FASP User Guide

FASP Developer Team

Contents

\mathbf{C}	ontei	nts	1			
1	Inti	roduction	3			
	1.1	General description	3			
	1.2	Roadmap: from basics to complex applications	3			
	1.3	How to use this guide	4			
	1.4	How to obtain FASP	4			
	1.5	Licensing	6			
	1.6	Building and installing the FASP library and examples	6			
2	B b	orief tutorial	11			
	2.1	Example 1:An AMG solver for the Poisson equation	11			
	2.2	Example 2: Conjugate gradient without preconditioning	17			
	2.3	Example 3: Conjugate gradient with preconditioner	20			
	2.4	How to change default parameters in FASP solver/preconditioner	23			
3	Dat	Data structures and basic usage				
	3.1	Vectors and sparse matrices	29			
	3.2	Block sparse matrices	34			
	3.3	I/O subroutines for sparse matrices	36			
	3.4	Sparse BLAS	39			
	3.5	Iterative methods	40			
	3.6	Geometric multigrid	42			
	3.7	Algebraic multigrid	43			
4	Mo	re advanced features	49			
	4.1	An OpenMP example	49			
	4.2	Predefined constants	50			
	4.3	Debugging and how to enable it	55			

0	9	CONTENTS
Ζ	L	CONTENIS

Bibliography 57

Chapter 1

Introduction

1.1 General description

The Fast Auxiliary Space Preconditioning (FASP) package provides C source files¹ to build a library of iterative solvers and preconditioners for the solution of large scale linear systems of equations. The components of the base FASP library include several ready-to-use, modern, and efficient iterative solvers used in applications ranging from simple examples of discretized scalar partial differential equations (PDEs) to numerical simulations of complex, multicomponent physical systems via the Auxiliary Space Preconditioning framework [19].

The main components of the FASP basic library are:

- Basic linear iterative methods;
- Standard Krylov subspace methods;
- Geometric and Algebraic Multigrid (G/AMG) methods;
- Incomplete factorization methods.

The FASP distribution also includes several examples for solving simple benchmark problems.

1.2 Roadmap: from basics to complex applications

A distinct feature of the FASP software project is that it is an open ended project. It contains a basic kernel of sources and is maintained by a team of developers with the expertise to build efficient solvers for a wide range of complex numerical models.

As typical for an open-source software, the further development of FASP project will also be based on the involvement of the community. While we have our own plans for expanding FASP's capabilities, we also count on the users' input in providing requests for, as well as, contributions to,

¹The C code is in the C99 standard.

the expansion of FASP in different application areas. Our team is ready to provide (or help with) the design and the implementation of efficient solvers based on the FASP kernel to best meet the goals and the requirements of our users.

The FASP software has been successfully used to build efficient solvers for several discretized PDEs and systems of PDEs: general scalar elliptic equations; linear elasticity; Brinkman equation; bi-harmonic equation; Stokes and Navier-Stokes equations; H(curl)/H(div) systems; Maxwell's system. The resulting solvers have been applied in simulations from fluid dynamics, underground water simulation, fluid-structure interactions; Oldryod-B and Johnson-Seglman models; black-oil model in reservoir simulation; magnetohydrodynamics (MHD).

Several of these benchmark problems are included as examples in the open-source distribution, others are under development or have more restrictive licensing.

1.3 How to use this guide

This user's guide describes how to use the existing solvers in FASP via a couple of simple tutorial problems. The user's guide is a self-contained document but does *not* provide any details about the algorithms or their implementation. Along with this guide, we provide a reference manual² for technical details on the implementation which includes references. We recommend that the users read these references to better understanding of the code. Furthermore, since FASP is under heavy development, please use this guide with caution because the code might have been changed before this document is updated.

1.4 How to obtain FASP

Downloading from SourceForge

The most updated version of FASP can be downloaded directly from

http://fasp.sf.net/download/faspsolver.zip

Downloading from BitBucket

FASP is also hosted on $BitBucket.org^3$ using Mercurial (Hg)⁴. A Hg client for GNU Linux, Mac OS X, or Windows can be downloaded from

http://mercurial.selenic.com/downloads/

²Available online at http://fasp.sourceforge.net. It is also available in "faspsolver/doc/doc.zip".

³Official website: https://bitbucket.org/

⁴Official website: http://mercurial.selenic.com/

There are also many other third-party clients which provides Hg services, for example: EasyMercurial⁵ (cross platform) and SourceTree⁶ (for Mac OS X only).

As a DVCS (Distributed Version Control System) source-control software, Hg is relatively new. But compared with other tools like Git, Hg is considered *friendlier* with a lower learning curve. This is despite the fact that Hg uses two distinct sets of commands and two distinct vocabularies for operations depending upon whether the repository is local or remote. Documentation for Hg is substantially better, including a book⁷. They've also had the advantage of trying the documentation on a fairly savvy group of developers (Mozilla) who gave them lots of feedback that helped polish the rough edges.

Linux or Mac OS X

First, you need to obtain a free copy of FASP kernel functions from our public Hg repository. If you are downloading FASP for the first time, you can clone the repository to your local machine:

```
"Download FASP kernel subroutines via HTTPS"

$ hg clone https://faspusers@bitbucket.org/fasp/faspsolver
```

If you have any problems when clone this repository, please send us an email to faspdev@gmail.com.

After a long pause⁸, you should have obtained "faspsolver" in your current directory successfully. If you have already cloned the repository before, you can just pull a new version and update your local version with it: Go to your local "faspsolver" directory and then

```
"Pull a new version from BitBucket"
$ hg pull

"Update you local version to the new version"
$ hg update
```

Windows OS

If you are using Windows, you may want to install TortoiseHg⁹. After installing it, the TortoiseHg menu has been merged into the right-click menu of Windows Explore. You could download FASP

⁵Official website: http://easyhg.org

⁶Official website: http://www.sourcetreeapp.com

⁷The hgbook, http://hgbook.red-bean.com/

⁸In fact, a very long pause. This is because the initial clone with copy all the history data which is about 400MB in total. Depending on the speed of your network, it could take 15 minutes to one hour.

⁹Official website: http://tortoisehg.bitbucket.org/

copy from BitBucket.org. Choose "TortoiseHg" --> "Clone" in the pop-up menu, the source address is

```
https://faspusers@bitbucket.org/fasp/faspsovler
```

Then press "Clone" and you will obtain "faspsolver" in the directory you set.

1.5 Licensing

This basic (kernel) FASP distribution is open-source and is licensed under GNU Lesser General Public License or LGPL. Other distributions may have different licensing (contact the developer team for details on this).

LICENSING: This software is free software distributed under the Lesser General Public License or LGPL, version 3.0 or any later versions. This software distributed in the hope that it will be useful, but WITHOUT ANY WARRANTY; without even the implied warranty of MERCHANTABILITY or FITNESS FOR A PARTICULAR PURPOSE. See the GNU Lesser General Public License http://www.gnu.org/licenses/ for more details.

1.6 Building and installing the FASP library and examples

FASP has been tested using the compilers and built-in libraries of several Linux distributions (Cent OS, Debian, Fedora, RedHat, Ubuntu) Mac OS X 10.6 and later (Leopard, Snow Leopard, Lion, Mavericks, Yosemite, El Capitan), and Windows (XP, Win 7) with several compliers, including gcc, g++, clang, icc, VC++. FASP also easily links to applications written in Fortran and this has been tested with gfortran, g95,ifort Fortran compilers.

FASP on Linux or Mac OS X

To build the FASP library for these operating systems: (1) go to the "faspsolver" directory; (2) modify the "FASP.mk.example" file to math your system and save it as "FASP.mk"; (3) then execute:

```
$ make config
$ make install
```

Note that, if you do not have "FASP.mk" present in the current directory, default settings will be used for building and installation FASP.

In order to be sure that everything went all right, you can go to the "faspsolver/test" directory and try to run a test problem as follows:

```
$ ./test.ex
```

If you need help with the options available, type

```
$ make help
```

and you will get the following screen

```
Fast Auxiliary Space Preconditioners (FASP)
                                                   Quick start:
To build FASP, copy "FASP.mk.example" to "FASP.mk" and put user-defined setting
there and then type "make config; make install".
More options:
$ make config
                      # Configure the building environment
$ make config CC=gcc  # Configure with a specific C compiler
$ make config debug=yes # Configure with compiler debug options ON
$ make config debug=all # Configure with FASP internal debug options ON
                       # Compile the library (after "make config")
$ make install
                      # install FASP libraries and related files
                     # Remove installed files by "make install"
$ make uninstall
$ make headers
                      # Generate function decorations automatically
                      # Generate the FASP documentation with Doxygen
$ make docs
$ make clean
                      # Remove obj files but retain configuration options
$ make distclean
                      # Clean and completely removes the build directory
                      # Show version information
$ make version
$ make help
                       # Show this screen
```

To uninstall FASP and clean up the working directory, you can run

```
$ make uninstall
$ make distclean
```

Windows 7

We provide a Visual Studio 2008 (VS08) distribution and a VS10 distribution of FASP for Windows users. For example, you can just open "faspsolver/vs08/faspsolver-vs08.sln" if you are using VS08 as your default developing environment. Then a single-click at the "Build Solution" on the menu or "F7" key will give you all the FASP libraries and the test programs in "faspsolver/test/". The way for building in VS10 is similar.

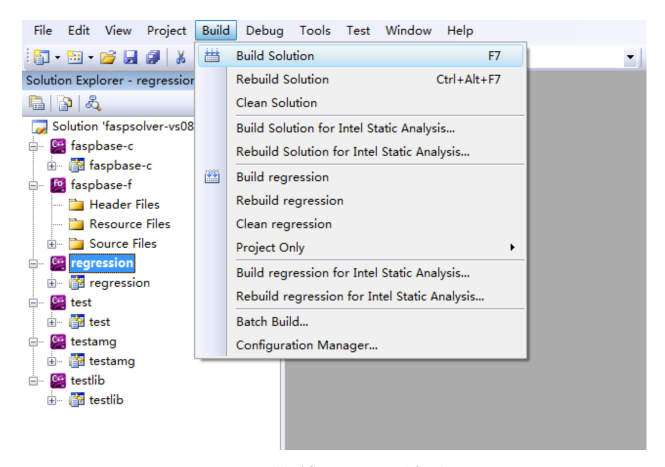


Figure 1.1: Build FASP using Visual Studio 2008.

You need a C/C++ complier and the Visual Studio to build FASP. For example, the build can be accomplished using either Microsoft Visual C++ or Intel C compiler.

If you are using other versions of Visual Studio (like VS05 or VS12), we advise NOT to convert the "VS08" files to your VS version because the FASP source files might be removed by the Visual Studio automatically. In such case, we recommend that you create your own version to build all the libraries and test programs.

If you need to build a VS FASP yourself, you need to create 5 projects:

- 1. "faspbase-c" contains all the ".c" and ".inl" files in the directory "./base/src/". You should add "./base/include" in Additional Directories. This project contains the core subroutines of faspsolver.
- 2. "faspbase-f" contains all the ".f" files in "./base/extra/sparsekit".

- 3. "testlib" contains all the ".c" files in "./test/src/". You should add "./test/include" in Additional Directories.
- 4. "test" is an executing program for test purpose in FASP. The source file is "./test/main/test.c".
- 5. "regression" is another executing program, which contains several methods to test the problems. The source file is "./test/main/regression.c".

NOTE: If you are using Visual C++, all the C files should be compiled as C++ code (by using the /TP compiling option).

After a successful build on VS, you will have two static libraries named "faspbase-c-vs08.lib" and "faspbase-f-vs08.lib". You can use the "lib" command to wrap them together as one single file (e.g. FASP.lib) for better portability. For example:

```
\texttt{C:} \\ \texttt{FASP} > \texttt{lib} \ / \texttt{ltcg} \ / \texttt{out:} \\ \texttt{FASP.lib} \ \texttt{faspbase-} \\ \texttt{c-vs08.lib} \ \texttt{faspbase-} \\ \texttt{f-vs08.lib}
```

Using a TCL based GUI for installation

For users who like more a GUI based installation, we provide a simple TCL Graphical User Interface (GUI) for building the FASP library. On a macine running Linux or Mac OS X with Tcl/Tk installed , you may invoke the GUI by typing

```
$ wish FASP_install.tcl
```

If all is OK, you should see on your screen the FASP window as shown in Figure 1.2. The rest of the building process is more or less straightforward: After choosing appropriate parameters, click "Config" first, followed by clicking "Install".

External libraries

There are a few *optional* external libraries that you might want to use, including memory allocation routines, direct solvers, ILU methods, discretization packages, etc. FASP has interfaces to several of them, for example, FASP can be linked to use UMFPack, SuperLU, MUMPS, SparseKit, dlmalloc.

FASP	FAST AUXILIARY SPACE PRECO	DNDITIONING
Quit Config Install HTML docs	Headers Uninstall	Help
Verbose output ?	○ Yes ○ No	
Build shared library ?	Yes O No	
Use Doxygen GUI (if found)?	Yes O No	
Build with OpenMP support ?	Yes O No	
Build with UMFPACK support ?	Yes O No	
Build type	Oebug Release	
The Debug build type assumes -g as a cortype assumes -O3 flag. Aditional compiler below.		
C flags		
F flags		
CXX flags		
Specify the path to SUITESPARSE (option use specific UMFPACK).	al, use it when you want to	
SUITESPARSE		

Figure 1.2: Install FASP using the TCL GUI on Mac OS X El Capitan.

