# Lis User Manual Version 1.2.71



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# 0 Changes from Version 1.1

- 1. Added support for eigensolvers.
- 2. Changed specifications of the following APIs:
  - (a) Changed names of lis\_output\_residual\_history() and lis\_get\_residual\_history() to lis\_solver\_output\_rhistory() and lis\_solver\_get\_rhistory(), respectively.
  - (b) Changed origin of Fortran subroutines lis\_vector\_set\_value() and lis\_vector\_get\_value() to 1.
  - (c) Changed origin of Fortran subroutine lis\_vector\_set\_size() to 1.
  - (d) Changed name of precision flag -precision to -f.

# 1 Introduction

Lis, a Library of Iterative Solvers for linear systems, is a numerical library written in C and Fortran for solving the systems of linear equations of the form

$$Ax = b$$

and the standard eigenvalue problems of the form

$$Ax = \lambda x$$

with real sparse matrices using iterative methods. The solvers available in Lis are listed in Table 1 and 2, and the preconditioners are listed in Table 3. The matrix storage formats are listed in Table 4.

Table 1: Linear Solvers

Table 1: Linear Solvers		
CG	CR	
$\operatorname{BiCG}$	BiCR[2]	
CGS	CRS[3]	
BiCGSTAB	BiCRSTAB[3]	
GPBiCG	GPBiCR[3]	
BiCGSafe[1]	BiCRSafe[4]	
BiCGSTAB(l)	TFQMR	
Jacobi	Orthomin(m)	
Gauss-Seidel	GMRES(m)	
SOR	FGMRES(m)[5]	
IDR(s)[13]	MINRES[14]	
GPBiCG BiCGSafe[1] BiCGSTAB(1) Jacobi Gauss-Seidel SOR	GPBiCR[3] BiCRSafe[4] TFQMR Orthomin(m) GMRES(m) FGMRES(m)[5]	

Table 2: Eigensolvers

Power Iteration
Inverse Iteration
Approximate Inverse Iteration
Rayleigh Quotient Iteration
Subspace Iteration
Lanczos Iteration
Conjugate Gradient[18, 19]
Conjugate Residual [20]

Table 3: Preconditioners

Table 4	4: Matrix	Storage	Formats
7	1	α.	

Jacobi	Compressed Row Storage	(CRS)
SSOR	Compressed Column Storage	(CCS)
ILU(k)	Modified Compressed Sparse Row	(MSR)
ILUT[6, 7]	Diagonal	(DIA)
Crout ILU[8, 7]	Ellpack-Itpack generalized diagonal	(ELL)
I+S[9]	Jagged Diagonal	(JDS)
SA-AMG[10]	Block Sparse Row	(BSR)
Hybrid[11]	Block Sparse Column	(BSC)
SAINV[12]	Variable Block Row	(VBR)
Additive Schwarz	Dense	(DNS)
User defined	Coordinate	(COO)

# 2 Installation

This section describes the instructions for installing and testing Lis. We assume Lis being installed on a Linux cluster.

# 2.1 Requirements

Installation of Lis requires a C compiler. Fortran interface requires a compiler which supports FOR-TRAN 77. AMG preconditioner requires a compiler which supports Fortran 90. For parallel computing environments, OpenMP or MPI-1 is used. Lis has been tested on the environments shown in Table 5 (see also Table 7).

Table 5: Major Tested Platforms

C Compilers	OS
Intel C/C++ Compiler 7.0, 8.0, 9.1, 10.1, 11.1,	Linux
Intel C++ Composer XE	Windows
IBM XL C/C++ V7.0, 9.0	AIX
	Linux
Sun WorkShop 6, Sun ONE Studio 7,	Solaris
Sun Studio 11, 12	
PGI C++ 6.0, 7.1, 10.5	Linux
gcc 3.3, 4.3	Linux
	Mac OS X
	Windows
Microsoft Visual C++ 2008, 2010	Windows
Fortran Compilers (Optional)	OS
Intel Fortran Compiler 8.1, 9.1, 10.1, 11.1,	Linux
Intel Fortran Composer XE	Windows
IBM XL Fortran V9.1, 11.1	AIX
	Linux
Sun WorkShop 6, Sun ONE Studio 7,	Solaris
Sun Studio 11, 12	
PGI Fortran 6.0, 7.1, 10.5	Linux
g77 3.3	Linux
gfortran 4.3, 4.4	Mac OS X
g95 0.91	Windows

# 2.2 Extracting Archive

Enter the following command to extract the archive files, where (\$VERSION) represents the version: >gunzip -c lis-(\$VERSION).tar.gz | tar xvf It creates a directory lis-(\$VERSION) along with its subfolders as shown in Figure 1.

# lis-(\$VERSION)

- + config
- | configuration files
- + include
- | header files
- + src
- | source files
- + test
  - test programs

Figure 1: Files contained in lis-(\$VERSION).tar.gz

# 2.3 Installing on UNIX and Compatible Systems

# 2.3.1 Configuring

Run the following script to generate a makefile:

• default: >./configure

• specifying the installation destination: >./configure --prefix=<install-dir>

Table 6 shows the options which can be specified for the configuration. Table 7 shows the computing environments which can be specified by TARGET.

Table 6: Major Configuration Options (see ./configure --help for complete list)

Use OpenMP
Use MPI
Use Fortran API
Use SA-AMG preconditioner
Use quadruple precision operation
Use gprof
Build shared libraries
Name of the installation destination directory
Computing environments
C compiler
Compilation options for C compilers
Fortran 90 compiler
Compilation options for the Fortran 90 compiler
Link options

# 2.3.2 Compiling

In the directory lis-(\$VERSION), run the following command to generate executable files: >make

To ensure that the library has been successfully built, enter as follows in the directory lis-(\$VERSION): >make check

It runs a test script using the executable files created in the lis-(\$VERSION)/test directory, which reads the data of the coefficient matrix and the right hand side vector from the file test/testmat.mtx and writes the solution of the system of linear equations Ax = b obtained with the BiCG method into test/sol.txt, and the residual history into test/res.txt. If all the elements of the solution are 1, then the result is correct. The result on SGI Altix 3700 is shown below.

Table 7: Examples of Targets (see configure for complete list)

<target></target>	Equivalent options		
cray_xt3	./configure CC=cc FC=ftn CFLAGS="-03 -B -fastsse -tp k8-64"		
	FCFLAGS="-03 -fastsse -tp k8-64 -Mpreprocess" FCLDFLAGS="-Mnomain"		
	ac_cv_sizeof_void_p=8 cross_compiling=yesenable-mpi		
	<pre>ax_f77_mangling="lower case, no underscore, extra underscore"</pre>		
fujitsu_pq	./configure CC=fcc FC=frt ac_cv_sizeof_void_p=8		
	CFLAGS="-03 -Kfast,ocl,preex" FFLAGS="-03 -Kfast,ocl,preex -Cpp"		
	FCFLAGS="-03 -Kfast,ocl,preex -Cpp -Am"		
	ax_f77_mangling="lower case, underscore, no extra underscore"		
hitachi	./configure CC=cc FC=f90 FCLDFLAGS="-lf90s" ac_cv_sizeof_void_p=8		
	CFLAGS="-Os -noparallel" FCFLAGS="-Oss -noparallel"		
	ax_f77_mangling="lower case, underscore, no extra underscore"		
ibm_bgl	./configure CC=blrts_xlc FC=blrts_xlf90		
	CFLAGS="-03 -qarch=440d -qtune=440 -qstrict		
	-I/bgl/BlueLight/ppcfloor/bglsys/include"		
	FFFLAGS="-03 -qarch=440d -qtune=440 -qsuffix=cpp=F -qfixed=72 -w		
	-I/bgl/BlueLight/ppcfloor/bglsys/include"		
	FCFLAGS="-03 -qarch=440d -qtune=440 -qsuffix=cpp=F90 -w		
	-I/bgl/BlueLight/ppcfloor/bglsys/include"		
	ac_cv_sizeof_void_p=4 cross_compiling=yesenable-mpi		
	ax_f77_mangling="lower case, no underscore, no extra underscore"		
nec_es	./configure CC=esmpic++ FC=esmpif90 AR=esar RANLIB=true		
	ac_cv_sizeof_void_p=8 ax_vector_machine=yes cross_compiling=yes		
	enable-mpienable-omp		
	ax_f77_mangling="lower case, no underscore, extra underscore"		
nec_sx9_cross	./configure CC=sxmpic++ FC=sxmpif90 AR=sxar RANLIB=true		
ac_cv_sizeof_void_p=8 ax_vector_machine=yes cross_compil:			
ax_f77_mangling="lower case, no underscore, extra unders			

```
matrix size = 100 x 100 (460 nonzero entries)
initial vector x = 0
precision: double
solver: BiCG 2
precon: none
storage: CRS
lis_solve: normal end

BiCG: number of iterations = 15 (double = 15, quad = 0)
BiCG: elapsed time = 5.178690e-03 sec.
BiCG: preconditioner = 1.277685e-03 sec.
BiCG: matrix creation = 1.254797e-03 sec.
BiCG: linear solver = 3.901005e-03 sec.
BiCG: relative residual 2-norm = 6.327297e-15
```

```
max number of threads = 32
number of threads = 2
matrix size = 100 x 100 (460 nonzero entries)
initial vector x = 0
precision : double
solver : BiCG 2
precon : none
storage : CRS
lis_solve : normal end

BiCG: number of iterations = 15 (double = 15, quad = 0)
BiCG: elapsed time = 8.960009e-03 sec.
BiCG: preconditioner = 2.297878e-03 sec.
BiCG: matrix creation = 2.072096e-03 sec.
BiCG: linear solver = 6.662130e-03 sec.
BiCG: relative residual 2-norm = 6.221213e-15
```

# number of processes = 2 matrix size = 100 x 100 (460 nonzero entries) initial vector x = 0 precision : double solver : BiCG 2 precon : none storage : CRS lis\_solve : normal end BiCG: number of iterations = 15 (double = 15, quad = 0) BiCG: elapsed time = 2.911400e-03 sec. BiCG: preconditioner = 1.560780e-04 sec. BiCG: matrix creation = 1.459997e-04 sec. BiCG: linear solver = 2.755322e-03 sec. BiCG: relative residual 2-norm = 6.221213e-15

# 2.3.3 Installing

lis\_config.h is the header file required to build the library, and lis.h and lisf.h are the header files required by C and Fortran compilers, respectively. liblis.a is the library file.

# 2.4 Installing on Windows Systems

Use one of the solution files or the project files for Microsoft Visual Studio in the directory lis-(\$VERSION)/win32. lis\_with\_fortran.sln is the solution file to be used with Intel Visual Fortran Compiler. lis\_with\_fortran\_mpi.sln is the solution file to be used with Visual Fortran and MPICH2. Header files are located in lis-(\$VERSION)/include. lis\_config\_win32.h is the header file required to build the library. lis.h and lisf.h are the header files required by C and Fortran compilers, respectively. The library files are generated in lis-(\$VERSION)/lib. The executable files of the test programs are generated in lis-(\$VERSION)/test.

# 2.5 Test Programs

### 2.5.1 test1

Usage: test1 matrix\_filename rhs\_setting solution\_filename residual\_filename [options]

This program inputs the data of the coefficient matrix from matrix\_filename and solves the system of linear equations Ax = b with the solver specified by options. It outputs the solution to solution\_filename and the residual history to residual\_filename. The Extended Matrix Market format, which is extended to allow vector data, is supported (see Appendix). One of the following values can be used for rhs\_setting:

0 Use the right hand side vector b included in the data file

1 Use  $b = (1, ..., 1)^T$ 

Use  $b = A \times (1, \dots, 1)^T$ 

rhs\_filename Filename for the right hand side vector

The PLAIN and MM formats are supported for rhs\_filename. test1f.F is the Fortran version of test1.c.

### 2.5.2 test2

Usage: test2 m n matrix\_type solution\_filename residual\_filename [options]

This program solves a discretized two dimensional Poisson equation Ax = b using the five point central difference scheme, with the coefficient matrix A of size mn in the storage format specified by matrix\_type and the solver specified by options. It outputs the solution to solution\_filename and the residual history to residual\_filename. The right hand side vector is set to make all the elements for the solution to be 1. The values m and n represent the numbers of lattice points in each dimension.

### 2.5.3 test3

Usage: test3 1 m n matrix\_type solution\_filename residual\_filename [options]

This program solves a discretized three dimensional Poisson equation Ax = b using the seven point central difference scheme, with the coefficient matrix A of size lmn in the storage format specified by matrix\_type and the solver specified by options. It outputs the solution to solution\_filename and the residual history to residual\_filename. The right hand side vector is set to make all the elements for the solution to be 1. The values l, m and m represent the numbers of lattice points in each dimension.

# 2.5.4 test4

This program solves the system of linear equations Ax = b with a specified solver and a preconditioner, where A is a tridiagonal matrix

$$\begin{pmatrix}
2 & -1 & & & & \\
-1 & 2 & -1 & & & & \\
& \ddots & \ddots & \ddots & & \\
& & -1 & 2 & -1 & \\
& & & -1 & 2
\end{pmatrix}$$

of size 12. The right hand side vector b is set to make all the elements of the solution x to be 1. test4f.F is the Fortran version of test4.c.

### 2.5.5 test5

Usage: test5 n gamma [options]

This program solves a system of linear equations Ax = b, where A is a Toeplitz matrix

$$\begin{pmatrix} 2 & 1 & & & & \\ 0 & 2 & 1 & & & \\ \gamma & 0 & 2 & 1 & & \\ & \ddots & \ddots & \ddots & \ddots & \\ & & \gamma & 0 & 2 & 1 \\ & & & \gamma & 0 & 2 \end{pmatrix}$$

with the solver specified by options. Note that the right hand vector is set to make all the elements of the solution to be 1. The value n is the size of the matrix A.

### 2.5.6 etest1

Usage: etest1 matrix\_filename solution\_filename residual\_filename [options]

This program inputs the matrix data from matrix\_filename and solves the eigenvalue problem  $Ax = \lambda x$  with the solver specified by options. It outputs the associated eigenvector to solution\_filename and the residual history to residual\_filename. The Matrix Market format is supported. etest1.F is the Fortran version of etest1.c.

### 2.5.7 etest2

Usage: etest2 m n matrix\_type solution\_filename residual\_filename [options]

This program solves the eigenvalue problem  $Ax = \lambda x$ , where the coefficient matrix A of size mn is derived from a discretized two dimensional Helmholtz equation using the five point central difference scheme, with the coefficient matrix in the storage format specified by matrix\_type and the solver specified by options. It outputs the associated eigenvector to solution\_filename and the residual history to residual\_filename. The values m and n represent the numbers of lattice points in each dimension.

# 2.5.8 etest3

Usage: etest3 1 m n matrix\_type solution\_filename residual\_filename [options]

This program solves the eigenvalue problem  $Ax = \lambda x$ , where the coefficient matrix A of size lmn is derived from a discretized three dimensional Helmholtz equation using the seven point central difference scheme, with the coefficient matrix in the storage format specified by matrix\_type and the solver specified by options. It outputs the associated eigenvector to solution\_filename and the residual history to residual\_filename. The values l, m and m represent the numbers of lattice points in each dimension.

### 2.5.9 etest4

Usage: etest4 n [options]

This program solves the eigenvalue problem  $Ax = \lambda x$  with a specified solver, where A is a tridiagonal

matrix

$$A = \begin{pmatrix} 2 & -1 & & & \\ -1 & 2 & -1 & & & \\ & \ddots & \ddots & \ddots & \\ & & -1 & 2 & -1 \\ & & & -1 & 2 \end{pmatrix}$$

of size  $n \times n$ , etest4f.F is the Fortran version of etest4.c

### 2.5.10 etest5

Usage: etest5 evalue\_filename evector\_filename

This program solves the eigenvalue problem  $Ax = \lambda x$  with subspace iteration, where A is a tridiagonal matrix

$$A = \begin{pmatrix} 2 & -1 & & & \\ -1 & 2 & -1 & & & \\ & \ddots & \ddots & \ddots & \\ & & -1 & 2 & -1 \\ & & & -1 & 2 \end{pmatrix}$$

of size  $12 \times 12$ . It outputs 2 extreme eigenvalues of smallest magnitude to evalue\_filename and the associated eigenvectors to evector\_filename in the extended Matrix Market format (see Appendix).

### 2.5.11 spmvtest1

Usage: spmvtest1 n iter

This program computes the multiply of a tridiagonal matrix derived from a discretized one dimensional Poisson equation using the three point central difference scheme

$$\begin{pmatrix} 2 & -1 \\ -1 & 2 & -1 \\ & \ddots & \ddots & \ddots \\ & & -1 & 2 & -1 \\ & & & -1 & 2 \end{pmatrix}$$

of size n and a vector  $(1, ..., 1)^T$ . FLOPS performance is measured as the average of *iter* iterations.

### 2.5.12 spmvtest2

Usage: spmvtest2 m n iter

This program computes the multiply of a sparse matrix, derived from a discretized two dimensional Poisson equation using the five point central difference scheme with available matrix storage formats, and a vector  $(1, ..., 1)^T$ . The values m and n represent the numbers of lattice points in the vertical and horizontal directions. FLOPS performance is measured as the average of *iter* iterations.

# 2.5.13 spmvtest3

Usage: spmvtest3 1 m n iter

This program computes the multiply of a sparse matrix, derived from a discretized three dimensional Poisson equation using the seven point central difference scheme with available matrix storage formats, and a vector  $(1, ..., 1)^T$ . The values l, m and n represent the numbers of lattice points in each dimension. FLOPS performance is measured as the average of *iter* iterations.

# 2.5.14 spmvtest4

Usage: spmvtest4 matrix\_filename\_list iter [block]

This program inputs the matrix data from the files listed in matrix\_filename\_list, and computes the multiplies of the matrices with available matrix storage formats and a vector  $(1, ..., 1)^T$ . FLOPS performance is measured twice as the average of *iter* iterations. If necessary, the block size of BSR and BSC can be specified by *block*.

# 2.5.15 spmvtest5

Usage: spmvtest5 matrix\_filename matrix\_type iter [block]

This program inputs the matrix data from matrix\_filename and compute the multiply of the matrix with matrix\_type and a vector  $(1, ..., 1)^T$ . FLOPS performance is measured twice as the average of *iter* iterations. If necessary, the block size of BSR and BSC can be specified by *block*.

# 2.6 Restrictions

The current version has the following restrictions:

### • Preconditioners

- If a preconditioner other than the Jacobi or SSOR is selected and the matrix A is not in the CRS format, a new matrix is created in the CRS format for preconditioning.
- SA-AMG preconditioner does not support BiCG.
- SA-AMG preconditioner does not support OpenMP version of Lis.
- Assembly of preconditioning matrices in SAINV is not parallelized.

# • Quadruple precision operations

- Jacobi, Gauss-Seidel, SOR, and  $\mathrm{IDR}(s)$  methods do not support quadruple precision operations.
- Conjugate gradient and conjugate residual methods for eigenproblems do not support quadruple precision operations.
- Jacobi, Gauss-Seidel and SOR do not support quadruple precision operations in the hybrid preconditioner.
- I+S and SA-AMG preconditioners do not support quadruple precision operations.

# • Matrix storage formats

- In the MPI environment, CRS is the only accepted format for user defined arrays.

# 3 Basic Operations

This section describes how to use the library. A program requires the following statements:

- Initialization
- Matrix creation
- Vector creation
- Solver creation
- Value assignment for matrix and vector
- Solver assignment for system of linear equations or eigenvalue problem
- Solver execution
- Finalization

In addition, it must include one of the following include statements:

- C #include "lis.h"
- Fortran #include "lisf.h"

When Lis is installed in \$(INSTALLDIR), lis.h and lisf.h are located in \$(INSTALLDIR)/include.

# 3.1 Initializing and Finalizing

The functions for initializing and finalizing the execution environment must be called at the top and the bottom of the program, respectively, as follows:

```
1: #include "lis.h"
2: int main(int argc, char* argv[])
3: {
4: lis_initialize(&argc, &argv);
5: ...
6: lis_finalize();
7: }
```

```
Fortran

1: #include "lisf.h"

2: call lis_initialize(ierr)

3: ...

4: call lis_finalize(ierr)
```

### Initializing

For initialization, the following functions are used:

- C lis\_initialize(int\* argc, char\*\* argv[])
- Fortran subroutine lis\_initialize(integer ierr)

This function initializes the MPI execution environment, and specifies options on the command line.

### Finalizing

For finalization, the following functions are used:

- C int lis\_finalize()
- Fortran subroutine lis\_finalize(integer ierr)

# 3.2 Operating Vectors

Assume that the size of a vector v is  $global\_n$ , and the size of each partial vector stored on nprocs processing elements is  $local\_n$ . If  $global\_n$  is divisible, then  $local\_n = global\_n / nprocs$ . For example, when the vector v is stored on two processing elements, as shown in Equation (3.1),  $global\_n$  and  $local\_n$  are 4 and 2, respectively.

$$v = \begin{pmatrix} 0\\ \frac{1}{2}\\ 3 \end{pmatrix} \text{ PE0}$$
PE1 (3.1)

In the case of creating the vector v in Equation (3.1), the serial and OpenMP versions create the vector v itself, while the MPI version creates partial vectors stored on a given number of processing elements.

