sets the mapping that assigns the function to each variable, if using the systems version.
If no assignment is made and the number of functions is $\mathrm{k}>1$, the mapping generated is $(0,1, \ldots, \mathrm{k}$ $1,0,1, \ldots, k-1, \ldots)$.

HYPRE_Int HYPRE_ParCSRHybridSetNodal (HYPRE_Solver solver, HYPRE_Int nodal)
(Optional) Sets whether to use the nodal systems version.
The default is 0 (the unknown based approach).
HYPRE_Int HYPRE_ParCSRHybridSetKeepTranspose(HYPRE_Solver solver, HYPRE_Int keepT) (Optional) Sets whether to store local transposed interpolation The default is 0 (don't store).

HYPRE_Int HYPRE_ParCSRHybridSetNonGalerkinTol (HYPRE_Solver solver, HYPRE_Int num_levels, HYPRE_Real *nongalerkin_tol)
(Optional) Sets whether to use non-Galerkin option The default is no non-Galerkin option num_levels sets the number of levels where to use it nongalerkin_tol contains the tolerances for <num_levels> levels
HYPRE_Int HYPRE_ParCSRHybridGetNumIterations(HYPRE_Solver solver, HYPRE_Int *num_its)
Retrieves the total number of iterations.
HYPRE_Int HYPRE_ParCSRHybridGetDSCGNumIterations(HYPRE_Solver solver, HYPRE_Int
*dscg_num_its)
Retrieves the number of iterations used by the diagonally scaled solver.
HYPRE_Int HYPRE_ParCSRHybridGetPCGNumIterations(HYPRE_Solver solver, HYPRE_Int
*pcg_num_its)
Retrieves the number of iterations used by the AMG preconditioned solver.
HYPRE_Int HYPRE_ParCSRHybridGetFinalRelativeResidualNorm(HYPRE_Solver solver, HYPRE_Real *norm)

Retrieves the final relative residual norm.
HYPRE_Int HYPRE_ParCSRHybridSetNumGridSweeps(HYPRE_Solver solver, HYPRE_Int *num_grid_sweeps)

HYPRE_Int HYPRE_ParCSRHybridGetSetupSolveTime(HYPRE_Solver solver, HYPRE_Real *time)

## ParCSR MGR Solver

Parallel multigrid reduction solver and preconditioner.
This solver or preconditioner is designed with systems of PDEs in mind. However, it can also be used for scalar linear systems, particularly for problems where the user can exploit information from the physics of the problem. In this way, the MGR solver could potentially be used as a foundation for a physics-based preconditioner.
HYPRE_Int HYPRE_MGRCreate(HYPRE_Solver *solver)
Create a solver object.
HYPRE_Int HYPRE_MGRDestroy (HYPRE_Solver solver)
Destroy a solver object.
HYPRE_Int HYPRE_MGRSetup(HYPRE_Solver solver, HYPRE_ParCSRMatrix A, HYPRE_ParVector b, HYPRE_ParVector x )
Setup the MGR solver or preconditioner.
If used as a preconditioner, this function should be passed to the iterative solver SetPrecond function.

## Parameters

- solver - [IN] object to be set up.
- A - [IN] ParCSR matrix used to construct the solver/preconditioner.
- $\mathbf{b}$ - right-hand-side of the linear system to be solved (Ignored by this function).
- $\mathbf{x}$ - approximate solution of the linear system to be solved (Ignored by this function).

HYPRE_Int HYPRE_MGRSolve(HYPRE_Solver solver, HYPRE_ParCSRMatrix A, HYPRE_ParVector b, HYPRE_ParVector x)
Solve the system or apply MGR as a preconditioner.
If used as a preconditioner, this function should be passed to the iterative solver SetPrecond function.

## Parameters

- solver - [IN] solver or preconditioner object to be applied.
- A - [IN] ParCSR matrix, matrix of the linear system to be solved
- $\mathbf{b}-[$ IN $]$ right hand side of the linear system to be solved
- $\mathbf{x}$ - [OUT] approximated solution of the linear system to be solved

HYPRE_Int HYPRE_MGRSetCpointsByContiguousBlock(HYPRE_Solver solver, HYPRE_Int block_size, HYPRE_Int max_num_levels, HYPRE_BigInt *idx_array, HYPRE_Int
*num_block_coarse_points, HYPRE_Int
**block_coarse_indexes)
Set the block data assuming that the physical variables are ordered contiguously, i.e.

$$
\mathrm{p} \_1, \mathrm{p} \_2, \ldots, \mathrm{p} \_\mathrm{n}, \mathrm{~s} \_1, \mathrm{~s} \_2, \ldots, \mathrm{~s} \_\mathrm{n}, \ldots
$$

## Parameters

- solver - [IN] solver or preconditioner object
- block_size - [IN] system block size
- max_num_levels - [IN] maximum number of reduction levels
- num_block_coarse_points - [IN] number of coarse points per block per level
- block_coarse_indexes - [IN] index for each block coarse point per level

HYPRE_Int HYPRE_MGRSetCpointsByBlock(HYPRE_Solver solver, HYPRE_Int block_size, HYPRE_Int max_num_levels, HYPRE_Int *num_block_coarse_points, HYPRE_Int **block_coarse_indexes)
Set the block data (by grid points) and prescribe the coarse indexes per block for each reduction level.

## Parameters

- solver - [IN] solver or preconditioner object
- block_size - [IN] system block size
- max_num_levels - [IN] maximum number of reduction levels
- num_block_coarse_points - [IN] number of coarse points per block per level
- block_coarse_indexes - [IN] index for each block coarse point per level

HYPRE_Int HYPRE_MGRSetCpointsByPointMarkerArray (HYPRE_Solver solver, HYPRE_Int block_size, HYPRE_Int max_num_levels, HYPRE_Int *num_block_coarse_points, HYPRE_Int **lvl_block_coarse_indexes, HYPRE_Int *point_marker_array)
Set the coarse indices for the levels using an array of tags for all the local degrees of freedom.
TODO: Rename the function to make it more descriptive.

## Parameters

- solver - [IN] solver or preconditioner object
- block_size - [IN] system block size
- max_num_levels - [IN] maximum number of reduction levels
- num_block_coarse_points - [IN] number of coarse points per block per level
- lvl_block_coarse_indexes - [IN] indices for the coarse points per level
- point_marker_array - [IN] array of tags for the local degrees of freedom

HYPRE_Int HYPRE_MGRSetNonCpointsToFpoints(HYPRE_Solver solver, HYPRE_Int nonCptToFptFlag)
(Optional) Set non C-points to F-points.
This routine determines how the coarse points are selected for the next level reduction. Options for nonCptToFptFlag are:

- 0 : Allow points not prescribed as $C$ points to be potentially set as $C$ points using classical AMG coarsening strategies (currently uses CLJP-coarsening).
- 1: Fix points not prescribed as $C$ points to be $F$ points for the next reduction

HYPRE_Int HYPRE_MGRSetMaxCoarseLevels(HYPRE_Solver solver, HYPRE_Int maxlev)
(Optional) Set maximum number of coarsening (or reduction) levels.
The default is 10 .
HYPRE_Int HYPRE_MGRSetBlockSize(HYPRE_Solver solver, HYPRE_Int bsize)
(Optional) Set the system block size.
This should match the block size set in the MGRSetCpointsByBlock function. The default is 1 .
HYPRE_Int HYPRE_MGRSetReservedCoarseNodes(HYPRE_Solver solver, HYPRE_Int reserved_coarse_size, HYPRE_BigInt *reserved_coarse_nodes)
(Optional) Defines indexes of coarse nodes to be kept to the coarsest level.
These indexes are passed down through the MGR hierarchy to the coarsest grid of the coarse grid (BoomerAMG) solver.

## Parameters

- solver - [IN] solver or preconditioner object
- reserved_coarse_size - [IN] number of reserved coarse points
- reserved_coarse_nodes - [IN] (global) indexes of reserved coarse points

HYPRE_Int HYPRE_MGRSetReservedCpointsLevelToKeep(HYPRE_Solver solver, HYPRE_Int level)
(Optional) Set the level for reducing the reserved Cpoints before the coarse grid solve.
This is necessary for some applications, such as phase transitions. The default is 0 (no reduction, i.e. keep the reserved cpoints in the coarse grid solve).

The default setup for the reduction is as follows: interp_type $=2$ restrict_type $=0$ F-relax method $=99$ Galerkin coarse grid
HYPRE_Int HYPRE_MGRSetRelaxType (HYPRE_Solver solver, HYPRE_Int relax_type)
(Optional) Set the relaxation type for F-relaxation.
Currently supports the following flavors of relaxation types as described in the BoomerAMGSetRelaxType: relax_type 0, 3-8, 13, 14, 18. Also supports AMG (options 1 and 2 ) and direct solver variants (9, 99, 199). See HYPRE_MGRSetLevelFRelaxType for details.

HYPRE_Int HYPRE_MGRSetFRelaxMethod(HYPRE_Solver solver, HYPRE_Int relax_method)
(Optional) Set the strategy for F-relaxation.
Options for relax_method are:

- 0 : Single-level relaxation sweeps for F-relaxation as prescribed by MGRSetRelaxType
- 1 : Multi-level relaxation strategy for F-relaxation ( $\mathrm{V}(1,0)$ cycle currently supported).