Chapter 2

B brief tutorial

In this chapter, we discuss several simple examples included with this FASP distribution and demonstrating how to use the FASP package for solving linear systems. We read the matrices from disk files (the files are also included in the fasp distribution). All the examples are found in "faspsolver/tutorial/".

Here we only discuss the C version of these examples, but the FASP distribution also includes F90 versions of some of the examples.

After you successfully build FASP (see §1.6), just go to the "faspsolver/tutorial/" directory and the tutorial examples should be ready to run.

In the description below, we display a typical output from runs of each of the examples. Note that the actual output depends on the solver parameters, and, on your computer it may be different than what you see here.

2.1 Example 1:An AMG solver for the Poisson equation

The first example is a standard one: We read from disk a symmetric positive definite matrix A and right-hand side b then we solve Ax = b using the classical AMG method [2, 13, 14]; see §3.7. In this example the matrix A included with the FASP distribution corresponds to a discretization with continuous piecewise linear finite elements of the Poisson equation

$$-\Delta u = f$$

(with the Dirichlet boundary conditions) on a triangulation of a bounded domain Ω .

```
/*! \file poisson-amg.c

2 *

3 * \brief The first test example for FASP: using AMG to solve

4 * the discrete Poisson equation from P1 finite element.
```

```
5
              C version.
6
      \note AMG example for FASP: C version
7
8
    * Solving the Poisson equation (P1 FEM) with AMG
9
10
11
12
   #include "fasp.h"
13
   #include "fasp_functs.h"
14
15
16
    * \fn int main (int argc, const char * argv[])
17
    \ast \brief This is the main function for the first example.
18
19
    * \author Chensong Zhang
20
21
    * \date 12/21/2011
22
23
     * Modified by Chensong Zhang on 09/22/2012
24
   int main (int argc, const char * argv[])
25
26
   {
27
       input_param
                       inparam; // parameters from input files
                       amgparam; // parameters for AMG
28
       AMG_param
29
       printf("\n======="");
30
       printf("\n|| FASP: AMG example -- C version ||");
31
       printf("\n======\n\n");
32
33
       // Step O. Set parameters: We can use ini/amg.dat
34
       fasp_param_set(argc, argv, &inparam);
35
       fasp_param_init(&inparam, NULL, &amgparam, NULL, NULL);
36
37
       // Set local parameters using the input values
38
39
       const int print_level = inparam.print_level;
40
       // Step 1. Get stiffness matrix and right-hand side
41
       // Read A and b -- P1 FE discretization for Poisson. The location
42
       // of the data files is given in "ini/amg.dat".
43
       dCSRmat A;
44
45
       dvector b, x;
46
       char filename1[512], *datafile1;
47
       char filename2 [512], *datafile2;
48
49
       // Read the stiffness matrix from matFE.dat
50
       strncpy(filename1, inparam.workdir, 128);
51
       datafile1="csrmat_FE.dat"; strcat(filename1, datafile1);
52
53
       // Read the RHS from rhsFE.dat
54
       strncpy(filename2, inparam.workdir, 128);
       datafile2="rhs_FE.dat"; strcat(filename2, datafile2);
55
56
       fasp_dcsrvec2_read(filename1, filename2,&A,&b);
```

```
58
        // Step 2. Print problem size and AMG parameters
59
        if (print_level>PRINT_NONE) {
60
            printf("A: m = %d, n = %d, nnz = %d n", A.row, A.col, A.nnz);
61
            printf("b: n = %d n", b.row);
62
63
            fasp_param_amg_print(&amgparam);
64
        }
65
66
        // Step 3. Solve the system with AMG as an iterative solver
67
        // Set the initial guess to be zero and then solve it
        // with AMG method as an iterative procedure
68
69
        fasp_dvec_alloc(A.row, &x);
70
        fasp_dvec_set(A.row, &x, 0.0);
71
72
        fasp\_solver\_amg(\&A, \&b, \&x, \&amgparam);
73
        // Step 4. Clean up memory
74
        fasp_dcsr_free(\&A);
75
        fasp_dvec_free(&b);
76
        fasp_dvec_free(&x);
77
78
79
        return FASP_SUCCESS;
80
81
82
                End of File
83
```

Since this is the first example, we will explain it in some detail:

- Line 1 tells the Doxygen documentation system that the filename is "poisson-amg.c". Line 3–5 tells the Doxygen what is the purpose of this file (function).
- Line 12–13 includes the main FASP header file "fasp.h" and FASP function decoration header "fasp_functs.h". These two headers shall be included in all files that requires FASP subroutines. Please also be noted that the function declarations in "fasp_functs.h" are automatically generated from the source files by an awk script and we do not recommend modifying this file, since your changes may be lost.
- Line 35 sets solver parameters using the default parameters or from the command line options; see more discussions in §2.4. In the "tutorial/ini/amg.dat" file, we can set the location of the data files, type of solvers, maximal number of iteration numbers, convergence tolerance, and many other parameters for iterative solvers.
- Line 44 defines a sparse matrix A in the compressed sparse row (CSR) format. Line 45 defines two vectors: the right-hand side b and the numerical solution x. We refer to §3.1 for definitions of vectors and general sparse matrices.

- Line 57 reads the matrix and the right-hand side from two disk files. Line 49–58 defines the filenames of them.
- Line 60–64 prints basic information of coefficient matrix, right-hand side, and solver parameters.
- Line 69–70 allocates memory for the solution vector x and set its initial value to be all zero.
- Line 72 solves Ax = b using the AMG method. Type of the AMG method and other parameters have been given in "amgparam" at Line 36; see §3.7.
- Line 75–77 frees up memory allocated for A, b, and x.

To run this example, type:

```
$ ./poisson-amg-c.ex
```

A sample output is as follows:

```
FASP: AMG example -- C version
fasp_dcsrvec2_read: reading file ../data/csrmat_FE.dat...
fasp_dcsrvec2_read: reading file ../data/rhs_FE.dat...
A: m = 3969, n = 3969, nnz = 27281
b: n = 3969
       Parameters in AMG_param
                                    2
AMG print level:
AMG max num of iter:
                                   1
AMG type:
                                   1.00e-06
AMG tolerance:
                                   20
AMG max levels:
AMG cycle type:
AMG coarse solver type:
AMG scaling of coarse correction:
AMG smoother type:
AMG smoother order:
                                   1
AMG num of presmoothing:
AMG num of postsmoothing:
                                   1
AMG coarsening type:
AMG interpolation type:
                                   1
AMG dof on coarsest grid:
                                   500
AMG strong threshold:
                                   0.3000
                                   0.2000
AMG truncation threshold:
                                   0.9000
AMG max row sum:
AMG aggressive levels:
                                   0
AMG aggressive path:
```

Level	Num of rows	Num of nonzeros	Avg. NNZ / row
0	3969	27281	6.87
1	1985	28523	14.37
2	541	7951	14.70
3	141	1803	12.79
	omplexity $=1.6$	72 Operator costs 0.0068 seconds.	emplexity = 2.403
Classica	l AMG setup cos		
Classica	l AMG setup cos	sts 0.0068 seconds.	·
Classica t Num	1 AMG setup cos		Conv. Factor
Classica Tt Num	1 AMG setup cos r / b 1.000000e+00 9.851978e-03	r	Conv. Factor
Classica t Num 0 1	1 AMG setup cos r / b 1.000000e+00 9.851978e-03 3.507451e-04	r	Conv. Factor
Classica It Num 0 1 2	1 AMG setup cos r / b 1.000000e+00 9.851978e-03 3.507451e-04	sts 0.0068 seconds. r $7.514358e+00$ $7.403129e-02$ $2.635624e-03$	Conv. Factor

We also provide a Fortran 90 example, which does the same thing as this C code except it gives less output, in "tutorial/main/poisson-amg.f90". Users who would like to call FASP solver from a Fortran based application can see how to do this example.

```
!> \file poisson-amg.f90
   the discrete Poisson equation from P1 finite element.
  ! >
   !>
          F90 version.
4
   !>
5
   !> \note AMG example for FASP: F90 version
6
7
   !> Solving the Poisson equation (P1 FEM) with AMG
8
9
   !>
10
   !> \author Chensong Zhang
   !> \date 12/21/2011
11
12
13
   program test
14
     implicit none
15
16
17
     double precision, dimension(:), allocatable :: u, b, a
                    dimension(:), allocatable :: ia, ja
18
     integer,
19
20
              :: iufile, n, nnz, i, prt_lvl, maxit
21
     double precision :: tol
```

```
22
      print*, ""
23
24
      write(*,"(A)") "|| FASP: AMG example -- F90 version ||"
25
      26
      print*, ""
27
28
      ! Step 0: user defined variables
29
30
      prt_lvl = 3
      maxit = 100
31
      \mathtt{tol} = 1.0\,\mathtt{d}{-6}
32
33
      iufile = 1
34
      ! Step 1: read A and b
35
36
      !===> Read data A from file
37
38
      open(unit=iufile,file='../data/csrmat_FE.dat')
39
40
      read(iufile,*) n
      allocate(ia(1:n+1))
41
      read(iufile,*) (ia(i),i=1,n+1)
42
43
      nnz=ia(n+1)-ia(1)
44
      \verb"allocate" (\verb"ja" (1:nnz")", \verb"a" (1:nnz")")
45
      read(iufile,*) (ja(i),i=1,nnz)
46
47
      read(iufile,*) (a(i),i=1,nnz)
48
      close(iufile)
49
50
      !===> Read data b from file
51
      open(unit=iufile,file='../data/rhs_FE.dat')
52
53
54
      read(iufile,*) n
      allocate(b(1:n))
55
      read(iufile,*) (b(i),i=1,n)
56
57
      close(iufile)
58
59
      !===> Shift the index to start from 0 (for C routines)
60
      forall (i=1:n+1) ia(i)=ia(i)-1
61
       \begin{array}{ll} \textbf{forall} & (\mathtt{i} \!=\! 1 \!:\! \mathtt{nnz}) & \mathtt{ja}(\mathtt{i}) \!=\! \mathtt{ja}(\mathtt{i}) \!-\! 1 \end{array} 
62
63
      ! Step 2: Solve the system
64
65
66
      !===> Initial guess
67
      allocate(u(1:n))
68
      u = 0.0 d0
69
      call fasp_fwrapper_amg(n,nnz,ia,ja,a,b,u,tol,maxit,prt_lvl)
70
71
      ! Step 3: Clean up memory
72
      deallocate(ia,ja,a)
73
      deallocate(b,u)
74
```

2.2 Example 2: Conjugate gradient without preconditioning

In the second example, we modify the previous example slightly and solve the Poisson equation using iterative methods (here by default we use the Conjugate Gradient method without preconditioning).

```
1
   /*! \file poisson-its.c
2
    * \brief The second test example for FASP: using ITS to solve
3
             the discrete Poisson equation from P1 finite element.
4
    * \note ITS example for FASP: C version
5
6
7
    * Solving the Poisson equation (P1 FEM) with iterative methods
8
9
   #include "fasp.h"
10
   #include "fasp_functs.h"
11
12
13
    * \fn int main (int argc, const char * argv[])
14
15
16
    * \brief This is the main function for the second example.
17
18
    * \author Feiteng Huang
    * \date 04/13/2012
19
20
21
    * Modified by Chensong Zhang on 09/22/2012
22
   int main (int argc, const char * argv[])
23
^{24}
25
                           inparam; // parameters from input files
       input_param
26
       itsolver_param
                           itparam; // parameters for itsolver
27
28
       printf("\n======="");
       printf("\n|| FASP: ITS example -- C version ||");
29
       printf("\n======\n\n");
30
31
       // Step O. Set parameters: We can use ini/its.dat
32
       fasp_param_set(argc, argv, &inparam);
33
34
       fasp_param_init(&inparam, &itparam, NULL, NULL, NULL);
35
36
       // Set local parameters
37
       const int print_level = inparam.print_level;
38
39
       // Step 1. Get stiffness matrix and right-hand side
```

```
// Read A and b -- P1 FE discretization for Poisson. The location
40
41
       // of the data files is given in "ini/its.dat".
42
       dCSRmat A;
       dvector b, x;
43
       char filename1[512], *datafile1;
44
       45
46
47
       // Read the stiffness matrix from matFE.dat
48
       strncpy(filename1, inparam.workdir, 128);
49
       datafile1="csrmat_FE.dat"; strcat(filename1, datafile1);
50
51
       // Read the RHS from rhsFE.dat
       strncpy(filename2, inparam.workdir, 128);
52
       datafile2="rhs_FE.dat"; strcat(filename2,datafile2);
53
54
55
       fasp_dcsrvec2_read(filename1,filename2,&A,&b);
56
       // Step 2. Print problem size and ITS parameters
57
       if (print_level>PRINT_NONE) {
58
           printf("A: m = %d, n = %d, nnz = %d\n", A.row, A.col, A.nnz);
59
            printf("b: n = %d\n", b.row);
60
           fasp_param_solver_print(&itparam);
61
       }
62
63
64
       // Step 3. Solve the system with ITS as an iterative solver
       // Set the initial guess to be zero and then solve it using standard
65
       // iterative methods, without applying any preconditioners
66
67
       fasp_dvec_alloc(A.row, &x);
       fasp_dvec_set(A.row,&x,0.0);
68
69
       fasp_solver_dcsr_itsolver(&A, &b, &x, NULL, &itparam);
70
71
72
       // Step 4. Clean up memory
       fasp_dcsr_free(\&A);
73
74
       fasp_dvec_free(&b);
75
       fasp_dvec_free(&x);
76
       return FASP_SUCCESS;
77
78
79
80
81
                End of File
82
```

This example is very similar to the first example and we briefly explain the differences:

- Line 67–68 allocates memory for the solution vector x and set its initial value to be all zero.
- Line 70 solves Ax = b using the general interface for Krylov subspace methods. Type the iterative method and other parameters have been specified in "itparam"; see §3.5 for details.