Programs to create the vector v are as follows, where the number of the processing elements for the MPI version is assumed to be two:

```
C (for serial and OpenMP versions)

1: int     i,n;
2: LIS_VECTOR     v;
3: n = 4;
4: lis_vector_create(0,&v);
5: lis_vector_set_size(v,0,n); /* or lis_vector_set_size(v,n,0); */
6:
7: for(i=0;i<n;i++)
8: {
9: lis_vector_set_value(LIS_INS_VALUE,i,(double)i,v);
10: }</pre>
C (for MPI version)
```

```
/*or int i,ln,is,ie;
1: int
                 i,n,is,ie;
2: LIS_VECTOR
                 v;
3: n = 4;
                                           /* ln = 2;
                                                                                        */
4: lis_vector_create(MPI_COMM_WORLD,&v);
5: lis_vector_set_size(v,0,n);
                                           /* lis_vector_set_size(v,ln,0);
                                                                                        */
6: lis_vector_get_range(v,&is,&ie);
7: for(i=is;i<ie;i++)
8: {
9:
       lis_vector_set_value(LIS_INS_VALUE,i,(double)i,v);
10: }
```

```
Fortran (for serial and OpenMP versions)

1: integer i,n
2: LIS_VECTOR v
3: n = 4
4: call lis_vector_create(0,v,ierr)
5: call lis_vector_set_size(v,0,n,ierr)
6:
7: do i=1,n
9: call lis_vector_set_value(LIS_INS_VALUE,i,DBLE(i),v,ierr)
10: enddo
```

```
Fortran (for MPI version)

1: integer i,n,is,ie

2: LIS_VECTOR v

3: n = 4

4: call lis_vector_create(MPI_COMM_WORLD,v,ierr)

5: call lis_vector_set_size(v,0,n,ierr)

6: call lis_vector_get_range(v,is,ie,ierr)

7: do i=is,ie-1

8: call lis_vector_set_value(LIS_INS_VALUE,i,DBLE(i),v,ierr);

9: enddo
```

### **Declaring Variables**

As the second line shows, the declaration is stated as follows:

```
LIS_VECTOR v;
```

### **Creating Vectors**

To create the vector v, the following functions are used:

- C int lis\_vector\_create(LIS\_Comm comm, LIS\_VECTOR \*v)
- Fortran subroutine lis\_vector\_create(LIS\_Comm comm, LIS\_VECTOR v, integer ierr)

For the example program above, **comm** must be replaced with the MPI communicator. For the serial and OpenMP versions, the value for **comm** is ignored.

### **Assigning Vector Sizes**

To assign the size of a vector v, the following functions are used:

- C int lis\_vector\_set\_size(LIS\_VECTOR v, int local\_n, int global\_n)
- Fortran subroutine lis\_vector\_create(integer local\_n, integer global\_n, LIS\_Comm comm, LIS\_VECTOR v, integer ierr)

Either local\_n or global\_n must be provided. This function creates a vector in one of the following ways: Creates partial vectors of size local\_n if local\_n is given, or create partial vectors of size global\_n, stored on a given number of processing elements, if global\_n is given.

In the case of the serial and OpenMP versions,  $local\_n = global\_n$ . It means that both  $lis\_vector\_set\_size(v,n,0)$  and  $lis\_vector\_set\_size(v,0,n)$  create a vector of size n.

For the MPI version,  $lis_vector_set_size(v,n,0)$  creates a partial vector of size  $n_p$  on the processing element p. On the other hand,  $lis_vector_set_size(v,0,n)$  creates a partial vector of size  $m_p$  on the processing element p. The value for  $m_p$  is determined by the library.

### **Assigning Elements**

To assign an element to the i-th row of the vector v, the following functions are used:

- C int lis\_vector\_set(int flag, int i, LIS\_SCALAR value, LIS\_VECTOR v)
- Fortran subroutine lis\_vector\_set\_value(int flag, int i, LIS\_SCALAR value, LIS\_VECTOR v, integer ierr)

For the MPI version, the i-th row of the global vector must be specified, instead of the i-th row of the partial vector. Either

```
\begin{split} & \texttt{LIS\_INS\_VALUE} \,:\, \texttt{v}[i] \,=\, value,\, \text{or} \\ & \texttt{LIS\_ADD\_VALUE} \,:\, \texttt{v}[i] \,=\, \texttt{v}[i] \,+\, value \end{split}
```

must be provided for flag.

### **Duplicating Vectors**

To create a vector which has the same information as for an existing vector, the following functions are used:

- C int lis\_vector\_duplicate(LIS\_VECTOR vin, LIS\_VECTOR \*vout)
- Fortran subroutine lis\_vector\_duplicate(LIS\_VECTOR vin, LIS\_VECTOR vout, integer ierr)

This function does not copy the elements of the vector. To copy the elements as well, the following functions must be added after the above functions:

- C int lis\_vector\_copy(LIS\_VECTOR vsrc, LIS\_VECTOR vdst)
- Fortran subroutine lis\_vector\_copy(LIS\_VECTOR vsrc, LIS\_VECTOR vdst, integer ierr)

### **Destroying Vectors**

To destroy a vector, the following functions are used:

- C int lis\_vector\_destroy(LIS\_VECTOR v)
- Fortran subroutine lis\_vector\_destroy(LIS\_VECTOR v, integer ierr)

# 3.3 Operating Matrices

Assume that the size of a matrix A is  $global\_n \times global\_n$ , and that the size of each row block of the matrix A stored on nprocs processing elements is  $local\_n \times global\_n$ . If  $global\_n$  is divisible, then  $local\_n = global\_n / nprocs$ . For example, when the row block of the matrix A is stored on two processing elements, as shown in Equation (3.2),  $global\_n$  and  $local\_n$  are 4 and 2, respectively.

$$A = \begin{pmatrix} 2 & 1 & & \\ 1 & 2 & 1 & \\ \hline & 1 & 2 & 1 \\ & & 1 & 2 \end{pmatrix} \text{ PE0}$$

$$(3.2)$$

A matrix in a specific storage format can be created in one of the following three ways:

### Method 1: Defining Arrays in Specific Storage Format with Library Functions

In the case of creating the matrix A in Equation (3.2) in the CRS format, the serial and OpenMP versions create the matrix A itself, and the MPI version creates on each processing element a partial matrix stored on given number of processing elements.

Programs to create the matrix A in the CRS format are as follows, where the number of the processing elements for the MPI version is assumed to be two:

```
C (for serial and OpenMP versions) -
 1: int
                  i,n;
 2: LIS_MATRIX
                  A;
 3: n = 4:
 4: lis_matrix_create(0,&A);
 5: lis_matrix_set_size(A,0,n); /* or lis_matrix_set_size(A,n,0); */
 6: for(i=0;i<n;i++) {
 7:
        if( i>0 ) lis_matrix_set_value(LIS_INS_VALUE,i,i-1,1.0,A);
 8:
        if( i<n-1 ) lis_matrix_set_value(LIS_INS_VALUE,i,i+1,1.0,A);</pre>
9:
        lis_matrix_set_value(LIS_INS_VALUE,i,i,2.0,A);
10:
     }
     lis_matrix_set_type(A,LIS_MATRIX_CRS);
11:
     lis_matrix_assemble(A);
```

```
- C (for MPI version) -
 1: int
                  i,n,gn,is,ie;
 2: LIS_MATRIX
                  A;
3: gn = 4;
                                              /* or n=2
4: lis_matrix_create(MPI_COMM_WORLD,&A);
5: lis_matrix_set_size(A,0,gn);
                                                    lis_matrix_set_size(A,n,0); */
 6: lis_matrix_get_size(A,&n,&gn);
7: lis_matrix_get_range(A,&is,&ie);
8: for(i=is;i<ie;i++) {
        if( i>0 ) lis_matrix_set_value(LIS_INS_VALUE,i,i-1,1.0,A);
9:
10:
        if( i<gn-1 ) lis_matrix_set_value(LIS_INS_VALUE,i,i+1,1.0,A);</pre>
        lis_matrix_set_value(LIS_INS_VALUE,i,i,2.0,A);
11:
12: }
13:
    lis_matrix_set_type(A,LIS_MATRIX_CRS);
    lis_matrix_assemble(A);
```

```
1: integer    i,n
2: LIS_MATRIX    A
3: n = 4
4: call lis_matrix_create(0,A,ierr)
5: call lis_matrix_set_size(A,0,n,ierr)
6: do i=1,n
7:    if( i>1 ) call lis_matrix_set_value(LIS_INS_VALUE,i,i-1,1.0d0,A,ierr)
```

```
8: if( i<n ) call lis_matrix_set_value(LIS_INS_VALUE,i,i+1,1.0d0,A,ierr)
9: call lis_matrix_set_value(LIS_INS_VALUE,i,i,2.0d0,A,ierr)
10: enddo
11: call lis_matrix_set_type(A,LIS_MATRIX_CRS,ierr)
```

- Fortran (for MPI version) -

12: call lis\_matrix\_assemble(A,ierr)

Fortran (for serial and OpenMP versions) -

```
1: integer
                  i,n,gn,is,ie
2: LIS_MATRIX
3: gn = 4
4: call lis_matrix_create(MPI_COMM_WORLD,A,ierr)
5: call lis_matrix_set_size(A,0,gn,ierr)
 6: call lis_matrix_get_size(A,n,gn,ierr)
7: call lis_matrix_get_range(A,is,ie,ierr)
8: do i=is,ie-1
        if( i>1 ) call lis_matrix_set_value(LIS_INS_VALUE,i,i-1,1.0d0,A,ierr)
9:
10:
        if( i<gn ) call lis_matrix_set_value(LIS_INS_VALUE,i,i+1,1.0d0,A,ierr)</pre>
11:
        call lis_matrix_set_value(LIS_INS_VALUE,i,i,2.0d0,A,ierr)
12: enddo
13: call lis_matrix_set_type(A,LIS_MATRIX_CRS,ierr)
14: call lis_matrix_assemble(A,ierr)
```

# **Declaring Variables**

As the second line shows, the declaration is stated as follows:

```
LIS_MATRIX A;
```

# **Creating Matrices**

To create the matrix A, the following functions are used:

- C int lis\_matrix\_create(LIS\_Comm comm, LIS\_MATRIX \*A)
- Fortran subroutine lis\_matrix\_create(LIS\_Comm comm, LIS\_MATRIX A, integer ierr)

comm must be replaced with the MPI communicator. For the serial and OpenMP versions, the value for comm is ignored.

# **Assigning Matrix Sizes**

To assign a size to the matrix A, the following functions are used:

- C int lis\_matrix\_set\_size(LIS\_MATRIX A, int local\_n, int global\_n)
- Fortran subroutine lis\_matrix\_set\_size(LIS\_MATRIX A, integer local\_n, integer global\_n, integer ierr)

Either  $local\_n$  or  $global\_n$  must be provided. This function creates matrices in one of the following two ways: Create partial matrices of size  $local\_n \times N$  if  $local\_n$  is given, or creating partial matrices of size  $global\_n \times global\_n$  stored on a given number of processing elements, if  $global\_n$  is given. N represents the total sum of  $local\_n$ .

In the case of the serial and OpenMP versions,  $local_n = global_n$ . It means that both  $lis_matrix_set_size(A,n,0)$  and  $lis_matrix_set_size(A,0,n)$  create a matrix of  $n \times n$ .

For the MPI version,  $lis_matrix_set_size(A,n,0)$  creates a partial matrix of size  $n_p \times N$  on the processing element p, where N is the total sum of  $n_p$ . On the other hand,  $lis_matrix_set_size(A,0,n)$  creates a partial matrix of size  $m_p \times n$  on the processing element p, where  $m_p$  is the number of the partial matrix, which is determined by the library.

### **Assigning Elements**

To assign an element to the cell at the i-th row and the j-th column of the matrix A, the following functions are used:

- C int lis\_matrix\_set\_value(int flag, int i, int j, LIS\_SCALAR value, LIS\_MATRIX A)
- Fortran subroutine lis\_matrix\_set\_value(integer flag, integer i, integer j, LIS\_SCALAR value, LIS\_MATRIX A, integer ierr)

For the MPI version, the i-th row and the j-th column of the global matrix must be specified, instead of the i-th row and the j-th column of the partial matrix. Either

```
LIS_INS_VALUE : A(i, j) = \text{value}, or
```

LIS\_ADD\_VALUE : A(i, j) = A(i, j) + value

must be provided for the parameter flag.

# **Assigning Storage Formats**

To assign a storage format to the matrix A, the following functions are used:

- C int lis\_matrix\_set\_type(LIS\_MATRIX A, int matrix\_type)
- Fortran subroutine lis\_matrix\_set\_type(LIS\_MATRIX A, int matrix\_type, integer ierr)

 $\mathtt{matrix\_type}$  of A is  $\mathtt{LIS\_MATRIX\_CRS}$  when the matrix is created. The following storage formats are accepted:

Storage formats		$matrix\_type$
Compressed Row Storage	(CRS)	{LIS_MATRIX_CRS 1}
Compressed Column Storage	(CCS)	{LIS_MATRIX_CCS 2}
Modified Compressed Sparse Row	(MSR)	{LIS_MATRIX_MSR 3}
Diagonal	(DIA)	{LIS_MATRIX_DIA 4}
Ellpack-Itpack generalized diagonal	(ELL)	{LIS_MATRIX_ELL 5}
Jagged Diagonal	(JDS)	{LIS_MATRIX_JDS 6}
Block Sparse Row	(BSR)	{LIS_MATRIX_BSR 7}
Block Sparse Column	(BSC)	{LIS_MATRIX_BSC 8}
Variable Block Row	(VBR)	{LIS_MATRIX_VBR 9}
Dense	(DNS)	{LIS_MATRIX_DNS 10}
Coordinate	(COO)	{LIS_MATRIX_COO 11}

# **Assembling Matrices**

After assigning elements, the following function must be used:

- C int lis\_matrix\_assemble(LIS\_MATRIX A)
- Fortran subroutine lis\_matrix\_assemble(LIS\_MATRIX A, integer ierr)

lis\_matrix\_assemble is assembled to the storage format specified with lis\_matrix\_set\_type.

# **Destroying Matrices**

To destroy a matrix, the following functions are used:

- C int lis\_matrix\_destroy(LIS\_MATRIX A)
- Fortran subroutine lis\_matrix\_destroy(LIS\_MATRIX A, integer ierr)

# Method 2: Defining Arrays in Specified Storage Format Directly

In the case of creating the matrix A in Equation (3.2) in the CRS format, the serial and OpenMP versions creates the matrix A itself, and the MPI version creates on each processing element a partial matrix stored on a given number of processing elements.

Programs to create the matrix A in the CRS format are as follows, where the number of the processing elements for the MPI version is assumed to be two:

```
C (for serial and OpenMP versions) -
 1: int
                  i,k,n,nnz;
 2: int
                  *ptr,*index;
3: LIS_SCALAR
                  *value;
4: LIS_MATRIX
                  A;
5: n = 4; nnz = 10; k = 0;
 6: lis_matrix_malloc_crs(n,nnz,&ptr,&index,&value);
7: lis_matrix_create(0,&A);
8: lis_matrix_set_size(A,0,n); /* or lis_matrix_set_size(A,n,0); */
9:
10: for(i=0;i<n;i++)
11: {
12:
        if( i>0 ) {index[k] = i-1; value[k] = 1; k++;}
13:
        index[k] = i; value[k] = 2; k++;
14:
        if( i \le n-1 ) {index[k] = i+1; value[k] = 1; k++;}
15:
        ptr[i+1] = k;
16: }
17:
     ptr[0] = 0;
18:
    lis_matrix_set_crs(nnz,ptr,index,value,A);
19: lis_matrix_assemble(A);
```

```
- C (for MPI version) -
 1: int
                  i,k,n,nnz,is,ie;
 2: int
                  *ptr,*index;
3: LIS_SCALAR
                  *value;
4: LIS_MATRIX
                 A;
5: n = 2; nnz = 5; k = 0;
6: lis_matrix_malloc_crs(n,nnz,&ptr,&index,&value);
7: lis_matrix_create(MPI_COMM_WORLD,&A);
8: lis_matrix_set_size(A,n,0);
9: lis_matrix_get_range(A,&is,&ie);
10: for(i=is;i<ie;i++)</pre>
11: {
        if( i>0 ) {index[k] = i-1; value[k] = 1; k++;}
12:
        index[k] = i; value[k] = 2; k++;
        if( i < n-1 ) {index[k] = i+1; value[k] = 1; k++;}
14:
15:
        ptr[i-is+1] = k;
16: }
17: ptr[0] = 0;
18: lis_matrix_set_crs(nnz,ptr,index,value,A);
19: lis_matrix_assemble(A);
```

# **Associating Arrays**

To associate the arrays required by the CRS format created by the user with the matrix A, the following functions are used:

- C int lis\_matrix\_set\_crs(int nnz, int row[], int index[], LIS\_SCALAR value[], LIS\_MATRIX A)
- Fortran subroutine lis\_matrix\_set\_crs(integer nnz, integer row(), integer index(), LIS\_SCALAR value(), LIS\_MATRIX A, integer ierr)

### Method 3: Reading Matrix and Vector Data from External Files

Programs to read the matrix A in Equation (3.2) in the CRS format and vector b in Equation (3.1) from external files are as follows:

```
C (for serial, OpenMP and MPI versions)

1: LIS_MATRIX A;
2: LIS_VECTOR b,x;
3: lis_matrix_create(LIS_COMM_WORLD,&A);
4: lis_vector_create(LIS_COMM_WORLD,&b);
5: lis_vector_create(LIS_COMM_WORLD,&x);
6: lis_matrix_set_type(A,LIS_MATRIX_CRS);
7: lis_input(A,b,x,"matvec.mtx");
```

```
Fortran (for serial, OpenMP and MPI versions)

1: LIS_MATRIX A

2: LIS_VECTOR b,x

3: call lis_matrix_create(LIS_COMM_WORLD,A,ierr)

4: call lis_vector_create(LIS_COMM_WORLD,b,ierr)

5: call lis_vector_create(LIS_COMM_WORLD,x,ierr)

6: call lis_matrix_set_type(A,LIS_MATRIX_CRS,ierr)

7: call lis_input(A,b,x,'matvec.mtx',ierr)
```

The content of the destination file matvec.mtx is as follows:

```
%%MatrixMarket matrix coordinate real general
```

```
4 4 10 1 0
1 2 1.0e+00
1 1 2.0e+00
2 3 1.0e+00
2 1 1.0e+00
2 2 2.0e+00
3 4 1.0e+00
3 2 1.0e+00
3 3 2.0e+00
4 4 2.0e+00
4 3 1.0e+00
1 0.0e+00
2 1.0e+00
3 2.0e+00
4 3.0e+00
```

### Reading from External Files

To input the matrix data for A from external files, the following functions are used:

```
• C int lis_input_matrix(LIS_MATRIX A, char *filename)
```

• Fortran subroutine lis\_input(LIS\_MATRIX A, character filename, integer ierr)

filename must be replaced with the file path. The following file formats are supported:

- Matrix Market format
- Harwell-Boeing format
- Lis format (original format)

To read the data for the matrix A and vectors b and x from external files, the following functions are used:

- C int lis\_input(LIS\_MATRIX A, LIS\_VECTOR b, LIS\_VECTOR x, char \*filename)
- Fortran subroutine lis\_input(LIS\_MATRIX A, LIS\_VECTOR b, LIS\_VECTOR x, character filename, integer ierr)

filename must be replaced with the file path. The following file formats are supported:

- Extended Matrix Market format (extended to allow vector data)
- Harwell-Boeing format
- Lis format (original format)

# 3.4 Solving Systems of Linear Equations

A program to solve the system of linear equations Ax = b with a specified solver is as follows:

```
C (for serial, OpenMP and MPI versions)

1: LIS_MATRIX A;
2: LIS_VECTOR b,x;
3: LIS_SOLVER solver;
4:
5: /* Create matrix and vector */
6:
7: lis_solver_create(&solver);
8: lis_solver_set_option("-i bicg -p none",solver);
9: lis_solver_set_option("-tol 1.0e-12",solver);
10: lis_solver(A,b,x,solver);
```

```
Fortran (for serial, OpenMP and MPI versions)

1: LIS_MATRIX A

2: LIS_VECTOR b,x

3: LIS_SOLVER solver

4:

5: /* Create matrix and vector */

6:

7: call lis_solver_create(solver,ierr)

8: call lis_solver_set_option('-i bicg -p none',solver,ierr)

9: call lis_solver_set_option('-tol 1.0e-12',solver,ierr)

10: call lis_solver(A,b,x,solver,ierr)
```

# **Creating Solvers**

To create a solver, the following functions are used:

- C int lis\_solver\_create(LIS\_SOLVER \*solver)
- Fortran subroutine lis\_solver\_create(LIS\_SOLVER solver, integer ierr)

# **Specifying Options**

To specify options, the following functions are used:

• C int lis\_solver\_set\_option(char \*text, LIS\_SOLVER solver)

• Fortran subroutine lis\_solver\_set\_option(character text, LIS\_SOLVER solver, integer ierr)

or

- C int lis\_solver\_set\_optionC(LIS\_SOLVER solver)
- Fortran subroutine lis\_solver\_set\_optionC(LIS\_SOLVER solver, integer ierr)

<code>lis\_solver\_set\_optionC</code> is a function which sets the options specified on the command line, and pass them to <code>solver</code> when the user's program is run.

The table below shows the available command line options, where  $-i \{cg|1\}$  means -i cg or -i 1 and -maxiter [1000] indicates that -maxiter defaults to 1,000.

