# STRUMPACK Users' Guide

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### 1 STRUMPACK Overview

STRUMPACK – STRUctured Matrix PACKage – is a C++ library for computations with dense and sparse matrices. It uses so-called *structured matrices*, i.e., matrices that exhibit some kind of low-rank property, in this case with Hierarchically Semi-Separable matrices (HSS), to speed up linear algebra operations. This version of STRUMPACK unifies two main components that were separate in previous versions: a package for dense matrix computations (STRUMPACK-dense) and a package (STRUMPACK-sparse) for sparse linear systems. The algorithms for solving dense linear systems are described in [6] while the algorithms for solving sparse linear systems are described in [4, 3]. STRUMPACK can be used as a general algebraic sparse direct solver (based on the multifrontal factorization method), or as an efficient preconditioner for sparse matrices obtained by discretization of partial differential equations. Included in STRUMPACK are also the GMRES and BiCGStab iterative Krylov solvers, that use the approximate, HSS-accelerated, sparse solver as a preconditioner for the efficient solution of sparse linear systems.

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Check the STRUMPACK website for more information and for the latest code:

http://portal.nersc.gov/project/sparse/strumpack/

## 2 Installation and Requirements

The STRUMPACK package uses the CMake build system (CMake version >= 2.8). The recommended way of building the STRUMPACK library is as follows:

```
> tar -xvzf strumpack-x.y.z.tar.gz
> cd strumpack-x.y.x
> mkdir build
> cd build
> cmake ... / -DCMAKE BUILD TYPE=Release \
             -DCMAKE INSTALL PREFIX=/path/to/install \
             -DCMAKE CXX COMPILER=<C++ compiler> \
                                                                                                                                                                                                                         # this and below are optional,
             -DCMAKE_C_COMPILER=<C compiler> \
                                                                                                                                                                                                                         # CMake will try to autodetect
             -DCMAKE Fortran COMPILER=<Fortran77 compiler> \
             -DSCALAPACK_LIBRARIES="/path/to/scalapack/libscalapack.a;/path/to/blacs/libblacs.a" \
             -DMETIS INCLUDES=/path/to/metis/incluce \
             -DMETIS LIBRARIES=/path/to/metis/libmetis.a \
             -DSTRUMPACK USE PARMETIS=ON \
                                                                                                                                                                                                                         # optional
             -DPARMETIS_INCLUDES=/path/to/parmetis/include \
             -DPARMETIS_LIBRARIES=/path/to/parmetis/libparmetis.a \
             -DSTRUMPACK USE SCOTCH=ON \
                                                                                                                                                                                                                         # optional
             -DSCOTCH INCLUDES=/path/to/scotch/include \
             -DSCOTCH LIBRARIES="/path/to/ptscotch/libscotch.a;...libscotcherr.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...libptscotch.a;...lib
> make test
                                           # optional, takes a while
> make install
```

The above will only work if you have the following dependencies, and CMake can find them:

• C++11, C and FORTRAN77 compilers. CMake looks for these compilers in the standard locations, if they are installed elsewhere, you can specify them as follows:

```
-DCMAKE_C_COMPILER=gcc \
-DCMAKE Fortran COMPILER=gfortran
```

- MPI (Message Passing Interface) library. You should not need to manually specify the MPI compiler wrappers. CMake will look for MPI options and libraries and set the appropriate compiler and linker flags.
- OpenMP v3.1 support is required in the C++ compiler to use the shared-memory parallelism in the code. OpenMP v3.1 introduces task parallelism, which is used extensively throughout the code. CMake will check whether your compiler supports OpenMP and sets the appropriate compiler and linker flags.
- BLAS, LAPACK and Scalapack libraries. For performance it is crucial to use optimized BLAS/LAPACK libraries like for instance Intel<sup>®</sup> MKL, AMD<sup>®</sup> ACML, Cray<sup>®</sup> LibSci or OpenBLAS. The default versions of the Intel<sup>®</sup> MKL and Cray<sup>®</sup> LibSci BLAS libraries will use multithreaded kernels, unless when they are called from within an OpenMP parallel region, in which case they run sequentially. This is the behavior STRUMPACK relies upon to achieve good performance when running in MPI+OpenMP hybrid mode. Scalapack depends on the Blacs communication library and on PBLAS (parallel Blas), both of which are typically included with the Scalapack installation (from Scalapack 2.0.2, the blacs library is included in the Scalapack library file). If CMake cannot locate these libraries, you can specify their path by setting the environment variable \$SCALAPACKDIR or by specifying the libraries manually:

```
> cmake ../ -DCMAKE_BUILD_TYPE=Release \
    -DSCALAPACK_LIBRARIES="/path/to/scalapack/libscalapack.a;/path/to/blacs/libblacs.a"
```

Or one can also directly modify the linker flags to add the ScaLAPACK and BLACS libraries:

```
> cmake ../ -DCMAKE_BUILD_TYPE=Release \
   -DCMAKE_EXE_LINKER_FLAGS="-L/usr/lib64/mpich/lib/ -lscalapack -lmpiblacs"
```

• METIS (≥ 5.1.0 required) for the nested dissection matrix reordering. Metis can be obtained from: http://glaros.dtc.umn.edu/gkhome/metis/metis/download.