To run this example, we can simply type:

```
$ ./poisson-its-c.ex
```

A sample output is as follows:

```
FASP: ITS example -- C version
                                               fasp_dcsrvec2_read: reading file ../data/csrmat_FE.dat...
fasp_dcsrvec2_read: reading file ../data/rhs_FE.dat...
A: m = 3969, n = 3969, nnz = 27281
b: n = 3969
        Parameters in itsolver_param
Solver print level:
                                           2
Solver type:
                                           1
Solver precond type:
                                           2
Solver max num of iter:
                                           500
                                           1.00\,\mathrm{e}\!-\!06
Solver tolerance:
Solver stopping type:
Calling PCG solver (CSR) ...
It Num
             ||r||/||b||
                                      ||r||
                                                       Conv. Factor
      0 |
           1.0000000e+00
                                  7.514358e+00
      1 \mid 5.078029 e - 01
                                  3.815813e+00
                                                           0.5078
      2 |
           3.728856 e - 01
                                  2.801996 e+00
                                                           0.7343
      3 \mid 3.359470e-01
                                  2.524426\,e+00
                                                           0.9009
      4 \mid 2.590574e-01
                               1.946650 e+00
                                                           0.7711
      5 \mid 2.380797e-01
                             1.789016 e+00
                                                           0.9190
      6 \mid 1.992579e-01
                             1.497295e+00
                                                           0.8369
      7 \mid 1.847971e-01
                              1.388631e+00
                                                           0.9274
      8 \mid 1.619777e-01
                                  1.217158\,\mathrm{e}{+00}
                                                           0.8765
     9 |
           1.513446\,\mathrm{e}\!-\!01
                                  1.137257\,\mathrm{e}{+00}
                                                           0.9344
    10 |
          1.364935\,\mathrm{e}\!-\!01
                                  1.025661\,\mathrm{e}{+00}
                                                           0.9019
    11 \mid 1.283425 e - 01
                               9.644117e-01
                                                           0.9403
    12 \mid 1.179652e-01
                               8.864327e-01
                                                           0.9191
    13 \mid 1.115146e-01
                                  8.379605\,\mathrm{e}{-01}
                                                           0.9453
    14 \mid 1.038726 e - 01
                                  7.805360\,\mathrm{e}{-01}
                                                           0.9315
    15 \mid 9.863412e-02
                                  7.411721 e - 01
                                                           0.9496
    16 \mid 9.277360 e - 02
                                  6.971341 \, \mathrm{e}{-01}
                                                           0.9406
    17 \mid 8.842679 e - 02
                                  6.644706 \, \mathrm{e}{-01}
                                                           0.9531
    18 \mid 8.378399e-02
                             | 6.295829 e - 01
                                                           0.9475
    19 \mid 8.011023e-02
                                  6.019770\,\mathrm{e}{-01}
                                                           0.9562
    20 \mid 7.633221 e - 02
                                  5.735875 e - 01
                                                           0.9528
    21 \quad | \quad 7.317756 \, \mathrm{e} \! - \! 02
                                  5.498824\,\mathrm{e}{-01}
                                                           0.9587
     22 |
            7.003292\,\mathrm{e}\!-\!02
                                  5.262524\,\mathrm{e}{-01}
                                                           0.9570
    23 \mid 6.728610 e - 02
                               | 5.056119e-01
                                                           0.9608
```

```
24
         6.461736\,\mathrm{e}\!-\!02
                                       4.855580\,\mathrm{e}\!-\!01
                                                                          0.9603
          6.219614 \, \mathrm{e}\!-\!02
                                        4.673640\,\mathrm{e}{-01}
                                                                          0.9625
26
          5.989276 \, \mathrm{e}{-02}
                                        4.500557\,\mathrm{e}{-01}
                                                                          0.9630
27 - 1
          5.773520\,\mathrm{e}\!-\!02
                                        4.338429\,\mathrm{e}{-01}
                                                                          0.9640
28
          5.571758\,\mathrm{e}\!-\!02
                                        4.186818\,\mathrm{e}\!-\!01
                                                                          0.9651
         5.377630\,\mathrm{e}{-02}
                                        4.040944\,\mathrm{e}{-01}
29
                                                                          0.9652
                                        3.906404e-01
30 |
         5.198586\,\mathrm{e}\!-\!02
                                                                          0.9667
31 |
         5.022413\,\mathrm{e}\!-\!02
                                        3.774021 \, \mathrm{e}{-01}
                                                                          0.9661
         4.861699\,\mathrm{e}\!-\!02
                                        3.653255\,\mathrm{e}\!-\!01
                                                                          0.9680
33 |
         4.700598 e - 02
                                        3.532197 e - 01
                                                                          0.9669
34
         4.554874\,\mathrm{e}\!-\!02
                                        3.422696\,\mathrm{e}{-01}
                                                                          0.9690
35
         4.406559 \, \mathrm{e}{-02}
                                        3.311246\,\mathrm{e}{-01}
                                                                          0.9674
36
         4.273253 e - 02
                                        3.211075 e - 01
                                                                          0.9697
37
         4.135901 \, \mathrm{e}{-02}
                                        3.107864 \, \mathrm{e} \! - \! 01
                                                                          0.9679
38
         4.013076 \, \mathrm{e}{-02}
                                        3.015569 \, \mathrm{e}{-01}
                                                                          0.9703
         3.885861 e - 02
39
                                        2.919975 e - 01
                                                                          0.9683
40
          3.776252 e - 02
                                        2.837611e-01
                                                                          0.9718
                                        2.764205\,\mathrm{e}\!-\!01
41
          3.678565\,\mathrm{e}\!-\!02
                                                                          0.9741
42
          3.648645\,\mathrm{e}\!-\!02
                                        2.741722\,\mathrm{e}\!-\!01
                                                                          0.9919
43
         3.725368 e - 02
                                        2.799375 e - 01
                                                                          1.0210
         3.922957 e - 02
                                        2.947850\,\mathrm{e}\!-\!01
                                                                          1.0530
44
45
         4.003513 e - 02
                                        3.008383 e - 01
                                                                          1.0205
46
         3.683219 e - 02
                                        2.767703 e - 01
                                                                          0.9200
          3.161285\,\mathrm{e}\!-\!02
                                        2.375503\,\mathrm{e}\!-\!01
47 l
                                                                          0.8583
48
          2.944107\,\mathrm{e}\!-\!02
                                        2.212307\,\mathrm{e}{-01}
                                                                          0.9313
         2.961834\,\mathrm{e}\!-\!02
49
                                        2.225628\,\mathrm{e}\!-\!01
                                                                          1.0060
         2.774118\,\mathrm{e}\!-\!02
                                                                          0.9366
50
                                        2.084571 \, \mathrm{e}{-01}
         2.513603\,\mathrm{e}\!-\!02
51 |
                                        1.888811 e - 01
                                                                          0.9061
          2.489908\,\mathrm{e}\!-\!02
                                        1.871006\,\mathrm{e}\!-\!01
                                                                          0.9906
          2.379644e-02
                                        1.788150 e - 01
53
                                                                          0.9557
          2.190590\,\mathrm{e}\!-\!02
54
                                        1.646088 e - 01
                                                                          0.9206
```

2.3 Example 3: Conjugate gradient with preconditioner

This example is a bit more involved and is a modification of the previous one. In this example, we wish to demonstrate how to use a the FASP library and run a preconditioned conjugate gradient (PCG) method.

```
/*! \file poisson-pcg.c
2
       \brief The third test example for FASP: using PCG to solve
               the discrete Poisson equation from P1 finite element.
3
4
               C version.
5
       \note PCG example for FASP: C version
6
7
8
       Solving the Poisson equation (P1 FEM) with PCG methods
9
10
11
   #include "fasp.h"
   #include "fasp_functs.h"
```

```
13
14
    /**
15
    * \fn int main (int argc, const char * argv[])
16
     * \brief This is the main function for the third example.
17
18
     * \author Feiteng Huang
19
20
     * \date 05/17/2012
21
     * Modified by Chensong Zhang on 09/22/2012
22
    */
23
24
    int main (int argc, const char * argv[])
25
                              inparam; // parameters from input files
26
        input_param
                             itparam; // parameters for itsolver
27
        itsolver_param
                              amgparam; // parameters for AMG
28
        AMG_param
        ILU_param
                              iluparam; // parameters for ILU
29
30
31
        printf("\n======="");
        printf("\n|| FASP: PCG example -- C version ||");
32
        printf("\n======\n\n");
33
34
35
        // Step O. Set parameters: We can use ini/pcg.dat
        fasp_param_set(argc, argv, &inparam);
36
        fasp_param_init(&inparam, &itparam, &amgparam, &iluparam, NULL);
37
38
        // Set local parameters
39
        const SHORT print_level = itparam.print_level;
40
        const SHORT pc_type = itparam.precond_type;
41
        const SHORT stop_type = itparam.stop_type;
42
        const INT maxit
                                = itparam.maxit;
43
        const REAL tol
44
                                 = itparam.tol;
45
        // Step 1. Get stiffness matrix and right-hand side
46
47
        // Read A and b -- P1 FE discretization for Poisson. The location
        // of the data files is given in "ini/pcg.dat".
48
        dCSRmat A;
49
        dvector b, x;
50
        char filename1[512], *datafile1;
51
        \begin{array}{ll} \textbf{char} & \texttt{filename2} \left[\, 5\, 1\, 2\, \right]\,, & *\texttt{datafile2}\,; \end{array}
52
53
54
        // Read the stiffness matrix from matFE.dat
55
        strncpy(filename1, inparam.workdir, 128);
        datafile1="csrmat_FE.dat"; strcat(filename1, datafile1);
56
57
58
        // Read the RHS from rhsFE.dat
59
        strncpy(filename2, inparam.workdir, 128);
60
        datafile2="rhs_FE.dat"; strcat(filename2, datafile2);
61
62
        {\tt fasp\_dcsrvec2\_read} \, (\, {\tt filename1} \, , {\tt filename2} \, , \& \, {\tt A} \, , \& \, {\tt b} \, ) \, ;
63
64
        // Step 2. Print problem size and PCG parameters
        if (print_level>PRINT_NONE) {
```

```
printf("A: m = %d, n = %d, nnz = %d n", A.row, A.col, A.nnz);
66
            printf("b: n = %d\n", b.row);
67
68
            fasp_param_solver_print(&itparam);
        }
69
70
        // Setp 3. Setup preconditioner
71
        // Preconditioner type is determined by pc_type
72
73
       precond *pc = fasp_precond_setup(pc_type, &amgparam, &iluparam, &A);
74
75
        // Step 4. Solve the system with PCG as an iterative solver
        // Set the initial guess to be zero and then solve it using PCG solver
76
77
        // Note that we call PCG interface directly. There is another way which
        // calls the abstract iterative method interface; see possion-its.c for
78
79
        // more details.
80
        fasp_dvec_alloc(A.row, &x);
81
        fasp_dvec_set(A.row, &x, 0.0);
82
        fasp_solver_dcsr_pcg(&A, &b, &x, pc, tol, maxit, stop_type, print_level);
83
84
        // Step 5. Clean up memory
85
       if (pc_type!=PREC_NULL) fasp_mem_free(pc->data);
86
87
        fasp_dcsr_free(\&A);
        fasp_dvec_free(&b);
88
        fasp_dvec_free(\&x);
89
90
        return FASP_SUCCESS;
91
92
93
94
               End of File
95
96
```

This example is very similar to the first example, and the details are as follows.

- Line 36 sets default parameters. In this example, we need parameters for iterative methods, AMG preconditioner, and ILU preconditioner.
- Line 73 sets up the desired preconditioner and prepare it for the preconditioned iterative methods.
- Line 83 calls PCG to solve Ax = b. One can also call the general iterative method interface as in the previous example.
- Line 86 cleans up auxiliary data associated with the preconditioner in use if necessary.