Specifying Linear Solvers (Default: -i bicg)

Method	Option	Auxiliary Options	
CG	-i {cg 1}		
$\operatorname{BiCG}$	-i {bicg 2}		
CGS	-i {cgs 3}		
BiCGSTAB	-i {bicgstab 4}		
BiCGSTAB(l)	-i {bicgstabl 5}	-ell [2]	Degree $l$
GPBiCG	-i {gpbicg 6}		
TFQMR	-i {tfqmr 7}		
Orthomin(m)	-i {orthomin 8}	-restart [40]	Restart value $m$
GMRES(m)	-i {gmres 9}	-restart [40]	Restart value $m$
Jacobi	-i {jacobi 10}		
Gauss-Seidel	-i {gs 11}		
SOR	-i {sor 12}	-omega [1.9]	Relaxation coefficient $\omega$ (0 < $\omega$ < 2)
BiCGSafe	-i {bicgsafe 13}		
$\operatorname{CR}$	-i {cr 14}		
$\operatorname{BiCR}$	-i {bicr 15}		
CRS	-i {crs 16}		
BiCRSTAB	-i {bicrstab 17}		
GPBiCR	-i {gpbicr 18}		
BiCRSafe	-i {bicrsafe 19}		
FGMRES(m)	-i {fgmres 20}	-restart [40]	Restart value $m$
IDR(s)	-i {idrs 21}	-irestart [2]	Restart value $s$
MINRES	-i {minres 22}		

# Specifying Preconditioners (Default: -p none)

Preconditioner	Option	Auxiliary Options	
None	-p {none 0}		
Jacobi	-p {jacobi 1}		
ILU(k)	-p {ilu 2}	-ilu_fill [0]	Fill level $k$
SSOR	-p {ssor 3}	-ssor_w [1.0]	Relaxation coefficient $\omega$ (0 < $\omega$ < 2)
Hybrid	-p {hybrid 4}	-hybrid_i [sor]	Linear equations solver
		-hybrid_maxiter [25]	Maximum number of iterations
		-hybrid_tol [1.0e-3]	Convergence criterion
		-hybrid_w [1.5]	Relaxation coefficient $\omega$ for SOR
			$(0 < \omega < 2)$
		-hybrid_ell [2]	Degree $l$ of BiCGSTAB(l)
		-hybrid_restart [40]	Restart values for GMRES and Orthomin
I+S	-p {is 5}	-is_alpha [1.0]	Parameter $\alpha$ for preconditioner
			of $I + \alpha S^{(m)}$ type
		-is_m [3]	Parameter $m$ for preconditioner
			of $I + \alpha S^{(m)}$ type
SAINV	-p {sainv 6}	-sainv_drop [0.05]	Drop criterion
SA-AMG	-p {saamg 7}	-saamg_unsym [false]	Selects unsymmetric version
			(Matrix structure must be symmetric)
		-saamg_theta [0.05 0.12]	Drop criterion $a_{ij}^2 \le \theta^2  a_{ii}   a_{jj} $
			(symmetric or unsymmetric)
Crout ILU	-p {iluc 8}	-iluc_drop [0.05]	Drop criterion
		-iluc_rate [5.0]	Ratio of maximum fill-in
ILUT	-p {ilut 9}	-ilut_drop [0.05]	Drop criterion
		-ilut_rate [5.0]	Ratio of maximum fill-in
Additive Schwarz	-adds true	-adds_iter [1]	Number of iterations

# Other Options

	other options		
Option			
-maxiter [1000]	Maximum number of iterations		
-tol [1.0e-12]	Convergence criterion		
-print [0]	Display of the residual		
	-print {none 0} None		
	-print {mem 1} Saves the residual history in memory		
	-print {out 2} Displays the residual history		
	-print {all 3} Saves the residual history and displays it on the screen		
-scale [0]	Scaling		
	(The result will overwrite the original matrix and vectors)		
	-scale {none 0} No scaling		
	-scale {jacobi 1} Jacobi scaling $D^{-1}Ax = D^{-1}b$		
	(D represents the diagonal of $A = (a_{ij})$ )		
	-scale {symm_diag 2} Diagonal scaling $D^{-1/2}AD^{-1/2}x = D^{-1/2}b$		
	$(D^{-1/2} \text{ represents a diagonal matrix with } 1/\sqrt{a_{ii}}$		
	as diagonal)		
-initx_zeros [true]	Behavior of the initial vector $x_0$		
	-initx_zeros {false 0} Given values		
	-initx_zeros {true 1} All elements are set to 0		
-omp_num_threads [t]	Number of threads		
-	(t represents the maximum number of threads)		
-storage [0]	Matrix storage format		
-storage_block [2]	Block size of BSR and BSC		

# Precision (Default: -f double)

Precision	Option	Auxiliary Options
DOUBLE	-f {double 0}	
QUAD	-f {quad 1}	

### Solving Systems of Linear Equations

To solve the system of linear equations Ax = b, the following functions are used:

- C int lis\_solve(LIS\_MATRIX A, LIS\_VECTOR b, LIS\_VECTOR x, LIS\_SOLVER solver)
- Fortran subroutine lis\_solve(LIS\_MATRIX A, LIS\_VECTOR b, LIS\_VECTOR x, LIS\_SOLVER solver, integer ierr)

# 3.5 Solving Eigenvalue Problems

A program to solve the eigenvalue problem  $Ax = \lambda x$  with a specified solver is as follows:

```
C (for serial, OpenMP and MPI versions)

1: LIS_MATRIX A;
2: LIS_VECTOR x;
3: LIS_REAL evalue;
4: LIS_ESOLVER esolver;
5:
6: /* Create matrix and vector */
7:
8: lis_esolver_create(&esolver);
9: lis_esolver_set_option("-e ii -i bicg -p none",esolver);
10: lis_esolver_set_option("-etol 1.0e-12 -tol 1.0e-12",esolver);
11: lis_esolve(A,x,evalue,esolver);
```

```
Fortran (for serial, OpenMP and MPI versions)

1: LIS_MATRIX A

2: LIS_VECTOR x

3: LIS_REAL evalue

4: LIS_ESOLVER esolver

5:

6: /* Create matrix and vector */

7:

8: call lis_esolver_create(esolver,ierr)

9: call lis_esolver_set_option('-e ii -i bicg -p none',esolver,ierr)

10: call lis_esolver_set_option('-etol 1.0e-12 -tol 1.0e-12',esolver,ierr)

11: call lis_esolve(A,x,evalue,esolver,ierr)
```

# Creating Eigensolvers

To create an eigensolver, the following functions are used:

- C int lis\_esolver\_create(LIS\_ESOLVER \*esolver)
- Fortran subroutine lis\_esolver\_create(LIS\_ESOLVER esolver, integer ierr)

# **Specifying Options**

To specify options, the following functions are used:

- C int lis\_esolver\_set\_option(char \*text, LIS\_ESOLVER esolver)
- Fortran subroutine lis\_esolver\_set\_option(character text, LIS\_ESOLVER esolver, integer ierr)

or

- C int lis\_esolver\_set\_optionC(LIS\_ESOLVER esolver)
- Fortran subroutine lis\_esolver\_set\_optionC(LIS\_ESOLVER esolver, integer ierr)

lis\_esolver\_set\_optionC is a function which sets the options specified on the command line, and pass them to esolver when the user's program is run.

The table below shows the available command line options, where -e {pi|1} means -e pi or -e 1 and -emaxiter [1000] indicates that -emaxiter defaults to 1,000.

Specifying Eigensolvers (Default: -i bicg)

Method	Option	Auxiliary Options	
Power Iteration	-e {pi 1}		
Inverse Iteration	-e {ii 2}	-i [bicg]	Linear solver
Approximate Inverse Iteration	-e {aii 3}		
Rayleigh Quotient Iteration	-e {rqi 4}	-i [bicg]	Linear solver
Subspace Iteration	-e {si 5}	-ss [2]	Size of subspace
		-m [O]	Mode number
Lanczos Iteration	-e {li 6}	-ss [2]	Size of subspace
		-m [O]	Mode number
Conjugate Gradient	-e {cg 7}		
Conjugate Residual	-e {cr 8}		

Specifying Preconditioners (Default: -p ilu)

Preconditioner	Option	Auxiliary Options	
None	-p {none 0}		
Jacobi	-p {jacobi 1}		
ILU(k)	-p {ilu 2}	-ilu_fill [0]	Fill level $k$
SSOR	-p {ssor 3}	-ssor_w [1.0]	Relaxation coefficient $\omega$ (0 < $\omega$ < 2)
Hybrid	-p {hybrid 4}	-hybrid_i [sor]	Linear equations solver
		-hybrid_maxiter [25]	Maximum number of iterations
		-hybrid_tol [1.0e-3]	Convergence criterion
		-hybrid_w [1.5]	Relaxation coefficient $\omega$ for SOR $(0 < \omega < 2)$
		-hybrid_ell [2]	Degree $l$ of BiCGSTAB(1)
		-hybrid_restart [40]	Restart values for GMRES and Orthomin
I+S	-p {is 5}	-is_alpha [1.0]	Parameter $\alpha$ for preconditioner
	_	-	of $I + \alpha S^{(m)}$ type
		-is_m [3]	Parameter $m$ for preconditioner
			of $I + \alpha S^{(m)}$ type
SAINV	-p {sainv 6}	-sainv_drop [0.05]	Drop criterion
SA-AMG	-p {saamg 7}	-saamg_unsym [false]	Selects unsymmetric version
			(Matrix structure must be symmetric)
		-saamg_theta [0.05 0.12]	Drop criterion $a_{ij}^2 \le \theta^2  a_{ii}   a_{jj} $
			(symmetric or unsymmetric)
Crout ILU	-p {iluc 8}	-iluc_drop [0.05]	Drop criterion
		-iluc_rate [5.0]	Ratio of maximum fill-in
ILUT	-p {ilut 9}	-ilut_drop [0.05]	Drop criterion
		-ilut_rate [5.0]	Ratio of maximum fill-in
Additive Schwarz	-adds true	-adds_iter [1]	Number of iterations

# Other Options

Option			
-emaxiter [1000]	Maximum number of iterations		
-etol [1.0e-12]	Convergence criterion		
-eprint [0]	Display of the residual		
	-eprint {none 0} None		
	-eprint {mem 1} Saves the residual history in memory		
	-eprint {out 2} Displays the residual history		
	-eprint {all 3} Saves the residual history and displays it on the screen		
-ie [ii]	Inner eigensolver used in Lanczos Iteration or Subspace Iteration		
	-ie {pi 1} Power Iteration (Subspace Iteration only)		
	-ie {ii 2} Inverse Iteration		
	-ie {aii 3} Approximate Inverse Iteration		
	-ie {rqi 4} Rayleigh Quotient Iteration		
-shift [0.0]	Amount of shift		
-initx_ones [true]	Behavior of the initial vector $x_0$		
	-initx_ones {false 0} Given values		
	-initx_ones {true 1} All elements are set to 1		
<pre>-omp_num_threads [t]</pre>	Number of threads		
	(t represents the maximum number of threads)		
-estorage [0]	Matrix storage format		
-estorage_block [2]	Block size of BSR and BSC		

Precision	Option	Auxiliary Options
DOUBLE	-ef {double 0}	
QUAD	-ef {quad 1}	

# Solving Eivenvalue Problems

To solve the eigenvalue problem  $Ax = \lambda x$ , the following functions are used:

- C int lis\_esolve(LIS\_MATRIX A, LIS\_VECTOR x, LIS\_REAL evalue, LIS\_ESOLVER esolver)
- Fortran subroutine lis\_esolve(LIS\_MATRIX A, LIS\_VECTOR x, LIS\_REAL evalue, LIS\_ESOLVER esolver, integer ierr)

# 3.6 Sample Programs

The following are the programs for solving the system of linear equations Ax = b, where the matrix A is a tridiagonal matrix

$$\begin{pmatrix}
2 & -1 & & & & \\
-1 & 2 & -1 & & & & \\
& \ddots & \ddots & \ddots & & \\
& & -1 & 2 & -1 & \\
& & & -1 & 2
\end{pmatrix}$$

of size 12. The the right hand side vector b is set to make all the elements of the solution x is 1. The program is located in the directory lis-(\$VERSION)/test.

```
- Test program: test4.c -
 1: #include <stdio.h>
 2: #include "lis.h"
 3: main(int argc, char *argv[])
 4: {
5:
        int i,n,gn,is,ie,iter;
        LIS_MATRIX A;
 6:
        LIS_VECTOR b,x,u;
7:
        LIS_SOLVER solver;
8:
9:
        n = 12;
10:
        lis_initialize(&argc,&argv);
11:
        lis_matrix_create(LIS_COMM_WORLD,&A);
12:
        lis_matrix_set_size(A,0,n);
13:
        lis_matrix_get_size(A,&n,&gn)
14:
        lis_matrix_get_range(A,&is,&ie)
        for(i=is;i<ie;i++)</pre>
15:
16:
17:
            if( i>0 ) lis_matrix_set_value(LIS_INS_VALUE,i,i-1,-1.0,A);
18:
            if( i<gn-1 ) lis_matrix_set_value(LIS_INS_VALUE,i,i+1,-1.0,A);</pre>
19:
            lis_matrix_set_value(LIS_INS_VALUE,i,i,2.0,A);
20:
        lis_matrix_set_type(A,LIS_MATRIX_CRS);
21:
        lis_matrix_assemble(A);
22:
23:
24:
        lis_vector_duplicate(A,&u);
25:
        lis_vector_duplicate(A,&b);
26:
        lis_vector_duplicate(A,&x);
27:
        lis_vector_set_all(1.0,u);
28:
        lis_matvec(A,u,b);
29:
30:
        lis_solver_create(&solver);
31:
        lis_solver_set_optionC(solver);
32:
        lis_solve(A,b,x,solver);
33:
        lis_solver_get_iters(solver,&iter);
34:
        printf("iter = %d\n",iter);
35:
        lis_vector_print(x);
36:
        lis_matrix_destroy(A);
37:
        lis_vector_destroy(u);
38:
        lis_vector_destroy(b);
39:
        lis_vector_destroy(x);
40:
        lis_solver_destroy(solver);
41:
        lis_finalize();
42:
        return 0;
43: }
}
```

```
Test program: test4f.F -
 1:
         implicit none
 3:#include "lisf.h"
 4:
 5:
         integer
                           i,n,gn,is,ie,iter,ierr
         LIS_MATRIX
 6:
                           Α
         LIS_VECTOR
7:
                           b,x,u
         LIS_SOLVER
 8٠
                            solver
         n = 12
9:
10:
         call lis_initialize(ierr)
11:
         call lis_matrix_create(LIS_COMM_WORLD, A, ierr)
12:
         call lis_matrix_set_size(A,0,n,ierr)
13:
         call lis_matrix_get_size(A,n,gn,ierr)
14:
         call lis_matrix_get_range(A,is,ie,ierr)
15:
         do i=is,ie-1
           if( i>1 ) call lis_matrix_set_value(LIS_INS_VALUE,i,i-1,-1.0d0,
16:
17:
                                                   A,ierr)
           if( i<gn ) call lis_matrix_set_value(LIS_INS_VALUE,i,i+1,-1.0d0,</pre>
18:
19:
                                                   A,ierr)
           call lis_matrix_set_value(LIS_INS_VALUE,i,i,2.0d0,A,ierr)
20:
21:
         enddo
22:
         call lis_matrix_set_type(A,LIS_MATRIX_CRS,ierr)
23:
         call lis_matrix_assemble(A,ierr)
24:
25:
         call lis_vector_duplicate(A,u,ierr)
26:
         call lis_vector_duplicate(A,b,ierr)
27:
         call lis_vector_duplicate(A,x,ierr)
28:
         call lis_vector_set_all(1.0d0,u,ierr)
29:
         call lis_matvec(A,u,b,ierr)
30:
31:
         call lis_solver_create(solver,ierr)
32:
         call lis_solver_set_optionC(solver,ierr)
33:
         call lis_solve(A,b,x,solver,ierr)
34:
         call lis_solver_get_iters(solver,iter,ierr)
35:
         write(*,*) 'iter = ',iter
36:
         call lis_vector_print(x,ierr)
37:
         call lis_matrix_destroy(A,ierr)
38:
         call lis_vector_destroy(b,ierr)
         call lis_vector_destroy(x,ierr)
39:
40:
         call lis_vector_destroy(u,ierr)
         call lis_solver_destroy(solver,ierr)
41:
42:
         call lis_finalize(ierr)
43:
44:
         stop
45:
         end
```

# 3.7 Compiling and Linking

Provided below is an example test4.c located in the directory lis-(\$VERSION)/test, compiled on SGI Altix 3700 using Intel C/C++ Compiler 8.1 (icc). Since the library includes Fortran 90 codes when the SA-AMG preconditioner is used, linking is processed by a Fortran 90 compiler.

```
Compiling
        >icc -c -I$(INSTALLDIR)/include test4.c
  Linking
        >icc -o test4 test4.o -llis
  Linking (with SA-AMG)
       >ifort -nofor_main -o test4 test4.o -llis
  for OpenMP verion —
  Compiling
       >icc -c -openmp -I$(INSTALLDIR)/include test4.c
  Linking
       >icc -openmp -o test4 test4.o -llis
  Linking (with SA-AMG)
        >ifort -nofor_main -openmp -o test4 test4.o -llis
  for MPI version —
  Compiling
        >icc -c -DUSE_MPI -I$(INSTALLDIR)/include test4.c
  Linking
       >icc -o test4 test4.o -llis -lmpi
  Linking (with SA-AMG)
       >ifort -nofor_main -o test4 test4.o -llis -lmpi
  - for OpenMP + MPI version —
  Compiling
        >icc -c -openmp -DUSE_MPI -I$(INSTALLDIR)/include test4.c
       >icc -openmp -o test4 test4.o -llis -lmpi
  Linking (with SA-AMG)
        >ifort -nofor_main -openmp -o test4 test4.o -llis -lmpi
   Provided below is an example test4f.F located in the directory lis-($VERSION)/test, compiled
on SGI Altix 3700 using Intel Fortran Compiler 8.1 (ifort). Since an #include statement is used in the
program, a compiler option -fpp is specified to use the preprocessor.
  for serial version -
  Compiling
       >ifort -c -fpp -I$(INSTALLDIR)/include test4f.F
  Linking
       >ifort -o test4 test4.o -llis
  - for OpenMP version -
  Compiling
        >ifort -c -fpp -openmp -I$(INSTALLDIR)/include test4f.F
        >ifort -openmp -o test4 test4.o -llis
```

```
for MPI version -
  Compiling
       >ifort -c -fpp -DUSE_MPI -I$(INSTALLDIR)/include test4f.F
  Linking
       >ifort -o test4 test4.o -llis -lmpi
  - for OpenMP + MPI version -
  Compiling
       >ifort -c -fpp -openmp -DUSE_MPI -I$(INSTALLDIR)/include test4f.F
  Linking
       >ifort -openmp -o test4 test4.o -llis -lmpi
3.8
      Running
The test programs test4 and test4f in the directory lis-($VERSION)/test are run as follows:
for serial version
      >./test4 -i bicgstab
for OpenMP version
```

The following results will be returned:

for OpenMP + MPI version

for MPI version

>env OMP\_NUM\_THREADS=2 ./test4 -i bicgstab

>mpirun -np 2 env OMP\_NUM\_THREADS=2 ./test4 -i bicgstab

>mpirun -np 2 ./test4 -i bicgstab

precision : double solver : BiCGSTAB 4 : none precon storage : CRS lis\_solve : normal end iter = 60 1.000000e+000 1 1.000000e+000 2 1.000000e+000 3 1.000000e+000 4 1.000000e+000 5 1.000000e+000 6 1.000000e+000 7 1.000000e+000 8 1.000000e+000 9 1.000000e+000

> 10 1.000000e+000 11 1.000000e+000

# 4 Quadruple Precision Operations

Double precision operations sometimes require a large number of iterations because of the rounding error. Lis supports "double-double", or quadruple precision operations by combining two double precision floating point numbers[15, 16]. To use the quadruple precision with the same interface as the double precision operations, both the matrix and vectors are assumed to be double precision. Lis also supports the performance acceleration of quadruple precision operations with SIMD instructions, such as Intel's Streaming SIMD Extensions (SSE) and IBM's Fused Multiply-Add (FMA)[24].

## 4.1 Using Quadruple Precision Operations

The test program test5.c solves a system of linear equations Ax = b, where A is a Toeplitz matrix

$$\begin{pmatrix} 2 & 1 & & & & \\ 0 & 2 & 1 & & & \\ \gamma & 0 & 2 & 1 & & \\ & \ddots & \ddots & \ddots & \ddots & \\ & & \gamma & 0 & 2 & 1 \\ & & & \gamma & 0 & 2 \end{pmatrix}.$$

The right hand vector is set to make all the elements of the solution to be 1. The value n is the size of the matrix A. test5 with -f option is run as follows:

## Double precision

```
By entering >./test5 200 2.0 -f double the following results will be returned:
```

```
n = 200, gamma = 2.000000
initial vector x = 0
precision : double
solver
         : BiCG 2
precon
          : none
         : CRS
storage
lis_solve : LIS_MAXITER(code=4)
BiCG: number of iterations
                              = 1001 (double = 1001, quad = 0)
BiCG: elapsed time
                              = 2.044368e-02 sec.
BiCG:
       preconditioner
                              = 4.768372e-06 sec.
BiCG:
         matrix creation
                              = 4.768372e-06 sec.
BiCG:
        linear solver
                              = 2.043891e-02 sec.
BiCG: relative residual 2-norm = 8.917591e+01
```

### Quadruple precision

By entering >./test5 200 2.0 -f quad the following results will be returned:

```
n = 200, gamma = 2.000000
initial vector x = 0
precision : quad
solver : BiCG 2
precon : none
storage : CRS
lis_solve : normal end
```

BiCG: number of iterations = 230 (double = 230, quad = 0)
BiCG: elapsed time = 2.267408e-02 sec.
BiCG: preconditioner = 4.549026e-04 sec.
BiCG: matrix creation = 5.006790e-06 sec.
BiCG: linear solver = 2.221918e-02 sec.

BiCG: relative residual 2-norm = 6.499145e-11

# 5 Matrix Storage Formats

This section describes the matrix storage formats supported by the library. Assume that the matrix row (column) number begins with 0 and that the number of the nonzero elements of the matrix A of  $n \times n$  is nnz.

# 5.1 Compressed Row Storage (CRS)

The CRS format uses three arrays ptr, index and value to store data.

- value is a double precision array with a length of nnz, which stores the nonzero elements of the matrix A along the row.
- index is an integer array with a length of nnz, which stores the column numbers of the nonzero elements stored in the array value.
- ptr is an integer array with a length of n + 1, which stores the starting points of the rows of the arrays value and index.

## 5.1.1 Creating Matrices (for Serial and OpenMP Versions)

The right diagram in Figure 2 shows how the matrix A in Figure 2 is stored in the CRS format. A program to create the matrix in the CRS format is as follows:

Figure 2: The data structure of the CRS format (for serial and OpenMP versions).

```
for serial and OpenMP versions
 1: int
                  n,nnz;
 2: int
                  *ptr,*index;
 3: LIS_SCALAR
                  *value:
 4: LIS_MATRIX
                  A:
 5: n = 4; nnz = 8;
         = (int *)malloc( (n+1)*sizeof(int) );
 6: ptr
7: index = (int *)malloc( nnz*sizeof(int) );
8: value = (LIS_SCALAR *)malloc( nnz*sizeof(LIS_SCALAR) );
9: lis_matrix_create(0,&A);
10: lis_matrix_set_size(A,0,n);
12: ptr[0] = 0; ptr[1] = 1; ptr[2] = 3; ptr[3] = 5; ptr[4] = 8;
13: index[0] = 0; index[1] = 0; index[2] = 1; index[3] = 1;
14: index[4] = 2; index[5] = 0; index[6] = 2; index[7] = 3;
15: value[0] = 11; value[1] = 21; value[2] = 22; value[3] = 32;
16: value[4] = 33; value[5] = 41; value[6] = 43; value[7] = 44;
17:
18:
    lis_matrix_set_crs(nnz,ptr,index,value,A);
    lis_matrix_assemble(A);
```

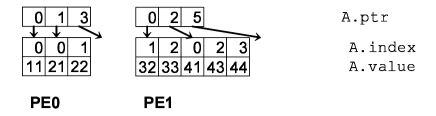


Figure 3: The data structure of the CRS format (for MPI version).