NOTE: This function will be removed in favor of HYPRE_MGRSetLevelFRelaxType!!
HYPRE_Int HYPRE_MGRSetLevelFRelaxMethod(HYPRE_Solver solver, HYPRE_Int *relax_method)
(Optional) This function is an extension of HYPRE_MGRSetFRelaxMethod.
It allows setting the F-relaxation strategy for each MGR level.
HYPRE_Int HYPRE_MGRSetLevelFRelaxType(HYPRE_Solver solver, HYPRE_Int *relax_type)
(Optional) Set the relaxation type for F-relaxation at each level.
This function takes precedence over, and will replace HYPRE_MGRSetFRelaxMethod and HYPRE_MGRSetRelaxType. Options for relax_type entries are:

- 0, 3-8, 13, 14, 18: (as described in BoomerAMGSetRelaxType)
- 1 : Multi-level relaxation strategy for F-relaxation ( $\mathrm{V}(1,0)$ cycle currently supported).
- 2 : AMG
- 9, 99, 199 : Gaussian Elimination variants (GE, GE with pivoting, direct inversion respectively)

HYPRE_Int HYPRE_MGRSetCoarseGridMethod(HYPRE_Solver solver, HYPRE_Int *cg_method)
(Optional) Set the strategy for coarse grid computation.
Options for cg_method are:

- 0 : Galerkin coarse grid computation using RAP.
- 5 : Galerkin coarse grid computation using RAI (injective prolongation).
- 1-4: Non-Galerkin coarse grid computation with dropping strategy.
- 1: inv(A_FF) approximated by its (block) diagonal inverse
- 2: CPR-like approximation with inv(A_FF) approximated by its diagonal inverse
- 3: CPR-like approximation with inv(A_FF) approximated by its block diagonal inverse
- 4: inv(A_FF) approximated by sparse approximate inverse

HYPRE_Int HYPRE_MGRSetLevelFRelaxNumFunctions(HYPRE_Solver solver, HYPRE_Int
*num_functions)
(Optional) Set the number of functions for F-relaxation V-cycle.
For problems like elasticity, one may want to perform coarsening and interpolation for block matrices. The number of functions corresponds to the number of scalar PDEs in the system.

HYPRE_Int HYPRE_MGRSetRestrictType(HYPRE_Solver solver, HYPRE_Int restrict_type)
(Optional) Set the strategy for computing the MGR restriction operator.
Options for restrict_type are:

- 0 : injection $[0 I]$
- 1: unscaled (not recommended)
- 2 : diagonal scaling (Jacobi)
- 3 : approximate inverse
- 4 : pAIR distance 1
- 5 : pAIR distance 2
- 12 : Block Jacobi
- 13 : CPR-like restriction operator
- 14 : (Block) Column-lumped restriction
- else : use classical modified interpolation

The default is injection.
HYPRE_Int HYPRE_MGRSetLevelRestrictType(HYPRE_Solver solver, HYPRE_Int *restrict_type)
(Optional) This function is an extension of HYPRE_MGRSetRestrictType.
It allows setting the restriction operator strategy for each MGR level.
HYPRE_Int HYPRE_MGRSetNumRestrictSweeps (HYPRE_Solver solver, HYPRE_Int nsweeps)
(Optional) Set number of restriction sweeps.
This option is for restrict_type $>2$.
HYPRE_Int HYPRE_MGRSetInterpType (HYPRE_Solver solver, HYPRE_Int interp_type)
(Optional) Set the strategy for computing the MGR interpolation operator.
Options for interp_type are:

- 0 : injection $[0 I]^{T}$
- 1 : L1-Jacobi
- 2 : diagonal scaling (Jacobi)
- 3 : classical modified interpolation
- 4 : approximate inverse
- 12 : Block Jacobi
- else : classical modified interpolation

The default is diagonal scaling.
HYPRE_Int HYPRE_MGRSetLevelInterpType(HYPRE_Solver solver, HYPRE_Int *interp_type)
(Optional) This function is an extension of HYPRE_MGRSetInterpType.
It allows setting the prolongation (interpolation) operator strategy for each MGR level.
HYPRE_Int HYPRE_MGRSetNumRelaxSweeps (HYPRE_Solver solver, HYPRE_Int nsweeps)
(Optional) Set number of relaxation sweeps.
This option is for the "single level" F-relaxation (relax_method $=0$ ).
HYPRE_Int HYPRE_MGRSetLevelNumRelaxSweeps (HYPRE_Solver solver, HYPRE_Int *nsweeps)
(Optional) This function is an extension of HYPRE_MGRSetNumRelaxSweeps.
It allows setting the number of single-level relaxation sweeps for each MGR level.
HYPRE_Int HYPRE_MGRSetNumInterpSweeps (HYPRE_Solver solver, HYPRE_Int nsweeps)
(Optional) Set number of interpolation sweeps.
This option is for interp_type $>2$.
HYPRE_Int HYPRE_MGRSetBlockJacobiBlockSize(HYPRE_Solver solver, HYPRE_Int blk_size)
(Optional) Set block size for block (global) smoother and interp/restriction.
This option is for interp_type/restrict_type $==12$, and smooth_type $=0$ or 1 .
HYPRE_Int HYPRE_MGRSetFSolver(HYPRE_Solver solver, HYPRE_PtrToParSolverFcn fine_grid_solver_solve, HYPRE_PtrToParSolverFcn fine_grid_solver_setup, HYPRE_Solver fsolver)
(Optional) Set the fine grid solver.

## Parameters

- solver - [IN] MGR solver/preconditioner object
- fine_grid_solver_solve - [IN] solve routine
- fine_grid_solver_setup - [IN] setup routine
- fine_grid_solver - [IN] fine grid solver object

HYPRE_Int HYPRE_MGRSetFSolverAtLevel(HYPRE_Solver solver, HYPRE_Solver fsolver, HYPRE_Int level)
(Optional) Set the F-relaxation solver at a given level.

## Parameters

- solver - [IN] MGR solver/preconditioner object
- fsolver - [IN] F-relaxation solver object
- level - [IN] MGR solver level

HYPRE_Int HYPRE_MGRBuildAff(HYPRE_ParCSRMatrix A, HYPRE_Int *CF_marker, HYPRE_Int debug_flag, HYPRE_ParCSRMatrix *A_ff)
(Optional) Extract A_FF block from matrix A.
TODO (VPM): Does this need to be exposed? Move to parcsr_mv?

HYPRE_Int HYPRE_MGRSetCoarseSolver(HYPRE_Solver solver, HYPRE_PtrToParSolverFcn coarse_grid_solver_solve, HYPRE_PtrToParSolverFcn coarse_grid_solver_setup, HYPRE_Solver coarse_grid_solver)
(Optional) Set the coarse grid solver.
Currently uses BoomerAMG. The default, if not set, is BoomerAMG with default options.

## Parameters

- solver - [IN] MGR solver/preconditioner object
- coarse_grid_solver_solve - [IN] solve routine for BoomerAMG
- coarse_grid_solver_setup - [IN] setup routine for BoomerAMG
- coarse_grid_solver - [IN] coarse grid solver object

HYPRE_Int HYPRE_MGRSetPrintLevel (HYPRE_Solver solver, HYPRE_Int print_level)
HYPRE_Int HYPRE_MGRSetFrelaxPrintLevel (HYPRE_Solver solver, HYPRE_Int print_level)
(Optional) Set the print level of the F-relaxation solver
HYPRE_Int HYPRE_MGRSetCoarseGridPrintLevel (HYPRE_Solver solver, HYPRE_Int print_level)
(Optional) Set the print level of the coarse grid solver
HYPRE_Int HYPRE_MGRSetTruncateCoarseGridThreshold(HYPRE_Solver solver, HYPRE_Real threshold)
(Optional) Set the threshold for dropping small entries on the coarse grid at each level.
No dropping is applied if threshold $=0.0$ (default).
HYPRE_Int HYPRE_MGRSetLogging(HYPRE_Solver solver, HYPRE_Int logging)
(Optional) Requests logging of solver diagnostics.
Requests additional computations for diagnostic and similar data to be logged by the user. Default is 0 , do nothing. The latest residual will be available if logging $>1$.
HYPRE_Int HYPRE_MGRSetMaxIter (HYPRE_Solver solver, HYPRE_Int max_iter)
(Optional) Set maximum number of iterations if used as a solver.
Set this to 1 if MGR is used as a preconditioner. The default is 20 .
HYPRE_Int HYPRE_MGRSetTol (HYPRE_Solver solver, HYPRE_Real tol)
(Optional) Set the convergence tolerance for the MGR solver.
Use tol $=0.0$ if MGR is used as a preconditioner. The default is 1.e-6.
HYPRE_Int HYPRE_MGRSetMaxGlobalSmoothIters(HYPRE_Solver solver, HYPRE_Int smooth_iter)
(Optional) Determines how many sweeps of global smoothing to do.
Default is 0 (no global smoothing).
HYPRE_Int HYPRE_MGRSetLevelSmoothIters(HYPRE_Solver solver, HYPRE_Int *smooth_iters)
(Optional) Determines how many sweeps of global smoothing to do on each level.
Default is 0 (no global smoothing).
HYPRE_Int HYPRE_MGRSetGlobalSmoothCycle(HYPRE_Solver solver, HYPRE_Int global_smooth_cycle)
(Optional) Set the cycle for global smoothing.
Options for global_smooth_cycle are:

- 1 : Pre-smoothing - Down cycle (default)
- 2 : Post-smoothing - Up cycle

HYPRE_Int HYPRE_MGRSetGlobalSmoothType(HYPRE_Solver solver, HYPRE_Int smooth_type)
(Optional) Determines type of global smoother.
Options for smooth_type are:

- 0 : block Jacobi (default)
- 1 : block Gauss-Seidel
- 2 : Jacobi
- 3 : Gauss-Seidel, sequential (very slow!)
- 4 : Gauss-Seidel, interior points in parallel, boundary sequential (slow!)
- 5 : hybrid Gauss-Seidel or SOR, forward solve
- 6 : hybrid Gauss-Seidel or SOR, backward solve
- 8 : Euclid (ILU)
- 16 : HYPRE_ILU
- 18 : L1-Jacobi

HYPRE_Int HYPRE_MGRSetLevelSmoothType(HYPRE_Solver solver, HYPRE_Int *smooth_type)
Sets the type of global smoother for each level in the multigrid reduction (MGR) solver.
This function allows the user to specify the type of global smoother to be used at each level of the multigrid reduction process. The types of smoothers available can be found in the documentation for HYPRE_MGRSetGlobalSmoothType. The smoother type for each level is indicated by the smooth_type array, which should have a size equal to max_num_coarse_levels.

Note: This function does not take ownership of the smooth_type array.