To run this example, we can simply type:

```
$ ./poisson-pcg-c.ex
```

A sample output is given as follows (note that the actual output depends on the solver parameters and might be different than what you see here):

```
{\tt FASP: \ PCG \ example -- \ C \ version}
fasp_dcsrvec2_read: reading file ../data/csrmat_FE.dat...
fasp_dcsrvec2_read: reading file ../data/rhs_FE.dat...
A: m = 3969, n = 3969, nnz = 27281
b: n = 3969
       Parameters in itsolver_param
Solver print level:
Solver type:
                                    1
Solver precond type:
                                    500
Solver max num of iter:
                                     1.00\,\mathrm{e}\!-\!06
Solver tolerance:
Solver stopping type:
  Level Num of rows Num of nonzeros Avg. NNZ / row
    0
               3969
                                 27281
                                                  6.87
               1985
                                                 14.37
    1
                                 28523
    2
                541
                                  7951
                                                 14.70
                                  1803
                                                 12.79
  Grid complexity = 1.672 | Operator complexity = 2.403
Classical AMG setup costs 0.0042 seconds.
It Num |
         ||r||/||b|| |
                                ||r||
                                            Conv. Factor
     0 |
         1.0000000e+00
                         | 7.514358e+00 |
     1 |
          1.156153 \, \mathrm{e}{-02}
                             8.687750\,\mathrm{e}{-02}
                                                   0.0116
         3.127181\,\mathrm{e}{-04}
                         2.349876e-03
                                                  0.0270
                         3.617014e-05
     3 \mid 4.813471e-06
                                                  0.0154
     4 \mid 5.312526e-08
                         | 3.992022 e - 07 |
                                                  0.0110
Number of iterations = 4 with relative residual 5.312526e-08.
```

2.4 How to change default parameters in FASP solver/preconditioner

In the previous examples, we have seen how to set solver parameters using the default settings. Here we discuss changing such parameters by reading them from a disk file. An example of such file is found in the FASP tutorial directory and is named "tutorial/ini/amg.dat".

```
$ ./poisson-amg-c.ex -ini ini/amg.dat
```

We take "tutorial/ini/amg.dat" as an example:

```
-%
                                                   %
2
   \% input parameters
   |\% lines starting with \% are comments
                                                   %
   \% must have spaces around the equal sign "="
5
6
   workdir = ../data/
                          \% work directory, no more than 128 characters
8
   print_level = 3
                         \% How much information to print out
9
10
                                                   -%
11
   \% parameters for multilevel iteration
12
13
   AMG_type
                             = SA
                                      \% C classic AMG
14
                                      \% SA smoothed aggregation
15
                                      \% UA unsmoothed aggregation
16
                                      \% V V-cycle | W W-cycle
17
   AMG_cycle_type
                             = v
                                      \% A AMLI-cycle | NA Nonlinear AMLI-cycleA
18
                                      \% tolerance for AMG
   AMG_tol
                            = 1e-8
19
                                      \% number of AMG iterations
                            = 100
20
   AMG_maxit
                            = 20
                                      \% max number of levels
21
   AMG_levels
   AMG_coarse_dof
                            = 500
                                      \% max number of coarse degrees of freedom
22
                           = OFF
                                      \% switch of scaling of the coarse grid correction
23
   AMG_coarse_scaling
                        = 2
24
   AMG_amli_degree
                                      \% degree of the polynomial used by AMLI cycle
25
   AMG_nl_amli_krylov_type = 6
                                      % Krylov method in NLAMLI cycle: 6 FGMRES | 7 GCG
26
27
   \% parameters for AMG smoothing
                                                   %
28
29
30
   AMG_smoother
                            = GS
                                      % GS | JACOBI | SGS
31
                                      \% SOR | SSOR | GSOR | SGSOR | POLY
32
                            = 0
                                      \% number of levels using ILU smoother
   AMG_ILU_levels
33
   AMG_Schwarz_levels
                            = 0
                                      \% number of levels using Schwarz smoother
34
   AMG_relaxation
                            = 1.1
                                      \% relaxation parameter for SOR smoother
35
   AMG_polynomial_degree = 3
                                      \% degree of the polynomial smoother
36
37
   AMG_presmooth_iter
                          = 2
                                      \% number of presmoothing sweeps
                                      \% number of postsmoothing sweeps
   AMG_postsmooth_iter
                            = 2
38
39
40
41
   \% parameters for classical AMG SETUP
                                                   %
42
43
44
   AMG_coarsening_type
                            = 1
                                      \% 1 Modified RS
45
                                      \% 3 Compatible Relaxation
                                      \% \ 4 \ {\tt Aggressive}
46
47
   AMG_interpolation_type
                            = 1
                                      \% 1 Direct | 2 Standard | 3 Energy-min
   AMG_strong_threshold = 0.6
                                      \% Strong threshold
48
   AMG\_truncation\_threshold = 0.4
                                      \% Truncation threshold
   AMG_max_row_sum = 0.9
                                      \% Max row sum
```

```
-%
52
   \% parameters for aggregation-type AMG SETUP
                                                     %
                                                     -%
54
55
                             = 0.08
                                       \% Strong coupled threshold
56
   AMG_strong_coupled
57
   AMG_max_aggregation
                             = 20
                                       \% Max size of aggregations
   AMG_tentative_smooth
                             = 0.67
                                       \% Smoothing factor for tentative prolongation
58
   AMG_smooth_filter
                             = OFF
                                       \% Switch for filtered matrix for smoothing
```

We now briefly discuss the parameters above: This example is very similar to the first example and we now briefly explain it:

- Line 7 sets the working directory, which should contain data files for the matrices (and right-hand side vectors when necessary).
- Line 8 sets the level of output for FASP routines. It should range from 0 to 10 with 0 means no output and 10 means output everything possible.
- Line 14–25 sets the basic parameters for multilevel iterations. For example, type of AMG, type of multilevel cycles, number of maximal levels, etc.
- Line 31–38 sets the type of smoothers, number of smoothing sweeps, etc.
- Line 44–50 sets the parameters for the setup phase of the classical AMG method (§3.7).
- Line 56–59 gives the parameters for the setup phase of the aggregation-base AMG methods (§3.7).

You can do a very simple experiment and change the AMG type from the classical AMG to smoothed aggregation AMG by revise Line 14 to

```
AMG_type = SA
```

Then you run "poisson-amg-c.ex" one more time and will get

```
| FASP: AMG example — C version | |

fasp_dcsrvec2_read: reading file ../data/csrmat_FE.dat...
fasp_dcsrvec2_read: reading file ../data/rhs_FE.dat...
A: m = 3969, n = 3969, nnz = 27281
b: n = 3969

Parameters in AMG_param

AMG print level: 3
AMG max num of iter: 100
```

```
AMG type:
AMG tolerance:
                                         1.00\,\mathrm{e}{-08}
AMG max levels:
AMG cycle type:
                                         1
AMG coarse solver type:
AMG scaling of coarse correction:
AMG smoother type:
{\tt AMG} \ {\tt smoother} \ {\tt order}:
AMG num of presmoothing:
AMG num of postsmoothing:
Aggregation type:
Aggregation number of pairs:
Aggregation quality bound:
                                       8.00
Calling SA AMG ...
  Level
           Num of rows
                           Num of nonzeros Avg. NNZ / row
    0
                 3969
                                     27281
                                                        6.87
    1
                  541
                                      6531
                                                      12.07
                                       421
                                                       10.27
  Grid complexity = 1.147 | Operator complexity = 1.255
Smoothed aggregation setup costs 0.0028\ \mbox{seconds}\,.
It Num | ||r||/||b|| |
                                    ||r||
                                                    Conv. Factor
     0 \mid 1.000000e+00
                                7.514358e+00
           4.345463 \, \mathrm{e}{-02}
                                                        0.0435
     1 |
                                3.265336 \, \mathrm{e}{-01}
          8.041967 \, \mathrm{e}{-03}
                                6.043022\,\mathrm{e}{-02}
                                                        0.1851
     3 \mid 3.808810 e - 03
                           | 2.862076 e - 02 |
                                                        0.4736
     4 \mid 1.838990 e - 03
                           | 1.381883 e - 02 |
                                                        0.4828
     5 \mid 8.675952e-04
                           | 6.519421 e - 03 |
                                                        0.4718
     6 \mid 4.089274 e - 04
                           | 3.072827 e - 03 |
                                                        0.4713
     7 \mid 1.939823 e - 04
                                1.457653\,\mathrm{e}{-03}
                                                        0.4744
          9.276723\,\mathrm{e}\!-\!05
                                6.970862\,\mathrm{e}{-04}
                                                        0.4782
                           3.360270\,\mathrm{e}{-04}
     9 \mid 4.471799e-05
                                                        0.4820
    10 \mid 2.171249 e - 05
                           | 1.631554e-04 |
                                                        0.4855
                           7.972239e-05
    11 \mid 1.060934e-05
                                                        0.4886
                           | 3.916668 e - 05
    12 \mid 5.212246 e - 06
                                                        0.4913
    13 \mid 2.572464 e - 06
                             1.933042e-05
                                                        0.4935
    14
          1.274466 \, \mathrm{e}{-06}
                                9.576797 e - 06
                                                        0.4954
    15 \mid 6.333891 e - 07
                                4.759512e-06
                                                        0.4970
                           2.371476e-06
    16 \mid 3.155926 e - 07
                                                        0.4983
    17 \mid 1.575755e-07
                           1.184079 e - 06
                                                        0.4993
    18 \mid 7.881043 e - 08
                           | 5.922098e-07 |
                                                        0.5001
                           | 2.965950 e - 07 |
    19 \mid 3.947044e-08
                                                        0.5008
    20 \mid 1.978978 \, \mathrm{e}{-08}
                                1.487075\,\mathrm{e}{-07}
                                                        0.5014
    21 \mid 9.931176 e - 09
                           7.462641e-08
                                                        0.5018
Number of iterations = 21 with relative residual 9.931176\,\mathrm{e}-09.
```

```
AMG solve costs 0.0053 seconds. AMG totally costs 0.0083 seconds.
```

You can compare this with the sample results in §2.1.

The input parameters allowed in FASP are not limited to the ones listed in this example. A list of possible iterative methods and preconditioners can be found in "base/include/fasp_const.h"; see §4.2. For more parameters and their ranges, we refer to the FASP Reference Manual.

Using "-ini [FILE]" is just one example of allowed command line option. To find out more what command line options are acceptable, you can type in a terminal window:

```
$ ./poisson-amg-c.ex -help
```

which will give you something like

```
FASP: AMG example — C version
                                         FASP command line options:
                    [CharValue] : Ini file name
 -\mathtt{ini}
 -print
                   [IntValue] : Print level
 -output
                   [IntValue] : Output to screen or a log file
 -\mathtt{solver}
                   [IntValue] : Solver type
                    [IntValue] : Preconditioner type
 -precond
                     [IntValue] : Max number of iterations
 -\mathtt{maxit}
                   [RealValue] : Tolerance for iterative solvers
 -\mathtt{tol}
                  [IntValue] : Max number of AMG iterations
 -\mathtt{amgmaxit}
 -{\tt amgtol}
                   [RealValue] : Tolerance for AMG methods
 -\mathtt{amgtype}
                   [IntValue] : AMG type
 -amgcycle [IntValue] : AMG cycle type
 -{\tt amgcoarsening} \qquad [\, {\tt IntValue} \,] \quad : \ {\tt AMG} \ \ {\tt coarsening} \ \ {\tt type}
 -{\tt amginterpolation} [IntValue] : AMG interpolation type
 -{\tt amgsmoother}
                   [IntValue] : AMG smoother type
 -{\tt amgsthreshold} [RealValue] : AMG strong threshold
 -amgscoupled [RealValue] : AMG strong coupled threshold
                           : Brief help messages
 -help
```

Chapter 3

Data structures and basic usage

In this chapter, we discuss the basic data structures and the important building blocks which are useful for constructing auxiliary space preconditioners for systems of PDEs in Chapter 4. In particular, we will discuss vectors, sparse matrices, iterative methods, and multigrid methods.

3.1 Vectors and sparse matrices

The data structures most often used for implementing iterative methods are sparse matrices and vectors. In this section, we first discuss the data structures for vectors and matrices in FASP; and then we discuss BLAS for sparse matrices. The definitions can be found in "base/include/fasp. h".

Vectors

The data structure for vectors is very simple. It only contains the length of the vector and an array which contains the entries of this vector.

```
330
331
     * \struct dvector
332
      * \brief Vector with n entries of REAL type
333
334
    typedef struct dvector{
335
336
         //! number of rows
         INT row;
337
338
339
         //! actual vector entries
         REAL *val;
340
341
    } dvector; /**< Vector of REAL type */
```

Sparse matrices

On the other hand, sparse matrices for PDE applications are very complicated. It depends on the particular applications, discretization methods, as well as solution algorithms. In FASP, there are several types of sparse matrices, COO, CSR, CSRL, BSR, and CSR Block, etc. The presentation closely follows ideas from Pissanetzky [12].

In this section, we use the following sparse matrix as an example to explain different formats for sparse matrices:

Example 3.1.1 Consider the following 4×5 matrix with 12 non-zero entries

$$\left(\begin{array}{cccccc}
1 & 1.5 & 0 & 0 & 12 \\
0 & 1 & 6 & 7 & 1 \\
3 & 0 & 6 & 0 & 0 \\
1 & 0 & 2 & 0 & 5
\end{array}\right)$$

(i) COO format

The coordinate (COO) format or IJ format is the simplest sparse matrix format.

```
192
193
     * \struct dCOOmat
     * \brief Sparse matrix of REAL type in COO (or IJ) format
194
195
196
     * Coordinate Format (I,J,A)
197
198
      * \note The starting index of A is 0.
      st \note Change I to rowind, J to colind. To avoid with complex.h confliction on I.
199
200
201
    typedef struct dCOOmat{
202
        //! row number of matrix A, m
203
        INT row;
204
205
        //! column of matrix A, n
206
        INT col;
207
208
209
        //! number of nonzero entries
        INT nnz;
210
211
        //! integer array of row indices, the size is nnz
212
        INT *rowind:
213
214
215
         //! integer array of column indices, the size is nnz
216
        INT *colind;
217
218
         //! nonzero entries of A
219
        REAL *val;
```

```
220 | } dCOOmat; /**< Sparse matrix of REAL type in COO format */
```

So it clear that the sparse matrix in Example 3.1.1 in COO format is stored as:

```
row = 4
col = 5
nnz = 12

I J val

0 0 1.0
0 1 1.5
0 4 12.0
1 1 1.0
1 2 6.0
1 3 7.0
1 4 1.0
......
```

Although the COO format is easy to understand or use, it wastes storage space and has little advantages in sparse BLAS operations.