## 5.1.2 Creating Matrices (for MPI Version)

Figure 3 shows how the matrix A in Figure 2 is stored in the CRS format on two processing elements. A program to create the matrix in the CRS format on two processing elements is as follows:

```
for MPI version -
 1: int
                  i,k,n,nnz,my_rank;
 2: int
                  *ptr,*index;
 3: LIS_SCALAR
                  *value;
 4: LIS_MATRIX
                  A;
 5: MPI_Comm_rank(MPI_COMM_WORLD,&my_rank);
 6: if( my_rank==0 ) {n = 2; nnz = 3;}
                     {n = 2; nnz = 5;}
7: else
          = (int *)malloc( (n+1)*sizeof(int) );
8: ptr
9: index = (int *)malloc( nnz*sizeof(int) );
10: value = (LIS_SCALAR *)malloc( nnz*sizeof(LIS_SCALAR) );
11: lis_matrix_create(MPI_COMM_WORLD,&A);
12: lis_matrix_set_size(A,n,0);
13: if( my_rank==0 ) {
14:
        ptr[0] = 0; ptr[1] = 1; ptr[2] = 3;
15:
        index[0] = 0; index[1] = 0; index[2] = 1;
16:
        value[0] = 11; value[1] = 21; value[2] = 22;}
17: else {
        ptr[0] = 0; ptr[1] = 2; ptr[2] = 5;
18:
        index[0] = 1; index[1] = 2; index[2] = 0; index[3] = 2; index[4] = 3;
19:
        value[0] = 32; value[1] = 33; value[2] = 41; value[3] = 43; value[4] = 44;}
20:
    lis_matrix_set_crs(nnz,ptr,index,value,A);
21:
    lis_matrix_assemble(A);
```

## 5.1.3 Associating Arrays

To associate the arrays required by the CRS format with the matrix A, the following functions are used:

- C int lis\_matrix\_set\_crs(int nnz, int row[], int index[], LIS\_SCALAR value[], LIS\_MATRIX A)
- Fortran subroutine lis\_matrix\_set\_crs(integer nnz, integer row(), integer index(), LIS\_SCALAR value(), LIS\_MATRIX A, integer ierr)

# 5.2 Compressed Column Storage (CCS)

The CSS format uses three arrays ptr, index and value to store data.

- value is a double precision array with a length of nnz, which stores the values for the nonzero elements of the matrix A along the column.
- index is an integer array with a length of nnz, which stores the row numbers of the nonzero elements stored in the array value.
- ptr is an integer array with a length of n + 1, which stores the starting points of the rows of the arrays value and index.

## 5.2.1 Creating Matrices (for Serial and OpenMP Versions)

The right diagram in Figure 4 shows how the matrix A in Figure 4 is stored in the CCS format. A program to create the matrix in the CCS format is as follows:

Figure 4: The data structure of the CCS format (for serial and OpenMP versions).

```
for serial and OpenMP versions —
 1: int
                 n,nnz;
 2: int
                 *ptr,*index;
3: LIS_SCALAR
                 *value;
4: LIS_MATRIX
                 A:
5: n = 4; nnz = 8;
6: ptr = (int *)malloc( (n+1)*sizeof(int) );
7: index = (int *)malloc( nnz*sizeof(int) );
8: value = (LIS_SCALAR *)malloc( nnz*sizeof(LIS_SCALAR) );
9: lis_matrix_create(0,&A);
10: lis_matrix_set_size(A,0,n);
11:
12: ptr[0] = 0; ptr[1] = 3; ptr[2] = 5; ptr[3] = 7; ptr[4] = 8;
13: index[0] = 0; index[1] = 1; index[2] = 3; index[3] = 1;
14: index[4] = 2; index[5] = 2; index[6] = 3; index[7] = 3;
15: value[0] = 11; value[1] = 21; value[2] = 41; value[3] = 22;
16: value[4] = 32; value[5] = 33; value[6] = 43; value[7] = 44;
17:
18:
    lis_matrix_set_ccs(nnz,ptr,index,value,A);
19: lis_matrix_assemble(A);
```

## 5.2.2 Creating Matrices (for MPI Version)

Figure 5 shows how the matrix A in Figure 4 is stored on two processing elements. A program to create the matrix in the CCS format on two processing elements is as follows:

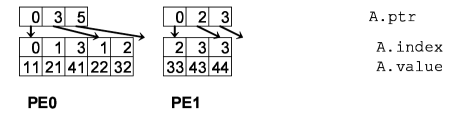


Figure 5: The data structure of the CCS format (for MPI version).

```
for MPI version -
 1: int
                  i,k,n,nnz,my_rank;
 2: int
                  *ptr,*index;
 3: LIS_SCALAR
                  *value;
 4: LIS_MATRIX
                  A:
5: MPI_Comm_rank(MPI_COMM_WORLD,&my_rank);
 6: if( my_rank==0 ) {n = 2; nnz = 3;}
                     {n = 2; nnz = 5;}
8: ptr
          = (int *)malloc( (n+1)*sizeof(int) );
9: index = (int *)malloc( nnz*sizeof(int) );
10: value = (LIS_SCALAR *)malloc( nnz*sizeof(LIS_SCALAR) );
11: lis_matrix_create(MPI_COMM_WORLD,&A);
12: lis_matrix_set_size(A,n,0);
13: if( my_rank==0 ) {
        ptr[0] = 0; ptr[1] = 3; ptr[2] = 5;
14:
15:
        index[0] = 0; index[1] = 1; index[2] = 3; index[3] = 1; index[4] = 2;
16:
        value[0] = 11; value[1] = 21; value[2] = 41; value[3] = 22; value[4] = 32}
17: else {
18:
        ptr[0] = 0; ptr[1] = 2; ptr[2] = 3;
19:
        index[0] = 2; index[1] = 3; index[2] = 3;
        value[0] = 33; value[1] = 43; value[2] = 44;}
20:
21:
     lis_matrix_set_ccs(nnz,ptr,index,value,A);
22:
     lis_matrix_assemble(A);
```

## 5.2.3 Associating Arrays

To associate the arrays required by the CCS format with the matrix A, the following functions are used:

- C int lis\_matrix\_set\_ccs(int nnz, int row[], int index[], LIS\_SCALAR value[],
- Fortran subroutine lis\_matrix\_set\_ccs(integer nnz, integer row(), integer index(), LIS\_SCALAR value(), LIS\_MATRIX A, integer ierr)

# 5.3 Modified Compressed Sparse Row (MSR)

The MSR format is a modified version of the CRS format. The MSR format is different in that it separates the diagonal elements before storing it. The MSR format uses two arrays index and value to store data. Assume that ndz represents the number of the zero elements of the diagonal.

- value is a double precision array with a length of nnz + ndz + 1, which stores the diagonal of the matrix A down to the n-th element. The n + 1-th element is not used. For the n + 2-th and after, the values of the nonzero elements except the diagonal of the matrix A are stored along the row.
- index is an integer array with a length of nnz + ndz + 1, which stores the starting points of the rows of the off-diagonal elements of the matrix A down to the n + 1-th element. For the n + 2-th and after, it stores the row numbers of the off-diagonal elements of the matrix A stored in the array value.

### 5.3.1 Creating Matrices (for Serial and OpenMP Versions)

The right diagram in Figure 6 shows how matrix A is stored in the MSR format. A program to create the matrix in the MSR format is as follows:

Figure 6: The data structure of the MSR format (for serial and OpenMP versions).

```
for serial and OpenMP versions -
 1: int
                  n,nnz,ndz;
                  *index;
 2: int
                  *value;
 3: LIS_SCALAR
 4: LIS_MATRIX
                  A:
5: n = 4; nnz = 8; ndz = 0;
 6: index = (int *)malloc( (nnz+ndz+1)*sizeof(int) );
7: value = (LIS_SCALAR *)malloc( (nnz+ndz+1)*sizeof(LIS_SCALAR) );
8: lis_matrix_create(0,&A);
9: lis_matrix_set_size(A,0,n);
10:
11: index[0] = 5; index[1] = 5; index[2] = 6; index[3] = 7;
12: index[4] = 9; index[5] = 0; index[6] = 1; index[7] = 0; index[8] =
13: value[0] = 11; value[1] = 22; value[2] = 33; value[3] = 44;
14: value[4] = 0; value[5] = 21; value[6] = 32; value[7] = 41; value[8] = 43;
15:
16:
    lis_matrix_set_msr(nnz,ndz,index,value,A);
    lis_matrix_assemble(A);
```

## 5.3.2 Creating Matrices (for MPI Version)

Figure 7 shows how the matrix A in Figure 6 is stored in the MSR format on two processing elements. A program to create the matrix in the MSR format on two processing element is as follows:

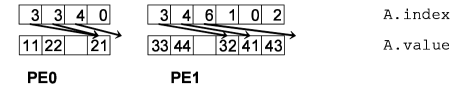


Figure 7: The data structure of the MSR format (for MPI version).

```
for MPI version -
 1: int
                  i,k,n,nnz,ndz,my_rank;
 2: int
                  *index;
 3: LIS_SCALAR
                  *value;
 4: LIS_MATRIX
                  A;
5: MPI_Comm_rank(MPI_COMM_WORLD,&my_rank);
 6: if( my_rank==0 ) {n = 2; nnz = 3; ndz = 0;}
                     {n = 2; nnz = 5; ndz = 0;}
7: else
8: index = (int *)malloc( (nnz+ndz+1)*sizeof(int) );
9: value = (LIS_SCALAR *)malloc( (nnz+ndz+1)*sizeof(LIS_SCALAR) );
10: lis_matrix_create(MPI_COMM_WORLD,&A);
11: lis_matrix_set_size(A,n,0);
12: if( my_rank==0 ) {
13:
        index[0] = 3; index[1] = 3; index[2] = 4; index[3] = 0;
14:
        value[0] = 11; value[1] = 22; value[2] = 0; value[3] = 21;}
15: else {
        index[0] = 3; index[1] = 4; index[2] = 6; index[3] = 1;
16:
        index[4] = 0; index[5] = 2;
17:
        value[0] = 33; value[1] = 44; value[2] = 0; value[3] = 32;
18:
        value[4] = 41; value[5] = 43;}
19:
    lis_matrix_set_msr(nnz,ndz,index,value,A);
20:
21:
    lis_matrix_assemble(A);
```

## 5.3.3 Associating Arrays

To associate the arrays required by the MSR format with the matrix A, the following functions are used:

- C int lis\_matrix\_set\_msr(int nnz, int ndz, int index[], LIS\_SCALAR value[], LIS\_MATRIX A)
- Fortran subroutine lis\_matrix\_set\_msr(integer nnz, integer ndz, integer index(), LIS\_SCALAR value(), LIS\_MATRIX A, integer ierr)

# 5.4 Diagonal (DIA)

DIA uses two arrays index and value to store data. Assume that nnd represents the number of the nonzero diagonal elements of the matrix A.

- value is a double precision array with a length of  $nnd \times n$ , which stores nonzero diagonal elements of the matrix A.
- index is an integer array with a length of nnd, which stores the offsets from the main diagonal.

For the OpenMP version, the following modifications have been made: DIA uses two arrays index and value to store data. Assume that nprocs represents the number of the threads.  $nnd_p$  is the number of the nonzero diagonal elements of the partial matrix into which the row block of the matrix A is divided. maxnnd is the maximum value  $nnd_p$ .

- value is a double precision array with a length of  $maxnnd \times n$ , which stores nonzero diagonal elements of the matrix A.
- index is an integer array with a length of  $nprocs \times maxnnd$ , which stores the offsets from the main diagonal.

## 5.4.1 Creating Matrices (for Serial Version)

The right diagram in Figure 8 shows how the matrix A in Figure 8 is stored in the DIA format. A program to create the matrix in the DIA format is as follows:

Figure 8: The data structure of the DIA format (for serial version).

```
for serial version -
 1: int
                  n,nnd;
 2: int
                  *index;
 3: LIS_SCALAR
                  *value;
 4: LIS_MATRIX
                  A;
 5: n = 4; nnd = 3;
 6: index = (int *)malloc( nnd*sizeof(int) );
7: value = (LIS_SCALAR *)malloc( n*nnd*sizeof(LIS_SCALAR) );
8: lis_matrix_create(0,&A);
9: lis_matrix_set_size(A,0,n);
11: index[0] = -3; index[1] = -1; index[2] = 0;
12: value[0] = 0; value[1] = 0; value[2] = 0; value[3] = 41;
13: value[4] = 0; value[5] = 21; value[6] = 32; value[7] = 43;
14: value[8] = 11; value[9] = 22; value[10] = 33; value[11] = 44;
15:
     lis_matrix_set_dia(nnd,index,value,A);
16:
     lis_matrix_assemble(A);
```

## 5.4.2 Creating Matrices (for OpenMP Version)

Figure 9 shows how the matrix A in Figure 8 is stored in the DIA format on two threads. A program to create the matrix in the DIA format on two threads is as follows:

-1	0		-3	-	0							A.index
0	21	11	22			0	41	32	43	33	44	A.value

Figure 9: The data structure of the DIA format (for OpenMP version).

```
for OpenMP version —
     1: int
                                                                         n, maxnnd, nprocs;
   2: int
                                                                          *index;
   3: LIS_SCALAR
                                                                          *value;
   4: LIS_MATRIX
                                                                        A;
   5: n = 4; maxnnd = 3; nprocs = 2;
    6: index = (int *)malloc( maxnnd*sizeof(int) );
   7: value = (LIS_SCALAR *)malloc( n*maxnnd*sizeof(LIS_SCALAR) );
   8: lis_matrix_create(0,&A);
   9: lis_matrix_set_size(A,0,n);
10:
11: index[0] = -1; index[1] = 0; index[2] = 0; index[3] = -3; index[4] = -1; index[5] = 0;
12: \ value[0] = 0; \ value[1] = 21; \ value[2] = 11; \ value[3] = 22; \ value[4] = 0; \ value[5] = 0; \\ value[5] = 0; \ value[6] = 0; \\ value[6] = 0; \ value[6] = 0; \\ value[6] = 0; \ value[6] = 0; \\ val
13: value[6] = 0; value[7] = 41; value[8] = 32; value[9] = 43; value[10] = 33; value[11] = 44;
14:
15: lis_matrix_set_dia(maxnnd,index,value,A);
16: lis_matrix_assemble(A);
```

## 5.4.3 Creating Matrices (for MPI Version)

Figure 10 shows how the matrix A in Figure 8 is stored in the DIA format on two processing elements. A program to create the matrix in the DIA format on two processing elements is as follows:

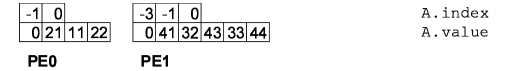


Figure 10: The data structure of the DIA format (for MPI version).

```
for MPI version -
 1: int
                  i,n,nnd,my_rank;
 2: int
                  *index;
3: LIS_SCALAR
                  *value;
 4: LIS_MATRIX
                  Α;
 5: MPI_Comm_rank(MPI_COMM_WORLD,&my_rank);
 6: if( my_rank==0 ) {n = 2; nnd = 2;}
7: else
                     {n = 2; nnd = 3;}
8: index = (int *)malloc( nnd*sizeof(int) );
9: value = (LIS_SCALAR *)malloc( n*nnd*sizeof(LIS_SCALAR) );
10: lis_matrix_create(MPI_COMM_WORLD,&A);
11: lis_matrix_set_size(A,n,0);
12: if( my_rank==0 ) {
13:
        index[0] = -1; index[1] = 0;
        value[0] = 0; value[1] = 21; value[2] = 11; value[3] = 22;}
14:
15: else {
16:
        index[0] = -3; index[1] = -1; index[2] = 0;
        value[0] = 0; value[1] = 41; value[2] = 32; value[3] = 43; value[4] = 33;
17:
18:
        value[5] = 44;
19:
    lis_matrix_set_dia(nnd,index,value,A);
20: lis_matrix_assemble(A);
```

#### 5.4.4 Associating Arrays

To associate the arrays required by the DIA format with the matrix A, the following functions are used:

- C int lis\_matrix\_set\_dia(int nnd, int index[], LIS\_SCALAR value[], LIS\_MATRIX A)
- Fortran subroutine lis\_matrix\_set\_dia(integer nnd, integer index(), LIS\_SCALAR value(), LIS\_MATRIX A, integer ierr)

## 5.5 Ellpack-Itpack generalized diagonal (ELL)

ELL uses two arrays index and value to store data. Assume that maxnzr is the maximum value for the number of the nonzero elements in the rows of the matrix A.

- value is a double precision array with a length of  $maxnzr \times n$ , which stores the nonzero elements of the rows of the matrix A along the column. The first column consists of the first nonzero elements of each row. If there is no nonzero elements to be stored, then 0 is stored.
- index is an integer array with a length of  $maxnzr \times n$ , which stores the column numbers of the nonzero elements stored in the array value. If the number of the nonzero elements in the *i*-th row is nnz, then index  $[nnz \times n + i]$  stores row number *i*.

## 5.5.1 Creating Matrices (for Serial and OpenMP Versions)

The right diagram in Figure 11 shows how the matrix A in Figure 11 is stored in the ELL format. A program to create the matrix in the ELL format is as follows:

Figure 11: The data structure of the ELL format (for serial and OpenMP versions).

```
for serial and OpenMP versions -
 1: int
                  n, maxnzr;
 2: int
                  *index:
 3: LIS_SCALAR
                  *value:
 4: LIS_MATRIX
5: n = 4; maxnzr = 3;
 6: index = (int *)malloc( n*maxnzr*sizeof(int) );
7: value = (LIS_SCALAR *)malloc( n*maxnzr*sizeof(LIS_SCALAR) );
8: lis_matrix_create(0,&A);
9: lis_matrix_set_size(A,0,n);
11: index[0] = 0; index[1] = 0; index[2] = 1; index[3] = 0; index[4] = 0; index[5] = 1;
12: index[6] = 2; index[7] = 2; index[8] = 0; index[9] = 1; index[10] = 2; index[11] = 3;
13: value[0] = 11; value[1] = 21; value[2] = 32; value[3] = 41; value[4] = 0; value[5] = 22;
14: value[6] = 33; value[7] = 43; value[8] = 0; value[9] = 0; value[10] = 0; value[11] = 44;
15:
    lis_matrix_set_ell(maxnzr,index,value,A);
16:
    lis_matrix_assemble(A);
```

## 5.5.2 Creating Matrices (for MPI Version)

Figure 12 shows how the matrix A in Figure 11 is stored in the ELL format. A program to create the matrix in the ELL format on two processing elements is as follows:

0 0 0 1	1 0 2 2 2 3	A.in
11 21 0 22	32 41 33 43 0 44	A.va
PE0	PE1	

Figure 12: The data structure of the ELL format (for MPI version).

```
for MPI version -
 1: int
                  i,n,maxnzr,my_rank;
 2: int
                  *index;
3: LIS_SCALAR
                  *value;
4: LIS_MATRIX
                 A;
 5: MPI_Comm_rank(MPI_COMM_WORLD,&my_rank);
 6: if( my_rank==0 ) {n = 2; maxnzr = 2;}
7: else
                     {n = 2; maxnzr = 3;}
8: index = (int *)malloc( n*maxnzr*sizeof(int) );
9: value = (LIS_SCALAR *)malloc( n*maxnzr*sizeof(LIS_SCALAR) );
10: lis_matrix_create(MPI_COMM_WORLD,&A);
11: lis_matrix_set_size(A,n,0);
12: if( my_rank==0 ) {
        index[0] = 0; index[1] = 0; index[2] = 0; index[3] = 1;
13:
14:
        value[0] = 11; value[1] = 21; value[2] = 0; value[3] = 22;}
15: else {
16:
        index[0] = 1; index[1] = 0; index[2] = 2; index[3] = 2; index[4] = 2;
17:
        index[5] = 3;
        value[0] = 32; value[1] = 41; value[2] = 33; value[3] = 43; value[4] = 0;
18:
19:
        value[5] = 44;
20: lis_matrix_set_ell(maxnzr,index,value,A);
21: lis_matrix_assemble(A);
```

#### 5.5.3 Associating Arrays

To associate an array required by the ELL format with the matrix A, the following functions are used:

- C int lis\_matrix\_set\_ell(int maxnzr, int index[], LIS\_SCALAR value[], LIS\_MATRIX A)
- Fortran subroutine lis\_matrix\_set\_ell(integer maxnzr, integer index(), LIS\_SCALAR value(), LIS\_MATRIX A, integer ierr)

# 5.6 Jagged Diagonal (JDS)

JDS first sorts the nonzero elements of the rows in decreasing order of size, and then stores them along the column. JDS uses four arrays perm, ptr, index and value to store date. Assume that maxnzr represents the maximum value of the number of the nonzero elements of the matrix A.

- perm is an integer array with a length of n, which stores the sorted row numbers.
- value is a double precision array with a length of nnz, which stores the jagged diagonal elements of the sorted matrix A. The first jagged diagonal consists of the first nonzero elements of each row. The next jagged diagonal consists of the second nonzero elements, and so on.
- index is an integer array with a length of nnz, which stores the row numbers of the nonzero elements stored in the array value.
- ptr is an integer array with a length of maxnzr + 1, which stores the starting points of the jagged diagonal elements.