Note: If smooth_type is a NULL pointer, a default global smoother (Jacobi) is used for all levels.

Note: This call is optional. It is intended for advanced users who need specific control over the smoothing process at different levels of the solver. If not called, the solver will proceed with default smoothing parameters.

## Param

HYPRE_Int HYPRE_MGRSetGlobalSmootherAtLevel (HYPRE_Solver solver, HYPRE_Solver smoother, HYPRE_Int level)
Sets the global smoother method for a specified MGR level using a HYPRE solver object.
This function enables solvers within hypre to be used as complex smoothers for a specific level within the multigrid reduction (MGR) scheme. Users can configure the solver options and pass the solver in as the smoother. Currently supported solver options via this interface are ILU and AMG.

Note: Unlike some other setup functions that might require an array to set options across multiple levels, this function focuses on a single level, identified by the level parameter.

Warning: The smoother passed to function takes precedence over the smoother type set for that level in the MGR hierarchy.

## Param

HYPRE_Int HYPRE_MGRGetNumIterations(HYPRE_Solver solver, HYPRE_Int *num_iterations)
(Optional) Return the number of MGR iterations.
HYPRE_Int HYPRE_MGRGetCoarseGridConvergenceFactor(HYPRE_Solver solver, HYPRE_Real *conv_factor)
(Optional) Return the relative residual for the coarse level system.
HYPRE_Int HYPRE_MGRSetPMaxElmts(HYPRE_Solver solver, HYPRE_Int P_max_elmts)
(Optional) Set the maximum number of nonzeros per row for interpolation operators.
HYPRE_Int HYPRE_MGRSetLevelPMaxElmts(HYPRE_Solver solver, HYPRE_Int *P_max_elmts)
(Optional) Set the maximum number of nonzeros per row for interpolation operators for each level.
HYPRE_Int HYPRE_MGRGetFinalRelativeResidualNorm(HYPRE_Solver solver, HYPRE_Real *res_norm)
(Optional) Return the norm of the final relative residual.

## ParCSR ILU Solver

(Parallel) Incomplete LU factorization.
HYPRE_Int HYPRE_ILUCreate(HYPRE_Solver *solver)
Create a solver object.
HYPRE_Int HYPRE_ILUDestroy (HYPRE_Solver solver)
Destroy a solver object.
HYPRE_Int HYPRE_ILUSetup(HYPRE_Solver solver, HYPRE_ParCSRMatrix A, HYPRE_ParVector b, HYPRE_ParVector x )
Setup the ILU solver or preconditioner.
If used as a preconditioner, this function should be passed to the iterative solver SetPrecond function.

## Parameters

- solver - [IN] object to be set up.
- A - [IN] ParCSR matrix used to construct the solver/preconditioner.
- $\mathbf{b}$ - right-hand-side of the linear system to be solved (Ignored by this function).
- $\mathbf{x}$ - approximate solution of the linear system to be solved (Ignored by this function).

HYPRE_Int HYPRE_ILUSolve(HYPRE_Solver solver, HYPRE_ParCSRMatrix A, HYPRE_ParVector b, HYPRE_ParVector x)
Solve the system or apply ILU as a preconditioner.
If used as a preconditioner, this function should be passed to the iterative solver SetPrecond function.

## Parameters

- solver - [IN] solver or preconditioner object to be applied.
- A - [IN] ParCSR matrix, matrix of the linear system to be solved
- $\mathbf{b}-[\mathrm{IN}]$ right hand side of the linear system to be solved
- $\mathbf{x}$ - [OUT] approximated solution of the linear system to be solved

HYPRE_Int HYPRE_ILUSetMaxIter(HYPRE_Solver solver, HYPRE_Int max_iter)
(Optional) Set maximum number of iterations if used as a solver.
Set this to 1 if ILU is used as a preconditioner. The default is 20 .
HYPRE_Int HYPRE_ILUSetIterativeSetupType(HYPRE_Solver solver, HYPRE_Int iter_setup_type)
(Optional) Set the algorithm type to compute the ILU factorization.
Options are:

- 0 : Non-iterative algorithm (default)
- 1: Asynchronous with in-place storage
- 2 : Asynchronous with explicit storage splitting
- 3 : Synchronous with explicit storage splitting
- 4 : Semi-synchronous with explicit storage splitting

Note: Iterative ILU is available only for zero fill-in and it depends on rocSPARSE.
HYPRE_Int HYPRE_ILUSetIterativeSetupOption(HYPRE_Solver solver, HYPRE_Int iter_setup_option)
(Optional) Set the compute option for iterative ILU in an additive fashion, i.e. ; multiple options can be turned on by summing their respective numeric codes as given below:

- 2: Use stopping tolerance to finish the algorithm
- 4: Compute correction norms
- 8: Compute residual norms
- 16: Save convergence history
- 32: Use rocSPARSE's internal COO format

The iterative ILU algorithm can terminate based on the maximum number of iterations (default) or a target tolerance (option 2). In the tolerance-based case, the max. number of iterations is still used to terminate the algorithm in case it does not converge to the requested tolerance. In addition, the tolerance-based mode uses residual norms by default (option 8). To use correction norms instead, enable option 4. Lastly, the convergence history for computing the triangular factors can be saved and printed out by enabling option 16.

Note: Iterative ILU is available only for zero fill-in, and it depends on rocSPARSE.

HYPRE_Int HYPRE_ILUSetIterativeSetupMaxIter (HYPRE_Solver solver, HYPRE_Int iter_setup_max_iter)
(Optional) Set the max.
number of iterations for the iterative ILU algorithm.
Note: Iterative ILU is available only for zero fill-in and it depends on rocSPARSE.
HYPRE_Int HYPRE_ILUSetIterativeSetupTolerance(HYPRE_Solver solver, HYPRE_Real iter_setup_tolerance)
(Optional) Set the stop tolerance for the iterative ILU algorithm.
Note: Iterative ILU is available only for zero fill-in and it depends on rocSPARSE.
HYPRE_Int HYPRE_ILUSetTriSolve(HYPRE_Solver solver, HYPRE_Int tri_solve)
(Optional) Set triangular solver type.
Options are:

- 0 : iterative
- 1 : direct (default)

HYPRE_Int HYPRE_ILUSetLowerJacobiIters(HYPRE_Solver solver, HYPRE_Int lower_jacobi_iterations)
(Optional) Set number of lower Jacobi iterations for the triangular L solves Set this to integer $>0$ when using iterative tri_solve (0).

The default is 5 iterations.
HYPRE_Int HYPRE_ILUSetUpperJacobiIters(HYPRE_Solver solver, HYPRE_Int upper_jacobi_iterations)
(Optional) Set number of upper Jacobi iterations for the triangular $U$ solves Set this to integer $>0$ when using iterative tri_solve (0).

The default is 5 iterations.
HYPRE_Int HYPRE_ILUSetTol (HYPRE_Solver solver, HYPRE_Real tol)
(Optional) Set the convergence tolerance for ILU.
Use tol $=0.0$ if ILU is used as a preconditioner. The default is 1.e-7.
HYPRE_Int HYPRE_ILUSetLevel0fFill(HYPRE_Solver solver, HYPRE_Int lfil)
(Optional) Set the level of fill $k$, for level-based $\operatorname{ILU}(\mathrm{k})$ The default is 0 (for ILU(0)).
HYPRE_Int HYPRE_ILUSetMaxNnzPerRow (HYPRE_Solver solver, HYPRE_Int nzmax)
(Optional) Set the max non-zeros per row in L and U factors (for ILUT) The default is 1000 .
HYPRE_Int HYPRE_ILUSetDropThreshold(HYPRE_Solver solver, HYPRE_Real threshold)
(Optional) Set the threshold for dropping in L and U factors (for ILUT).
Any fill-in less than this threshold is dropped in the factorization. The default is $1.0 \mathrm{e}-2$.
HYPRE_Int HYPRE_ILUSetDropThresholdArray (HYPRE_Solver solver, HYPRE_Real *threshold)
(Optional) Set the array of thresholds for dropping in ILUT.
B, E, and F correspond to upper left, lower left and upper right of $2 \times 2$ block decomposition respectively. Any fill-in less than threshold is dropped in the factorization.

- threshold[0] : threshold for matrix B.
- threshold[1] : threshold for matrix E and F.
- threshold[2] : threshold for matrix S (Schur Complement). The default is $1.0 \mathrm{e}-2$.

HYPRE_Int HYPRE_ILUSetNSHDropThreshold(HYPRE_Solver solver, HYPRE_Real threshold)
(Optional) Set the threshold for dropping in Newton-Schulz-Hotelling iteration (NSH-ILU).
Any entries less than this threshold are dropped when forming the approximate inverse matrix. The default is $1.0 \mathrm{e}-2$.

HYPRE_Int HYPRE_ILUSetNSHDropThresholdArray (HYPRE_Solver solver, HYPRE_Real *threshold)
(Optional) Set the array of thresholds for dropping in Newton-Schulz-Hotelling iteration (for NSH-ILU).
Any fill-in less than thresholds is dropped when forming the approximate inverse matrix.

- threshold[0] : threshold for Minimal Residual iteration (initial guess for NSH).
- threshold[1] : threshold for Newton-Schulz-Hotelling iteration.

The default is $1.0 \mathrm{e}-2$.
HYPRE_Int HYPRE_ILUSetSchurMaxIter (HYPRE_Solver solver, HYPRE_Int ss_max_iter)
(Optional) Set maximum number of iterations for Schur System Solve.
For GMRES-ILU, this is the maximum number of iterations for GMRES. The Krylov dimension for GMRES is set equal to this value to avoid restart. For NSH-ILU, this is the maximum number of iterations for NSH solve. The default is 5 .