CMake looks for the Metis inlude files the library in the default locations as well as in \$metisdir/include and \$metisdir/lib. Using the Bash shell, the METISDIR environment variable can be set as export METISDIR=/usr/local/metis/. Alternatively, you can specify the location of the header and library as follows:

```
> cmake ../ -DCMAKE_BUILD_TYPE=Release \
 -DMETIS_INCLUDES=/usr/local/metis/include \
 -DMETIS_LIBRARIES=/usr/local/metis/lib/libmetis.a
```

• PARMETIS (optional) for parallel nested dissection. ParMetis can be download from

http://glaros.dtc.umn.edu/gkhome/metis/parmetis/download

The steps to make sure CMake can find ParMetis are similar as for Metis. Enable with -dstrumpack\_use\_parmetis. The CMake variables are \$parmetisdir or parmetis\_includes and parmetis\_libraries.

• SCOTCH and PT-SCOTCH (≥ 5.1.12) (optional) for matrix reordering. Scotch can be downloaded from:

http://www.labri.fr/perso/pelegrin/scotch/

Configuring CMake to find (PT-)Scotch is similar to Metis. Enable with -DSTRUMPACK\_USE\_SCOTCH For (PT-)Scotch the CMake variables are \$SCOTCHDIR or SCOTCH\_INCLUDES and SCOTCH\_LIBRARIES. Make sure to specify all libraries: libscotch, libscotcherr, libptscotch and libptscotcherr.

- getopt\_long: This is a GNU extension to the POSIX getopt() C library function.
- TCMalloc, TBB Malloc or jemalloc: This is optional, but recommended, as it can lead to dramatic performance improvements for multithreaded code that performs frequent memory allocations. Link with the one of these libraries, e.g.:

```
-DCMAKE EXE LINKER FLAGS="-ltcmalloc"
```

to replace the default memory allocator (C++ new) with a more scalable implementation. See also Section 9.

The code was tested on GNU/Linux with the GNU and Intel<sup>®</sup> compilers and the OpenBLAS, Intel<sup>®</sup> MKL<sup>®</sup> and Cray<sup>®</sup> LibSci<sup>®</sup> numerical libraries. If you encounter issues on other platforms or with other BLAS/LAPACK implementations, please let us know. Successful compilation will create a libstrumpack.a file

## 3 Algorithm

The algorithm used in STRUMPACK is described in detail in [4], and is based on the work by Jianlin Xia [8]. Here we summarize the main algorithm features. Section 5 has more information on the low-rank compression strategy and how to tune this to get a good preconditioner for your specific problem. There are three main steps in the algorithm: matrix reordering, factorization and solve.

Matrix reordering: There are three distinct matrix reordering steps: one for stability, one to limit fill-in and one to reduce HSS-ranks. First, the matrix is reordered and possibly scaled for numerical stability by the MC64 code [2]. For many matrices, this reordering can safely be disabled. By default, MC64 is used to maximize the product of the diagonal values of the matrix, and to scale the rows and columns of the matrix. Alternatively, MC64 can be used to maximize the smallest diagonal value or to maximize the sum of the diagonals. Next, a nested dissection reordering is applied to limit fill-in. Both (Par)Metis and (PT-)Scotch are supported. We expose one user tunable parameter which controls the size of the smallest separators. Finally, when HSS compression is used, there is an extra reordering step to reduce the HSS-ranks. This reordering uses Metis and does not require user tuning.

Factorization: Before the actual numerical factorization, there is a symbolic factorization step to construct the elimination tree. After that, the multifrontal factorization procedure traverses this elimination tree from bottom (smallest separators) to top (root separator). With each node of the elimination tree a dense matrix is associated, referred to as a frontal matrix, or simply front. These fronts can possibly be compressed as Hierarchically Semi-Separable (HSS) matrices. This compression will only pay off for fronts that are large enough, which are typically the frontal matrices at the nodes in the elimination tree close to the root. Without any HSS compression, the solver acts as a standard multifrontal direct solver. HSS approximations are constructed using a randomized sampling algorithm.

Solve: Once the matrix is factorized, the factors can be used to efficiently solve a linear system of equations by doing a forward and a backward solve sweeps. When no HSS compression is used, this is a direct solver. The multifrontal solve procedure is then used within an iterative refinement loop, with typically only 1 or very few iterations. However, when the factors are compressed using HSS, a single multifrontal solve is only approximate and the solve is by default used as a preconditioner for GMRES(30). The required number of GMRES iterations will depend strongly on the quality of the HSS approximation.

## 4 Using STRUMPACK Sparse

This section gives an overview on the basic usage of the sparse solvers in STRUMPACK. Many STRUMPACK options can be set from the command line. Running with --help or -h, will give you a list of supported runtime options.

An example Makefile is available in the examples/ directory. This Makefile is generated by the cmake command, see Section 2.

The STRUMPACK package is written in C++, and offers a simple C++ interface. See Section 8 if you prefer a C interface. STRUMPACK-sparse has three different solver classes, all interaction happens through objects of these classes:

#### • StrumpackSparseSolver<scalar,integer=int>

This class represents the sparse solver for a single computational node, optionally using OpenMP parallelism. Use this if you are running the code sequentially, on a (multicore) laptop or desktop or on a single node of a larger cluster. This class is defined in StrumpackSparseSolver.hpp, so include this header if you intend to use it.

#### • StrumpackSparseSolverMPI<scalar,integer=int>

This solver has (mostly) the same interface as StrumpackSparseSolver<scalar,integer> but the numerical factorization and multifrontal solve phases run in parallel using MPI and ScaLAPACK. However, the inputs (sparse matrix, right-hand side vector) need to be available completely on every MPI process. The reordering phase uses Metis or Scotch (not ParMetis or PTScotch) and the symbolic factorization is threaded, but not distributed. The (multifrontal) solve is done in parallel, but the right-hand side vectors need to be available completely on every processor. Make sure to call MPI\_Init[\_thread] before instantiating an object of this class and include the header file StrumpackSparseSolverMPI.hpp. We do not recommend this solver, instead, use StrumpackSparseSolverMPIDist whenever possible.