For the OpenMP version, the following modifications have been made: JDS uses four arrays perm, ptr, index and value to store data. Assume that nprocs is the number of the threads.  $maxnzr_p$  is the number of the nonzero diagonal elements of the partial matrix into which the row block of the matrix A is divided. maxmaxnzr is the maximum value of  $maxnzr_p$ .

- perm is an integer array with a length of n, which stores the sorted row numbers.
- value is a double precision array with a length of nnz, which stores jagged diagonal elements of the sorted matrix A. The first jagged diagonal consists of the first nonzero elements of each row. The next jagged diagonal consist of the second nonzero elements of each row, and so on.
- index is an integer array with a length of nnz, which stores the row numbers of the nonzero elements stored in the array value.
- ptr is an integer array with a length of  $nprocs \times (maxmaxnzr+1)$ , which stores the starting points of the jagged diagonal elements.

# 5.6.1 Creating Matrices (for Serial Version)

The right diagram in Figure 13 shows how the matrix A in Figure 13 is stored in the JDS format. A program to create the matrix in the JDS format is as follows:

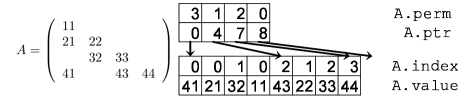


Figure 13: The data structure of the JDS format (for serial version).

```
for serial version -
 1: int
                 n,nnz,maxnzr;
 2: int
                 *perm,*ptr,*index;
3: LIS_SCALAR
                 *value;
4: LIS_MATRIX
                 A;
5: n = 4; nnz = 8; maxnzr = 3;
 6: perm = (int *)malloc( n*sizeof(int) );
7: ptr = (int *)malloc( (maxnzr+1)*sizeof(int) );
8: index = (int *)malloc( nnz*sizeof(int) );
9: value = (LIS_SCALAR *)malloc( nnz*sizeof(LIS_SCALAR) );
10: lis_matrix_create(0,&A);
11: lis_matrix_set_size(A,0,n);
13: perm[0] = 3; perm[1] = 1; perm[2] = 2; perm[3] = 0;
14: ptr[0] = 0; ptr[1] = 4; ptr[2] = 7; ptr[3] = 8;
15: index[0] = 0; index[1] = 0; index[2] = 1; index[3] = 0;
16: index[4] = 2; index[5] = 1; index[6] = 2; index[7] = 3;
17: value[0] = 41; value[1] = 21; value[2] = 32; value[3] = 11;
18: value[4] = 43; value[5] = 22; value[6] = 33; value[7] = 44;
19:
20: lis_matrix_set_jds(nnz,maxnzr,perm,ptr,index,value,A);
21: lis_matrix_assemble(A);
```

## 5.6.2 Creating Matrices (for OpenMP Version)

Figure 14 shows how the matrix A in Figure 13 is stored in the JDS format on two threads. A program to create the matrix in the JDS format on two threads is as follows:

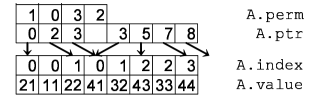


Figure 14: The data structure of the JDS format (for OpenMP version).

```
for OpenMP version -
 1: int
                  n,nnz,maxmaxnzr,nprocs;
 2: int
                   *perm,*ptr,*index;
 3: LIS_SCALAR
                   *value;
 4: LIS_MATRIX
                  Α;
 5: n = 4; nnz = 8; maxmaxnzr = 3; nprocs = 2;
 6: perm = (int *)malloc( n*sizeof(int) );
 7: ptr = (int *)malloc( nprocs*(maxmaxnzr+1)*sizeof(int) );
 8: index = (int *)malloc( nnz*sizeof(int) );
 9: value = (LIS_SCALAR *)malloc( nnz*sizeof(LIS_SCALAR) );
10: lis_matrix_create(0,&A);
11: lis_matrix_set_size(A,0,n);
13: perm[0] = 1; perm[1] = 0; perm[2] = 3; perm[3] = 2;
14: ptr[0] = 0; ptr[1] = 2; ptr[2] = 3; ptr[3] = 0;
15: ptr[4] = 3; ptr[5] = 5; ptr[6] = 7; ptr[7] = 8;
16: index[0] = 0; index[1] = 0; index[2] = 1; index[3] = 0;
17: index[4] = 1; index[5] = 2; index[6] = 2; index[7] = 3;
18: value[0] = 21; value[1] = 11; value[2] = 22; value[3] = 41;
19: value[4] = 32; value[5] = 43; value[6] = 33; value[7] = 44;
20:
    lis_matrix_set_jds(nnz,maxmaxnzr,perm,ptr,index,value,A);
22: lis_matrix_assemble(A);
```

## 5.6.3 Creating Matrices (for MPI Version)

Figure 15 shows how the matrix A in Figure 13 is stored in the JDS format on two processing elements. A program to create the matrix in the JDS format on two processing elements is as follows:

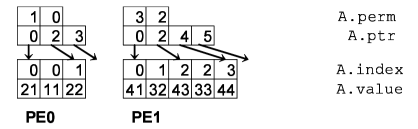


Figure 15: The data structure of the JDS format (for MPI version).

```
for MPI version -
 1: int
                  i,n,nnz,maxnzr,my_rank;
 2: int
                  *perm,*ptr,*index;
 3: LIS_SCALAR
                  *value;
 4: LIS_MATRIX
 5: MPI_Comm_rank(MPI_COMM_WORLD,&my_rank);
 6: if( my_rank==0 ) {n = 2; nnz = 3; maxnzr = 2;}
                     {n = 2; nnz = 5; maxnzr = 3;}
7: else
8: perm = (int *)malloc( n*sizeof(int) );
9: ptr = (int *)malloc( (maxnzr+1)*sizeof(int) );
10: index = (int *)malloc( nnz*sizeof(int) );
11: value = (LIS_SCALAR *)malloc( nnz*sizeof(LIS_SCALAR) );
12: lis_matrix_create(MPI_COMM_WORLD,&A);
13: lis_matrix_set_size(A,n,0);
14: if( my_rank==0 ) {
15:
        perm[0] = 1; perm[1] = 0;
        ptr[0] = 0; ptr[1] = 2; ptr[2] = 3;
16:
        index[0] = 0; index[1] = 0; index[2] = 1;
17:
        value[0] = 21; value[1] = 11; value[2] = 22;}
18:
19: else {
        perm[0] = 3; perm[1] = 2;
20:
21:
        ptr[0] = 0; ptr[1] = 2; ptr[2] = 4; ptr[3] = 5;
        index[0] = 0; index[1] = 1; index[2] = 2; index[3] = 2; index[4] = 3;
22:
        value[0] = 41; value[1] = 32; value[2] = 43; value[3] = 33; value[4] = 44;}
23:
   lis_matrix_set_jds(nnz,maxnzr,perm,ptr,index,value,A);
25:
    lis_matrix_assemble(A);
```

## 5.6.4 Associating Arrays

To associate an array required by the JDS format with the matrix A, the following functions are used:

- C int lis\_matrix\_set\_jds(int nnz, int maxnzr, int perm[], int ptr[], int index[], LIS\_SCALAR value[], LIS\_MATRIX A)
- Fortran subroutine lis\_matrix\_set\_jds(integer nnz, integer maxnzr, integer ptr(), integer index(), LIS\_SCALAR value(), LIS\_MATRIX A, integer ierr)

# 5.7 Block Sparse Row (BSR)

BSR breaks down the matrix A into partial matrices called blocks, with a size of  $r \times c$ . BSR stores nonzero blocks, in which at least one nonzero element exists, with the same step as that for the CRS format. Assume that nr = n/r and nnzb are the numbers of nonzero blocks of A. BSR uses three arrays bptr, bindex and value to store matrices.

- value is a double precision array with a length of  $nnzb \times r \times c$ , which stores all the elements of the nonzero blocks.
- bindex is an integer array with a length of nnzb, which stores block column numbers of the nonzero blocks.
- bptr is an integer array with a length of nr + 1, which stores the starting points of the block rows in the array bindex.

### 5.7.1 Creating Matrices (for Serial and OpenMP Versions)

The right diagram in Figure 16 shows how the matrix A in Figure 16 is stored in the BSR format. A program to create the matrix in the BSR format is as follows:

$$A = \begin{pmatrix} 11 & & & & \\ 21 & 22 & & & \\ \hline & 32 & 33 & \\ 41 & & 43 & 44 \end{pmatrix} \begin{pmatrix} 0 & 1 & 3 & & & \\ \hline & 0 & 0 & 1 & & \\ \hline & 1 & 21 & 0 & 22 & 0 & 41 & 32 & 0 & 33 & 43 & 0 & 44 \\ \hline & 11 & 21 & 0 & 22 & 0 & 41 & 32 & 0 & 33 & 43 & 0 & 44 \\ \hline \end{pmatrix} & A. \text{bindex}$$

Figure 16: The data structure of the BSR format (for serial and OpenMP versions).

```
for serial and OpenMP versions —
 1: int
                 n,bnr,bnc,nr,nc,bnnz;
 2: int
                  *bptr,*bindex;
3: LIS_SCALAR
                  *value;
 4: LIS_MATRIX
                 A;
5: n = 4; bnr = 2; bnc = 2; bnnz = 3; nr = (n-1)/bnr+1; nc = (n-1)/bnc+1;
         = (int *)malloc( (nr+1)*sizeof(int) );
7: bindex = (int *)malloc( bnnz*sizeof(int) );
8: value = (LIS_SCALAR *)malloc( bnr*bnc*bnnz*sizeof(LIS_SCALAR) );
9: lis_matrix_create(0,&A);
10: lis_matrix_set_size(A,0,n);
11:
12: bptr[0] = 0; bptr[1] = 1; bptr[2] = 3;
13: bindex[0] = 0; bindex[1] = 0; bindex[2] = 1;
14: value[0] = 11; value[1] = 21; value[2] = 0; value[3] = 22;
             = 0; value[5] = 41; value[6] = 32; value[7] = 0;
15: value[4]
16: value[8] = 33; value[9] = 43; value[10] = 0; value[11] = 44;
17:
18:
    lis_matrix_set_bsr(bnr,bnc,bnnz,bptr,bindex,value,A);
19:
    lis_matrix_assemble(A);
```

## 5.7.2 Creating Matrices (for MPI Version)

Figure 17 shows how the matrix A in Figure 16 is stored in the BSR format on two processing elements. A program to create the matrix in the BSR format on two processing elements is as follows:

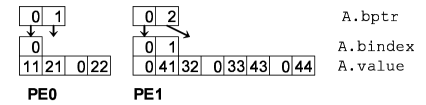


Figure 17: The data structure of the BSR format (for MPI version).

```
for MPI version -
 1: int
                  n,bnr,bnc,nr,nc,bnnz,my_rank;
 2: int
                  *bptr,*bindex;
3: LIS_SCALAR
                  *value;
 4: LIS_MATRIX
                  A;
 5: MPI_Comm_rank(MPI_COMM_WORLD,&my_rank);
 6: if( my_rank==0 ) {n = 2; bnr = 2; bnc = 2; bnnz = 1; nr = (n-1)/bnr+1; nc = (n-1)/bnc+1;}
                     {n = 2; bnr = 2; bnc = 2; bnnz = 2; nr = (n-1)/bnr+1; nc = (n-1)/bnc+1;}
7: else
8: bptr
          = (int *)malloc( (nr+1)*sizeof(int) );
9: bindex = (int *)malloc( bnnz*sizeof(int) );
10: value = (LIS_SCALAR *)malloc( bnr*bnc*bnnz*sizeof(LIS_SCALAR) );
11: lis_matrix_create(MPI_COMM_WORLD,&A);
12: lis_matrix_set_size(A,n,0);
13: if( my_rank==0 ) {
14:
        bptr[0] = 0; bptr[1] = 1;
15:
        bindex[0] = 0;
        value[0] = 11; value[1] = 21; value[2] = 0; value[3] = 22;}
16:
17: else {
18:
        bptr[0] = 0; bptr[1] = 2;
        bindex[0] = 0; bindex[1] = 1;
19:
20:
        value[0] = 0; value[1] = 41; value[2] = 32; value[3] = 0;
        value[4] = 33; value[5] = 43; value[6] = 0; value[7] = 44;}
21:
22: lis_matrix_set_bsr(bnr,bnc,bnnz,bptr,bindex,value,A);
23: lis_matrix_assemble(A);
```

#### 5.7.3 Associating Arrays

To associate the arrays required by the BSR format with the matrix A, the following functions are used:

- C int lis\_matrix\_set\_bsr(int bnr, int bnc, int bnnz, int bptr[], int bindex[], LIS\_SCALAR value[], LIS\_MATRIX A)
- Fortran subroutine lis\_matrix\_set\_bsr(integer bnr, integer bnc, integer bnnz, integer bptr(), integer bindex(), LIS\_SCALAR value(), LIS\_MATRIX A, integer ierr)

# 5.8 Block Sparse Column (BSC)

BSC breaks down the matrix A into partial matrices called blocks, with a size of  $r \times c$ . BSC stores nonzero blocks, in which at least one nonzero block exists, in the same step as that for the CCS format. Assume that nc = n/c and nnzb are the numbers of nonzero blocks of A. BSC uses three arrays bptr, bindex and value to store matrices.

- value is a double precision array with a length of  $nnzb \times r \times c$ , which stores all the elements of the nonzero blocks.
- bindex is an integer array with a length of nnzb, which stores the block row numbers of the nonzero blocks.
- bptr is an integer array with a length of nc+1, which stores the starting points of the block columns in the array bindex.

#### 5.8.1 Creating Matrices (for Serial and OpenMP Versions)

The right diagram in Figure 18 shows how the matrix A in Figure 18 is stored in the BSC format. A program to create the matrix in the BSC format is as follows:

$$A = \begin{pmatrix} 11 & & & & \\ 21 & 22 & & & \\ \hline & 32 & 33 & \\ 41 & & 43 & 44 \end{pmatrix} \begin{pmatrix} 0 & 1 & 3 & & & \\ \hline & 0 & 1 & 1 & & \\ \hline & 0 & 1 & 1 & & \\ \hline & 11 & 21 & 0 & 22 & 0 & 41 & 32 & 0 & 33 & 43 & 0 & 44 \\ \hline & 11 & 21 & 0 & 22 & 0 & 41 & 32 & 0 & 33 & 43 & 0 & 44 \\ \hline \end{pmatrix} & A. \text{ bindex}$$

Figure 18: The data structure of the BSC format (for serial and OpenMP versions).

```
for serial and OpenMP versions —
 1: int
                  n,bnr,bnc,nr,nc,bnnz;
 2: int
                  *bptr,*bindex;
3: LIS_SCALAR
                  *value;
 4: LIS_MATRIX
                  A;
5: n = 4; bnr = 2; bnc = 2; bnnz = 3; nr = (n-1)/bnr+1; nc = (n-1)/bnc+1;
         = (int *)malloc( (nc+1)*sizeof(int) );
7: bindex = (int *)malloc( bnnz*sizeof(int) );
8: value = (LIS_SCALAR *)malloc( bnr*bnc*bnnz*sizeof(LIS_SCALAR) );
9: lis_matrix_create(0,&A);
10: lis_matrix_set_size(A,0,n);
11:
12: bptr[0] = 0; bptr[1] = 1; bptr[2] = 3;
13: bindex[0] = 0; bindex[1] = 1; bindex[2] = 1;
14: value[0] = 11; value[1] = 21; value[2] = 0; value[3] = 22;
             = 0; value[5] = 41; value[6] = 32; value[7] = 0;
15: value[4]
16: value[8] = 33; value[9] = 43; value[10] = 0; value[11] = 44;
17:
18:
    lis_matrix_set_bsc(bnr,bnc,bnnz,bptr,bindex,value,A);
19:
    lis_matrix_assemble(A);
```

## 5.8.2 Creating Matrices (for MPI Version)

Figure 19 shows how the matrix A in Figure 18 is stored in the BSC format on two processing elements. A program to create the matrix in the BSC format on two processing elements is as follows:

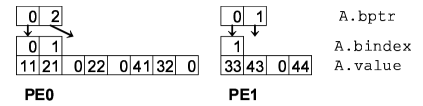


Figure 19: The data structure of the BSC format (for MPI version).

```
for MPI version -
 1: int
                  n,bnr,bnc,nr,nc,bnnz,my_rank;
 2: int
                  *bptr,*bindex;
3: LIS_SCALAR
                  *value;
 4: LIS_MATRIX
                  A;
 5: MPI_Comm_rank(MPI_COMM_WORLD,&my_rank);
 6: if( my_n = 0 ) {n = 2; bnr = 2; bnc = 2; bnnz = 2; nr = (n-1)/bnr+1; nc = (n-1)/bnc+1;}
                     {n = 2; bnr = 2; bnc = 2; bnnz = 1; nr = (n-1)/bnr+1; nc = (n-1)/bnc+1;}
7: else
8: bptr
          = (int *)malloc( (nr+1)*sizeof(int) );
9: bindex = (int *)malloc( bnnz*sizeof(int) );
10: value = (LIS_SCALAR *)malloc( bnr*bnc*bnnz*sizeof(LIS_SCALAR) );
11: lis_matrix_create(MPI_COMM_WORLD,&A);
12: lis_matrix_set_size(A,n,0);
13: if( my_rank==0 ) {
14:
        bptr[0] = 0; bptr[1] = 2;
15:
        bindex[0] = 0; bindex[1] = 1;
        value[0] = 11; value[1] = 21; value[2] = 0; value[3] = 22;
16:
        value[4] = 0; value[5] = 41; value[6] = 32; value[7] = 0;}
17:
18: else {
        bptr[0] = 0; bptr[1] = 1;
19:
20:
        bindex[0] = 1;
        value[0] = 33; value[1] = 43; value[2] = 0; value[3] = 44;}
21:
22: lis_matrix_set_bsc(bnr,bnc,bnnz,bptr,bindex,value,A);
23: lis_matrix_assemble(A);
```

#### 5.8.3 Associating Arrays

To associate the arrays required by the BSC format with the matrix A, the following functions are used:

- C int lis\_matrix\_set\_bsc(int bnr, int bnc, int bnnz, int bptr[], int bindex[], LIS\_SCALAR value[], LIS\_MATRIX A)
- Fortran subroutine lis\_matrix\_set\_bsc(integer bnr, integer bnc, integer bnnz, integer bptr(), integer bindex(), LIS\_SCALAR value(), LIS\_MATRIX A, integer ierr)

## 5.9 Variable Block Row (VBR)

VBR is the generalized version of BSR. The division points of the rows and columns are given by the arrays row and col. VBR stores the nonzero blocks (the blocks in which at least one nonzero block exists) in the same step as that for the CRS format. Assume that nr and nc are the numbers of row and column divisions, respectively, and that nnzb denotes the number of the nonzero blocks of A, and nnz denotes the total number of the elements of the nonzero blocks. VBR uses six arrays bptr, bindex, row, col, ptr and value to store matrices.

- row is an integer array with a length of nr + 1, which stores the starting row number of the block rows.
- $\bullet$  col is an integer array with a length of nc+1, which stores the starting column number of the block columns.
- bindex is an integer array with a length of nnzb, which stores the block column numbers of the nonzero blocks.
- bptr is an integer array with a length of nr + 1, which stores the starting points of the block rows in the array bindex.
- value is a double precision array with a length of nnz, which stores all the elements of the nonzero blocks.
- ptr is an integer array with a length of nnzb + 1, which stores the starting points of the nonzero blocks in the array value.