NOTE: In FASP, the indices always start from 0, instead of from 1. This is often the source of problems related to vectors and matrices.

(ii) CSR format

The most commonly used data structure for sparse matrices nowadays is probably the so-called compressed sparse row (CSR) format, according to Saad [15]. The compressed row storage format of a matrix $A \in \mathbb{R}^{n \times m}$ (n rows and m columns) consists of three arrays, as follows:

- 1. An integer array of row pointers of size n+1;
- 2. An integer array of column indexes of size nnz;
- 3. An array of actual matrix entries.

In FASP, we define:

```
/**

132  /**

133  * \struct dCSRmat

134  * \brief Sparse matrix of REAL type in CSR format

135  *

136  * CSR Format (IA, JA, A) in REAL

137  *

138  * \note The starting index of A is 0.
```

```
139
    typedef struct dCSRmat{
140
141
         //! row number of matrix A, m
142
143
         INT row;
144
         //! column of matrix A, n
145
146
         INT col;
147
         //! number of nonzero entries
148
         INT nnz;
149
150
         //! integer array of row pointers, the size is m+1
151
152
         INT *IA;
153
         //! integer array of column indexes, the size is nnz
154
155
156
         //! nonzero entries of A
157
         REAL *val;
158
159
    } dCSRmat; /**< Sparse matrix of REAL type in CSR format */
160
```

The matrix (only nonzero elements) is stored in the array val row after row, in a way that i-th row begins at val(IA(i)) and ends at val(IA(i+1)-1). In the same way, JA(IA(i)) to JA(IA(i+1)-1) will contain the column indexes of the non-zeros in row i. Thus IA is of size n+1 (number of rows in val plus one), JA and val are of size equal to the number of non-zeroes. The total number of non-zeroes is equal to IA(n+1)-1.

NOTE: When the sparse matrix A is a boolean (i.e. all entries are either 0 or 1), the actual non-zeroes are not stored because it is understood that, if it is nonzero, it could only be 1 and there is no need to store it.

The matrix in Example 3.1.1 in CSR format is represented in the following way:

• IA is of size 5 and

$$IA = \parallel 0 \parallel 3 \parallel 7 \parallel 9 \parallel 12 \parallel$$

• JA is of size IA(5) - 1 = 12

• val is of the same size as JA and

$$val = || 1. | 1.5 | 12. || 1. | 7. | 6. | 1. || 3. | 6. || 2. | 5. | 1. ||$$

Here we use double vertical bars to separate rows and single vertical bars to separate values.

NOTE: The indices in JA and entries of val does NOT have to be ordered as seen in this example. Sometimes they are sorted in ascending order in each row. More often, the diagonal entries are stored in the first position in each row and the rest are sorted in ascending order.

Below is a "non-numeric" example.

Example 3.1.2 Consider the following sparse matrix:

$$\begin{pmatrix}
a_{11} & 0 & a_{13} & 0 \\
0 & a_{22} & a_{23} & a_{24} \\
0 & a_{32} & 0 & a_{34} \\
a_{41} & a_{42} & a_{43} & 0
\end{pmatrix}$$

For this matrix, we have that the number of non-zeros nnz = 10. Furthermore, the three arrays of in the CSR format are:

$$IA = \left\| \begin{array}{c|c|c} 0 & 2 & 5 & 7 \end{array} \right\|,$$

$$JA = \left\| \begin{array}{c|c|c} 0 & 2 & 1 & 2 & 3 & 1 & 3 & 0 & 1 & 2 \end{array} \right\|,$$

and

$$val = || a_{11} | a_{13} || a_{22} | a_{23} | a_{24} || a_{32} | a_{34} || a_{41} | a_{42} | a_{43} ||.$$

NOTE: The CSR format presents challenges to sparse matrix-vector product mainly because of the high cache missing rate due to indirect memory access and irregular access pattern. In order to reduce the cache missing rate, we introduce an improved data format, CSRL.

(iii) CSRL format

CSRL matrix format [10] groups rows with same number of nonzeros together and improves cache hitting rate.

```
253
254
      * \struct dCSRLmat
255
      * \brief Sparse matrix of REAL type in CSRL format
256
    typedef struct dCSRLmat{
257
258
         //! number of rows
259
         INT row:
260
261
262
         //! number of cols
         INT col;
263
264
265
         //! number of nonzero entries
266
         INT nnz;
```

```
267
268
         //! number of different values in i-th row, i=0:nrows-1
269
270
         //! nz_diff[i]: the i-th different value in 'nzrow'
271
         INT *nz_diff;
272
273
         //! row index of the matrix (length-grouped): rows with same nnz are together
274
275
         INT *index;
276
         //! j in {start[i],...,start[i+1]-1} means nz_diff[i] nnz in index[j]-row
277
278
         INT *start;
279
         //! column indices of all the nonzeros
280
281
         INT * ja;
282
283
         //! values of all the nonzero entries
        REAL *val;
284
285
      dCSRLmat; /**< Sparse matrix of REAL type in CSRL format */
286
```

3.2 Block sparse matrices

For PDE applications, we often need to solve systems of partial differential equations. Many iterative methods and preconditioners could take advantages of the structure of PDE systems and improve efficiency. So we often need to use semi-structured (block) sparse data structures to store the coefficient matrix arising from PDE systems.

Depending on different applications and different solving algorithms, we can use two types of block matrices: dBSRmat (or BSR Block Compressed Sparse Row) and block_dCSRmat (CSR Block or Block of CSR matrices).

For more details as well as other specialized block matrices, readers are referred to the header file "base/include/fasp_block.h".

As an example, we consider the following matrix, which have been used in §3.1 for the CSR format. We add structure to this matrix and divide it as a 2×2 block matrix:

Example 3.2.1

$$\begin{pmatrix}
a_{11} & 0 & a_{13} & 0 \\
0 & a_{22} & a_{23} & a_{24} \\
\hline
0 & a_{32} & 0 & a_{34} \\
a_{41} & a_{42} & a_{43} & 0
\end{pmatrix}$$

(i) BSR format

This format is a standard data structure for storing block sparse matrices which has been used by the Intel MKL library.

```
33
34
    * \struct dBSRmat
    * \brief Block sparse row storage matrix of REAL type
35
    * \note This data structure is adapted from the Intel MKL library. Refer to:
37
    * http://software.intel.com/sites/products/documentation/hpc/mkl/lin/index.htm
38
39
    * \note Some of the following entries are capitalized to stress that they are
40
            for blocks!
41
42
43
   typedef struct dBSRmat {
44
45
        //! number of rows of sub-blocks in matrix A, M
46
        INT ROW;
47
       //! number of cols of sub-blocks in matrix A, N \,
48
       INT COL;
49
50
       //! number of nonzero sub-blocks in matrix A, NNZ
51
52
53
       //! dimension of each sub-block
54
       INT nb; // NOTE: for the moment, allow nb*nb full block
55
56
        //! storage manner for each sub-block
57
        INT storage_manner; // 0: row-major order, 1: column-major order
58
59
60
        //! A real array that contains the elements of the non-zero blocks of
61
        //! a sparse matrix. The elements are stored block-by-block in row major
62
       //! order. A non-zero block is the block that contains at least one non-zero
63
       //! element. All elements of non-zero blocks are stored, even if some of
       //! them is equal to zero. Within each nonzero block elements are stored
64
65
        //! in row-major order and the size is (NNZ*nb*nb).
       REAL *val;
66
67
       //! integer array of row pointers, the size is ROW+1
68
69
70
       //! Element i of the integer array columns is the number of the column in the
71
72
        //! block matrix that contains the i-th non-zero block. The size is NNZ.
73
       INT *JA;
74
   } dBSRmat; /**< Matrix of REAL type in BSR format */
```

For the matrix in Example 3.2.1, we have that the number of block rows ROW = 2, the number of block columns COL = 2, and the number of block nonzeros NNZ = 4. The block size is nb = 2.

We can choose different storage manners for storing the small blocks. Suppose that we set it to be 0, i.e. row-major format. Then the three arrays of in the BSR format are:

$$IA = | 0 | 8 | 16 |,$$

$$JA = | 0 | 1 | 0 | 1 |,$$

and

$$val = || a_{11} | 0 | 0 | a_{22} || a_{13} | 0 | a_{23} | a_{24} ||$$
$$|| 0 | a_{32} | a_{41} | a_{42} || 0 | a_{34} | a_{43} | 0 ||.$$

We immediately notice that this format might be not be the best choice for this particular matrix due to all the blocks are nonzero blocks, i.e., contain nonzero entries. However, for PDE applications, this does not usually happen.

(ii) CSR Block format

This format is simple and is derived from the dCSRmat data structure. The following definition explains itself.

```
77
      \struct block_dCSRmat
78
      \brief Block REAL CSR matrix format
79
80
81
       \note The starting index of A is 0.
82
   typedef struct block_dCSRmat {
83
84
        //! row number of blocks in A, m
85
        INT brow;
86
87
        //! column number of blocks A, n
88
        INT bcol;
89
90
        //! blocks of dCSRmat, point to blocks[brow][bcol]
91
92
        dCSRmat **blocks;
93
   } block_dCSRmat; /**< Matrix of REAL type in Block CSR format */
94
```

3.3 I/O subroutines for sparse matrices

In FASP, we provided several functions for reading, writing, and printing different formats of sparse matrices in plain text or binary formats. These functions can be found in "base/src/io.c" and we list the available functions as follows:

```
/*----*/
834
835
    void fasp_dcsrvec1_read (const char *filename,
836
837
                              dCSRmat *A,
838
                              dvector *b);
839
    void fasp_dcsrvec2_read (const char *filemat,
840
                              const char *filerhs,
841
842
                              dCSRmat *A,
843
                              dvector *b );
844
845
    void fasp_dcsr_read (const char *filename,
846
                          dCSRmat *A);
847
    void fasp_dcoo_read (const char *filename,
848
                          dCSRmat *A);
849
850
    void fasp_dcoo1_read (const char *filename,
851
852
                           dCOOmat *A);
853
    void fasp_dcoo_shift_read (const char *filename,
854
855
                                dCSRmat *A);
856
    void fasp_dmtx_read (const char *filename,
857
858
                          dCSRmat *A);
859
860
    void fasp_dmtxsym_read (const char *filename,
                             dCSRmat *A);
861
862
863
    void fasp_dstr_read (const char *filename,
                          dSTRmat *A);
864
865
    void fasp_dbsr_read (const char *filename,
866
                          dBSRmat *A);
867
868
869
    void fasp_dvecind_read (const char *filename,
                             dvector *b);
870
871
872
    void fasp_dvec_read (const char *filename,
873
                          dvector *b);
874
875
    void fasp_ivecind_read (const char *filename,
876
                             ivector *b);
877
878
    void fasp_ivec_read (const char *filename,
879
                          ivector *b);
880
    void fasp_dcsrvec1_write (const char *filename,
881
882
                               dCSRmat *A,
883
                               dvector *b);
884
885
    void fasp_dcsrvec2_write (const char *filemat,
886
                             const char *filerhs,
```

```
dCSRmat *A,
887
888
                                dvector *b );
889
    void fasp_dcoo_write (const char *filename,
890
                            dCSRmat *A);
891
892
893
    void fasp_dstr_write (const char *filename,
894
                            dSTRmat *A);
895
    void fasp_dbsr_write (const char *filename,
896
897
                            dBSRmat *A);
898
899
    void fasp_dvec_write (const char *filename,
                            dvector *vec);
900
901
902
    void fasp_dvecind_write (const char *filename,
903
                              dvector *vec);
904
905
    void fasp_ivec_write (const char *filename,
906
                           ivector *vec);
907
908
    void fasp_dvec_print (INT n,
909
                            dvector *u);
910
    void fasp_ivec_print (INT n,
911
912
                            ivector *u);
913
    void fasp_dcsr_print (dCSRmat *A);
914
915
    void fasp_dcoo_print (dCOOmat *A);
916
917
    void fasp_dbsr_print (dBSRmat *A);
918
919
    void fasp_dbsr_write_coo (const char *filename,
920
921
                                const dBSRmat *A);
922
    void fasp_dcsr_write_coo (const char *filename,
923
                                const dCSRmat *A);
924
925
    void fasp_dstr_print (dSTRmat *A);
926
927
928
    void fasp_matrix_read (const char *filename,
929
                            void *A);
930
931
    void fasp_matrix_read_bin (const char *filename,
932
                                 void *A);
933
934
    void fasp_matrix_write (const char *filename,
935
                              void *A,
936
                              INT flag);
937
938
    void fasp_vector_read (const char *filerhs,
939
                           void *b);
```

3.4. SPARSE BLAS

```
940
941 void fasp_vector_write (const char *filerhs,
942 void *b,
943
944
945 void fasp_hb_read (const char *input_file,
946 dCSRmat *A,
947 dvector *b);
```

NOTE: The above function decorations are taken from "base/include/fasp_functs.h". This header file is automatically generated based on the source codes. Users are discouraged from changing it by hand; their changes may be lost.