HYPRE_Int HYPRE_ILUSetType(HYPRE_Solver solver, HYPRE_Int ilu_type)
Set the type of ILU factorization.
Options for ilu_type are:

- 0 : BJ with $\operatorname{ILU}(\mathrm{k})($ default, with $\mathrm{k}=0)$
- 1: BJ with ILUT
- 10 : GMRES with ILU(k)
- 11 : GMRES with ILUT
- 20 : NSH with ILU(k)
- 21 : NSH with ILUT
- 30 : RAS with ILU(k)
- 31 : RAS with ILUT
- 40 : (nonsymmetric permutation) DDPQ-GMRES with ILU(k)
- 41 : (nonsymmetric permutation) DDPQ-GMRES with ILUT
- 50 : GMRES with RAP-ILU(0) using MILU(0) for P

HYPRE_Int HYPRE_ILUSetLocalReordering (HYPRE_Solver solver, HYPRE_Int reordering_type)
Set the type of reordering for the local matrix.
Options for reordering_type are:

- 0 : No reordering
- 1 : RCM (default)

HYPRE_Int HYPRE_ILUSetPrintLevel (HYPRE_Solver solver, HYPRE_Int print_level)
(Optional) Set the print level to print setup and solve information.

- 0 : no printout (default)
- 1 : print setup information
- 2 : print solve information
- 3 : print both setup and solve information

HYPRE_Int HYPRE_ILUSetLogging(HYPRE_Solver solver, HYPRE_Int logging)
(Optional) Requests logging of solver diagnostics.
Requests additional computations for diagnostic and similar data to be logged by the user. Default is 0 , do nothing. The latest residual will be available if logging $>1$.
HYPRE_Int HYPRE_ILUGetNumIterations (HYPRE_Solver solver, HYPRE_Int *num_iterations)
(Optional) Return the number of ILU iterations.
HYPRE_Int HYPRE_ILUGetFinalRelativeResidualNorm(HYPRE_Solver solver, HYPRE_Real *res_norm)
(Optional) Return the norm of the final relative residual.
HYPRE_ParCSRMatrix GenerateLaplacian(MPI_Comm comm, HYPRE_BigInt nx, HYPRE_BigInt ny, HYPRE_BigInt nz, HYPRE_Int P, HYPRE_Int Q, HYPRE_Int R, HYPRE_Int p, HYPRE_Int q, HYPRE_Int r, HYPRE_Real *value)

HYPRE_ParCSRMatrix GenerateLaplacian27pt(MPI_Comm comm, HYPRE_BigInt nx, HYPRE_BigInt ny, HYPRE_BigInt nz, HYPRE_Int P, HYPRE_Int Q, HYPRE_Int R, HYPRE_Int p, HYPRE_Int q, HYPRE_Int r, HYPRE_Real *value)

HYPRE_ParCSRMatrix GenerateLaplacian9pt(MPI_Comm comm, HYPRE_BigInt nx, HYPRE_BigInt ny, HYPRE_Int P, HYPRE_Int Q, HYPRE_Int p, HYPRE_Int q, HYPRE_Real *value)

HYPRE_ParCSRMatrix GenerateDifConv(MPI_Comm comm, HYPRE_BigInt nx, HYPRE_BigInt ny, HYPRE_BigInt nz, HYPRE_Int P, HYPRE_Int Q, HYPRE_Int R, HYPRE_Int p, HYPRE_Int q, HYPRE_Int r, HYPRE_Real *value)

HYPRE_ParCSRMatrix GenerateRotate7pt (MPI_Comm comm, HYPRE_BigInt nx, HYPRE_BigInt ny, HYPRE_Int P, HYPRE_Int Q, HYPRE_Int p, HYPRE_Int q, HYPRE_Real alpha, HYPRE_Real eps)

HYPRE_ParCSRMatrix GenerateVarDifConv(MPI_Comm comm, HYPRE_BigInt nx, HYPRE_BigInt ny, HYPRE_BigInt nz, HYPRE_Int P, HYPRE_Int Q, HYPRE_Int R, HYPRE_Int p, HYPRE_Int q, HYPRE_Int r, HYPRE_Real eps, HYPRE_ParVector *rhs_ptr)

HYPRE_ParCSRMatrix GenerateRSVarDifConv(MPI_Comm comm, HYPRE_BigInt nx, HYPRE_BigInt ny, HYPRE_BigInt nz, HYPRE_Int P, HYPRE_Int Q, HYPRE_Int R, HYPRE_Int p, HYPRE_Int q, HYPRE_Int r, HYPRE_Real eps, HYPRE_ParVector *rhs_ptr, HYPRE_Int type)
float *hypre_GenerateCoordinates(MPI_Comm comm, HYPRE_BigInt nx, HYPRE_BigInt ny, HYPRE_BigInt nz, HYPRE_Int P, HYPRE_Int Q, HYPRE_Int R, HYPRE_Int p, HYPRE_Int q, HYPRE_Int r, HYPRE_Int coorddim)

HYPRE_Int HYPRE_BoomerAMGSetPostInterpType(HYPRE_Solver solver, HYPRE_Int post_interp_type)
(Optional) Switches on use of Jacobi interpolation after computing an original interpolation
HYPRE_Int HYPRE_BoomerAMGSetJacobiTruncThreshold(HYPRE_Solver solver, HYPRE_Real jacobi_trunc_threshold)
(Optional) Sets a truncation threshold for Jacobi interpolation.
HYPRE_Int HYPRE_BoomerAMGSetNumCRRelaxSteps(HYPRE_Solver solver, HYPRE_Int num_CR_relax_steps)
(Optional) Defines the number of relaxation steps for CR The default is 2 .
HYPRE_Int HYPRE_BoomerAMGSetCRRate(HYPRE_Solver solver, HYPRE_Real CR_rate)
(Optional) Defines convergence rate for CR The default is 0.7 .
HYPRE_Int HYPRE_BoomerAMGSetCRStrongTh(HYPRE_Solver solver, HYPRE_Real CR_strong_th)
(Optional) Defines strong threshold for CR The default is 0.0 .
HYPRE_Int HYPRE_BoomerAMGSetCRUseCG(HYPRE_Solver solver, HYPRE_Int CR_use_CG)
(Optional) Defines whether to use CG
HYPRE_Int HYPRE_BoomerAMGSetISType(HYPRE_Solver solver, HYPRE_Int IS_type)
(Optional) Defines the Type of independent set algorithm used for CR

## ParCSR LOBPCG Eigensolver

These routines should be used in conjunction with the generic interface in Eigensolvers.
HYPRE_Int HYPRE_ParCSRSetupInterpreter (mv_InterfaceInterpreter *i)
Load interface interpreter.
Vector part loaded with hypre_ParKrylov functions and multivector part loaded with mv_TempMultiVector functions.

HYPRE_Int HYPRE_ParCSRSetupMatvec (HYPRE_MatvecFunctions *mv)
Load Matvec interpreter with hypre_ParKrylov functions.
HYPRE_Int HYPRE_ParCSRMultiVectorPrint (void *x_, const char *fileName)
void *HYPRE_ParCSRMultiVectorRead(MPI_Comm comm, void *ii_, const char *fileName)

## Functions

HYPRE_Int HYPRE_SchwarzCreate(HYPRE_Solver *solver)
HYPRE_Int HYPRE_SchwarzDestroy (HYPRE_Solver solver)
HYPRE_Int HYPRE_SchwarzSetup(HYPRE_Solver solver, HYPRE_ParCSRMatrix A, HYPRE_ParVector b, HYPRE_ParVector x)

HYPRE_Int HYPRE_SchwarzSolve(HYPRE_Solver solver, HYPRE_ParCSRMatrix A, HYPRE_ParVector b, HYPRE_ParVector x )

```
HYPRE_Int HYPRE_SchwarzSetVariant (HYPRE_Solver solver, HYPRE_Int variant)
HYPRE_Int HYPRE_SchwarzSetOverlap(HYPRE_Solver solver, HYPRE_Int overlap)
HYPRE_Int HYPRE_SchwarzSetDomainType(HYPRE_Solver solver, HYPRE_Int domain_type)
HYPRE_Int HYPRE_SchwarzSetRelaxWeight (HYPRE_Solver solver, HYPRE_Real relax_weight)
HYPRE_Int HYPRE_SchwarzSetDomainStructure(HYPRE_Solver solver, HYPRE_CSRMatrix domain_structure)
HYPRE_Int HYPRE_SchwarzSetNumFunctions(HYPRE_Solver solver, HYPRE_Int num_functions)
HYPRE_Int HYPRE_SchwarzSetDofFunc (HYPRE_Solver solver, HYPRE_Int *dof_func)
HYPRE_Int HYPRE_SchwarzSetNonSymm(HYPRE_Solver solver, HYPRE_Int use_nonsymm)
HYPRE_Int HYPRE_ParCSRCGNRCreate(MPI_Comm comm, HYPRE_Solver *solver)
HYPRE_Int HYPRE_ParCSRCGNRDestroy(HYPRE_Solver solver)
HYPRE_Int HYPRE_ParCSRCGNRSetup (HYPRE_Solver solver, HYPRE_ParCSRMatrix A, HYPRE_ParVector b, HYPRE_ParVector x)
HYPRE_Int HYPRE_ParCSRCGNRSolve(HYPRE_Solver solver, HYPRE_ParCSRMatrix A, HYPRE_ParVector b, HYPRE_ParVector x)
HYPRE_Int HYPRE_ParCSRCGNRSetTol (HYPRE_Solver solver, HYPRE_Real tol)
HYPRE_Int HYPRE_ParCSRCGNRSetMinIter(HYPRE_Solver solver, HYPRE_Int min_iter)
HYPRE_Int HYPRE_ParCSRCGNRSetMaxIter (HYPRE_Solver solver, HYPRE_Int max_iter)
HYPRE_Int HYPRE_ParCSRCGNRSetStopCrit(HYPRE_Solver solver, HYPRE_Int stop_crit)
HYPRE_Int HYPRE_ParCSRCGNRSetPrecond(HYPRE_Solver solver, HYPRE_PtrToParSolverFcn precond, HYPRE_PtrToParSolverFcn precondT, HYPRE_PtrToParSolverFcn precond_setup, HYPRE_Solver precond_solver)
HYPRE_Int HYPRE_ParCSRCGNRGetPrecond(HYPRE_Solver solver, HYPRE_Solver *precond_data)
HYPRE_Int HYPRE_ParCSRCGNRSetLogging(HYPRE_Solver solver, HYPRE_Int logging)
HYPRE_Int HYPRE_ParCSRCGNRGetNumIterations (HYPRE_Solver solver, HYPRE_Int *num_iterations)
HYPRE_Int HYPRE_ParCSRCGNRGetFinalRelativeResidualNorm(HYPRE_Solver solver, HYPRE_Real *norm)
```


### 8.7 Krylov Solvers

## group KrylovSolvers

A basic interface for Krylov solvers.
These solvers support many of the matrix/vector storage schemes in hypre. They should be used in conjunction with the storage-specific interfaces, particularly the specific Create() and Destroy() functions.