## 5.9.1 Creating Matrices (for Serial and OpenMP Versions)

The right diagram in Figure 20 shows how the matrix A in Figure 20 is stored in the VBR format. A program to create the matrix in the VBR format is as follows:

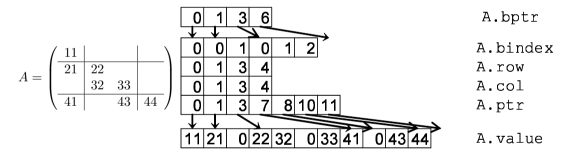


Figure 20: The data structure of the VBR format (for serial and OpenMP versions).

```
for serial and OpenMP versions
 1: int
                 n,nnz,nr,nc,bnnz;
 2: int
                  *row,*col,*ptr,*bptr,*bindex;
 3: LIS_SCALAR
                  *value;
 4: LIS_MATRIX
                 A;
5: n = 4; nnz = 11; bnnz = 6; nr = 3; nc = 3;
          = (int *)malloc( (nr+1)*sizeof(int) );
 6: bptr
          = (int *)malloc( (nr+1)*sizeof(int) );
7: row
8: col
          = (int *)malloc( (nc+1)*sizeof(int) );
9: ptr
          = (int *)malloc( (bnnz+1)*sizeof(int) );
10: bindex = (int *)malloc( bnnz*sizeof(int) );
11: value = (LIS_SCALAR *)malloc( nnz*sizeof(LIS_SCALAR) );
12: lis_matrix_create(0,&A);
13: lis_matrix_set_size(A,0,n);
15: bptr[0] = 0; bptr[1] = 1; bptr[2] = 3; bptr[3] = 6;
16: row[0] = 0; row[1] = 1; row[2] = 3; row[3] = 4;
17: col[0] = 0; col[1] = 1; col[2] = 3; col[3] = 4;
18: bindex[0] = 0; bindex[1] = 0; bindex[2] = 1; bindex[3] = 0;
19: bindex[4] = 1; bindex[5] = 2;
             = 0; ptr[1]
                                             = 3; ptr[3]
20: ptr[0]
                             = 1; ptr[2]
                                                             = 7;
21: ptr[4]
              = 8; ptr[5]
                              = 10; ptr[6]
                                             = 11;
22: value[0] = 11; value[1] = 21; value[2]
                                            = 0; value[3]
23: value[4] = 32; value[5] = 0; value[6] = 33; value[7]
24: value[8] = 0; value[9] = 43; value[10] = 44;
25:
26:
    lis_matrix_set_vbr(nnz,nr,nc,bnnz,row,col,ptr,bptr,bindex,value,A);
    lis_matrix_assemble(A);
```

## 5.9.2 Creating Matrices (for MPI Version)

Figure 21 shows how the matrix A in Figure 20 is stored in the VBR format on two processing elements. A program to create the matrix in the VBR format on two processing elements is as follows:

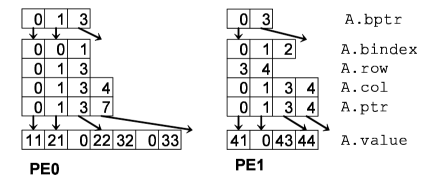


Figure 21: The data structure of the VBR format (for MPI version).

```
for MPI version -
 1: int
                 n,nnz,nr,nc,bnnz,my_rank;
 2: int
                 *row,*col,*ptr,*bptr,*bindex;
 3: LIS_SCALAR
                 *value:
 4: LIS_MATRIX
                 A;
5: MPI_Comm_rank(MPI_COMM_WORLD,&my_rank);
 6: if( my_rank==0 ) {n = 2; nnz = 7; bnnz = 3; nr = 2; nc = 3;}
                   {n = 2; nnz = 4; bnnz = 3; nr = 1; nc = 3;}
7: else
 8: bptr
          = (int *)malloc( (nr+1)*sizeof(int) );
9: row
          = (int *)malloc( (nr+1)*sizeof(int) );
          = (int *)malloc( (nc+1)*sizeof(int) );
          = (int *)malloc( (bnnz+1)*sizeof(int) );
12: bindex = (int *)malloc( bnnz*sizeof(int) );
13: value = (LIS_SCALAR *)malloc( nnz*sizeof(LIS_SCALAR) );
14: lis_matrix_create(MPI_COMM_WORLD,&A);
15: lis_matrix_set_size(A,n,0);
16: if( my_rank==0 ) {
       bptr[0] = 0; bptr[1] = 1; bptr[2] = 3;
17:
18:
       row[0] = 0; row[1] = 1; row[2] = 3;
       col[0] = 0; col[1] = 1; col[2] = 3; col[3] = 4;
19:
       bindex[0] = 0; bindex[1] = 0; bindex[2] = 1;
20:
               = 0; ptr[1]
                              = 1; ptr[2] = 3; ptr[3]
21:
       value[0] = 11; value[1] = 21; value[2] = 0; value[3] = 22;
22:
23:
       value[4] = 32; value[5] = 0; value[6] = 33;}
24: else {
25:
       bptr[0] = 0; bptr[1] = 3;
       row[0] = 3; row[1] = 4;
26:
27:
       col[0] = 0; col[1] = 1; col[2] = 3; col[3] = 4;
28:
       bindex[0] = 0; bindex[1] = 1; bindex[2] = 2;
               = 0; ptr[1] = 1; ptr[2]
29:
                                                = 3; ptr[3]
                                                                = 4;
       ptr[0]
       value[0] = 41; value[1] = 0; value[2] = 43; value[3] = 44;}
30:
    lis_matrix_set_vbr(nnz,nr,nc,bnnz,row,col,ptr,bptr,bindex,value,A);
31:
32: lis_matrix_assemble(A);
```

### 5.9.3 Associating Arrays

To associate the arrays required by the VBR format with the matrix A, the following functions are used:

- C int lis\_matrix\_set\_vbr(int nnz, int nr, int nc, int bnnz, int row[], int col[], int ptr[], int bptr[], int bindex[], LIS\_SCALAR value[], LIS\_MATRIX A)
- Fortran subroutine lis\_matrix\_set\_vbr(integer nnz, integer nr, integer nc, integer bnnz, integer row(), integer col(), integer ptr(), integer bptr(), integer bindex(), LIS\_SCALAR value(), LIS\_MATRIX A, integer ierr)

# 5.10 Coordinate (COO)

COO uses three arrays row, col and value to store data.

- value is a double precision array with a length of nnz, which stores the nonzero elements.
- row is an integer array with a length of nnz, which stores the row numbers of the nonzero elements.
- $\bullet$  col is an integer array with a length of nnz, which stores the column numbers of the nonzero elements.

## 5.10.1 Creating Matrices (for Serial and OpenMP Versions)

The right diagram in Figure 22 shows how the matrix A in Figure 22 is stored in the COO format. A program to create the matrix in the COO format is as follows:

Figure 22: The data structure of the COO format (for serial and OpenMP versions).

```
for serial and OpenMP versions –
 1: int
                  n,nnz;
 2: int
                  *row,*col;
3: LIS_SCALAR
                  *value:
 4: LIS_MATRIX
                  A;
5: n = 4; nnz = 8;
6: row = (int *)malloc( nnz*sizeof(int) );
7: col = (int *)malloc( nnz*sizeof(int) );
8: value = (LIS_SCALAR *)malloc( nnz*sizeof(LIS_SCALAR) );
9: lis_matrix_create(0,&A);
10: lis_matrix_set_size(A,0,n);
12: row[0] = 0; row[1] = 1; row[2] = 3; row[3] = 1;
13: row[4] = 2; row[5] = 2; row[6] = 3; row[7] = 3;
14: col[0] = 0; col[1] = 0; col[2] = 0; col[3] = 1;
15: col[4] = 1; col[5] = 2; col[6] = 2; col[7] = 3;
16: value[0] = 11; value[1] = 21; value[2] = 41; value[3] = 22;
17: value[4] = 32; value[5] = 33; value[6] = 43; value[7] = 44;
18:
19:
    lis_matrix_set_coo(nnz,row,col,value,A);
    lis_matrix_assemble(A);
```

## 5.10.2 Creating Matrices (for MPI Version)

Figure 23 shows how the matrix A in Figure 22 is stored in the COO format on two processing elements. A program to create the matrix in the COO format on two processing elements is as follows:

0	1	1	3	2	2	3	3	A.row
0	0	1	0	1	2	2	3	A.col
11	21	22	41	32	33	43	44	A.value
PE	Ξ0		PI	≣1				

Figure 23: The data structure of the COO format (for MPI version).

```
for MPI version -
 1: int
                  n,nnz,my_rank;
 2: int
                  *row,*col;
 3: LIS_SCALAR
                  *value;
 4: LIS_MATRIX
                  Α;
5: MPI_Comm_rank(MPI_COMM_WORLD,&my_rank);
 6: if( my_rank==0 ) {n = 2; nnz = 3;}
                     {n = 2; nnz = 5;}
7: else
8: row
          = (int *)malloc( nnz*sizeof(int) );
         = (int *)malloc( nnz*sizeof(int) );
9: col
10: value = (LIS_SCALAR *)malloc( nnz*sizeof(LIS_SCALAR) );
11: lis_matrix_create(MPI_COMM_WORLD,&A);
12: lis_matrix_set_size(A,n,0);
13: if( my_rank==0 ) {
        row[0] = 0; row[1] = 1; row[2] = 1;
15:
        col[0] = 0; col[1] = 0; col[2] = 1;
16:
        value[0] = 11; value[1] = 21; value[2] = 22;}
17: else {
        row[0] = 3; row[1] = 2; row[2] = 2; row[3] = 3; row[4] = 3;
18:
        col[0] = 0; col[1] = 1; col[2] = 2; col[3] = 2; col[4] = 3;
19:
        value[0] = 41; value[1] = 32; value[2] = 33; value[3] = 43; value[4] = 44;}
20.
21: lis_matrix_set_coo(nnz,row,col,value,A);
22: lis_matrix_assemble(A);
```

## 5.10.3 Associating Arrays

To associate the arrays required by the COO format with the matrix A, the following functions are used:

- C int lis\_matrix\_set\_coo(int nnz, int row[], int col[], LIS\_SCALAR value[], LIS\_MATRIX A)
- Fortran subroutine lis\_matrix\_set\_coo(integer nnz, integer row(), integer col(), LIS\_SCALAR value(), LIS\_MATRIX A, integer ierr)

## 5.11 Dense (DNS)

DNS uses one array value to store data.

• value is a double precision array with a length of  $n \times n$ , which stores the elements with priority given to the columns.

## 5.11.1 Creating Matrices (for Serial and OpenMP Versions)

The right diagram in Figure 24 shows how the matrix A in Figure 24 is stored in the DNS format. A program to create the matrix in the DNS format is as follows:

Figure 24: The data structure of the DNS format (for serial and OpenMP versions).

```
for serial and OpenMP versions —
 1: int
2: LIS_SCALAR
                  *value;
3: LIS_MATRIX
                  A;
4: n = 4;
5: value = (LIS_SCALAR *)malloc( n*n*sizeof(LIS_SCALAR) );
6: lis_matrix_create(0,&A);
7: lis_matrix_set_size(A,0,n);
9: value[0] = 11; value[1] = 21; value[2] = 0; value[3] = 41;
10: value[4] = 0; value[5] = 22; value[6] = 32; value[7] = 0;
11: value[8] = 0; value[9] = 0; value[10] = 33; value[11] = 43;
12: value[12] = 0; value[13] = 0; value[14] = 0; value[15] = 44;
13:
14: lis_matrix_set_dns(value,A);
15: lis_matrix_assemble(A);
```

## 5.11.2 Creating Matrices (for MPI Version)

Figure 25 shows how the matrix A in Figure 24 is stored in the DNS format on two processing elements. A program to create the matrix in the DNS format on two processing elements is as follows:

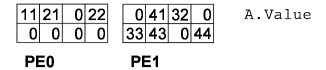


Figure 25: The data structure of the DNS format (for MPI version).

```
for MPI version -
 1: int
                  n,my_rank;
                  *value;
 2: LIS_SCALAR
3: LIS_MATRIX
                  A;
 4: MPI_Comm_rank(MPI_COMM_WORLD,&my_rank);
 5: if( my_rank==0 ) {n = 2;}
 6: else
                     {n = 2;}
7: value = (LIS_SCALAR *)malloc( n*n*sizeof(LIS_SCALAR) );
8: lis_matrix_create(MPI_COMM_WORLD,&A);
9: lis_matrix_set_size(A,n,0);
10: if( my_rank==0 ) {
11:
        value[0] = 11; value[1] = 21; value[2] = 0; value[3] = 22;
        value[4] = 0; value[5] = 0; value[6] = 0; value[7] = 0;}
12:
13: else {
14:
        value[0] = 0; value[1] = 41; value[2] = 32; value[3] = 0;
        value[4] = 33; value[5] = 43; value[6] = 0; value[7] = 44;}
15:
16:
    lis_matrix_set_dns(value,A);
    lis_matrix_assemble(A);
17:
```

#### 5.11.3 Associating Arrays

To associate the arrays required by the DNS format with the matrix A, the following functions are used:

- C int lis\_matrix\_set\_dns(LIS\_SCALAR value[], LIS\_MATRIX A)
- Fortran subroutine lis\_matrix\_set\_dns(LIS\_SCALAR value(), LIS\_MATRIX A, integer ierr)

# 6 Functions

This section describes the functions which can be employed by users. The return values of the functions in C and the values of ierr in Fortran are as follows:

### Return Values

LIS\_SUCCESS(0) Normal termination

LIS\_ILL\_OPTION(1) Illegal option
LIS\_BREAKDOWN(2) Breakdown

LIS\_OUT\_OF\_MEMORY(3) Insufficient working memory

LIS\_MAXITER(4) Did not converge within the maximum number of iterations

LIS\_NOT\_IMPLEMENTED(5) Not implemented

LIS\_ERR\_FILE\_IO(6) File I/O error

## 6.1 Operating Vector Elements

Assume that the size of the vector v is  $global\_n$  and that the size of the partial vectors stored on nprocs processing elements is  $local\_n$ .  $global\_n$  and  $local\_n$  are called the global size and the local size, respectively.

## 6.1.1 lis\_vector\_create

C int lis\_vector\_create(LIS\_Comm comm, LIS\_VECTOR \*v)
Fortran subroutine lis\_vector\_create(LIS\_Comm comm, LIS\_VECTOR v, integer ierr)

## Description

Create vector v

# Input

LIS\_Comm MPI communicator

### Output

v Vector

ierr Return code

### Note

For the serial and OpenMP versions, the value for **comm** is ignored.

## 6.1.2 lis\_vector\_destroy

```
C int lis_vector_destroy(LIS_VECTOR v)
Fortran subroutine lis_vector_destroy(LIS_VECTOR v, integer ierr)
```

## Description

Destroy vector v

## Input

v Vector to be destroyed

Output

ierr Return code

## 6.1.3 lis\_vector\_duplicate

```
C int lis_vector_duplicate(void *vin, LIS_VECTOR *vout)
Fortran subroutine lis_vector_duplicate(LIS_VECTOR vin, LIS_VECTOR vout, integer ierr)
```

## Description

Create vector  $v_{out}$  which has the same information as  $v_{in}$ 

#### Input

vin Source vector

Output

vout Destination vector

ierr Return code

## Note

The function lis\_vector\_duplicate does not copy the values, but only reserves an area. To copy the values as well, the function lis\_vector\_copy must be used after this function.

#### 6.1.4 lis\_vector\_set\_size

## Description

Assign size of vector v

## Input

v Vector

local\_n Size of partial vector

global\_n Size of global vector

Output

ierr Return code

#### Note

Either  $local\_n$  or  $global\_n$  must be provided. This function can create a vector in one of the following ways: Creates partial vectors of size  $local\_n$  if  $local\_n$  is given, or creates partial vectors stored on a given number of processing elements, if  $global\_n$  is given.

In the case of the serial and OpenMP versions,  $local\_n = global\_n$ . It means that both  $lis\_vector\_set\_size(v,n,0)$  and  $lis\_vector\_set\_size(v,0,n)$  create a vector of size n.

## 6.1.5 lis\_vector\_get\_size

```
C int lis_vector_get_size(LIS_VECTOR v, int *local_n, int *global_n)
Fortran subroutine lis_vector_get_size(LIS_VECTOR v, integer local_n,
    integer global_n, integer ierr)
```

## Description

Get size of vector v

## Input

v Vector

## Output

local\_n Size of partial vector

global\_n Size of global vector

ierr Return code

## Note

In the case of the serial and OpenMP versions,  $local_n = global_n$ .

## 6.1.6 lis\_vector\_get\_range

```
C int lis_vector_get_range(LIS_VECTOR v, int *is, int *ie)
Fortran subroutine lis_vector_get_range(LIS_VECTOR v, integer is, integer ie, integer ierr)
```

### Description

Get location of partial vector v in global vector

## Input

v Partial vector

## Output

is Location where partial vector  $\boldsymbol{v}$  starts in global vector

ie 1+ location where partial vector v ends in global vector

ierr Return code

#### Note

For the serial and OpenMP versions, a vector of size n results in is = 0 and ie = n.

#### 6.1.7 lis\_vector\_set\_value

#### Description

Assign scalar value to i-th row of vector v

## Input

flag LIS\_INS\_VALUE : v[i] = value

LIS\_ADD\_VALUE : v[i] = v[i] + value

i Location where value is assigned

value Scalar value to be assigned

v Destination vector

## Output

v Vector with scalar value assigned to i-th row

ierr Return code

#### Note

For the MPI version, the i-th row of the global vector must be specified instead of the i-th row of the partial vector.

## 6.1.8 lis\_vector\_get\_value

```
C int lis_vector_get_value(LIS_VECTOR v, int i, LIS_SCALAR *value)
Fortran subroutine lis_vector_get_value(LIS_VECTOR v, integer i, LIS_SCALAR value, integer ierr)
```

### Description

Get value of *i*-th row of vector v

### Input

i Location where value should be assigned

v Destination vector

Output

value Value of *i*-th row

ierr Return code

#### Note

For the MPI version, the i-th row of the global vector must be specified instead of the i-th row of the partial vector.

#### 6.1.9 lis\_vector\_set\_values

## Description

Assign scalar values value[i] to the index[i]-th row of vector v

## Input

flag LIS\_INS\_VALUE : v[index[i]] = value[i]

LIS\_ADD\_VALUE : v[index[i]] = v[index[i]] + value[i]

Number of elements of array which stores scalar values to be as-

signed

index Array which stores location where scalar values should be assigned

value Array which stores scalar values to be assigned

v Destination vector

Output

v Vector with scalar value[i] assigned to its index[i]-th row

ierr Return code

### Note

For the MPI version, the index[i]-th row of the global vector must be specified instead of the index[i]-th row of the partial vector.

## 6.1.10 lis\_vector\_get\_values

```
C int lis_vector_get_values(LIS_VECTOR v, int start, int count, LIS_SCALAR value[])
Fortran subroutine lis_vector_get_values(LIS_VECTOR v, integer start, integer count, LIS_SCALAR value(), integer ierr)
```

## Description

Get scalar values of start + i-th row of vector v, where i = 0, 1, ..., count - 1

## Input

start Starting location

count Number of values to get

v Destination vector

Output

value Vector to store scalar values

ierr Return code

## Note

For the MPI version, the start+i-th row of the global vector must be specified instead of the start+i-th row of the partial vector.

#### 6.1.11 lis\_vector\_scatter

C int lis\_vector\_scatter(LIS\_SCALAR value[], LIS\_VECTOR v)
Fortran subroutine lis\_vector\_scatter(LIS\_SCALAR value(), LIS\_VECTOR v, integer ierr)

## Description

Assign scalar values of i-th row of vector v, where  $i = 0, 1, ..., global_n - 1$ 

#### Input

value Array which stores scalar values to be assigned

Output

v Destination vector

ierr Return code

Note

## 6.1.12 lis\_vector\_gather

C int lis\_vector\_gather(LIS\_VECTOR v, LIS\_SCALAR value[])
Fortran subroutine lis\_vector\_gather(LIS\_VECTOR v, LIS\_SCALAR value(), integer ierr)

## Description

Get scalar values of *i*-th row of vector v, where  $i = 0, 1, ..., global_n - 1$ 

## Input

v Source vector

Output

value Vector to store scalar values

ierr Return code

Note

## 6.1.13 lis\_vector\_copy

C int lis\_vector\_copy(LIS\_VECTOR x, LIS\_VECTOR y)
Fortran subroutine lis\_vector\_copy(LIS\_VECTOR x, LIS\_VECTOR y, integer ierr)

## Description

Copy vector x

## Input

x Source vector

Output

y Destination vector

ierr Return code

## 6.1.14 lis\_vector\_set\_all

C int lis\_vector\_set\_all(LIS\_SCALAR value, LIS\_VECTOR x)
Fortran subroutine lis\_vector\_set\_all(LIS\_SCALAR value, LIS\_VECTOR x, integer ierr)

## Description

Assign scalar value to all elements of vector v

## Input

value Scalar value to be assigned

v Destination vector

Output

v Vector with value assigned to all elements

# 6.2 Operating Matrix Elements

Assume that the size of the matrix A is  $global\_n \times global\_n$  and that the size of each partial matrix stored on nprocs processing elements is  $local\_n \times global\_n$ . Here,  $global\_n$  and  $local\_n$  are called the number of the rows of the global matrix and the number of the rows of the partial matrix, respectively.

#### 6.2.1 lis\_matrix\_create

C int lis\_matrix\_create(LIS\_Comm comm, LIS\_MATRIX \*A)
Fortran subroutine lis\_matrix\_create(LIS\_Comm comm, LIS\_MATRIX A, integer ierr)

## Description

Create matrix A

### Input

LIS\_Comm MPI communicator

## Output

A Matrix

ierr Return code

#### Note

For the sequential and the OpenMP versions, the value for comm is ignored.

# 6.2.2 lis\_matrix\_destroy

C int lis\_matrix\_destroy(LIS\_MATRIX A)
Fortran subroutine lis\_matrix\_destroy(LIS\_MATRIX A, integer ierr)

# Description

Destroy matrix A

## Input

A Matrix to be destroyed

## Output

## 6.2.3 lis\_matrix\_duplicate

#### Description

Create matrix  $A_{out}$  which has the same information as the original  $A_{in}$ 

Input

Ain Source matrix

Output

Aout Destination matrix

ierr Return code

#### Note

The function <code>lis\_matrix\_duplicate</code> does not copy the values of the elements of the matrix, but only reserves an area. To copy the values of the elements as well, the function <code>lis\_matrix\_copy</code> must be used.

#### 6.2.4 lis\_matrix\_malloc

## Description

Allocate memory for matrix A

## Input

A Matrix

nnz\_row Average number of nonzero elements

nnz Array of numbers of nonzero elements in each row

Output

ierr Return code

#### Note

Either nnz\_row or nnz must be provided.

#### 6.2.5 lis\_matrix\_set\_value

#### Description

Assign value to (i, j) element of matrix A

#### Input

flag LIS\_INS\_VALUE : A(i, j) = value

LIS\_ADD\_VALUE : A(i,j) = A(i,j) + value

i Row number of matrix

j Column number of matrix

value Value to be assigned

A Matrix

### Output

A Matrix

ierr Return code

#### Note

For the MPI version, the i-th row and the j-th column of the global matrix must be specified, instead of the i-th row and the j-th column of the partial matrix.

The function lis\_matrix\_set\_value stores the assigned value in a temporary internal format. For this reason, when lis\_matrix\_set\_value is used, the function lis\_matrix\_assemble must be called.

## 6.2.6 lis\_matrix\_assemble

```
C int lis_matrix_assemble(LIS_MATRIX A)
Fortran subroutine lis_matrix_assemble(LIS_MATRIX A, integer ierr)
```

## Description

Build matrix A in specified storage format

## Input

A Matrix

## Output

A Matrix built in specified storage format

#### 6.2.7 lis\_matrix\_set\_size

## Description

Assign size of matrix A

### Input

A Matrix

local\_n Number of rows of partial matrix
global\_n Number of rows of global matrix

Output

ierr Return code

#### Note

Either  $local\_n$  or  $global\_n$  must be provided. This function can create matrices in one of the following two ways: Create partial matrices of size  $local\_n \times N$  if  $local\_n$  is given, or create partial matrices of size  $global\_n \times global\_n$  is stored in a given number of processing elements, if  $global\_n$  is given. N represents the total sum of  $local\_n$ .

In case of the serial and OpenMP versions,  $local_n$  is equal to  $global_n$ . It means that both  $lis_matrix_set_size(A,n,0)$  and  $lis_matrix_set_size(A,0,n)$  create a matrix of  $n \times n$ .

For the MPI version,  $lis_matrix_set_size(A,n,0)$  creates on the processing element p a partial matrix of size  $n_p \times N$ , where N is the total sum of  $n_p$ . On the other hand,  $lis_matrix_set_size(A,0,n)$  creates on the processing element p a partial matrix of size  $m_p \times n$ , where  $m_p$  is the number of the partial matrix, which is determined by the library.