3.4 Sparse BLAS

The matrix-vector multiplication: y = Ax can be performed in the following simple way:

```
1
 2
     * \fn void fasp_blas_dcsr_mxv (dCSRmat *A, REAL *x, REAL *y)
 3
 4
      * \brief Matrix-vector multiplication y = A*x
 5
 6
      * \param A Pointer to dCSRmat matrix A
 7
      * \param x
                    Pointer to array x
     * \param y Pointer to array y
 8
 9
      * \author Chensong Zhang
10
     * \date 07/01/2009
11
12
13
    void fasp_blas_dcsr_mxv (dCSRmat *A,
14
                                   REAL *x.
                                   REAL *y)
15
16
    {
17
         const INT
                      m = A \rightarrow row;
         const INT *ia = A->IA, *ja = A->JA;
18
19
         const REAL *aj = A->val;
20
21
         {\tt INT \ i} \;,\;\; {\tt k} \;,\;\; {\tt beg} \;,\;\; {\tt end} \;;
         register REAL tmp;
22
23
24
         for (i=0; i< m; ++i) {
              \mathtt{tmp} \; = \; 0.0;
25
              \mathtt{beg} \, = \, \mathtt{ia}\,[\,\mathtt{i}\,]\,; \ \mathtt{end} \, = \, \mathtt{ia}\,[\,\mathtt{i}\,+1\,];
^{26}
              27
28
              y[i] = tmp;
         }
29
    }
```

This is only a simple example for sparse matrix-vector multiplication (SpMV) kernel. Since we need many types of sparse matrices, there are various of versions of SpMV for different data structures. See the Reference Manual for more details.

3.5 Iterative methods

In FASP, there are a couple of standard preconditioned iterative methods [15] implemented, including preconditioned CG, BiCGstab, GMRES, Variable Restarting GMRES, Flexible GMRES, etc. In this section, we use the CSR matrix format as example to introduce how to call these iterative methods. To learn more details, we refer to the Reference Manual.

We first notice the abstract interface for the iterative methods. The following code segment is taken from "base/src/itsolver_csr.c":

```
20
21
    * \fn INT fasp_solver_dcsr_itsolver (dCSRmat *A, dvector *b, dvector *x,
22
                                           precond *pc, itsolver_param *itparam)
23
      \brief Solve Ax=b by preconditioned Krylov methods for CSR matrices
24
25
      \param A
                       Pointer to the coeff matrix in dCSRmat format
26
27
      \param b
                       Pointer to the right hand side in dvector format
      \param x
                       Pointer to the approx solution in dvector format
28
29
     * \param pc
                       Pointer to the preconditioning action
      \param itparam Pointer to parameters for iterative solvers
30
31
32
                       Iteration number if converges; ERROR otherwise.
     * \return
33
     * \author Chensong Zhang
34
     * \date 09/25/2009
35
36
      \note This is an abstract interface for iterative methods.
37
38
   INT fasp_solver_dcsr_itsolver (dCSRmat *A,
39
40
                                    dvector *b,
41
                                    dvector *x,
42
                                    precond *pc,
43
                                    itsolver_param *itparam)
```

The names of the input arguments explain themselves mostly and they are explained in the Reference Manual in detail.

We briefly discuss how to call this function; and, once you understand PCG, you can easily call other iterative methods.

```
// ILU setup for whole matrix
ILU_data LU;
if ((status = fasp_ilu_dcsr_setup(A,&LU,iluparam)) < 0 ) goto FINISHED;
```

```
466
467
         // check iludata
468
         if ( (status = fasp_mem_iludata_check(&LU)) < 0 ) goto FINISHED;</pre>
469
         // set preconditioner
470
471
         precond pc;
472
         pc.data = \&LU;
473
         pc.fct = fasp_precond_ilu;
474
475
         // call iterative solver
         status = fasp_solver_dcsr_itsolver(A,b,x,&pc,itparam);
476
```

Now we explain this code segment a little bit:

- Line 463–465 performs the setup phase for ILU method. The particular type of ILU method is determined by "iluparam"; see §2.4. Line 7 performs a simple memory check for ILU.
- Line 471–473 defines the preconditioner data structure "pc", which contains two parts: one is the actual preconditioning action "pc.fct", the other is the auxiliary data which is needed to perform the preconditioning action "pc.data".
- Line 476 calls iterative methods. "A" is the matrix in dCSRmat format; "b" and "x" are the right-hand side and the solution vectors, respectively. Similar to ILU setup, the type of iterative methods is determined by "itparam".

Apparently, we now left with no choice but introducing "itparam".

```
/**
  * \struct itsolver_param
  * \brief Parameters passed to iterative solvers
  */
typedef struct {

    SHORT itsolver_type; /**< solver type: see message.h */
    SHORT precond_type; /**< preconditioner type: see message.h */
    SHORT stop_type; /**< stopping criteria type */
    INT maxit; /**< max number of iterations */
    REAL tol; /**< convergence tolerance */
    INT restart; /**< number of steps for restarting: for GMRES etc */
    SHORT print_level; /**< print level: 0--10 */

} itsolver_param; /**< Parameters for iterative solvers */</pre>
```

Possible "itsolver_type" includes:

```
/**

* \brief Definition of solver types for iterative methods

*/

#define SOLVER_DEFAULT

0 /**< Use default solver in FASP */
```

```
#define SOLVER_CG
                                           1 /**< Conjugate Gradient */</pre>
#define SOLVER_BiCGstab
                                           2 /**< Bi-Conjugate Gradient Stabilized */</pre>
#define SOLVER_MinRes
                                            3 /**< Minimal Residual */
#define SOLVER_GMRES
                                            4 /** Generalized Minimal Residual */
#define SOLVER_VGMRES
                                             5 /**< Variable Restarting GMRES */
#define SOLVER_VFGMRES 6  /**< Variable Restarting Flexible GMRE:
#define SOLVER_GCG 7  /**< Generalized Conjugate Gradient */
#define SOLVER_GCR 8  /**< Generalized Conjugate Residual */</pre>
                                            6 /**< Variable Restarting Flexible GMRES */
#define SOLVER_SCG 11 /**< Conjugate Gradient with safe net */
#define SOLVER_SBiCGstab 12 /**< BiCGstab with safe net */
#define SOLVER_SMinRes 13 /**< MinRes with safe net */
#define SOLVER_SGMRES
                                          14 /**< GMRes with safe net */
#define SOLVER.SGMRES 14 /**< GMMes with safe net */
#define SOLVER.SVGMRES 15 /**< Variable-restart GMRES with safe net */
#define SOLVER.SVFGMRES 16 /**< Variable-restart FGMRES with safe net */
                                          16 /**< Variable-restart FGMRES with safe net */
#define SOLVER_SGCG
                                          17 /**< GCG with safe net */
#define SOLVER_AMG 21 /**< AMG as an iterative solver */
#define SOLVER_FMG 22 /**< Full AMG as an solver */
```

3.6 Geometric multigrid

The geometric multigrid method (GMG) is one of the most efficient solving techniques for discrete algebraic systems arising from many types of partial differential equations [1, 16]. GMG utilizes a hierarchy of grids or discretizations and reduces the error at a number of frequencies simultaneously. Because of its plausible linear complexity—i.e., the low computational cost of solving a linear system with N unknowns is O(N)—the GMG method is one of the most popular Poisson solvers. Although the GMG's applicability is limited as it requires explicit information on the hierarchy of the discrete system, when it can be applied, GMG is far more efficient than its algebraic version, the algebraic multigrid (AMG) method.

We now give a simple example on calling the geometric multigrid for solving the Poisson's equation in 2D (discretized by the standard five-point finite difference stencil). Consider the Poisson equation

$$\begin{cases}
-\Delta u &= f & \text{in } \Omega \\
u &= 0 & \text{on } \partial\Omega,
\end{cases}$$

where $\Omega = (0,1)^2 \subset \mathbb{R}^2$. The main reason why we choose this simplest possible setting is to emphasize that, even for a simple problem, the new heterogeneous architectures present challenges for numerical implementation. Another reason is to allow us to use explicit stencils and to avoid the bottleneck of sparse matrix-vector production. The standard central finite difference method is applied to discretize the Poisson's equation. In other words, the Laplace operator is discretized by the classical second-order central difference scheme. After discretization, we end up with a system

of linear equations:

$$\mathbf{A}\vec{u} = \vec{f}$$
.

We use the five-point central difference scheme in 2D. Consider a uniform square mesh of $\Omega = [0,1]^2$ with size $h = \frac{1}{n}$ and in which $x_i = ih$, $y_j = jh$ (i, j = 0, 1, ..., n). Let $u_{i,j}$ be the numerical approximation of $u(x_i, y_j)$. The five-point central difference scheme for solving the Poisson's equation in 2D can be written as follows:

$$-u_{i-1,j} - u_{i,j-1} + 4u_{i,j} - u_{i+1,j} - u_{i,j+1} = h^2 f(x_i, y_i)$$
 $i, j = 1, 2, \dots, n-1$.

The sample code for this solver can be found in "test/main/testgmg.c" and a piece of the source code is listed as follows:

```
case 2: // 2 dimension
254
255
                   u = (REAL *) malloc((nx+1)*(ny+1)*sizeof(REAL));
256
257
                   fasp_array_set((nx+1)*(ny+1), u, 0.0);
258
                   b = (REAL *) malloc((nx+1)*(ny+1)*sizeof(REAL));
259
                   for (i = 0; i \le nx; i++) {
260
                        for (j = 0; j \le ny; j++) {
261
                             {\tt b\,[\,j*(\,nx+1)+i\,]\,\,=\,\,h*h*f2d\,(\,i\,\,,\,\,\,j\,\,,\,\,\,nx\,\,,\,\,\,ny\,)\,\,;}
262
263
264
265
                   {\tt switch} (method) {
266
267
268
                        case 1: // V-cycle
                             fasp_poisson_gmg_2D(u, b, nx, ny, maxlevel, rtol, prtlvl);
269
270
271
                        case 2: // FMG
272
                             fasp_poisson_fgmg_2D(u, b, nx, ny, maxlevel, rtol, prtlvl);
273
274
275
                        case 3: // PCG
276
                             fasp_poisson_pcg_gmg_2D(u, b, nx, ny, maxlevel, rtol, prtlvl);
277
278
                             break;
279
280
                   }
281
282
                   break:
```

3.7 Algebraic multigrid

The classical algebraic multigrid method [14] is an important component in many of our auxiliary space preconditioners. Because of its user-friendly and scalability, AMG becomes increasingly

popular in scientific and engineering computing, especially when GMG is difficult or not possible to be applied. Various of new AMG techniques [17, 18, 4, 7, 5, 9, 6, 20, 3, 11, 8] have emerged in recent years.

The following code segment is part of "base/src/amg.c" and it is a good example which shows how to call different AMG methods (classical AMG, smoothed aggregation, un-smoothed aggregation) and different multilevel iterative methods (V-cycle, W-cycle, AMLI-cycle, Nonlinear AMLI-cycle, etc).

```
42
        const SHORT
                        max_levels = param->max_levels;
                       prtlvl
43
        const SHORT
                                     = param->print_level;
        const SHORT amg_type
                                     = param->AMG_type;
44
45
        const SHORT cycle_type = param->cycle_type;
46
        const INT
                        nnz = A \rightarrow nnz, m = A \rightarrow row, n = A \rightarrow col;
47
        // local variables
48
49
        AMG_data *
                       mgl = fasp_amg_data_create(max_levels);
50
        REAL
                       AMG_start , AMG_end;
51
52
53
    #if DEBUG_MODE > 0
        printf("### DEBUG: %s ..... [Start]\n", __FUNCTION__);
54
55
    #endif
56
        if ( prtlvl > PRINT_NONE ) fasp_gettime(&AMG_start);
57
58
        // check matrix data
59
60
        if ( m != n ) {
             printf("### ERROR: A is not a square matrix!\n");
61
             {\tt fasp\_chkerr} \left( \, {\tt ERROR\_MAT\_SIZE} \; , \; \; {\tt \_\_FUNCTION\_\_} \right) \; ;
62
63
        }
64
        if (nnz \ll 0)
65
             printf("### ERROR: A has no nonzero entries!\n");
66
             fasp_chkerr(ERROR_MAT_SIZE, __FUNCTION__);
67
68
        }
69
        // Step 0: initialize mgl[0] with A, b and x
70
        mgl[0].A = fasp_dcsr_create(m, n, nnz);
71
72
        fasp_dcsr_cp(A, \&mgl[0].A);
73
        mgl[0].b = fasp_dvec_create(n);
74
75
        fasp_dvec_cp(b, &mgl[0].b);
76
        mgl[0].x = fasp_dvec_create(n);
77
78
        fasp_dvec_cp(x, \&mgl[0].x);
79
80
        // Step 1: AMG setup phase
        switch (amg_type) {
81
82
83
             case SA_AMG: // Smoothed Aggregation AMG setup
```

```
if ( prtlvl > PRINT_NONE ) printf("\nCalling SA AMG ...\n");
84
85
                 status = fasp_amg_setup_sa(mgl, param); break;
86
             case UA_AMG: // Unsmoothed Aggregation AMG setup
87
                 if ( prtlvl > PRINT_NONE ) printf("\nCalling UA AMG ...\n");
88
89
                 status = fasp_amg_setup_ua(mgl, param); break;
90
             default: // Classical AMG setup
91
92
                 if ( prtlvl > PRINT_NONE ) printf("\nCalling classical AMG ...\n");
93
                 status = fasp_amg_setup_rs(mgl, param);
94
95
         }
96
         // Step 2: AMG solve phase
97
         if ( status == FASP_SUCCESS ) { // call a multilevel cycle
98
99
             switch (cycle_type) {
100
101
                 case AMLI_CYCLE: // AMLI-cycle
102
                      fasp_amg_solve_amli(mgl, param); break;
103
104
                 case NL_AMLI_CYCLE: // Nonlinear AMLI-cycle
105
                      fasp_amg_solve_nl_amli(mgl, param); break;
106
107
                 default: // V,W-cycles (determined by param)
108
                      fasp_amg_solve(mgl, param); break;
109
110
             }
111
112
             fasp_dvec_cp(\&mgl[0].x, x);
113
114
115
116
         else { // call a backup solver
117
118
             if ( prtlvl > PRINT_MIN ) {
119
                 printf("### WARNING: AMG setup failed!\n");
120
                 printf("### WARNING: Use a backup solver instead.\n");
121
122
             {\tt fasp\_solver\_dcsr\_spgmres \ (A\,,\ b\,,\ x\,,\ NULL\,,\ param->tol\,,\ param->maxit\,,}
123
124
                                          20, 1, prtlv1);
125
126
         }
```