## Krylov Solvers

typedef HYPRE_Int (*HYPRE_PtrToModifyPCFcn)(HYPRE_Solver, HYPRE_Int, HYPRE_Real)

## HYPRE_MODIFYPC

## PCG Solver

HYPRE_Int HYPRE_PCGSetup(HYPRE_Solver solver, HYPRE_Matrix A, HYPRE_Vector b, HYPRE_Vector x)

Prepare to solve the system.
The coefficient data in $b$ and $x$ is ignored here, but information about the layout of the data may be used.
HYPRE_Int HYPRE_PCGSolve(HYPRE_Solver solver, HYPRE_Matrix A, HYPRE_Vector b, HYPRE_Vector x)

Solve the system.
HYPRE_Int HYPRE_PCGSetTol(HYPRE_Solver solver, HYPRE_Real tol)
(Optional) Set the relative convergence tolerance.
HYPRE_Int HYPRE_PCGSetAbsoluteTol (HYPRE_Solver solver, HYPRE_Real a_tol)
(Optional) Set the absolute convergence tolerance (default is 0 ).
If one desires the convergence test to check the absolute convergence tolerance only, then set the relative convergence tolerance to 0.0 . (The default convergence test is $\langle C * r, r>\leq \max$ (relative _tolerance $\left.{ }^{2} *<C * b, b\right\rangle$, absolute _tolerance ${ }^{2}$ ).)

HYPRE_Int HYPRE_PCGSetResidualTol(HYPRE_Solver solver, HYPRE_Real rtol)
(Optional) Set a residual-based convergence tolerance which checks if $\left\|r_{\text {old }}-r_{n e w}\right\|<r t o l\|b\|$.
This is useful when trying to converge to very low relative and/or absolute tolerances, in order to bail-out before roundoff errors affect the approximation.

HYPRE_Int HYPRE_PCGSetAbsoluteTolFactor(HYPRE_Solver solver, HYPRE_Real abstolf)
HYPRE_Int HYPRE_PCGSetConvergenceFactorTol (HYPRE_Solver solver, HYPRE_Real cf_tol)
HYPRE_Int HYPRE_PCGSetStopCrit(HYPRE_Solver solver, HYPRE_Int stop_crit)
HYPRE_Int HYPRE_PCGSetMaxIter (HYPRE_Solver solver, HYPRE_Int max_iter)
(Optional) Set maximum number of iterations.
HYPRE_Int HYPRE_PCGSetTwoNorm(HYPRE_Solver solver, HYPRE_Int two_norm)
(Optional) Use the two-norm in stopping criteria.
HYPRE_Int HYPRE_PCGSetRelChange(HYPRE_Solver solver, HYPRE_Int rel_change)
(Optional) Additionally require that the relative difference in successive iterates be small.
HYPRE_Int HYPRE_PCGSetRecomputeResidual (HYPRE_Solver solver, HYPRE_Int recompute_residual)
(Optional) Recompute the residual at the end to double-check convergence.
HYPRE_Int HYPRE_PCGSetRecomputeResidualP(HYPRE_Solver solver, HYPRE_Int recompute_residual_p)
(Optional) Periodically recompute the residual while iterating.

HYPRE_Int HYPRE_PCGSetFlex (HYPRE_Solver solver, HYPRE_Int flex)
(Optional) Setting this to 1 allows use of Polak-Ribiere Method (flexible) this incrceases robustness, but adds an additional dot product per iteration

HYPRE_Int HYPRE_PCGSetSkipBreak(HYPRE_Solver solver, HYPRE_Int skip_break)
(Optional) Skips subnormal alpha, gamma and iprod values in CG.
If set to 0 (default): will break if values are below HYPRE_REAL_MIN If set to 1 : will break if values are below HYPRE_REAL_TRUE_MIN (requires C11 minimal or will check to HYPRE_REAL_MIN) If set to 2 : will break if values are $<=0$. If set to 3 or larger: will not break at all

HYPRE_Int HYPRE_PCGSetPrecond(HYPRE_Solver solver, HYPRE_PtrToSolverFcn precond, HYPRE_PtrToSolverFen precond_setup, HYPRE_Solver precond_solver)
(Optional) Set the preconditioner to use.
HYPRE_Int HYPRE_PCGSetPreconditioner(HYPRE_Solver solver, HYPRE_Solver precond) (Optional) Set the preconditioner to use in a generic fashion.
This function does not require explicit input of the setup and solve pointers of the preconditioner object. Instead, it automatically extracts this information from the aforementioned object.
HYPRE_Int HYPRE_PCGSetLogging(HYPRE_Solver solver, HYPRE_Int logging)
(Optional) Set the amount of logging to do.
HYPRE_Int HYPRE_PCGSetPrintLevel (HYPRE_Solver solver, HYPRE_Int level)
(Optional) Set the amount of printing to do to the screen.
HYPRE_Int HYPRE_PCGGetNumIterations(HYPRE_Solver solver, HYPRE_Int *num_iterations)
Return the number of iterations taken.
HYPRE_Int HYPRE_PCGGetFinalRelativeResidualNorm(HYPRE_Solver solver, HYPRE_Real *norm)
Return the norm of the final relative residual.
HYPRE_Int HYPRE_PCGGetResidual (HYPRE_Solver solver, void *residual)
Return the residual.
HYPRE_Int HYPRE_PCGGetTol(HYPRE_Solver solver, HYPRE_Real *tol)
HYPRE_Int HYPRE_PCGGetResidualTol(HYPRE_Solver solver, HYPRE_Real *rtol)
HYPRE_Int HYPRE_PCGGetAbsoluteTolFactor (HYPRE_Solver solver, HYPRE_Real *abstolf)
HYPRE_Int HYPRE_PCGGetConvergenceFactorTol (HYPRE_Solver solver, HYPRE_Real *cf_tol)
HYPRE_Int HYPRE_PCGGetStopCrit(HYPRE_Solver solver, HYPRE_Int *stop_crit)
HYPRE_Int HYPRE_PCGGetMaxIter (HYPRE_Solver solver, HYPRE_Int *max_iter)
HYPRE_Int HYPRE_PCGGetTwoNorm(HYPRE_Solver solver, HYPRE_Int *two_norm)
HYPRE_Int HYPRE_PCGGetRelChange(HYPRE_Solver solver, HYPRE_Int *rel_change)
HYPRE_Int HYPRE_PCGGetSkipBreak(HYPRE_Solver solver, HYPRE_Int *skip_break)
HYPRE_Int HYPRE_PCGGetFlex (HYPRE_Solver solver, HYPRE_Int *flex)
HYPRE_Int HYPRE_PCGGetPrecond(HYPRE_Solver solver, HYPRE_Solver *precond_data_ptr)

HYPRE_Int HYPRE_PCGGetLogging(HYPRE_Solver solver, HYPRE_Int *level)
HYPRE_Int HYPRE_PCGGetPrintLevel (HYPRE_Solver solver, HYPRE_Int *level)
HYPRE_Int HYPRE_PCGGetConverged(HYPRE_Solver solver, HYPRE_Int * converged)

## GMRES Solver

HYPRE_Int HYPRE_GMRESSetup(HYPRE_Solver solver, HYPRE_Matrix A, HYPRE_Vector b, HYPRE_Vector x )
Prepare to solve the system.
The coefficient data in $b$ and $x$ is ignored here, but information about the layout of the data may be used.
HYPRE_Int HYPRE_GMRESSolve(HYPRE_Solver solver, HYPRE_Matrix A, HYPRE_Vector b, HYPRE_Vector $x$ )
Solve the system.
HYPRE_Int HYPRE_GMRESSetTol (HYPRE_Solver solver, HYPRE_Real tol)
(Optional) Set the relative convergence tolerance.
HYPRE_Int HYPRE_GMRESSetAbsoluteTol (HYPRE_Solver solver, HYPRE_Real a_tol)
(Optional) Set the absolute convergence tolerance (default is 0 ).
If one desires the convergence test to check the absolute convergence tolerance only, then set the relative convergence tolerance to 0.0 . (The convergence test is $\|r\| \leq \max$ (relative _tolerance $*\|b\|$, absolute _tolerance).)
HYPRE_Int HYPRE_GMRESSetConvergenceFactorTol (HYPRE_Solver solver, HYPRE_Real cf_tol)
HYPRE_Int HYPRE_GMRESSetStopCrit(HYPRE_Solver solver, HYPRE_Int stop_crit)
HYPRE_Int HYPRE_GMRESSetMinIter(HYPRE_Solver solver, HYPRE_Int min_iter)
HYPRE_Int HYPRE_GMRESSetMaxIter(HYPRE_Solver solver, HYPRE_Int max_iter) (Optional) Set maximum number of iterations.

HYPRE_Int HYPRE_GMRESSetKDim(HYPRE_Solver solver, HYPRE_Int k_dim)
(Optional) Set the maximum size of the Krylov space.
HYPRE_Int HYPRE_GMRESSetRelChange(HYPRE_Solver solver, HYPRE_Int rel_change)
(Optional) Additionally require that the relative difference in successive iterates be small.
HYPRE_Int HYPRE_GMRESSetSkipRealResidualCheck(HYPRE_Solver solver, HYPRE_Int skip_real_r_check)
(Optional) By default, hypre checks for convergence by evaluating the actual residual before returnig from GMRES (with restart if the true residual does not indicate convergence).