#### 6.2.8 lis\_matrix\_get\_size

#### Description

Get size of matrix A

## Input

A Matrix

#### Output

local\_n Number of rows of partial matrix

global\_n Number of rows of global matrix

ierr Return code

#### Note

In case of the serial and OpenMP versions,  $local_n = global_n$ .

# ${\bf 6.2.9 \quad lis\_matrix\_get\_range}$

```
C int lis_matrix_get_range(LIS_MATRIX A, int *is, int *ie)
Fortran subroutine lis_matrix_get_range(LIS_MATRIX A, integer is, integer ie,
    integer ierr)
```

## Description

Get location of partial matrix A in global matrix

# Input

A Partial matrix

# Output

is Location where partial matrix A starts in global matrix

ie 1+ location where partial matrix A ends in global matrix

ierr Return code

## Note

For the serial and OpenMP versions, a matrix of  $n \times n$  results in is = 0 and ie = n.

## 6.2.10 lis\_matrix\_set\_type

C int lis\_matrix\_set\_type(LIS\_MATRIX A, int matrix\_type)
Fortran subroutine lis\_matrix\_set\_type(LIS\_MATRIX A, int matrix\_type, integer ierr)

## Description

Assign storage format

## Input

A Matrix

matrix\_type Storage format

Output

ierr Return code

#### Note

 $\mathtt{matrix\_type}$  of A is LIS\_MATRIX\_CRS when the matrix is created. The table below shows the available storage formats for  $\mathtt{matrix\_type}$ .

storage format		matrix_type
Compressed Row Storage	(CRS)	LIS_MATRIX_CRS
Compressed Column Storage	(CCS)	LIS_MATRIX_CCS
Modified Compressed Sparse Row	(MSR)	LIS_MATRIX_MSR
Diagonal	(DIA)	LIS_MATRIX_DIA
Ellpack-Itpack generalized diagonal	(ELL)	LIS_MATRIX_ELL
Jagged Diagonal	(JDS)	LIS_MATRIX_JDS
Block Sparse Row	(BSR)	LIS_MATRIX_BSR
Block Sparse Column	(BSC)	LIS_MATRIX_BSC
Variable Block Row	(VBR)	LIS_MATRIX_VBR
Dense	(DNS)	LIS_MATRIX_DNS
Coordinate	(COO)	LIS_MATRIX_COO

## 6.2.11 lis\_matrix\_get\_type

## Description

Get storage format

### Input

A Matrix

## Output

matrix\_type Storage format ierr Return code

#### 6.2.12 lis\_matrix\_set\_blocksize

C int lis\_matrix\_set\_blocksize(LIS\_MATRIX A, int bnr, int bnc, int row[],
 int col[])

Fortran subroutine lis\_matrix\_set\_blocksize(LIS\_MATRIX A, integer bnr, integer bnc, integer row[], integer col[], integer ierr)

### Description

Assign block size for BSR, BSC, and VBR

### Input

A Matrix

bnr Row block size for BSR (BSC) or number of row blocks for VBR

bnc Column block size for BSR (BSC) or number of column blocks for

VBR

row Array of row division information about VBR

col Array of column division information about VBR

Output

ierr Return code

### 6.2.13 lis\_matrix\_convert

C int lis\_matrix\_convert(LIS\_MATRIX Ain, LIS\_MATRIX Aout)
Fortran subroutine lis\_matrix\_convert(LIS\_MATRIX Ain, LIS\_MATRIX Aout, integer ierr)

## Description

Convert matrix  $A_{in}$  into  $A_{out}$  of type specified with lis\_matrix\_set\_type

## Input

Ain Source matrix

Output

Aout Matrix converted into specified type

ierr Return code

## Note

The specification of the converted storage format is set to Aout by using lis\_matrix\_set\_type. The specification of the block size of BSR, BSC, and VBR is set to Aout by using lis\_matrix\_set\_blocksize.

In converting the storage type of the source matrix into a specified type, the conversions indicated by one in the table below are performed directly, and the other conversions are made via the indicated types. The conversions with no indication are made via the CRS type.

Src \Dst	CRS	CCS	MSR	DIA	ELL	JDS	BSR	BSC	VBR	DNS	COO
CRS	1	1	1	1	1	1	1	CCS	1	1	1
COO	1	1	1	CRS	CRS	CRS	CRS	CCS	CRS	CRS	1

## 6.2.14 lis\_matrix\_copy

C int lis\_matrix\_copy(LIS\_MATRIX Ain, LIS\_MATRIX Aout)
Fortran subroutine lis\_matrix\_copy(LIS\_MATRIX Ain, LIS\_MATRIX Aout, integer ierr)

## Description

Copy values of matrix elements

Input

Ain Source matrix

Output

Aout Destination matrix

ierr Return code

## 6.2.15 lis\_matrix\_get\_diagonal

C int lis\_matrix\_get\_diagonal(LIS\_MATRIX A, LIS\_VECTOR d)
Fortran subroutine lis\_matrix\_get\_diagonal(LIS\_MATRIX A, LIS\_VECTOR d, integer ierr)

## Description

Store diagonal elements of matrix A to vector d

Input

A Matrix

Output

d Vector which stores diagonal elements of matrix

#### 6.2.16 lis\_matrix\_set\_crs

### Description

Set matrix A in CRS format

### Input

nnz Number of nonzero elements

ptr, index, value Arrays for CRS format

A Matrix

## Output

A Matrix and associated arrays

#### Note

When lis\_matrix\_set\_crs is used, the function lis\_matrix\_assemble must be called.

#### 6.2.17 lis\_matrix\_set\_ccs

```
C int lis_matrix_set_ccs(int nnz, int ptr[], int index[], LIS_SCALAR value[], LIS_MATRIX A)

Fortran subroutine lis_matrix_set_ccs(integer nnz, integer row(), integer index(), LIS_SCALAR value(), LIS_MATRIX A, integer ierr)
```

### Description

Set matrix A in CCS format

## Input

nnz Number of nonzero elements

ptr, index, value Arrays for CCS format

A Matrix

## Output

A Matrix and associated arrays

### Note

When lis\_matrix\_set\_ccs is used, the function lis\_matrix\_assemble must be called.

#### 6.2.18 lis\_matrix\_set\_msr

### Description

Set matrix A in MSR format

### Input

nnz Number of nonzero elements

ndz Number of nonzero elements in diagonal

index, value Arrays for MSR format

A Matrix

### Output

A Matrix and associated arrays

#### Note

When lis\_matrix\_set\_msr is used, the function lis\_matrix\_assemble must be called.

### 6.2.19 lis\_matrix\_set\_dia

### Description

Set matrix A in DIA format

## Input

nnd Number of nonzero diagonal elements

index, value Arrays for DIA format

A Matrix

### Output

A Matrix and associated arrays

#### Note

When lis\_matrix\_set\_dia is used, the function lis\_matrix\_assemble must be called.

#### 6.2.20 lis\_matrix\_set\_ell

### Description

Set matrix A in ELL format

### Input

maxnzr Maximum number of nonzero elements in each row

index, value Arrays for ELL format

A Matrix

#### Output

A Matrix and associated arrays

#### Note

When lis\_matrix\_set\_ell is used, the function lis\_matrix\_assemble must be called.

## 6.2.21 lis\_matrix\_set\_jds

### Description

Set matrix A in JDS format

## Input

nnz Number of nonzero elements

maxnzr Maximum number of nonzero elements in each row

perm, ptr, index, value Arrays for JDS format

A Matrix

## Output

A Matrix and associated arrays

## Note

When lis\_matrix\_set\_jds is used, the function lis\_matrix\_assemble must be called.

#### 6.2.22 lis\_matrix\_set\_bsr

C int lis\_matrix\_set\_bsr(int bnr, int bnc, int bnnz, int bptr[], int bindex[], LIS\_SCALAR value[], LIS\_MATRIX A)

Fortran subroutine lis\_matrix\_set\_bsr(integer bnr, integer bnc, integer bnnz, integer bptr(), integer bindex(), LIS\_SCALAR value(), LIS\_MATRIX A, integer ierr)

### Description

Set matrix A in BSR format

#### Input

bnr Row block size

bnc Column block size

bnnz Number of nonzero blocks

bptr, bindex, value Arrays for BSR format

A Matrix

## Output

A Matrix and associated arrays

#### Note

When lis\_matrix\_set\_bsr is used, the function lis\_matrix\_assemble must be called.

#### 6.2.23 lis\_matrix\_set\_bsc

## Description

Set matrix A in BSC format

integer ierr)

#### Input

bnr Row block size

bnc Column block size

bnnz Number of nonzero blocks

bptr, bindex, value Arrays for BSC format

A Matrix

### Output

A Matrix and associated arrays

#### Note

When lis\_matrix\_set\_bsc is used, the function lis\_matrix\_assemble must be called.

#### 6.2.24 lis\_matrix\_set\_vbr

```
C int lis_matrix_set_vbr(int nnz, int nr, int nc, int bnnz, int row[], int col[], int ptr[], int bptr[], int bindex[], LIS_SCALAR value[], LIS_MATRIX A)

Fortran subroutine lis_matrix_set_vbr(integer nnz, integer nr, integer nc, integer bnnz, integer row(), integer col(), integer ptr(), integer bptr(), integer bindex(), LIS_SCALAR value(), LIS_MATRIX A, integer ierr)
```

#### Description

Set matrix A in VBR format

#### Input

nnz Number of all nonzero elements

nr Number of row blocks

nc Number of column blocks

bnnz Number of nonzero blocks

row, col, ptr, bptr, bindex, value Arrays for VBR format

A Matrix

### Output

A Matrix and associated arrays

#### Note

When lis\_matrix\_set\_vbr is used, the function lis\_matrix\_assemble must be called.

## 6.2.25 lis\_matrix\_set\_coo

#### Description

Set matrix A in COO format

### Input

nnz Number of nonzero elements

row, col, value Arrays for COO format

A Matrix

### Output

A Matrix and associated arrays

#### Note

When lis\_matrix\_set\_coo is used, the function lis\_matrix\_assemble must be called.

## 6.2.26 lis\_matrix\_set\_dns

C int lis\_matrix\_set\_dns(LIS\_SCALAR value[], LIS\_MATRIX A)
Fortran subroutine lis\_matrix\_set\_dns(LIS\_SCALAR value(), LIS\_MATRIX A, integer ierr)

# Description

Set matrix A in DNS format

## Input

value Array for DNS format

A Matrix

Output

A Matrix and associated arrays

# Note

When lis\_matrix\_set\_dns is used, the function lis\_matrix\_assemble must be called.

# 6.3 Operating Vectors and Matrices

## 6.3.1 lis\_vector\_scale

C int lis\_vector\_scale(LIS\_SCALAR alpha, LIS\_VECTOR x)
Fortran subroutine lis\_vector\_scale(LIS\_SCALAR alpha, LIS\_VECTOR x, integer ierr)

## Description

Multiply vector x by  $\alpha$ 

## Input

alpha Scalar value  $\alpha$ 

x Vector to be multiplied

Output

**x** Vector multiplied by  $\alpha$ 

ierr Return code

#### 6.3.2 lis\_vector\_dot

## Description

Calculate inner product  $x^T y$ 

## Input

x Vector

y Vector

Output

val Inner product value

## 6.3.3 lis\_vector\_nrm1

C int lis\_vector\_nrm1(LIS\_VECTOR x, LIS\_SCALAR \*val)
Fortran subroutine lis\_vector\_nrm1(LIS\_VECTOR x, LIS\_SCALAR val, integer ierr)

# Description

Calculate 1-norm of vector x

### Input

x Vector

## Output

val 1-norm of vector

ierr Return code

## 6.3.4 lis\_vector\_nrm2

C int lis\_vector\_nrm2(LIS\_VECTOR x, LIS\_SCALAR \*val)
Fortran subroutine lis\_vector\_nrm2(LIS\_VECTOR x, LIS\_SCALAR val, integer ierr)

## Description

Calculate 2-norm of vector x

## Input

x Vector

# Output

val 2-norm of vector

## 6.3.5 lis\_vector\_nrmi

C int lis\_vector\_nrmi(LIS\_VECTOR x, LIS\_SCALAR \*val)
Fortran subroutine lis\_vector\_nrmi(LIS\_VECTOR x, LIS\_SCALAR val, integer ierr)

# Description

Calculate infinity norm of vector x

Input

x Vector

Output

val infinity norm of vector

## 6.3.6 lis\_vector\_axpy

C int lis\_vector\_axpy(LIS\_SCALAR alpha, LIS\_VECTOR x, LIS\_VECTOR y)
Fortran subroutine lis\_vector\_axpy(LIS\_SCALAR alpha, LIS\_VECTOR x, LIS\_VECTOR y, integer ierr)

## Description

Calculate  $y = \alpha x + y$ 

## Input

alpha Scalar value

x, y Vectors

## Output

y  $\alpha x + y$  (vector y is overwritten)

ierr Return code

# 6.3.7 lis\_vector\_xpay

C int lis\_vector\_xpay(LIS\_VECTOR x, LIS\_SCALAR alpha, LIS\_VECTOR y)
Fortran subroutine lis\_vector\_xpay(LIS\_VECTOR x, LIS\_SCALAR alpha, LIS\_VECTOR y, integer ierr)

## Description

Calculate  $y = x + \alpha y$ 

## Input

alpha Scalar value

x, y Vectors

## Output

y  $x + \alpha y$  (vector y is overwritten)

## 6.3.8 lis\_vector\_axpyz

C int lis\_vector\_axpyz(LIS\_SCALAR alpha, LIS\_VECTOR x, LIS\_VECTOR y, LIS\_VECTOR z)

Fortran subroutine lis\_vector\_axpyz(LIS\_SCALAR alpha, LIS\_VECTOR x, LIS\_VECTOR y, LIS\_VECTOR z, integer ierr)

## Description

Calculate  $z = \alpha x + y$ 

## Input

alpha Scalar value

x, y Vectors

## Output

z  $x + \alpha y$ 

ierr Return code

## 6.3.9 lis\_matrix\_scaling

## Description

Scale matrix A

## Input

A Matrix

b Vector

action LIS\_SCALE\_JACOBI : Jacobi scaling  $D^{-1}Ax = D^{-1}b$ , where D rep-

resents the diagonal of  $A = (a_{ij})$ 

LIS\_SCALE\_SYMM\_DIAG : Diagonal scaling  $D^{-1/2}AD^{-1/2}x=D^{-1/2}b$ , where  $D^{-1/2}$  represents a diagonal matrix with  $1/\sqrt{a_{ii}}$  as di-

agonal

# Output

d Vector which stores diagonal elements of  $D^{-1}$  or  $D^{-1/2}$ 

## 6.3.10 lis\_matvec

C void lis\_matvec(LIS\_MATRIX A, LIS\_VECTOR x, LIS\_VECTOR y)
Fortran subroutine lis\_matvec(LIS\_MATRIX A, LIS\_VECTOR x, LIS\_VECTOR y)

# Description

Calculate matrix vector product y = Ax

## Input

A Matrix

x Vector

## Output

 $\mathbf{y}$  Ax

## 6.3.11 lis\_matvect

C void lis\_matvect(LIS\_MATRIX A, LIS\_VECTOR x, LIS\_VECTOR y)
Fortran subroutine lis\_matvect(LIS\_MATRIX A, LIS\_VECTOR x, LIS\_VECTOR y)

## Description

Calculate transposed matrix vector product  $y = A^T x$ 

# Input

A Matrix

x Vector

Output

 $\mathbf{y}$   $A^Tx$ 

# 6.4 Solving Systems of Linear Equations

## 6.4.1 lis\_solver\_create

C int lis\_solver\_create(LIS\_SOLVER \*solver)
Fortran subroutine lis\_solver\_create(LIS\_SOLVER solver, integer ierr)

## Description

Create solver

## Input

None

## Output

solver Solver

ierr Return code

## Note

solver has the information on the solver, the preconditioner, etc.

# 6.4.2 lis\_solver\_destroy

C int lis\_solver\_destroy(LIS\_SOLVER solver)
Fortran subroutine lis\_solver\_destroy(LIS\_SOLVER solver, integer ierr)

## Description

Destroy solver

## Input

solver to be destroyed

# Output

## 6.4.3 lis\_solver\_set\_option

## Description

Set options for solver

## Input

text Command line options

## Output

solver Solver

ierr Return code

## Note

The table below shows the available command line options, where  $-i \{cg|1\}$  means -i cg or -i 1 and -maxiter [1000] indicates that -maxiter defaults to 1,000.

Specifying Linear Solvers (Default: -i bicg)

Method	Option	Auxiliary Options	<u> </u>
CG	-i {cg 1}		
$\operatorname{BiCG}$	-i {bicg 2}		
CGS	-i {cgs 3}		
BiCGSTAB	-i {bicgstab 4}		
BiCGSTAB(l)	-i {bicgstabl 5}	-ell [2]	Degree $l$
GPBiCG	-i {gpbicg 6}		
TFQMR	-i {tfqmr 7}		
Orthomin(m)	-i {orthomin 8}	-restart [40]	Restart value $m$
GMRES(m)	-i {gmres 9}	-restart [40]	Restart value $m$
Jacobi	-i {jacobi 10}		
Gauss-Seidel	-i {gs 11}		
SOR	-i {sor 12}	-omega [1.9]	Relaxation coefficient $\omega$ (0 < $\omega$ < 2)
BiCGSafe	-i {bicgsafe 13}		
$\operatorname{CR}$	-i {cr 14}		
BiCR	-i {bicr 15}		
CRS	-i {crs 16}		
BiCRSTAB	-i {bicrstab 17}		
GPBiCR	-i {gpbicr 18}		
BiCRSafe	-i {bicrsafe 19}		_
FGMRES(m)	-i {fgmres 20}	-restart [40]	Restart value m
IDR(s)	-i {idrs 21}	-irestart [2]	Restart value $s$
MINRES	-i {minres 22}		

# Specifying Preconditioners (Default: -p none)

Preconditioner	Option	Auxiliary Options	
None	-p {none 0}		
Jacobi	-p {jacobi 1}		
ILU(k)	-p {ilu 2}	-ilu_fill [0]	Fill level $k$
SSOR	-p {ssor 3}	-ssor_w [1.0]	Relaxation coefficient $\omega$ (0 < $\omega$ < 2)
Hybrid	-p {hybrid 4}	-hybrid_i [sor]	Linear equations solver
		-hybrid_maxiter [25]	Maximum number of iterations
		-hybrid_tol [1.0e-3]	Convergence criterion
		-hybrid_w [1.5]	Relaxation coefficient $\omega$ for SOR
			$(0 < \omega < 2)$
		-hybrid_ell [2]	Degree $l$ of BiCGSTAB(1)
		-hybrid_restart [40]	Restart values for GMRES and Orthomin
I+S	-p {is 5}	-is_alpha [1.0]	Parameter $\alpha$ for preconditioner
			of $I + \alpha S^{(m)}$ type
		-is_m [3]	Parameter $m$ for preconditioner
			of $I + \alpha S^{(m)}$ type
SAINV	-p {sainv 6}	-sainv_drop [0.05]	Drop criterion
SA-AMG	-p {saamg 7}	-saamg_unsym [false]	Selects unsymmetric version
			(Matrix structure must be symmetric)
		-saamg_theta [0.05 0.12]	Drop criterion $a_{ij}^2 \leq \theta^2  a_{ii}   a_{jj} $
			(symmetric or unsymmetric)
Crout ILU	-p {iluc 8}	-iluc_drop [0.05]	Drop criterion
		-iluc_rate [5.0]	Ratio of maximum fill-in
ILUT	-p {ilut 9}	-ilut_drop [0.05]	Drop criterion
		-ilut_rate [5.0]	Ratio of maximum fill-in
Additive Schwarz	-adds true	-adds_iter [1]	Number of iterations

# Other Options

	0 01101	S P t l c l c c c c c c c c c c c c c c c c	
Option			
-maxiter [1000]	Maximum number of iterations		
-tol [1.0e-12]	Convergence criterion		
-print [0]	Display of the residual		
	-print {none 0}	None	
	-print {mem 1}	Saves the residual history in memory	
	-print {out 2}	Displays the residual history	
	-print {all 3}	Saves the residual history and displays it on the screen	
-scale [0]	Scaling		
	(The result will overwrite	te the original matrix and vectors)	
	-scale {none 0}	No scaling	
	-scale {jacobi 1}	Jacobi scaling $D^{-1}Ax = D^{-1}b$	
		(D represents the diagonal of $A = (a_{ij})$ )	
	-scale {symm_diag 2}	Diagonal scaling $D^{-1/2}AD^{-1/2}x = D^{-1/2}b$	
		$(D^{-1/2} \text{ represents a diagonal matrix with } 1/\sqrt{a_{ii}}$	
		as diagonal)	
-initx_zeros [true]	Behavior of the initial vector $x_0$		
	-initx_zeros {false	0} Given values	
	-initx_zeros {true 1	All elements are set to 0	
-omp_num_threads [t]	Number of threads		
	(t represents the maximum number of threads)		
-storage [0]	Matrix storage format		
-storage_block [2]	Block size of BSR and I	BSC	

# Precision (Default: -f double)

Precision	Option	Auxiliary Options
	-f {double 0}	<i>v</i> 1
QUAD	-f {quad 1}	

# 6.4.4 lis\_solver\_set\_optionC

```
C int lis_solver_set_optionC(LIS_SOLVER solver)
Fortran subroutine lis_solver_set_optionC(LIS_SOLVER solver, integer ierr)
```

# Description

Set options for solver on command line

## Input

None

# Output

solver Solver

## 6.4.5 lis\_solve

# Description

Solve system of linear equations Ax = b with specified solver

# Input

A Coefficient matrix

b Right hand side vector

x Initial vector

solver Solver

Output

x Solution

solver Number of iterations, execution time, etc.

ierr Return code (0)

## 6.4.6 lis\_solve\_kernel

```
C int lis_solve_kernel(LIS_MATRIX A, LIS_VECTOR b, LIS_VECTOR x,
LIS_SOLVER solver, LIS_PRECON, precon)

Fortran subroutine lis_solve_kernel(LIS_MATRIX A, LIS_VECTOR b, LIS_VECTOR x,
LIS_SOLVER solver, LIS_PRECON precon, integer ierr)
```

## Description

Solve system of linear equations Ax = b with specified solver and predefined preconditioner

## Input

A Coefficient matrix

b Right hand side vector

x Initial vector

solver Solver

precon Preconditioner

Output

x Solution

solver Number of iterations, execution time, etc.

ierr Return code (0)

## 6.4.7 lis\_solver\_get\_status

## Description

Get status from solver

## Input

solver Solver

## Output

status Number of iterations

ierr Return code

## 6.4.8 lis\_solver\_get\_iters

## Description

Get number of iterations from solver

# Input

solver Solver

## Output

iters Number of iterations

## 6.4.9 lis\_solver\_get\_itersex

## Description

Get number of iterations from solver

#### Input

solver Solver

## Output

iters Number of iterations

iters\_double Number of double precision iterations

iters\_quad Number of quadruple precision iterations

ierr Return code

# 6.4.10 lis\_solver\_get\_time

# Description

Get execution time from solver

## Input

solver Solver

## Output

times Time in seconds for execution

## 6.4.11 lis\_solver\_get\_timeex

```
C int lis_solver_get_timeex(LIS_SOLVER solver, double *times, double *ptimes, double *p_c_times, double *p_i_times)

Fortran subroutine lis_solver_get_timeex(LIS_SOLVER solver, real*8 times, real*8 itimes, real*8 ptimes, real*8 p_c_times, real*8 p_i_times, integer ierr)
```

### Description

Get execution time from solver

## Input

solver Solver

Output

times Total time in seconds

itimes Time in seconds for iteration

ptimes Time in seconds for preconditioning

p\_c\_times Time in seconds for creating preconditioner

p\_i\_times Time in seconds for iteration in preconditioner

ierr Return code

## 6.4.12 lis\_solver\_get\_residualnorm

C int lis\_solver\_get\_residualnorm(LIS\_SOLVER solver, LIS\_REAL \*residual)
Fortran subroutine lis\_solver\_get\_residualnorm(LIS\_SOLVER solver, LIS\_REAL residual,
integer ierr)

## Description

Calculate relative redidual norm  $||b - Ax||_2/||b||_2$  from solution x

### Input

solver Solver

Output

residual norm  $||b - Ax||_2/||b||_2$ 

# 6.4.13 lis\_solver\_get\_rhistory

```
C int lis_solver_get_rhistory(VECTOR v)
Fortran subroutine lis_solver_get_rhistory(LIS_VECTOR v, integer ierr)
```

## Description

Store residual norm history of solver

## Input

None

## Output

v Vector

ierr Return code

#### Note

The vector v must be created in advance with the function lis\_vector\_create. When the vector v is shorter than the residual history, it stores the residual history in order to the vector v.