#### • StrumpackSparseSolverMPIDist<scalar,integer=int>

This solver is fully distributed. The numerical factorization and solve as well as the symbolic factorization are distributed. The input is now a block-row distributed sparse matrix and a correspondingly distributed right-hand side. For matrix reordering, ParMetis or PT-Scotch are used. Include the header file StrumpackSparseSolverMPIDist.hpp and call MPI\_Init[\_thread]. Unfortunately, there is no distributed version of the MC64 reordering code, so if this reordering (and scaling) step is enabled, the code will gather the distributed sparse matrix on a single node and then apply MC64 sequentially.

The three solver classes StrumpackSparseSolver, StrumpackSparseSolverMPI and

StrumpackSparseSolverMPIDist depend on two template parameters <scalar,integer>: the type of a scalar and an integer type. The scalar type can be float, double, std::complex<float> or std::complex<double>. It is recommended to first try to simply use the default integer=int type, unless you run into 32 bit integer overflow problems. In that case one can switch to for instance int64\_t (a signed integer type).

#### 4.1 StrumpackSparseSolver Example

The following shows the typical way to use a (sequential or multithreaded) STRUMPACK sparse solver:

```
#include "StrumpackSparseSolver.hpp"
using namespace strumpack; // all strumpack code is in the strumpack namespace,
int main(int argc, char* argv[]) {
 int N = ...;
                             // construct an NxN CSR matrix with nnz nonzeros
 int* row ptr = ...;
                             // N+1 integers
 int* col ind = ...;
                             // nnz integers
 double* val = ...;
                             // nnz scalars
 double* x = new double[N]; // will hold the solution vector
 double* b = ...;
                             // set a right-hand side b
 StrumpackSparseSolver<double> sp;
                                                 // create solver object
  sp.options().set_rel_tol(1e-10);
                                                 // set options
  sp.options().set_gmres_restart(10);
                                                 // ...
                                                 // enable HSS compression, see section 5
  sp.options().enable_HSS();
```

$$A = \begin{bmatrix} 8.2 & 0.1 & & & & & & \\ 0 & & -4.8 & & & & \\ 6.2 & 1.1 & & 2.6 & & & \\ & & -1.0 & & & & \\ & & & 99.9 & 4.0 \end{bmatrix} \qquad \begin{array}{c} \mathsf{row\_ptr} = [0, \, 3, \, 5, \, 8, \, 9, \, 11] \\ \mathsf{col\_ind} = [0, \, 1, \, 4 \, | \, 0, \, 2 \, | \, 0, \, 1, \, 3 \, | \, 2 \, | \, 3, \, 4] \\ \mathsf{values} = [8.2, \, 0.1, \, 3.1 \, | \, 0, \, -4.8 \, | \, 6.2, \, 1.1, \, 2.6 \, | \, -1.0 \, | \, 99.9, \, 4.0] \\ \end{array}$$

Figure 1: Illustration of a small  $5 \times 5$  sparse matrix with 11 nonzeros and its Compressed Sparse Row (CSR) or Yale format representation. We always use 0-based indexing! Let N=5 denote the number of rows. The row\_ptr array has N+1 elements, with element i denoting the start of row i in the col\_ind and values arrays. Element row\_ptr[N] = nnz, i.e., the total number of nonzero elements in the matrix. The values array holds the actual matrix values, ordered by row. The corresponding elements in col\_ind give the column indices for each nonzero. There can be explicit zero elements in the matrix. The nonzero values and corresponding column indices need not be sorted by column (within a row).

```
sp.options().set_from_command_line(argc, argv); // parse command line options
sp.set_csr_matrix(N, row_ptr, col_ind, val); // set the matrix (copy)
sp.reorder(); // reorder matrix
sp.factor(); // numerical factorization
sp.solve(b, x); // solve Ax=b
... // check residual/error and cleanup
}
```

The main steps are: create solver object, set options (parse options from the command line), set matrix, reorder, factor and finally solve. The matrix should be in the Compressed Sparse Row (CSR) format, also called Yale format, with 0 based indices. Figure 1 illustrates the CSR format. In the basic scenario, it is not necessary to explicitly call reorder and factor, since trying to solve with a StrumpackSparseSolver object that is not factored yet, will internally call the factor routine, which will call reorder if necessary.

The above code should be linked with -lstrumpack and with the Metis, ParMetis, Scotch, PT-Scotch, BLAS, LAPACK, and ScaLAPACK libraries.

### 4.2 StrumpackSparseSolverMPI Example

Usage of the StrumpackSparseSolverMPI<scalar,integer=int> solver is very similar:

```
#include "StrumpackSparseSolverMPI.hpp"
using namespace strumpack;
int main(int argc, char* argv[]) {
 int thread_level, rank;
 // StrumpackSparseSolverMPI uses OpenMP so we should ask for MPI_THREAD_FUNNELED at least
 MPI_Init_thread(&argc, &argv, MPI_THREAD_FUNNELED, &thread_level);
 MPI_Comm_rank(MPI_COMM_WORLD, &rank);
 if (thread level != MPI THREAD FUNNELED && rank == 0)
    std::cout << "MPI,.implementation.does.not.support,MPI THREAD FUNNELED" << std::endl;</pre>
    // define the same CSR matrix as for StrumpackSparseSolver
   int N = ...;
                         // construct an NxN CSR matrix with nnz nonzeros
   int* row_ptr = ...;
                               // N+1 integers
   int* col ind = ...;
                               // nnz integers
```

The only difference here is the use of StrumpackSparseSolverMPI instead of StrumpackSparseSolver and the calls to MPI Init thread, Cblacs exit and MPI Finalize.