This option allows users to skip the evaluation and the check of the actual residual for badly conditioned problems where restart is not expected to be beneficial.

HYPRE_Int HYPRE_GMRESSetPrecond(HYPRE_Solver solver, HYPRE_PtrToSolverFcn precond, HYPRE_PtrToSolverFcn precond_setup, HYPRE_Solver precond_solver)
(Optional) Set the preconditioner to use.

HYPRE_Int HYPRE_GMRESSetLogging(HYPRE_Solver solver, HYPRE_Int logging) (Optional) Set the amount of logging to do.

HYPRE_Int HYPRE_GMRESSetPrintLevel(HYPRE_Solver solver, HYPRE_Int level) (Optional) Set the amount of printing to do to the screen.

HYPRE_Int HYPRE_GMRESGetNumIterations(HYPRE_Solver solver, HYPRE_Int *num_iterations) Return the number of iterations taken.

HYPRE_Int HYPRE_GMRESGetFinalRelativeResidualNorm(HYPRE_Solver solver, HYPRE_Real *norm)

Return the norm of the final relative residual.
HYPRE_Int HYPRE_GMRESGetResidual (HYPRE_Solver solver, void *residual)
Return the residual.
HYPRE_Int HYPRE_GMRESGetSkipRealResidualCheck(HYPRE_Solver solver, HYPRE_Int
*skip_real_r_check)
HYPRE_Int HYPRE_GMRESGetTol (HYPRE_Solver solver, HYPRE_Real *tol)
HYPRE_Int HYPRE_GMRESGetAbsoluteTol (HYPRE_Solver solver, HYPRE_Real *tol)
HYPRE_Int HYPRE_GMRESGetConvergenceFactorTol (HYPRE_Solver solver, HYPRE_Real *cf_tol)
HYPRE_Int HYPRE_GMRESGetStopCrit (HYPRE_Solver solver, HYPRE_Int *stop_crit)
HYPRE_Int HYPRE_GMRESGetMinIter(HYPRE_Solver solver, HYPRE_Int *min_iter)
HYPRE_Int HYPRE_GMRESGetMaxIter (HYPRE_Solver solver, HYPRE_Int *max_iter)
HYPRE_Int HYPRE_GMRESGetKDim(HYPRE_Solver solver, HYPRE_Int *k_dim)
HYPRE_Int HYPRE_GMRESGetRelChange(HYPRE_Solver solver, HYPRE_Int *rel_change)
HYPRE_Int HYPRE_GMRESGetPrecond (HYPRE_Solver solver, HYPRE_Solver *precond_data_ptr)
HYPRE_Int HYPRE_GMRESGetLogging(HYPRE_Solver solver, HYPRE_Int * level)
HYPRE_Int HYPRE_GMRESGetPrintLevel (HYPRE_Solver solver, HYPRE_Int *level)
HYPRE_Int HYPRE_GMRESGetConverged(HYPRE_Solver solver, HYPRE_Int * converged)

FlexGMRES Solver

HYPRE_Int HYPRE_FlexGMRESSetup(HYPRE_Solver solver, HYPRE_Matrix A, HYPRE_Vector b, HYPRE_Vector x )
Prepare to solve the system.
The coefficient data in $b$ and $x$ is ignored here, but information about the layout of the data may be used.
HYPRE_Int HYPRE_FlexGMRESSolve(HYPRE_Solver solver, HYPRE_Matrix A, HYPRE_Vector b, HYPRE_Vector x )
Solve the system.
HYPRE_Int HYPRE_FlexGMRESSetTol (HYPRE_Solver solver, HYPRE_Real tol)
(Optional) Set the convergence tolerance.

HYPRE_Int HYPRE_FlexGMRESSetAbsoluteTol (HYPRE_Solver solver, HYPRE_Real a_tol)
(Optional) Set the absolute convergence tolerance (default is 0 ).
If one desires the convergence test to check the absolute convergence tolerance only, then set the relative convergence tolerance to 0.0 . (The convergence test is $\|r\| \leq \max$ (relative _tolerance $*\|b\|$, absolute _tolerance).)

HYPRE_Int HYPRE_FlexGMRESSetConvergenceFactorTol (HYPRE_Solver solver, HYPRE_Real cf_tol)
HYPRE_Int HYPRE_FlexGMRESSetMinIter(HYPRE_Solver solver, HYPRE_Int min_iter)
HYPRE_Int HYPRE_FlexGMRESSetMaxIter(HYPRE_Solver solver, HYPRE_Int max_iter) (Optional) Set maximum number of iterations.

HYPRE_Int HYPRE_FlexGMRESSetKDim(HYPRE_Solver solver, HYPRE_Int k_dim) (Optional) Set the maximum size of the Krylov space.

HYPRE_Int HYPRE_FlexGMRESSetPrecond(HYPRE_Solver solver, HYPRE_PtrToSolverFcn precond, HYPRE_PtrToSolverFen precond_setup, HYPRE_Solver precond_solver)
(Optional) Set the preconditioner to use.
HYPRE_Int HYPRE_FlexGMRESSetLogging (HYPRE_Solver solver, HYPRE_Int logging)
(Optional) Set the amount of logging to do.
HYPRE_Int HYPRE_FlexGMRESSetPrintLevel (HYPRE_Solver solver, HYPRE_Int level) (Optional) Set the amount of printing to do to the screen.

HYPRE_Int HYPRE_FlexGMRESGetNumIterations(HYPRE_Solver solver, HYPRE_Int *num_iterations) Return the number of iterations taken.

HYPRE_Int HYPRE_FlexGMRESGetFinalRelativeResidualNorm(HYPRE_Solver solver, HYPRE_Real *norm)
Return the norm of the final relative residual.
HYPRE_Int HYPRE_FlexGMRESGetResidual (HYPRE_Solver solver, void *residual) Return the residual.

HYPRE_Int HYPRE_FlexGMRESGetTol (HYPRE_Solver solver, HYPRE_Real *tol)
HYPRE_Int HYPRE_FlexGMRESGetConvergenceFactorTol (HYPRE_Solver solver, HYPRE_Real *cf_tol)
HYPRE_Int HYPRE_FlexGMRESGetStopCrit(HYPRE_Solver solver, HYPRE_Int *stop_crit)
HYPRE_Int HYPRE_FlexGMRESGetMinIter (HYPRE_Solver solver, HYPRE_Int *min_iter)
HYPRE_Int HYPRE_FlexGMRESGetMaxIter (HYPRE_Solver solver, HYPRE_Int *max_iter)
HYPRE_Int HYPRE_FlexGMRESGetKDim(HYPRE_Solver solver, HYPRE_Int *k_dim)
HYPRE_Int HYPRE_FlexGMRESGetPrecond(HYPRE_Solver solver, HYPRE_Solver *precond_data_ptr)
HYPRE_Int HYPRE_FlexGMRESGetLogging(HYPRE_Solver solver, HYPRE_Int *level)
HYPRE_Int HYPRE_FlexGMRESGetPrintLevel (HYPRE_Solver solver, HYPRE_Int *level)
HYPRE_Int HYPRE_FlexGMRESGetConverged(HYPRE_Solver solver, HYPRE_Int *converged)
HYPRE_Int HYPRE_FlexGMRESSetModifyPC(HYPRE_Solver solver, HYPRE_PtrToModifyPCFcn modify_pc)
(Optional) Set a user-defined function to modify solve-time preconditioner attributes.

## LGMRES Solver

HYPRE_Int HYPRE_LGMRESSetup(HYPRE_Solver solver, HYPRE_Matrix A, HYPRE_Vector b, HYPRE_Vector x )

Prepare to solve the system.
The coefficient data in $b$ and $x$ is ignored here, but information about the layout of the data may be used.
HYPRE_Int HYPRE_LGMRESSolve(HYPRE_Solver solver, HYPRE_Matrix A, HYPRE_Vector b, HYPRE_Vector x)
Solve the system.

Details on LGMRES may be found in A. H. Baker, E.R. Jessup, and T.A. Manteuffel, "A technique for accelerating the convergence of restarted GMRES." SIAM Journal on Matrix Analysis and Applications, 26 (2005), pp. 962-984. LGMRES(m,k) in the paper corresponds to LGMRES(Kdim+AugDim, AugDim).

HYPRE_Int HYPRE_LGMRESSetTol (HYPRE_Solver solver, HYPRE_Real tol)
(Optional) Set the convergence tolerance.
HYPRE_Int HYPRE_LGMRESSetAbsoluteTol (HYPRE_Solver solver, HYPRE_Real a_tol)
(Optional) Set the absolute convergence tolerance (default is 0 ).
If one desires the convergence test to check the absolute convergence tolerance only, then set the relative convergence tolerance to 0.0 . (The convergence test is $\|r\| \leq \max$ (relative _tolerance $*\|b\|$, absolute _tolerance).)

HYPRE_Int HYPRE_LGMRESSetConvergenceFactorTol (HYPRE_Solver solver, HYPRE_Real cf_tol)
HYPRE_Int HYPRE_LGMRESSetMinIter (HYPRE_Solver solver, HYPRE_Int min_iter)
HYPRE_Int HYPRE_LGMRESSetMaxIter (HYPRE_Solver solver, HYPRE_Int max_iter) (Optional) Set maximum number of iterations.

HYPRE_Int HYPRE_LGMRESSetKDim(HYPRE_Solver solver, HYPRE_Int k_dim)
(Optional) Set the maximum size of the approximation space (includes the augmentation vectors).
HYPRE_Int HYPRE_LGMRESSetAugDim(HYPRE_Solver solver, HYPRE_Int aug_dim)
(Optional) Set the number of augmentation vectors (default: 2).
HYPRE_Int HYPRE_LGMRESSetPrecond(HYPRE_Solver solver, HYPRE_PtrToSolverFcn precond, HYPRE_PtrToSolverFen precond_setup, HYPRE_Solver precond_solver)
(Optional) Set the preconditioner to use.
HYPRE_Int HYPRE_LGMRESSetLogging(HYPRE_Solver solver, HYPRE_Int logging)
(Optional) Set the amount of logging to do.
HYPRE_Int HYPRE_LGMRESSetPrintLevel (HYPRE_Solver solver, HYPRE_Int level)
(Optional) Set the amount of printing to do to the screen.
HYPRE_Int HYPRE_LGMRESGetNumIterations (HYPRE_Solver solver, HYPRE_Int *num_iterations)
Return the number of iterations taken.