## 6.4.14 lis\_solver\_get\_solver

#### Description

Get solver number from solver

## Input

solver Solver

## Output

nsol Solver number lierr Return code

#### Note

The number of the solver is as follows:

Solver	Number	Solver	Number
CG	1	SOR	12
$\operatorname{BiCG}$	2	BiCGSafe	13
CGS	3	CR	14
BiCGSTAB	4	BiCR	15
BiCGSTAB(l)	5	CRS	16
GPBiCG	6	BiCRSTAB	17
TFQMR	7	GPBiCR	18
Orthomin(m)	8	BiCRSafe	19
GMRES(m)	9	FGMRES(m)	20
Jacobi	10	IDR(s)	21
Gauss-Seidel	11	MINRES	22

# ${\bf 6.4.15}\quad {\bf lis\_get\_solvername}$

C int lis\_get\_solvername(int nsol, char \*name)
Fortran subroutine lis\_get\_solvername(integer nsol, character name, integer ierr)

## Description

Get solver name from solver number

## Input

nsol Solver number

### Output

name Solver name ierr Return code

# 6.5 Solving Eigenvalue Problems

# ${\bf 6.5.1} \quad {\bf lis\_esolver\_create}$

C int lis\_esolver\_create(LIS\_ESOLVER \*esolver)
Fortran subroutine lis\_esolver\_create(LIS\_ESOLVER esolver, integer ierr)

### Description

Create eigensolver

## Input

None

## Output

esolver Eigensolver ierr Return code

## Note

esolver has the information on the eigensolver, the preconditioner, etc.

## 6.5.2 lis\_esolver\_destroy

C int lis\_esolver\_destroy(LIS\_ESOLVER esolver)
Fortran subroutine lis\_esolver\_destroy(LIS\_ESOLVER esolver, integer ierr)

## Description

Destroy eigensolver

## Input

esolver Eigensolver to be destoyed

# Output

### 6.5.3 lis\_esolver\_set\_option

### Description

Set options for eigensolver

### Input

text Command line options

### Output

esolver Eigensolver ierr Return code

### Note

The table below shows the available command line options, where -e {pi|1} means -e pi or -e 1 and -emaxiter [1000] indicates that -emaxiter defaults to 1,000.

Specifying Eigensolvers (Default: -e pi)

Specifying Eigensorvers (Delaute. C pr)				
Method	Option	Auxiliary Options		
Power Iteration	-e {pi 1}			
Inverse Iteration	-e {ii 2}	-i [bicg]	Linear solver	
Approximate Inverse Iteration	-e {aii 3}			
Rayleigh Quotient Iteration	-e {rqi 4}	-i [bicg]	Linear solver	
Subspace Iteration	-e {si 5}	-ss [2]	Size of subspace	
		-m [O]	Mode number	
Lanczos Iteration	-e {li 6}	-ss [2]	Size of subspace	
		-m [O]	Mode number	
Conjugate Gradient	-e {cg 7}			
Conjugate Residual	-e {cr 8}			

# ${\bf Specifying\ Preconditioners}\ ({\rm Default:\ -p\ ilu})$

Preconditioner	Option	Auxiliary Options	
None	-p {none 0}		
Jacobi	-p {jacobi 1}		
ILU(k)	-p {ilu 2}	-ilu_fill [0]	Fill level $k$
SSOR	-p {ssor 3}	-ssor_w [1.0]	Relaxation coefficient $\omega$ (0 < $\omega$ < 2)
Hybrid	-p {hybrid 4}	-hybrid_i [sor]	Linear equations solver
		-hybrid_maxiter [25]	Maximum number of iterations
		-hybrid_tol [1.0e-3]	Convergence criterion
		-hybrid_w [1.5]	Relaxation coefficient $\omega$ for SOR
			$(0 < \omega < 2)$
		-hybrid_ell [2]	Degree $l$ of BiCGSTAB(l)
		-hybrid_restart [40]	Restart values for GMRES and Orthomin
I+S	-p {is 5}	-is_alpha [1.0]	Parameter $\alpha$ for preconditioner
			of $I + \alpha S^{(m)}$ type
		-is_m [3]	Parameter $m$ for preconditioner
			of $I + \alpha S^{(m)}$ type
SAINV	-p {sainv 6}	-sainv_drop [0.05]	Drop criterion
SA-AMG	-p {saamg 7}	-saamg_unsym [false]	Selects unsymmetric version
			(Matrix structure must be symmetric)
		-saamg_theta [0.05 0.12]	Drop criterion $a_{ij}^2 \le \theta^2  a_{ii}   a_{jj} $
			(symmetric or unsymmetric)
Crout ILU	-p {iluc 8}	-iluc_drop [0.05]	Drop criterion
		-iluc_rate [5.0]	Ratio of maximum fill-in
ILUT	-p {ilut 9}	-ilut_drop [0.05]	Drop criterion
		-ilut_rate [5.0]	Ratio of maximum fill-in
Additive Schwarz	-adds true	-adds_iter [1]	Number of iterations

# Other Options

Option				
-emaxiter [1000]	Maximum number of iterations			
-etol [1.0e-12]	Convergence criterion			
-eprint [0]	Display of the residual			
	-eprint {none 0} None			
	-eprint {mem 1} Saves the residual history in memory			
	-eprint {out 2} Displays the residual history			
	-eprint {all 3} Saves the residual history and displays it on the screen			
-ie [ii]	Inner eigensolver used in Lanczos Iteration or Subspace Iteration			
	-ie {pi 1} Power Iteration (Subspace Iteration only)			
	-ie {ii 2} Inverse Iteration			
	-ie {aii 3} Approximate Inverse Iteration			
	-ie {rqi 4} Rayleigh Quotient Iteration			
-shift [0.0]	Amount of shift			
-initx_ones [true]	Behavior of the initial vector $x_0$			
	-initx_ones {false 0} Given values			
	-initx_ones {true 1} All elements are set to 1			
-omp_num_threads [t]	Number of threads			
	(t represents the maximum number of threads)			
-estorage [0]	Matrix storage format			
-estorage_block [2]	Block size of BSR and BSC			

# Precision (Default: -ef double)

Precision	Option	Auxiliary Options
DOUBLE	-ef {double 0}	
QUAD	-ef {quad 1}	

### 6.5.4 lis\_esolver\_set\_optionC

```
C int lis_esolver_set_optionC(LIS_ESOLVER esolver)
Fortran subroutine lis_esolver_set_optionC(LIS_ESOLVER esolver, integer ierr)
```

#### Description

Set options for eigensolver on command line

### Input

None

### Output

esolver Eigensolver ierr Return code

### 6.5.5 lis\_esolve

#### Description

Solve eigenvalue problem  $Ax = \lambda x$  with specified eigensolver

### Input

A Matrix

x Initial vector

esolver Eigensolver

Output

evalue Eigenvalue of mode specified by -m [0] option

x Associated eigenvector

esolver Number of iterations, execution time, etc.

ierr Return code (0)

### 6.5.6 lis\_esolver\_get\_status

### Description

Get status from eigensolver

### Input

esolver Eigensolver

## Output

status Number of iterations

ierr Return code

### 6.5.7 lis\_esolver\_get\_iters

### Description

Get number of iterations from eigensolver

# Input

esolver Eigensolver

### Output

iters Number of iterations

### 6.5.8 lis\_esolver\_get\_itersex

### Description

Get number of iterations from eigensolver

### Input

esolver Eigensolver

### Output

iters Number of iterations

iters\_double Number of double precision iterations

iters\_quad Number of quadruple precision iterations

ierr Return code

# 6.5.9 lis\_esolver\_get\_time

# Description

Get execution time from eigensolver

# Input

esolver Eigensolver

## Output

times Time in seconds for execution

### 6.5.10 lis\_esolver\_get\_timeex

C int lis\_esolver\_get\_timeex(LIS\_ESOLVER esolver, double \*times, double \*times, double \*ptimes, double \*pc\_times, double \*p\_i\_times)

Fortran subroutine lis\_esolver\_get\_timeex(LIS\_ESOLVER esolver, real\*8 times, real\*8 itimes, real\*8 ptimes, real\*8 p\_c\_times, real\*8 p\_i\_times, integer ierr)

#### Description

Get execution time from eigensolver

### Input

esolver Eigensolver

Output

times Total time in seconds

itimes Time in seconds for iteration

ptimes Time in seconds for preconditioning

p\_c\_times Time in seconds for creating preconditioner

p\_i\_times Time in seconds for iteration in preconditioner

ierr Return code

### $\bf 6.5.11 \quad lis\_esolver\_get\_residual norm$

### Description

Calculate relative residual norm  $||\lambda x - Ax||_2/\lambda$  from eigenvector x

#### Input

esolver Eigensolver

Output

residual norm  $||\lambda x - Ax||_2/\lambda$ 

# 6.5.12 lis\_esolver\_get\_rhistory

```
C int lis_esolver_get_rhistory(VECTOR v)
Fortran subroutine lis_esolver_get_rhistory(LIS_VECTOR v, integer ierr)
```

### Description

Store residual norm history of eigensolver

### Input

None

### Output

v Vector

ierr Return code

#### Note

The vector v must be created in advance with the function lis\_vector\_create. When the vector v is shorter than the residual history, it stores the residual history in order to the vector v.

### 6.5.13 lis\_esolver\_get\_evalues

#### Description

Store eigenvalues in vector

### Input

esolver Eigensolver

## Output

v Vector which stores eigenvalues

ierr Return code

#### Note

The vector v must be created in advance with the function lis\_vector\_create.

### 6.5.14 lis\_esolver\_get\_evectors

### Description

Store eigenvectors in matrix

## Input

esolver Eigensolver

### Output

A Matrix in CRS format which stores eigenvectors

ierr Return code

#### Note

The matrix A must be created in advance with the function  $lis_matrix_create$ .

### 6.5.15 lis\_esolver\_get\_esolver

### Description

Get eigensolver number from eigensolver

### Input

esolver Eigensolver

Output

nesol Eigensolver number

ierr Return code

#### Note

The number of the eigensolver is as follows:

Method	Number
Power Iteration	1
Inverse Iteration	2
Approximate Inverse Iteration	3
Rayleigh Quotient Iteration	4
Subspace Iteration	5
Lanczos Iteration	6
Conjugate Gradient	7
Conjugate Residual	8

### 6.5.16 lis\_get\_esolvername

C int lis\_get\_esolvername(int nesol, char \*ename)
Fortran subroutine lis\_get\_esolvername(integer nesol, character ename, integer ierr)

### Description

Get eigensolver name from eigensolver number

### Input

nesol Eigensolver number

Output

name Eigensolver name

# 6.6 Operating External Files

# 6.6.1 lis\_input

### Description

Read matrix and vector from file

### Input

filename Source file

### Output

A Matrix in specified storage format

b Right hand side vector

x Solution

ierr Return code

#### Note

The supported file formats are shown below:

- Matrix Market format (extended to allow vector data to be read)
- Harwell-Boeing format

## 6.6.2 lis\_input\_vector

```
C int lis_input_vector(LIS_VECTOR v, char *filename)
Fortran subroutine lis_input_vector(LIS_VECTOR v, character filename, integer ierr)
```

### Description

Read vector from file

## Input

filename Source file

### Output

v Vector

ierr Return code

#### Note

The following formats are supported:

- PLAIN format
- MM format

### 6.6.3 lis\_input\_matrix

#### Description

Read matrix from file

### Input

filename Source file

#### Output

A Matrix in specified storage format

x Solution

ierr Return code

#### Note

The supported file formats are shown below:

- Matrix Market format (extended to allow vector data to be read)
- Harwell-Boeing format

### 6.6.4 lis\_output

C int lis\_output(LIS\_MATRIX A, LIS\_VECTOR b, LIS\_VECTOR x, int format, char \*filename)

Fortran subroutine lis\_output(LIS\_MATRIX A, LIS\_VECTOR b, LIS\_VECTOR x, integer format, character path, integer ierr)

# Description

Write matrix and vector into file

### Input

A Matrix

b Right hand side vector (If no vector is written to an external file,

then NULL must be input.)

x Solution (If no vector is written to an external file, then NULL must

be input.)

format File format

LIS\_FMT\_MM Matrix Market format

filename Destination file

Output

### 6.6.5 lis\_output\_vector

#### Description

Write vector into file

### Input

v Vector

format File format

LIS\_FMT\_PLAIN PLAIN format
LIS\_FMT\_MM MM format

LIS\_FMT\_LIS Lis format(ASCII)

filename Destination file

Output

ierr Return code

### 6.6.6 lis\_output\_matrix

#### Description

Write matrix into file

# Input

A Matrix

format File format

LIS\_FMT\_MM Matrix Market format

filename Destination file

Output

### 6.7 Other Functions

### 6.7.1 lis\_initialize

```
C int lis_initialize(int* argc, char** argv[])
Fortran subroutine lis_initialize(integer ierr)
```

#### Description

Initialize execution environment

### Input

argc Number of command line arguments

argv Command line argument

Output

ierr Return code

#### 6.7.2 lis\_finalize

```
C void lis_finalize()
Fortran subroutine lis_finalize(integer ierr)
```

### Description

Finalize execution environment

#### Input

None

### Output

ierr Return code

### 6.7.3 lis\_wtime

```
C double lis_wtime()
Fortran function lis_wtime()
```

### Description

Measure elapsed time

### Input

None

#### Output

Elapsed time in seconds from given point is returned as double number

#### Note

To measure the processing time, call <code>lis\_wtime</code> to get the starting time, call it again to get the ending time, and calculate the difference.

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# A File Formats

This section describes the file formats available for the library.

#### A.1 Extended Matrix Market Format

The Matrix Market format[32] is not designed to store vector data. For this library, it has been extended to handle vector data. Assume that the number of the nonzero elements for the matrix  $A = (a_{ij})$  of  $M \times N$  is L and that  $a_{ij} = A(I, J)$ . The format is as follows:

```
%%MatrixMarket matrix coordinate real general <-- Header
%
                                                  | Comment lines with 0 or more lines
%
                                                <-+
MNLBX
                                                <-- Numbers of rows, columns, and
I1 J1 A(I1,J1)
                                                       nonzero elements (0 or 1) (0 or 1)
I2 J2 A(I2,J2)
                                                  | Row and column number values
                                                  | The index is one origin
IL JL A(IL, JL)
                                                <-+
I1 B(I1)
                                                <-+
I2 B(I2)
                                                  | Exists only when B=1
                                                  | Row number value
IM B(IM)
                                                <-+
I1 X(I1)
                                                <-+
I2 X(I2)
                                                  | Exists only when X=1
                                                  | Row number value
IM X(IM)
```

The extended Matrix Market format for the matrix A and the vector b in Equation (A.1) is as follows:

$$A = \begin{pmatrix} 2 & 1 & & \\ 1 & 2 & 1 & & \\ & 1 & 2 & 1 \\ & & 1 & 2 \end{pmatrix} \qquad b = \begin{pmatrix} 0 & & \\ 1 & & \\ 2 & & \\ 3 & & \end{pmatrix}$$
 (A.1)

 $\M$ MatrixMarket matrix coordinate real general

```
4 4 10 1 0
1 2 1.00e+00
1 1 2.00e+00
2 3
    1.00e+00
2 1
    1.00e+00
2 2
    2.00e+00
3 4 1.00e+00
3 2 1.00e+00
3 3 2.00e+00
4 4 2.00e+00
4 3 1.00e+00
1 0.00e+00
2 1.00e+00
3 2.00e+00
4 3.00e+00
```

# A.2 Harwell-Boeing Format

The Harwell-Boeing format inputs and outputs the matrix in the CCS storage format. Assume that the array value stores the values of the nonzero elements of the matrix A, the array index stores the row indices of the nonzero elements and the array ptr stores pointers to the top of each column in the arrays value and index. The format is as follows:

```
Line 1 (A72,A8)
  1 - 72 Title
 73 - 80 Key
Line 2 (5I14)
  1 - 14 Total number of lines excluding header
  15 - 28 Number of lines for ptr
  29 - 42 Number of lines for index
  43 - 56 Number of lines for value
  57 - 70 Number of lines for right hand side vectors
Line 3 (A3,11X,4I14)
  1 - 3 Matrix type
           Col.1: R Real matrix
                   C Complex matrix (Not supported)
                   P Pattern only (Not supported)
            Col.2: S Symmetric
                   U Unsymmetric
                   H Hermitian (Not supported)
                   Z Skew symmetric (Not supported)
                   R Rectangular (Not supported)
            Col.3: A Assembled
                   E Elemental matrices (Not supported)
  4 - 14 Blank space
  15 - 28 Number of rows
  29 - 42 Number of columns
  43 - 56 Number of nonzero elements
  57 - 70 0
Line 4 (2A16,2A20)
  1 - 16 Format for ptr
  17 - 32 Format for index
  33 - 52 Format for value
  53 - 72 Format for right hand side vectors
Line 5 (A3,11X,2I14) Only presents if there are right hand side vectors
          Right hand side vector type
   1
           F for full storage
           M for same format as matrix (Not supported)
          G if a starting vector is supplied
         X if an exact solution is supplied
   4 - 14 Blank space
  15 - 28 Number of right hand side vectors
  29 - 42 Number of nonzero elements
```

The Harwell-Boeing format for the matrix A and the vector b in Equation (A.1) is as follows:

```
1 - - - - - 10 - - - - - 20 - - - - - 30 - - - - - 40 - - - - - 50 - - - - - - 60 - - - - - 70 - - - - - 80
Harwell-Boeing format sample
                                                                    2
            8
                          1
                                        1
                                                      4
RUA
                                        4
                                                     10
                                                                     4
(11i7)
                                (3e26.18)
                                                   (3e26.18)
                (13i6)
F
                          1
                           9
             3
     1
    1
          2
                1
                      2
                            3
                                  2
                                        3
                                              4
                                                    3
                                                          4
  2.0000000000000000E+00 1.00000000000000E+00 1.0000000000000E+00
  2.00000000000000000E+00
                           1.00000000000000000E+00 1.000000000000000E+00
  2.00000000000000000E+00
                           1.00000000000000000E+00 1.000000000000000E+00
  2.0000000000000000E+00
                           1.00000000000000000E+00 2.000000000000000E+00
  0.0000000000000000E+00
  3.0000000000000000E+00
```

# A.3 Extended Matrix Market Format for Vectors

The Matrix Market format [32] has been extended to store vector data. Assume that the vector  $b = (b_i)$  is a vector of order N and that  $b_i = B(I)$ . The format is as follows:

```
      %//MatrixMarket vector coordinate real general
      <-- Header</td>

      %
      <-+</td>

      %
      | Comment lines with 0 or more lines

      %
      <-+</td>

      N
      <-- Number of rows</td>

      I1 B(I1)
      <-+</td>

      I2 B(I2)
      | Row number value

      . . .
      | The index is one origin

      IN B(IN)
      <-+</td>
```

The extended Matrix Market format for the vector b in Equation (A.1) is as follows:

```
%%MatrixMarket vector coordinate real general
4
1 0.00e+00
2 1.00e+00
3 2.00e+00
4 3.00e+00
```

### A.4 PLAIN Format for Vectors

The PLAIN format for vectors is designed to write vector values in order. Assume that the vector  $b = (b_i)$  is a vector of order N and that  $b_i = B(I)$ . The format is as follows:

The PLAIN format for the vector b in Equation (A.1) is as follows:

0.00e+00 1.00e+00 2.00e+00 3.00e+00