#### 4.3 StrumpackSparseSolverMPIDist Example

Finally, we illustrate the usage of StrumpackSparseSolverMPIDist<scalar,integer=int> solver. This interface takes a block-row distributed compressed sparse row matrix as input. This matrix format is illustrated in Figure 2.

```
#include "StrumpackSparseSolverMPI.hpp"
using namespace strumpack;
int main(int argc, char* argv[]) {
 int thread level, rank, P;
 MPI Init thread(&argc, &argv, MPI THREAD FUNNELED, &thread level);
 MPI Comm rank(MPI COMM WORLD, &rank);
 MPI_Comm_size(MPI_COMM_WORLD, &P);
   // define a block-row distributed CSR matrix
   int* dist = new int[P];
   // set dist such that processor p owns rows [dist[p], dist[p+1]) of the sparse matrix
   for (int p=0; p<P; p++) dist[p] = .. ;</pre>
   // local n is the number of rows of the input matrix assigned to me
   int local_n = dist[rank+1] - dist[rank];
   int* row_ptr = new int[local_n+1];
    .. // set the sparse matrix row pointers in row_ptr
   int local nnz = row ptr[local n+1] - row ptr[0];
   int* col ind = new int[local nnz];
   .. // set the sparse matrix column indices in col_ind
   double* val = new double[local_nnz];
    .. // set the matrix nonzero value in val
   double* x = new double[local_n];
                                               // local part of solution
              = new double[local_n];
   double* b
                                               // local part of rhs
   for (int i=0; i<local_n; i++) b[i] = ...;  // set the rhs</pre>
   StrumpackSparseSolverMPIDist<double> sp(MPI_COMM_WORLD);
   sp.options().set_reordering_method(ReorderingStrategy::PARMETIS);
   sp.options().set_from_command_line(argc, argv);
    sp.set_distributed_csr_matrix(local_n, row_ptr, col_ind, val, dist);
```

$$A = \begin{bmatrix} 8.2 & 0.1 & & & & & \\ \hline 0 & & -4.8 & & & \\ \hline 6.2 & 1.1 & & 2.6 & & \\ \hline & & -1.0 & & & \\ \hline & & & 99.9 & 4.0 \end{bmatrix} \begin{matrix} P_0 \\ P_2 \end{matrix}$$
 
$$\text{dist} = \begin{bmatrix} 0, 1, 3, 5 \end{bmatrix} \qquad P_2$$
 
$$\text{row\_ptr} = \begin{bmatrix} 0, 3 \end{bmatrix} \qquad \text{row\_ptr} = \begin{bmatrix} 0, 2, 5 \end{bmatrix} \qquad \text{row\_ptr} = \begin{bmatrix} 0, 1, 3 \end{bmatrix}$$
 
$$\text{col\_ind} = \begin{bmatrix} 0, 1, 4 \end{bmatrix} \qquad \text{col\_ind} = \begin{bmatrix} 0, 2 \mid 0, 1, 3 \end{bmatrix} \qquad \text{col\_ind} = \begin{bmatrix} 2 \mid 3, 4 \end{bmatrix}$$
 
$$\text{values} = \begin{bmatrix} 8.2, 0.1, 3.1 \end{bmatrix} \qquad \text{values} = \begin{bmatrix} 0, -4.8 \mid 6.2, 1.1, 2.6 \end{bmatrix} \qquad \text{values} = \begin{bmatrix} -1.0 \mid 99.9, 4.0 \end{bmatrix}$$

Figure 2: Illustration of a small  $5 \times 5$  sparse matrix with 11 nonzeros and its block-row distributed compressed sparse row representation. We always use 0-based indexing! Process  $P_0$  owns row 0, process  $P_1$  has rows 1 and 2 and process  $P_2$  has rows 3 and 4. This distribution of rows over the processes is represented by the dist array. Process p owns rows [dist[p],dist[p+1]). If N=5 is the number of rows in the entire matrix and P is the total number of processes, then dist[P]=N. The (same) dist array is stored on every process. Each process holds a CSR representation of only its local rows of the matrix, see Figure 1.

```
sp.solve(b, x);
... // check residual/error, cleanup
}
Cblacs_exit(1);
MPI_Finalize();
}
```

### 4.4 Initializing the Solver Object

Let

```
typedef strumpack::StrumpackSparseSolver<scalar,integer> Sp;
typedef strumpack::StrumpackSparseSolverMPI<scalar,integer> SpMPI;
typedef strumpack::StrumpackSparseSolverMPIDist<scalar,integer> SpMPIDist;
```

Each of the solver classes has two constructors:

```
Sp::StrumpackSparseSolver(bool verbose=true, bool root=true);
Sp::StrumpackSparseSolver(int argc, char* argv[], bool verbose=true, bool root=true);
```

```
SpMPI::StrumpackSparseSolverMPIDist(MPI_Comm comm, bool verbose=true);
SpMPI::StrumpackSparseSolverMPIDist(MPI_Comm comm, int argc, char* argv[], bool verbose=true);
```

```
SpMPIDist::StrumpackSparseSolverMPIDist(MPI_Comm comm, bool verbose=true);
SpMPIDist::StrumpackSparseSolverMPIDist(MPI_Comm comm, int argc, char* argv[], bool verbose=true);
```

where argc and argv contain the command line options and the verbose option can be set to false to suppress output of the solver. Note that since SpMPIDist is a subclass of SpMPI, which is a subclass of Sp, all public members of Sp are also members of SpMPI and SpMPIDist. The public interface to the SpMPI class is exactly the same as that for the Sp class.