HYPRE_Int HYPRE_LGMRESGetFinalRelativeResidualNorm(HYPRE_Solver solver, HYPRE_Real *norm)
Return the norm of the final relative residual.
HYPRE_Int HYPRE_LGMRESGetResidual (HYPRE_Solver solver, void *residual)
Return the residual.
HYPRE_Int HYPRE_LGMRESGetTol (HYPRE_Solver solver, HYPRE_Real *tol)
HYPRE_Int HYPRE_LGMRESGetConvergenceFactorTol (HYPRE_Solver solver, HYPRE_Real *cf_tol)
HYPRE_Int HYPRE_LGMRESGetStopCrit (HYPRE_Solver solver, HYPRE_Int *stop_crit)
HYPRE_Int HYPRE_LGMRESGetMinIter (HYPRE_Solver solver, HYPRE_Int *min_iter)
HYPRE_Int HYPRE_LGMRESGetMaxIter (HYPRE_Solver solver, HYPRE_Int *max_iter)
HYPRE_Int HYPRE_LGMRESGetKDim(HYPRE_Solver solver, HYPRE_Int *k_dim)
HYPRE_Int HYPRE_LGMRESGetAugDim(HYPRE_Solver solver, HYPRE_Int *k_dim)
HYPRE_Int HYPRE_LGMRESGetPrecond(HYPRE_Solver solver, HYPRE_Solver *precond_data_ptr)
HYPRE_Int HYPRE_LGMRESGetLogging(HYPRE_Solver solver, HYPRE_Int *level)
HYPRE_Int HYPRE_LGMRESGetPrintLevel (HYPRE_Solver solver, HYPRE_Int *level)
HYPRE_Int HYPRE_LGMRESGetConverged(HYPRE_Solver solver, HYPRE_Int * converged)

## COGMRES Solver

HYPRE_Int HYPRE_COGMRESSetup(HYPRE_Solver solver, HYPRE_Matrix A, HYPRE_Vector b, HYPRE_Vector x )

Prepare to solve the system.
The coefficient data in $b$ and $x$ is ignored here, but information about the layout of the data may be used.
HYPRE_Int HYPRE_COGMRESSolve(HYPRE_Solver solver, HYPRE_Matrix A, HYPRE_Vector b, HYPRE_Vector x )

Solve the system.
HYPRE_Int HYPRE_COGMRESSetTol (HYPRE_Solver solver, HYPRE_Real tol)
(Optional) Set the convergence tolerance.
HYPRE_Int HYPRE_COGMRESSetAbsoluteTol (HYPRE_Solver solver, HYPRE_Real a_tol)
(Optional) Set the absolute convergence tolerance (default is 0 ).
If one desires the convergence test to check the absolute convergence tolerance only, then set the relative convergence tolerance to 0.0 . (The convergence test is $\|r\| \leq \max$ (relative _tolerance $*\|b\|$, absolute _tolerance).)

HYPRE_Int HYPRE_COGMRESSetConvergenceFactorTol (HYPRE_Solver solver, HYPRE_Real cf_tol)
HYPRE_Int HYPRE_COGMRESSetMinIter(HYPRE_Solver solver, HYPRE_Int min_iter)
HYPRE_Int HYPRE_COGMRESSetMaxIter(HYPRE_Solver solver, HYPRE_Int max_iter)
(Optional) Set maximum number of iterations.

HYPRE_Int HYPRE_COGMRESSetKDim(HYPRE_Solver solver, HYPRE_Int k_dim)
(Optional) Set the maximum size of the Krylov space.
HYPRE_Int HYPRE_COGMRESSetUnroll (HYPRE_Solver solver, HYPRE_Int unroll) (Optional) Set number of unrolling in mass funcyions in COGMRES Can be 4 or 8 . Default: no unrolling.

HYPRE_Int HYPRE_COGMRESSetCGS(HYPRE_Solver solver, HYPRE_Int cgs)
(Optional) Set the number of orthogonalizations in COGMRES (at most 2).
HYPRE_Int HYPRE_COGMRESSetPrecond(HYPRE_Solver solver, HYPRE_PtrToSolverFcn precond, HYPRE_PtrToSolverFen precond_setup, HYPRE_Solver precond_solver)
(Optional) Set the preconditioner to use.
HYPRE_Int HYPRE_COGMRESSetLogging(HYPRE_Solver solver, HYPRE_Int logging)
(Optional) Set the amount of logging to do.
HYPRE_Int HYPRE_COGMRESSetPrintLevel (HYPRE_Solver solver, HYPRE_Int level)
(Optional) Set the amount of printing to do to the screen.
HYPRE_Int HYPRE_COGMRESGetNumIterations(HYPRE_Solver solver, HYPRE_Int *num_iterations) Return the number of iterations taken.

HYPRE_Int HYPRE_COGMRESGetFinalRelativeResidualNorm(HYPRE_Solver solver, HYPRE_Real *norm)

Return the norm of the final relative residual.
HYPRE_Int HYPRE_COGMRESGetResidual (HYPRE_Solver solver, void *residual)
Return the residual.
HYPRE_Int HYPRE_COGMRESGetTol (HYPRE_Solver solver, HYPRE_Real *tol)
HYPRE_Int HYPRE_COGMRESGetConvergenceFactorTol (HYPRE_Solver solver, HYPRE_Real *cf_tol)
HYPRE_Int HYPRE_COGMRESGetMinIter(HYPRE_Solver solver, HYPRE_Int *min_iter)
HYPRE_Int HYPRE_COGMRESGetMaxIter (HYPRE_Solver solver, HYPRE_Int *max_iter)
HYPRE_Int HYPRE_COGMRESGetKDim(HYPRE_Solver solver, HYPRE_Int *k_dim)
HYPRE_Int HYPRE_COGMRESGetUnroll (HYPRE_Solver solver, HYPRE_Int *unroll)
HYPRE_Int HYPRE_COGMRESGetCGS (HYPRE_Solver solver, HYPRE_Int *cgs)
HYPRE_Int HYPRE_COGMRESGetPrecond(HYPRE_Solver solver, HYPRE_Solver *precond_data_ptr)
HYPRE_Int HYPRE_COGMRESGetLogging(HYPRE_Solver solver, HYPRE_Int *level)
HYPRE_Int HYPRE_COGMRESGetPrintLevel(HYPRE_Solver solver, HYPRE_Int *level)
HYPRE_Int HYPRE_COGMRESGetConverged(HYPRE_Solver solver, HYPRE_Int *converged)
HYPRE_Int HYPRE_COGMRESSetModifyPC(HYPRE_Solver solver, HYPRE_PtrToModifyPCFcn modify_pc)
(Optional) Set a user-defined function to modify solve-time preconditioner attributes.

## BiCGSTAB Solver

HYPRE_Int HYPRE_BiCGSTABDestroy(HYPRE_Solver solver)
HYPRE_Int HYPRE_BiCGSTABSetup(HYPRE_Solver solver, HYPRE_Matrix A, HYPRE_Vector b, HYPRE_Vector x )
Prepare to solve the system.
The coefficient data in $b$ and $x$ is ignored here, but information about the layout of the data may be used.
HYPRE_Int HYPRE_BiCGSTABSolve(HYPRE_Solver solver, HYPRE_Matrix A, HYPRE_Vector b, HYPRE_Vector x )

Solve the system.
HYPRE_Int HYPRE_BiCGSTABSetTol(HYPRE_Solver solver, HYPRE_Real tol)
(Optional) Set the convergence tolerance.
HYPRE_Int HYPRE_BiCGSTABSetAbsoluteTol (HYPRE_Solver solver, HYPRE_Real a_tol)
(Optional) Set the absolute convergence tolerance (default is 0 ).
If one desires the convergence test to check the absolute convergence tolerance only, then set the relative convergence tolerance to 0.0 . (The convergence test is $\|r\| \leq \max$ (relative _tolerance $*\|b\|$, absolute _tolerance).)

HYPRE_Int HYPRE_BiCGSTABSetConvergenceFactorTol (HYPRE_Solver solver, HYPRE_Real cf_tol)
HYPRE_Int HYPRE_BiCGSTABSetStopCrit(HYPRE_Solver solver, HYPRE_Int stop_crit)
HYPRE_Int HYPRE_BiCGSTABSetMinIter (HYPRE_Solver solver, HYPRE_Int min_iter)
HYPRE_Int HYPRE_BiCGSTABSetMaxIter (HYPRE_Solver solver, HYPRE_Int max_iter) (Optional) Set maximum number of iterations.