The code above is very simple and we only wish to point out that:

- Line 42–45 reads some of the parameters from "AMG_param", which can be defined from a input file; see §2.4.
- Line 50–78 initializes the "AMG_data" with a copy of the coefficient matrix, the right-hand side, and the initial solution (it will store the final solution eventually).

- Line 81–95 calls three different AMG setup methods, determined by "amg_type".
- Line 98–115 calls three different multilevel iterative methods, determined by "cycle_type".

Parameters for AMG

There are a couple of controlling parameters for algebraic multigrid methods in FASP. Basically, there are four types of parameters for AMG—They control multilevel iterations, smoothing, classical AMG setup, and aggregation AMG setup. The following is a sample from "test/ini/input.dat" and a brief explanation of each parameter is given.

```
-%
55
                                                     %
56
   \% parameters for multilevel iteration
57
                                                     -%
58
59
   AMG_type
                              = C
                                       \% C classic AMG
60
                                       \% SA smoothed aggregation
61
                                       \% UA unsmoothed aggregation
                                       \% V V-cycle | W W-cycle
62
   AMG_cycle_type
                              = V
                                       \% A AMLI-cycle | NA Nonlinear AMLI-cycleA
63
   AMG_tol
                              = 1e-6
                                       \% tolerance for AMG
64
   AMG_maxit
                                       \% number of AMG iterations
                              = 1
65
   AMG_levels
                              = 20
                                       \% max number of levels
66
   AMG_coarse_dof
                              = 500
                                       \% max number of coarse degrees of freedom
67
                                       \% coarsest solver: 0 iterative |
   AMG_coarse_solver
                              = 0
68
                                       \% 31 SuperLU | 32 UMFPack | 33 MUMPS
69
   AMG_coarse_scaling
                              = OFF
                                       \% switch of scaling of the coarse grid correction
70
   AMG_amli_degree
                              = 2
                                       \% degree of the polynomial used by AMLI cycle
71
                                       \% Krylov method in NLAMLI cycle: 6 FGMRES | 7 GCG
   AMG_nl_amli_krylov_type = 6
72
73
                                                     -%
74
   \% parameters for AMG smoothing
                                                     %
75
                                                     -%
76
   %-
77
                                       % GS | JACOBI | SGS SOR | SSOR
78
   AMG_smoother
                              = GS
                                       \% GSOR | SGSOR | POLY | L1DIAG | CG
79
   AMG_smooth_order
                              = CF
                                       \% NO: natural order | CF: CF order
80
   AMG_ILU_levels
                              = 0
                                       \% number of levels using ILU smoother
81
                                       \% number of levels using Schwarz smoother
   {\tt AMG\_Schwarz\_levels}
                              = 0
82
   AMG_relaxation
                              = 1.0
                                       \% relaxation parameter for SOR smoother
   AMG_polynomial_degree
                              = 3
                                       \% degree of the polynomial smoother
84
   AMG_presmooth_iter
                              = 1
                                       \% number of presmoothing sweeps
85
86
   AMG_postsmooth_iter
                              = 1
                                       \% number of postsmoothing sweeps
87
                                                     -%
88
                                                     %
89
   \% parameters for classical AMG SETUP
90
   %
                                                     -%
91
                                       \% 1 Modified RS
92
   AMG_coarsening_type
                              = 1
93
                                       \% 3 Compatible Relaxation
94
                                       \% 4 Aggressive
```

```
{\tt AMG\_interpolation\_type} \quad = \quad 1 \qquad \qquad \% \ 1 \ {\tt Direct} \ | \ 2 \ {\tt Standard} \ | \ 3 \ {\tt Energy-min}
    AMG_strong_threshold
                               = 0.3
                                         \% Strong threshold
97
    AMG_truncation_threshold = 0.1
                                         \% Truncation threshold
   AMG_max_row_sum
                              = 0.9
                                         \% Max row sum
98
99
100
                                                       -%
                                                       %
    \% parameters for aggregation—type AMG SETUP
101
                                                       -%
102
103
                                         \% 1 Matching | 2 VMB
104
    AMG_aggregation_type
                               = 2
    AMG_pair_number
                               = 2
                                         \% Number of pairs in matching
105
106
    AMG_strong_coupled
                               = 0.08
                                         \% Strong coupled threshold
107
    AMG_max_aggregation
                               = 20
                                         \% Max size of aggregations
    AMG_tentative_smooth
                              = 0.67
                                         \% Smoothing factor for tentative prolongation
108
                               = OFF
                                         \% Switch for filtered matrix for smoothing
109
    AMG_smooth_filter
110 AMG_quality_bound
                               = 8.0
                                         \% quality of aggregation: 8.0~{
m sysmm} | 10.0~{
m unsymm}
```

NOTE: Here we can not discuss the details of these parameters as a full discussion requires more understand of the underlying algorithms which we have completely omitted. So to learn more about, we refer to the Reference Manual.

Chapter 4

More advanced features

In this chapter, we discuss a few more advanced features of FASP. We will discuss parallel versions of FASP and its build-in features for debugging purposes. These features will be helpful for people who would like to develop on the top of FASP. For users who only wish to call a few standard solvers, they can skip this chapter.

4.1 An OpenMP example

OpenMP¹ (Open Multiprocessing) is an API that supports multi-platform shared memory multi-processing programming in C, C++, and Fortran, on most processor architectures and operating systems. It consists of a set of compiler directives, library routines, and environment variables that influence run-time behavior. Some preliminary OpenMP support has been included since the very beginning of FASP. We consistently improves and expands OpenMP support as multiprocessor architectures become the dominant desktop computing environment.

NOTE: By default, OpenMP is disabled in FASP. In order to turn it on, you need to modify FASP.mk slightly as follows.

To enable OpenMP support, you need to uncomment one line in "FASP.mk" and set "openmp" to be "yes".

```
# You may use multithread version after you enable OpenMP support. To
# setup the environment, you need
# >> export OMP_NUM_THREADS=4 (for bash)
# >> setenv OMP_NUM_THREADS 4 (for tcsh)
# If you want to compile with OpenMP support, uncomment the next line:
# #
```

¹Official website: http://openmp.org/

```
44  # openmp=yes
45  #
```

After you build FASP with "openmp=yes", OpenMP is turned on and the number of threads is determined by the environment variable OMP_NUM_THREADS. For example, to use 8 threads in sh/bash, you need to set:

```
$ export OMP_NUM_THREADS=8
```