#### 4.5 Sparse Matrix Format

The sparse matrix should be specified in compressed sparse row format [7]:

Internally, the matrix is copied, so it will not be modified. Previous versions of STRUMPACK also supported the CSC format, but this is now deprecated. If the sparsity pattern of the matrix is symmetric (the values do not have to be symmetric), then you can set symmetric\_pattern=true. This saves some work in the setup phase of the solver.

For the SpMPIDist solver the input is a block-row distributed compressed sparse row matrix (as illustrated in the example above):

```
void SpMPIDist::set_distributed_csr_matrix
    (integer local_rows, integer* row_ptr, integer* col_ind,
    scalar* values, integer* dist, bool symmetric_pattern=false);
```

Alternatively, you can also specify a sequential CSR matrix to the Spmpidist solver:

For this routine, the matrix only needs to be specified completely on the root process. Other processes can pass NULL for the arrays.

#### 4.6 Setting and Parsing Options

The solver class has an object of type SPOptions<scalar>, which can be accessed through:

```
SPOptions<scalar>& Sp::options();
```

The SPOptions<scalar> class is defined in SPOptions.hpp. The complete public interface for the SPOptions<scalar> class is given in Section 4.10.1. The following subsections describe some of the options available from SPOptions<scalar> in more detail.

#### 4.7 Reordering

There are three types of matrix reordering: for numerical stability, to reduce fill-in and to reduce the HSS-ranks. These reorderings are all performed when calling

```
ReturnCode Sp::reorder();
```

The return value is of type ReturnCode (defined in strumpack\_parameters.hpp) and can be

```
enum class ReturnCode {
   SUCCESS,     /*!< Operation completed successfully. */
   MATRIX_NOT_SET,     /*!< The input matrix was not set.     */
   REORDERING_ERROR     /*!< The matrix reordering failed.     */
};</pre>
```

#### 4.7.1 Reordering for numerical stability

The reordering for numerical stability is performed using the MC64 code. For many matrices, this reordering is not necessary and can safely be disabled! MC64 supports 5 different modes

0: no reordering for stability, this disables MC64

- 1: currently not supported
- 2: maximize the smallest diagonal value
- **3:** maximize the smallest diagonal value, different strategy
- 4: maximize sum of diagonal values
- 5: maximize product of diagonal values and apply row and column scaling

which can be selected via

```
void SPOptions::set_mc64job(int job);
void SPOptions::set_mc64job(MC64Job job);
int SPOptions::mc64job() const;
```

where mc64() queries the currently selected strategy (the default is 5: maximum product of diagonal values plus row and column scaling). Instead of entering a number, you can specify one of the enumerations in MC64Job:

The command line option

```
--sp_mc64job [0-5]
```

can also be used.

#### 4.7.2 Nested dissection reordering

The STRUMPACK sparse solver supports both (Par)Metis and (PT-)Scotch for the matrix reordering. The following functions can set the preferred method or check the currently selected method:

```
void SPOptions::set_reordering_method(ReorderingStrategy m);
ReorderingStrategy SPOptions::reordering_method() const;
```

The options for MatrixReorderingStrategy are

When the solver is an object of Sp, PARMETIS or PTSCOTCH are not supported. When the solver is parallel, either an SpMPI or SpMPIDist object, and METIS, SCOTCH or RCM are chosen, then the graph of the complete matrix will be gathered onto the root process and the root process will call the (sequential) Metis, Scotch or RCM reordering routine. For large graphs this might fail due to insufficient memory.

The GEOMETRIC option is only allowed for regular grids. In this case, the dimensions of the grid should be specified in the function

```
ReturnCode Sp::reorder(int nx=1, int ny=1, int nz=1);
```

For instance for a regular 2d  $2000 \times 4000$  grid, you can call this as sp.reorder(2000, 4000). In the general algebraic case, the grid dimensions don't have to be provided. The reordering method can also be specified via the command line option

--sp\_reordering\_method [metis|parmetis|scotch|ptscotch|geometric|rcm]

#### 4.8 Factorization

Compute the factorization by calling

```
ReturnCode Sp::factor();
```

where the possible return values are the same as for Sp::reorder(). If Sp::reorder() was not called already, it is called automatically. When HSS compression is not enabled, this will compute an exact LU factorization of the (permuted) sparse input matrix. If HSS compression is enabled (with SPOptions::enable\_HSS() or --sp\_enable\_HSS, see Section 5), the factorization is only approximate.

#### 4.9 Solve

Solve the linear system Ax = b by calling

```
ReturnCode Sp::solve(scalar* b, scalar* x, bool use_initial_guess=false);
```

By default (bool use\_initial\_guess=false) the input in x is ignored. If bool use\_initial\_guess=true, x is used as initial guess for the iterative solver (if an iterative solver is used, for instance iterative refinement or GMRES). If the Sp::factor() was not called, it is called automatically. The return values are the same as for Sp::reorder().

The iterative solver can be chosen through:

```
void SPOptions::set_Krylov_solver(KrylovSolver s);
```

where KrylovSolver can take the following values:

with KrylovSolver::AUTO being the default value. The KrylovSolver::AUTO setting will use iterative refinement when HSS compression is not enabled, and preconditioned GMRES when HSS compression is enabled, see Section 5. To use the solver as a preconditioner, or a single (approximate) solve, set the solver to KrylovSolver::DIRECT. When calling SpMPIDist::solve, the right-hand side and solution vectors should only point to the local parts!

#### 4.10 All Options for the Sparse Solver

The HSS specific options are stored in an object of type HSSOptions<scalar>, inside the SPOptions object. These options are described in Section 5.