HYPRE_Int HYPRE_BiCGSTABSetPrecond(HYPRE_Solver solver, HYPRE_PtrToSolverFcn precond, HYPRE_PtrToSolverFen precond_setup, HYPRE_Solver precond_solver)
(Optional) Set the preconditioner to use.
HYPRE_Int HYPRE_BiCGSTABSetLogging (HYPRE_Solver solver, HYPRE_Int logging)
(Optional) Set the amount of logging to do.
HYPRE_Int HYPRE_BiCGSTABSetPrintLevel (HYPRE_Solver solver, HYPRE_Int level)
(Optional) Set the amount of printing to do to the screen.
HYPRE_Int HYPRE_BiCGSTABGetNumIterations (HYPRE_Solver solver, HYPRE_Int *num_iterations)
Return the number of iterations taken.
HYPRE_Int HYPRE_BiCGSTABGetFinalRelativeResidualNorm(HYPRE_Solver solver, HYPRE_Real *norm)

Return the norm of the final relative residual.
HYPRE_Int HYPRE_BiCGSTABGetResidual(HYPRE_Solver solver, void *residual)
Return the residual.
HYPRE_Int HYPRE_BiCGSTABGetPrecond(HYPRE_Solver solver, HYPRE_Solver *precond_data_ptr)

## CGNR Solver

HYPRE_Int HYPRE_CGNRDestroy(HYPRE_Solver solver)
HYPRE_Int HYPRE_CGNRSetup(HYPRE_Solver solver, HYPRE_Matrix A, HYPRE_Vector b, HYPRE_Vector x )

Prepare to solve the system.
The coefficient data in $b$ and $x$ is ignored here, but information about the layout of the data may be used.
HYPRE_Int HYPRE_CGNRSolve(HYPRE_Solver solver, HYPRE_Matrix A, HYPRE_Vector b, HYPRE_Vector x )

Solve the system.
HYPRE_Int HYPRE_CGNRSetTol(HYPRE_Solver solver, HYPRE_Real tol)
(Optional) Set the convergence tolerance.
HYPRE_Int HYPRE_CGNRSetStopCrit(HYPRE_Solver solver, HYPRE_Int stop_crit)
HYPRE_Int HYPRE_CGNRSetMinIter(HYPRE_Solver solver, HYPRE_Int min_iter)
HYPRE_Int HYPRE_CGNRSetMaxIter(HYPRE_Solver solver, HYPRE_Int max_iter)
(Optional) Set maximum number of iterations.
HYPRE_Int HYPRE_CGNRSetPrecond(HYPRE_Solver solver, HYPRE_PtrToSolverFcn precond, HYPRE_PtrToSolverFcn precondT, HYPRE_PtrToSolverFen precond_setup, HYPRE_Solver precond_solver)
(Optional) Set the preconditioner to use.
Note that the only preconditioner available in hypre for use with CGNR is currently BoomerAMG. It requires to use Jacobi as a smoother without CF smoothing, i.e. relax_type needs to be set to 0 or 7 and relax_order needs to be set to 0 by the user, since these are not default values. It can be used with a relaxation weight for Jacobi, which can significantly improve convergence.

HYPRE_Int HYPRE_CGNRSetLogging(HYPRE_Solver solver, HYPRE_Int logging)
(Optional) Set the amount of logging to do.
HYPRE_Int HYPRE_CGNRGetNumIterations(HYPRE_Solver solver, HYPRE_Int *num_iterations)
Return the number of iterations taken.
HYPRE_Int HYPRE_CGNRGetFinalRelativeResidualNorm(HYPRE_Solver solver, HYPRE_Real *norm) Return the norm of the final relative residual.

HYPRE_Int HYPRE_CGNRGetPrecond(HYPRE_Solver solver, HYPRE_Solver *precond_data_ptr)

### 8.8 Eigensolvers

## group Eigensolvers

A basic interface for eigensolvers
These eigensolvers support many of the matrix/vector storage schemes in hypre. They should be used in conjunction with the storage-specific interfaces.

## LOBPCG Eigensolver

HYPRE_Int HYPRE_LOBPCGCreate (mv_InterfaceInterpreter *interpreter, HYPRE_MatvecFunctions *mvfunctions, HYPRE_Solver *solver)

LOBPCG constructor.
HYPRE_Int HYPRE_LOBPCGDestroy (HYPRE_Solver solver)
LOBPCG destructor.
HYPRE_Int HYPRE_LOBPCGSetPrecond(HYPRE_Solver solver, HYPRE_PtrToSolverFcn precond, HYPRE_PtrToSolverFen precond_setup, HYPRE_Solver precond_solver)
(Optional) Set the preconditioner to use.
If not called, preconditioning is not used.
HYPRE_Int HYPRE_LOBPCGGetPrecond(HYPRE_Solver solver, HYPRE_Solver *precond_data_ptr)
HYPRE_Int HYPRE_LOBPCGSetup(HYPRE_Solver solver, HYPRE_Matrix A, HYPRE_Vector b, HYPRE_Vector $x$ )
Set up $A$ and the preconditioner (if there is one).
HYPRE_Int HYPRE_LOBPCGSetupB(HYPRE_Solver solver, HYPRE_Matrix B, HYPRE_Vector x) (Optional) Set up B.

If not called, $B=I$.
HYPRE_Int HYPRE_LOBPCGSetupT (HYPRE_Solver solver, HYPRE_Matrix T, HYPRE_Vector x) (Optional) Set the preconditioning to be applied to $\mathrm{Tx}=\mathrm{b}$, not $\mathrm{Ax}=\mathrm{b}$.

HYPRE_Int HYPRE_LOBPCGSolve(HYPRE_Solver solver, mv_MultiVectorPtr y, mv_MultiVectorPtr x, HYPRE_Real *lambda)

Solve A x = lambda B x, y'x $=0$.
HYPRE_Int HYPRE_LOBPCGSetTol (HYPRE_Solver solver, HYPRE_Real tol)
(Optional) Set the absolute convergence tolerance.
HYPRE_Int HYPRE_LOBPCGSetRTol (HYPRE_Solver solver, HYPRE_Real tol)
(Optional) Set the relative convergence tolerance.
HYPRE_Int HYPRE_LOBPCGSetMaxIter (HYPRE_Solver solver, HYPRE_Int max_iter)
(Optional) Set maximum number of iterations.
HYPRE_Int HYPRE_LOBPCGSetPrecondUsageMode(HYPRE_Solver solver, HYPRE_Int mode)
Define which initial guess for inner PCG iterations to use: mode $=0$ : use zero initial guess, otherwise use RHS.

HYPRE_Int HYPRE_LOBPCGSetPrintLevel (HYPRE_Solver solver, HYPRE_Int level)
(Optional) Set the amount of printing to do to the screen.
utilities_FortranMatrix *HYPRE_LOBPCGResidualNorms (HYPRE_Solver solver)
utilities_FortranMatrix *HYPRE_LOBPCGResidualNormsHistory (HYPRE_Solver solver)
utilities_FortranMatrix *HYPRE_LOBPCGEigenvaluesHistory (HYPRE_Solver solver)
HYPRE_Int HYPRE_LOBPCGIterations(HYPRE_Solver solver)
void hypre_LOBPCGMultiOperatorB (void *data, void *x, void *y)
void lobpcg_MultiVectorByMultiVector (mv_MultiVectorPtr x, mv_MultiVectorPtr y, utilities_FortranMatrix *xy)

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[^0]:    HYPRE_Int HYPRE_ParCSRCOGMRESSetup(HYPRE_Solver solver, HYPRE_ParCSRMatrix A, HYPRE_ParVector b, HYPRE_ParVector x)

    HYPRE_Int HYPRE_ParCSRCOGMRESSolve(HYPRE_Solver solver, HYPRE_ParCSRMatrix A, HYPRE_ParVector b, HYPRE_ParVector x)

    HYPRE_Int HYPRE_ParCSRCOGMRESSetKDim(HYPRE_Solver solver, HYPRE_Int k_dim)
    HYPRE_Int HYPRE_ParCSRCOGMRESSetUnroll (HYPRE_Solver solver, HYPRE_Int unroll)
    HYPRE_Int HYPRE_ParCSRCOGMRESSetCGS (HYPRE_Solver solver, HYPRE_Int cgs)
    HYPRE_Int HYPRE_ParCSRCOGMRESSetTol (HYPRE_Solver solver, HYPRE_Real tol)
    HYPRE_Int HYPRE_ParCSRCOGMRESSetAbsoluteTol (HYPRE_Solver solver, HYPRE_Real a_tol)
    HYPRE_Int HYPRE_ParCSRCOGMRESSetMinIter(HYPRE_Solver solver, HYPRE_Int min_iter)
    HYPRE_Int HYPRE_ParCSRCOGMRESSetMaxIter(HYPRE_Solver solver, HYPRE_Int max_iter)
    HYPRE_Int HYPRE_ParCSRCOGMRESSetPrecond(HYPRE_Solver solver, HYPRE_PtrToParSolverFcn precond, HYPRE_PtrToParSolverFcn precond_setup, HYPRE_Solver precond_solver)

    HYPRE_Int HYPRE_ParCSRCOGMRESGetPrecond(HYPRE_Solver solver, HYPRE_Solver *precond_data)
    HYPRE_Int HYPRE_ParCSRCOGMRESSetLogging(HYPRE_Solver solver, HYPRE_Int logging)
    HYPRE_Int HYPRE_ParCSRCOGMRESSetPrintLevel (HYPRE_Solver solver, HYPRE_Int print_level)
    HYPRE_Int HYPRE_ParCSRCOGMRESGetNumIterations(HYPRE_Solver solver, HYPRE_Int *num_iterations)

    HYPRE_Int HYPRE_ParCSRCOGMRESGetFinalRelativeResidualNorm(HYPRE_Solver solver, HYPRE_Real *norm)

    HYPRE_Int HYPRE_ParCSRCOGMRESGetResidual (HYPRE_Solver solver, HYPRE_ParVector *residual)
    Returns the residual.

    ## ParCSR FlexGMRES Solver

    These routines should be used in conjunction with the generic interface in Krylov Solvers.
    HYPRE_Int HYPRE_ParCSRFlexGMRESCreate(MPI_Comm comm, HYPRE_Solver *solver)
    Create a solver object.
    HYPRE_Int HYPRE_ParCSRFlexGMRESDestroy (HYPRE_Solver solver)
    Destroy a solver object.
    HYPRE_Int HYPRE_ParCSRFlexGMRESSetup(HYPRE_Solver solver, HYPRE_ParCSRMatrix A, HYPRE_ParVector b, HYPRE_ParVector x)

    HYPRE_Int HYPRE_ParCSRFlexGMRESSolve(HYPRE_Solver solver, HYPRE_ParCSRMatrix A, HYPRE_ParVector b, HYPRE_ParVector x)

    HYPRE_Int HYPRE_ParCSRFlexGMRESSetKDim(HYPRE_Solver solver, HYPRE_Int k_dim)