4.2 Predefined constants

FASP also many predefined constants used in the source files. Using these macros makes the source codes more readable. These constants are defined in "base/include/fasp_const.h" and a printout of this file is below:

```
1
    /*! \file fasp_const.h
2
       \brief Definition of all kinds of messages, including error messages,
3
               solver types, etc.
4
       \note This is internal use only. Do NOT change.
5
6
7
    * Created by Chensong Zhang on 03/20/2010.
8
    * Modified by Chensong Zhang on 12/06/2011.
9
10
    * Modified by Chensong Zhang on 12/25/2011.
     * Modified by Chensong Zhang on 04/22/2012.
11
12
       Modified by Ludmil Zikatanov on 02/15/2013: CG \rightarrow SMOOTHER_CG.
       Modified by Chensong Zhang on 02/16/2013: GS \rightarrow SMOOTHER_GS, etc.
13
    st Modified by Chensong Zhang on 04/09/2013: Add safe Krylov methods.
14
     st Modified by Chensong Zhang on 09/22/2013: Clean up Doxygen.
15
    * Modified by Chensong Zhang on 09/17/2013: Filename changed from message.h.
16
17
18
19
20
                                       /*-- allow multiple inclusions --*/
21
   #ifndef __FASP_MESSAGES__
   #define __FASP_MESSAGES__
23
24
    * \brief Definition of return status and error messages
25
26
   #define FASP_SUCCESS
                                   0 /**< return from function successfully */</pre>
27
28
29
   #define ERROR_OPEN_FILE
                                  -10 /**< fail to open a file */
30
   #define ERROR_WRONG_FILE
                                  -11 /**< input contains wrong format */
  #define ERROR_INPUT_PAR
                                  -13 /**< wrong input argument */
31
   #define ERROR_REGRESS
                                  -14 /**< regression test fail */
   #define ERROR_MAT_SIZE
                              -15 /**< wrong problem size */
```

```
#define ERROR_NUM_BLOCKS -18 /**< wrong number of blocks */
   #define ERROR_MISC
                               -19 /**< other error */
36
  //---
37 | #define ERROR_ALLOC_MEM -20 /**< fail to allocate memory */
38 #define ERROR_DATA_STRUCTURE -21 /**< problem with data structures */
   \verb|#define ERROR_DATA_ZERODIAG| -22 /** < \verb|matrix has zero diagonal entries */|
39
   #define ERROR_DUMMY_VAR -23 /**< unexpected input data */
40
  //---
41
   #define ERROR_AMG_INTERP_TYPE -30 /**< unknown interpolation type */
   #define ERROR_AMG_SMOOTH_TYPE -31 /**< unknown smoother type */
43
   #define ERROR_AMG_COARSE_TYPE -32 /**< unknown coarsening type */
44
   #define ERROR_AMG_COARSEING -33 /**< coarsening step failed to complete */
45
46
   //----
   #define ERROR_SOLVER_TYPE -40 /**< unknown solver type */
47
  #define ERROR_SOLVER_PRECTYPE -41 /**< unknown precond type */
48
49 #define ERROR_SOLVER_STAG -42 /**< solver stagnates */
50 #define ERROR_SOLVER_SOLSTAG -43 /**< solver's solution is too small */
51 #define ERROR_SOLVER_TOLSMALL -44 /**< solver's tolerance is too small */
52 #define ERROR_SOLVER_ILUSETUP -45 /**< ILU setup error */
53 #define ERROR_SOLVER_MISC -46 /**< misc solver error during run time */
  #define ERROR_SOLVER_MAXIT -48 /**< maximal iteration number exceeded */
54
   #define ERROR_SOLVER_EXIT -49 /**< solver does not quit successfully */
   //----
56
   #define ERROR_QUAD_TYPE -60 /**< unknown quadrature type */
#define ERROR_QUAD_DIM -61 /**< unsupported quadrature dim */
57
58
59
   #define ERROR_LIC_TYPE -80 /**< wrong license type */
60
61
   #define ERROR_UNKNOWN -99 /**< an unknown error type */
63
64
65
    * \brief Definition of logic type
66
                                 1 /**< logic TRUE */
   #define TRUE
67
   #define FALSE
                                  0 /**< logic FALSE */
68
69
70
    * \brief Definition of switch
71
72
   #define ON
                                   1 /**< turn on certain parameter */</pre>
73
   #define OFF
                                   0 /**< turn off certain parameter */</pre>
74
75
   /**
76
   * \brief Print level for all subroutines -- not including DEBUG output
77
78
79
   #define PRINT_NONE
                                  0 /**< silent: no printout at all */</pre>
80
   #define PRINT MIN
                                 1 /**< quiet: print error, important warnings */</pre>
81
  #define PRINT_SOME
                                 2 /**< some: print less important warnings */
82 #define PRINT_MORE
                                 4 /**< more: print some useful debug info */
   #define PRINT_MOST
                                 8 /**< most: maximal printouts, no files */
83
   #define PRINT_ALL
                                 10 /**< all: all printouts, including files */
84
85
86
```

```
* \brief Definition of matrix format
 87
    **/
 88
                                   0 /**< matrix-free format: only mxv action */</pre>
 89 #define MAT_FREE
 90 #define MAT CSR
                                   1 /**< compressed sparse row */</pre>
                                   2 /**< block-wise compressed sparse row */</pre>
91 #define MAT_BSR
    #define MAT_STR
                                   3 /**< structured sparse matrix */</pre>
 92
93 #define MAT_bCSR
                                   4 /**< block matrix of CSR */
 94 #define MAT_bBSR
                                  5 /**< block matrix of BSR for bordered systems */
 95 #define MAT_CSRL
                                   6 /**< modified CSR to reduce cache missing */
   #define MAT_SymCSR
                                  7 /**< symmetric CSR format */
96
97
98
99
     * \brief Definition of solver types for iterative methods
100
    #define SOLVER_DEFAULT 0 /**< Use default solver in FASP */
101
102 //----
103 #define SOLVER_CG
                                  1 /**< Conjugate Gradient */</pre>
#define SOLVER_BiCGstab
                                  2 /**< Bi-Conjugate Gradient Stabilized */</pre>
                                  3 /** Minimal Residual */
    #define SOLVER_MinRes
105
106 #define SOLVER_GMRES
                                   4 /**< Generalized Minimal Residual */
#define SULVER_VGMRES 5 /**< Variable Restarting GMRES */
108 #define SULVER_VFGMRES 6 /**< Variable Restarting Flexible GMRES */
109 #define SULVER GCG 7 /**< C
    #define SOLVER_GCR
                                   8 /** Generalized Conjugate Residual */
110
    //-----
111
    #define SOLVER_SCG 11 /**< Conjugate Gradient with safe net */
112
#define SOLVER_SBiCGstab 12 /**< BiCGstab with safe net */
#define SOLVER_SMinRes 13 /**< MinRes with safe net */
115 #define SOLVER_SGMRES
                                  14 /**< GMRes with safe net */
116 #define SOLVER_SVGMRES
                                  15 /**< Variable-restart GMRES with safe net */
#define SOLVER_SVFGMRES 16 /**< Variable-restart FGMF #define SOLVER_SGCG 17 /**< GCG with safe net */
                                  16 /**< Variable-restart FGMRES with safe net */
119 //---
120 #define SOLVER_AMG 21 /**< AMG as an iterative solver */
122 //---
   #define SOLVER_SUPERLU 31 /**< SuperLU Direct Solver */
#define SOLVER_UMFPACK 32 /**< UMFPack Direct Solver */
123
124
                                  33 /**< MUMPS Direct Solver */
    #define SOLVER_MUMPS
125
126
127
128
    * \brief Definition of iterative solver stopping criteria types
129
130 #define STOP_REL_RES
                                   1 /**< relative residual ||r||/||b|| */
    #define STOP_REL_PRECRES
                                  2 /**< relative B-residual ||r||_B/||b||_B */
131
    #define STOP_MOD_REL_RES
                                   3 /**< modified relative residual ||r||/||x|| */</pre>
133
134
135
    * \brief Definition of preconditioner type for iterative methods
136
137 #define PREC_NULL
                                   0 /**< with no precond */</pre>
    #define PREC_DIAG
                                   1 /**< with diagonal precond */</pre>
138
139 #define PREC_AMG 2 /**< with AMG precond */
```

```
140#define PREC_FMG3/**< with full AMG precond */</th>141#define PREC_ILU4/**< with ILU precond */</td>
142 #define PREC_SCHWARZ
                                  5 /**< with Schwarz preconditioner */
143
144 /**
    * \brief Type of ILU methods
145
146 */
                                  1 /**< ILUk */
147 #define ILUk
148 #define ILUt
                                  2 /**< ILUt */
                                  3 /**< ILUtp */
149 #define ILUtp
150
151
152
    * \brief Type of Schwarz smoother
153
4 #define SCHWARZ_FORWARD 1 /**< Forward ordering */
155 #define SCHWARZ_BACKWARD
                                  2 /**< Backward ordering */
156 #define SCHWARZ SYMMETRIC
                                  3 /** Symmetric smoother */
157
158
159
    * \brief Definition of AMG types
160
161#define CLASSIC_AMG1 /**< classic AMG */</td>162#define SA_AMG2 /**< smoothed aggregation</td>
                                  2 /**< smoothed aggregation AMG */
163 #define UA_AMG
                                  3 /**< unsmoothed aggregation AMG */</pre>
164
165
166
    * \brief Definition of aggregation types
167 */
168 #define PAIRWISE
                                  1 /**< pairwise aggregation */</pre>
169 #define VMB
                                   2 /**< VMB aggregation */
170
171
172
    * \brief Definition of cycle types
173 | */
175 #define W_CYCLE 1 /**< V-cycle */
176 #define AMLI_CYCLE
                                  3 /**< AMLI-cycle */
   #define AMLI_CYCLE 3 /**< AMLI-cycle */
#define NL_AMLI_CYCLE 4 /**< Nonlinear AMLI-cycle */
177
178
179
180
   * \brief Definition of standard smoother types
181 */
182 #define SMOOTHER_JACOBI 1 /**< Jacobi smoother */
183 #define SMOOTHER_GS
                                  2 /**< Gauss-Seidel smoother */
                                  3 /**< Symmetric Gauss-Seidel smoother */
184 #define SMOOTHER_SGS
185 #define SMOOTHER_CG
                                  4 /**< CG as a smoother */
5 /**< SOR smoother */
186 #define SMOOTHER_SOR
                                  6 /**< SSOR smoother */
187 #define SMOOTHER_SSOR
188 #define SMOOTHER_GSOR
                                 7 /**< GS + SOR smoother */
                                  8 /**< SGS + SSOR smoother */
189 #define SMOOTHER_SGSOR
190 #define SMOOTHER_POLY
                                  9 /**< Polynomial smoother */
#define SMOOTHER_LIDIAG 10 /**< L1 norm diagonal scaling smoother */
192
```

```
193 /**
    *\ \brief Definition of coarsening types
194
196 #define COARSE RS
                                  1 /**< Classical */
197 #define COARSE_RSP
                                  2 /**< Classical, with positive offdiags */
198 #define COARSE_CR
                                  3 /**< Compatible relaxation */</pre>
199 #define COARSE_AC
                                  4 /**< Aggressive coarsening */
200 #define COARSE_MIS
                                  5 /**< Aggressive coarsening based on MIS */
201
202
    *\ \brief Definition of interpolation types
203
204
205 #define INTERP_DIR 1 /**< Direct interpolation */
206 #define INTERP_STD
                                   2 /**< Standard interpolation */</pre>
207 #define INTERP_ENG
                                  3 /**< energy minimization interpolation */</pre>
209 /**
    * \brief Type of vertices (DOFs) for coarsening
210
211
212 #define GOPT
                                   -5 /**< Cannot fit in aggregates */
213 #define UNPT
                                  -1 /**< Undetermined points */
214 #define FGPT
                                   0 /**< Fine grid points */
                                  1 /**< Coarse grid points */
215 #define CGPT
216 #define ISPT
                                   2 /**< Isolated points */
217
218
219
   * \brief Definition of smoothing order
220 */
221 #define NO_ORDER
                                  0 /**< Natural order smoothing */</pre>
222 #define CF ORDER
                                   1 /**< C/F order smoothing */</pre>
223
224
    st \brief Type of ordering for smoothers
225
226
227 #define USERDEFINED 0 /**< User defined order */
228 #define CPFIRST
                                  1 /**< C-points first order */
229 #define FPFIRST
                                  -1 /**< F-points first order */
                                 12 /**< Ascending order */
230 #define ASCEND
                       21 /**< Descending order */
    #define DESCEND
231
232
233 /**
234 * \brief Some global constants
235 */
236 #define BIGREAL
                               1e+20 /**< A large real number */
237 #define SMALLREAL
                               1e-20 /**< A small real number */
    #define SMALLREAL2
                               1e-40 /**< An extremely small real number */
238
                              20 /**< Maximal refinement level */
239 #define MAX_REFINE_LVL
240 #define MAX_AMG_LVL
                                 20 /**< Maximal AMG coarsening level */
241 #define MIN_CDOF
                                 20 /**< Minimal number of coarsest variables */
#define MIN_CRATE 0.9 /**< Minimal coarsening ratio */
243 #define MAX_CRATE 20.0 /**< Maximal coarsening ratio */
244 #define MAX_RESTART 20 /**< Maximal restarting number */
245 #define MAX_STAG 20 /**< Maximal number of stagnation times */
```

```
#define STAG_RATIO
246
                                    1e-4 /**< Stagnation tolerance = tol*STAGRATIO */</pre>
                                          /**< Smallest size for OpenMP version */
247
    #define OPENMP_HOLDS
                                    2000
248
    #endif
                                           /* end if for __FASP_MESSAGES__ */
249
250
251
                  End of File
252
253
```

4.3 Debugging and how to enable it

NOTE: The default FSP build is a REALEASE version and such version is compiled with optimization and no warnings are displayed during the build. How to build the FASP library with debugging enabled is described below.

There is a built-in debug feature which is intended to help developers and users to locate malfunctions and bugs in FASP (and hopefully fix them). In order to turn this feature on, you need to add the debug option during the config stage by

```
$ make config debug=all
```

When this debug feature is turned on, there will be a lot more information printed when you run FASP. If you just want to enable the debugging and warnings during the compile stage, you can do so by using

```
$ make config debug=yes
```

Bibliography

- J. Bramble. Multigrid methods, volume 294 of Pitman Research Notes in Mathematics Series. Longman Scientific & Technical, Harlow, 1993.
- [2] A. Brandt, S. McCormick, and J. Ruge. Algebraic multigrid (AMG) for automatic multigrid solution with application to geodetic computations, Report. Inst. Comp. Studies Colorado State Univ, 109:110, 1982.
- [3] J. Brannick and L. Zikatanov. Algebraic multigrid methods based on compatible relaxation and energy minimization. Lecture Notes in Computational Science and Engineering, 55:15, 2007.
- [4] M. Brezina, A. Cleary, R. Falgout, V. Henson, J. Jones, T. A. Manteuffel, S. F. McCormick, and J. W. Ruge. Algebraic multigrid based on element interpolation (AMGe). SIAM Journal on Scientific Computing, 22(5):1570–1592, 2000.
- [5] T. Chartier, R. Falgout, V. Henson, J. Jones, T. Manteuffel, S. McCormick, J. Ruge, and P. Vassilevski. Spectral AMGe (ρAMGe). SIAM J. Sci. Comput., 25(1):1–26, 2003.
- [6] R. Falgout and P. Vassilevski. On generalizing the algebraic multigrid framework. SIAM J. Numer. Anal., 42(4):1669–1693 (electronic), 2004.
- [7] V. Henson and P. Vassilevski. Element-free AMGe: general algorithms for computing interpolation weights in AMG. SIAM J. Sci. Comput., 23(2):629–650, 2001.
- [8] X. Hu, P. S. Vassilevski, and J. Xu. Comparative convergence analysis of nonlinear amli-cycle multigrid. SIAM J. Num. Anal, 51(2):1349-1369, 2013.
- [9] O. E. Livne. Coarsening by compatible relaxation. Numer. Linear Algebra Appl., 11(2-3):205-227, 2004.
- [10] J. Mellor-crummey and J. Garvin. Optimizing Sparse Matrix-Vector Product Computations Using Unroll and Jam. International Journal of High Performance Computing Applications, 18(2):225—-236, 2004.
- [11] A. Muresan and Y. Notay. Analysis of aggregation-based multigrid. SIAM Journal on Scientific Computing, 30(2):1082–1103, 2008.
- [12] S. Pissanetzky. Sparse matrix technology. Academic Press Inc. [Harcourt Brace Jovanovich Publishers], London, 1984.
- [13] J. Ruge and K. Stuben. Efficient solution of finite difference and finite element equations. In D. J. Paddon and H. Holstein, editors, Multigrid Methods for Integral and Differential Equations, volume 3, pages 169–212. Clarendon Press, 1985.
- [14] J. Ruge and K. Stüben. Algebraic multigrid. Multigrid methods, 3:73–130, 1987.
- [15] Y. Saad. Iterative methods for sparse linear systems. Society for Industrial and Applied Mathematics, Philadelphia, PA, second edition, 2003.

58 BIBLIOGRAPHY

[16] U. Trottenberg, C. Oosterlee, and A. Schüller. Multigrid. With contributions by A. Brandt, P. Oswald and K. Stuben, 2001.

- [17] P. Vaněk, J. Mandel, and M. Brezina. Algebraic multigrid by smoothed aggregation for second and fourth order elliptic problems. *Computing*, 56(3):179–196, 1996.
- [18] W. Wan, T. Chan, and B. Smith. An energy-minimizing interpolation for robust multigrid methods. SIAM Journal on Scientific Computing, 21(4):1632–1649, 2000.
- [19] J. Xu. Fast Poisson-Based Solvers for Linear and Nonlinear PDEs Jinchao Xu. In *Proceedings of the International Congress of Mathematicians*, pages 2886–2912, 2010.
- [20] J. Xu and L. Zikatanov. On an energy minimizing basis for algebraic multigrid methods. *Computing and Visualization in Science*, 7(3):121–127, 2004.