#### 4.10.1 SPOptions<scalar> Interface

The complete public interface to the options class is as follows, wher the real type is the real part of a scalar, i.e., decltype(std::real(scalar(0))).

```
template<typename scalar> class SPOptions {
public:
 SPOptions();
 SPOptions(int argc, char* argv[]);
 /* print statistics?
 void set_verbose(bool verbose);
                                                        bool verbose() const;
  /* maximum iterations in iterative solver
 void set_maxit(int maxit);
                                                        int maxit() const;
                                                                   */
  /* relative residual stopping criterion for iterative solver
 void set_rel_tol(real rel_tol);
                                                        real rel_tol() const;
 /* absolute residual stopping criterion for iterative solver
                                                                   */
 void set_abs_tol(real abs_tol);
                                                        real abs_tol() const;
 /* type of iterative solver to use, see section 4.9
                                                                   */
 void set_Krylov_solver(KrylovSolver s);
                                                        KrylovSolver Krylov_solver() const;
 /* GMRES restart
                                                                   */
 void set_gmres_restart(int m);
                                                        int gmres_restart() const;
 /* type of Gram-Schmidt used in GMRES
                                                                   */
 void set_GramSchmidt_type(GramSchmidtType t);
                                                        GramSchmidtType GramSchmidt_type() const;
 /* nested-dissection code, see section 4.7.2
                                                        ReorderingStrategy reordering_method() const;
 void set_reordering_method(ReorderingStrategy m);
 /* stop nested-dissection when domains are smaller than nd_param */
 void set_nd_param(int nd_param);
                                                        int nd_param() const;
 /* use the internal (undocumented) metis routine METIS_NodeNDP */
                                                        bool use METIS NodeNDP() const;
 void enable METIS NodeNDP();
 /* do not use METIS_NodeNDP, use METIS_NodeND instead
                                                                   */
 void disable_METIS_NodeNDP();
                                                                   */
 /* use METIS_NodeND instead of METIS_NodeDNP
 void enable_METIS_NodeND();
                                                        bool use_METIS_NodeND() const;
 /* do not use METIS_NodeND, use METIS_NodeNDP instead
                                                                   */
 void disable_METIS_NodeND();
 /* build the supernodal tree using the MUMPS SYMQAMD code
 void enable MUMPS SYMQAMD();
                                                        bool use MUMPS SYMQAMD() const;
 /* do not use MUMPS_SYMQAMD, use fundamental supernodes
                                                                    */
 void disable_MUMPS_SYMQAMD();
 /* when using MUMPS_SYMQAMS, enable aggressive amalgamation
 void enable_agg_amalg();
                                                        bool use_agg_amalg() const;
 /* when using MUMPS SYMQAMS, disable aggressive amalgamation
 void disable_agg_amalg();
 /st set the job to be used for static pivoting, see section 4.7.1 */
 void set_mc64job(int job);
                                                        int mc64job() const;
 void set_mc64job(MC64Job job);
  /* not used at the moment
 void enable_assembly_tree_log();
                                                        bool log_assembly_tree() const;
 void disable assembly tree log();
 /* enable HSS compression, see section 5
 void enable_HSS();
                                                        bool use_HSS() const;
 /* disable HSS compression
 void disable_HSS();
  /* set the minimum size of a separator for HSS compression
 void set_HSS_min_sep_size(int s);
                                                        int HSS_min_sep_size() const;
```

```
/* set level to 1 to enable length 2 connections in the separator
  * before computing separator reordering to reduce HSS ranks.
  * Set to to disable length 2 connections.
 void set separator ordering level(int l);
                                                        int separator ordering level() const;
 /* best not to touch this
 void enable indirect sampling();
 void disable indirect sampling();
                                                        bool indirect sampling() const;
                                                        bool replace tiny pivots() const;
 void enable replace tiny pivots();
 void disable_replace_tiny_pivots(;
 /* get the HSS specific options, see section 5.
                                                                    */
 const HSS::HSSOptions<scalar>& HSS options() const;
 HSS::HSSOptions<scalar>& HSS options();
 /* parse the options in argc/argv set in the constructor
 void set from command line();
 /* parse the options in argc/argv
 void set from command line(int argc, char* argv[]);
 /* print out message listing all command line options
 void describe options() const;
};
```

This uses the following (scoped) enumeration for the Gram-Schmidt type used in GMRES:

```
enum class GramSchmidtType {
   CLASSICAL, /*!< Classical Gram-Schmidt is faster, more scalable. */
   MODIFIED /*!< Modified Gram-Schmidt is slower, but stable. */
};</pre>
```

#### 4.10.2 Command Line Options

To get a list of all available options, make sure to pass "int argc, char\* argv[]" when initializing the StrumpackSparseSolver or when calling SPOptions::set\_from\_command\_line and run the application with --help or -h. Some default values listed here are for double precision and might be different when running in single precision.

```
STRUMPACK options:
```

```
--sp_maxit int (default 5000)
--sp_rel_tol real (default 1e-06)
--sp_abs_tol real (default 1e-10)
--sp_Krylov_solver auto|direct|refinement|pgmres|gmres|pbicgstab|bicgstab
--sp gmres restart int (default 30)
--sp GramSchmidt type modified|classical
--sp reordering method natural|metis|scotch|parmetis|ptscotch|rcm|geometric
--sp_nd_param int (default 8)
--sp_enable_METIS_NodeNDP (default true)
--sp disable METIS NodeNDP (default false)
--sp_enable_METIS_NodeND (default false)
--sp disable METIS NodeND (default true)
--sp enable MUMPS SYMQAMD (default false)
--sp_disable_MUMPS_SYMQAMD (default true)
--sp_enable_agg_amalg (default false)
--sp_disable_agg_amalg (default true)
```



Figure 3: Illustration of a Hierarchically Semi-Separable (HSS) matrix. Gray blocks are dense matrices. Off-diagonal blocks, on different levels of the HSS hierarchy, are low-rank. The low-rank factors of off-diagonal blocks of different levels are related.

```
--sp_mc64job 0-5 (default 0)
--sp_enable_hss (default false)
--sp_disable_hss (default true)
--sp_hss_min_sep_size int (default 256)
--sp_separator_ordering_level (default 1)
--sp_enable_indirect_sampling
--sp_disable_indirect_sampling
--sp_enable_replace_tiny_pivots
--sp_disable_replace_tiny_pivots
--sp_verbose or -v (default true)
--sp_quiet or -q (default false)
--help or -h
```

## 5 HSS Preconditioning

(no argument)

--sp enable hss

The sparse multifrontal solver can optionally use Hierarchically Semi-Separable, rank-structured matrices to compress the fill-in. In the multifrontal method, computations are performed on dense matrices called frontal matrices. A frontal matrix can be approximated as an HSS matrix, but this will only be beneficial (compared to storing the frontal as a standard dense matrix and operating on it with BLAS/LAPACK routines) if the frontal matrix is large enough.

Figure 3 illustrates the HSS matrix format. The matrix is partitioned as a  $2 \times 2$  block matrix, with the partitioning recursively applied on the diagonal blocks, until diagonal blocks are smaller than a specified leaf size. The off-diagonal block on each level of the hierarchy are approximated by a low-rank product. This low-rank storage format asymptotically reduces memory usage and floating point operations, while introducing approximation errors. HSS compression is not used by default in the STRUMPACK sparse solver (the default is to perform exact LU factorization), but can be turned on/off via the command line:

```
--sp_disable_hss (no argument)

or via the C++ API as follows

void SPOptions<scalar>::enable_HSS();
void SPOptions<scalar>::disable_HSS();
bool SPOptions<scalar>::use_HSS(); // check whether HSS compression is enabled
```

When HSS compression is enabled, the default STRUMPACK behavior is to use the HSS enabled approximate LU factorization as a preconditioner within GMRES. This behavior can also be changed, see Section 4.9.

However, HSS compression has a considerable overhead and only pays off for sufficiently large matrices. Therefore STRUMPACK has a tuning parameter to specify the minimum size a dense matrix needs to be to be considered a candidate for HSS compression. The minimum dense matrix size for HSS compression is set via the command line via

```
--sp_hss_min_sep_size int (default 256)
or via the C++ API as follows
```

```
void SPOptions<scalar>::set_HSS_min_sep_size(int s);
int SPOptions<scalar>::HSS_min_sep_size() const;  // get the current value
```

The routine set\_HSS\_min\_sep\_size(int s) refers to the size of the top-left block of the front only. This top-left block is the part that corresponds to a separator, as given by the nested dissection reordering algorithm. This top-left block is also referred to as the block containing the fully-summed variable. Factorization (LU in the dense case, ULV in the HSS case) is only applied to this top-left block. Tuning the value for the minimum separator size can have a big impact on performance and memory usage!

The above options affect the use of HSS within the multifrontal solver. There are more, HSS specific, options which are stored in an object of type HSS::HSSOptions<scalar>. An object of this type is stored in the SPOptions<scalar> object stored in the StrumpackSparseSolver. It can be accessed via the HSS\_options() routine as follows:

```
StrumpackSparseSolver<double> sp; // create solver object
sp.options().enable_HSS(); // enable HSS compression in the multifrontal solver
sp.options().HSS_options().set_leaf_size(256); // set the HSS leaf size
```

In STRUMPACK, HSS matrices are constructed using a randomized sampling algorithm [5]. To construct an HSS approximation for a matrix A, sampling of the rows and columns of A is computed by multiplication with a tall and skinny random matrix R as follows:  $S^r = AR$  and  $S^c = A^*R$ . Ideally, the number of columns in the matrix R is d = r + p, with r the maximum off-diagonal block rank in the HSS matrix and p a small oversampling parameter. Unfortunately, the HSS rank is not known a-priori, so it needs to determined adaptively. The adaptive sampling scheme used in STRUMPACK starts with an initial number of random vector  $d_0$ , and increases this in steps of  $\Delta d$ , until the compression quality reaches the desired user specified tolerance, or until the maximum rank is reached. The compression tolerances can greatly impact performance. They can be set using:

```
--hss_rel_tol real (default 0.01)
--hss_abs_tol real (default 1e-08)
```

or via the C++ API

```
void HSSOptions<scalar>::set_rel_tol(real rel_tol);
void HSSOptions<scalar>::set_abs_tol(real abs_tol);
real HSSOptions<scalar>::rel_tol() const; // get the current value
real HSSOptions<scalar>::abs_tol() const;
```

#### 5.1 HSSOptions<scalar> Interface

Other options are available to tune for instance the initial number of random vectors  $d_0$ , the increment  $\Delta d$ , the random number generator or the random number distribution. The complete public interface for the HSSOptions<scalar> class is:

```
void set abs tol(real abs tol);
                                          real abs tol() const;
/* size of the smallest blocks in the HSS hierarchy
void set leaf size(int leaf size);
                                         int leaf size() const;
/* initial number of random vectors used in the
   adaptive randomized compression algorithm
void set d0(int d0);
                                          int d0() const;
/* number of random vectors added in each step of the
   adaptive randomized HSS compression algorithm
void set dd(int dd);
                                          int dd() const;
/* currently not used
void set q(int q);
                                          int q() const;
/* maximum rank in the HSS representation
void set max rank(int max rank);
                                         int max rank() const;
/* random engine/generator to use, see below
void set random engine(random::RandomEngine random engine);
random::RandomEngine random engine() const;
/* the random number distribution, see below
void set random distribution
(random::RandomDistribution random distribution);
random::RandomDistribution random distribution() const;
/* the compression algorithm to use
void set_compression_algorithm(CompressionAlgorithm a);
CompressionAlgorithm compression algorithm() const;
/* for expert users
void set_user_defined_random(bool user_defined_random);
bool user defined random() const;
/* for expert users
void set_synchronized_compression(bool sync);
bool synchronized compression() const;
/* currently not used
void set_log_ranks(bool log_ranks);
                                          bool log_ranks() const;
/* print statistics?
                                                        */
void set_verbose(bool verbose);
                                          bool verbose() const;
/* parse options in argc/argv
void set_from_command_line(int argc, char* argv[]);
/* print description of command line options
void describe_options() const;
```

#### 5.2 HSS Command Line Options

The HSS specific command line options are:

```
HSS Options:
--hss_rel_tol real (default 0.01)
--hss_abs_tol real (default 1e-08)
--hss_leaf_size int (default 128)
--hss_d0 int (default 128)
--hss_dd int (default 32)
--hss_q int (default 0)
--hss_max_rank int (default 5000)
--hss_random_distribution normal|uniform (default normal(0,1))
--hss_random_engine linear|mersenne (default minstd_rand)
--hss_compression_algorithm original|stable (default stable)
--hss_user_defined_random (default false)
```

```
--hss_enable_sync (default true)
--hss_disable_sync (default false)
--hss_log_ranks (default false)
--hss_verbose or -v (default false)
--hss_quiet or -q (default true)
--help or -h
```

## 6 HSS Approximation of Dense Matrices

The HSS code can be found in the src/HSS/ subdirectory. All HSS code is in a namespace called HSS. The class for a sequential/multithreaded HSS matrix is HSSMatrix<scalar>, while the distributed memory HSS class is HSSMatrix<scalar>. For examples of the usage of these classes, see the test code in test/test\_HSS\_seq.cpp and test/test\_HSS\_mpi.cpp respectively. There is also one sequential example in examples/MLkernel.cpp, which uses HSS compression for kernel matrices as used in certain machine learning applications, see for instance [1]. This part of the code will be better documented in future STRUMPACK releases.

## 7 Examples

A number of examples are available in the examples/ folder. This folder, with the sources, is copied to the build directory when running cmake. However, the examples are not part of the CMake build system, so they will not be compiled and linked when running make. A simple Makefile is generated in the examples/ folder in the build directory, with the compiler setting based on what CMake has discovered. However, this Makefile is rather simplistic and in certain cases it might need some manual edits. However, it can serve as an example for how to compile code using STRUMPACK. Check the README file in the examples/ directory for more details.

#### 8 C Interface

The C interface is defined in the header file StrumpackSparseSolver.h and is very similar to the C++ interface. For example usage see the programs sexample.c, dexample.c, cexample.c and zexample.c in the examples/directory, for simple single and double precision real and complex example programs. Note that since the STRUMPACK code is written in C++, even when using the C interface you should link with a C++ aware linker or link with the standard C++ library. For instance when using the GNU toolchain, link with g++ instead of gcc or link with gcc and include -lstdc++.

## 9 Advanced Usage Tips

- To keep track of the number of floating point operations performed in the STRUMPACK Sparse Solver, you can run CMake with -DSTRUMPACK\_COUNT\_FLOPS=ON. Then, when running, do not set the quiet flag in the StrumpackSparseSolver constructor or on the command line and the solver will print some statistics. This will also enable a counter for data movement in the solve phase, from which the (approximately) attained bandwidth usage is derived. This is done because the solve phase is typically bandwidth limited, while the factorization is flop limited.
- There is also some support for PAPI. Run CMake with -DSTRUMPACK\_USE\_PAPI=ON and specify the PAPI include folders and libraries via -DPAPI\_INCLUDES=.. and -DPAPI\_LIBRARIES=...
- We have added timers all throughout the code. These can be enabled with -DSTRUMPACK\_TASK\_TIMERS=ON. Running the code will generate a file time.log. A script to visualize these timings is provided.

- If you compile with MKL or OpenBLAS, you can take advantage of some extra optimized routines by specifying -D HAVE MKL or -D HAVE OPENBLAS respectively.
- The code is not completely thread safe at the moment: do not call solve on the same StrumpackSparseSolve object from different threads simultaneously.
- For comments, feature requests or bug reports: {pghysels,xsli,qichavez}@lbl.gov

## 10 FAQ

 Help, I get this compilation error: catastrophic error: cannot open source file "chrono" #include <chrono>

You need a C++11 capable compiler, and also a C++11 enabled standard library. For instance suppose you are using the Intel 15.0 C++ compiler with GCC 4.4 headers. The Intel 15.0 C++ compiler supports the C++11 standard, but the GCC 4.4 headers do not implement the C++11 standard library. You should install/load a newer GCC version (or just the headers). On cray machines, this can be done with module unload gcc; module load gcc/4.9.3 for instance.

• When running make test, many of the tests fail!

The parallel execution in ctest is invoked by the MPIEXEC command as discovered by CMake. On many HPC clusters, this does not run unless it is executed from within a batch script. In this case all parallel tests will fail. At the moment, a small number of tests still fail. This is normal behavior.

## 11 Acknowledgements

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