Parallel processing and matrix solver interface: mtxp

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Notice

• Version number of the software in the following will change owing to updates of the software.

1 Parallel matrix solver library: PETSc

1.1 Download by git

1. Make a directory (if already /opt directory exists, skip first two lines)

2. Download by git

```
git clone -b release https://gitlab.com/petsc/petsc.git petsc
```

1.2 Configure

1. Create a configure script "gfortran.py" by your favarite editor

gfortran.py

```
#!/usr/bin/env python
# Build PETSc with gfortran
configure_options = [
  '--with-cc=clang',
  '--with-cxx=clang++',
  '--with-fc=gfortran-mp-10',
  '--with-shared-libraries=0',
  '--with-cxx-dialect=C++11',
  '--download-mpich=1',
  '--download-hypre=0',
  '--download-fblaslapack=1',
  '--download-spooles=1',
  '--download-superlu=1',
  '--download-metis=1',
  '--download-parmetis=1',
  '--download-superlu_dist=1',
  '--download-blacs=1',
  '--download-scalapack=1',
  '--download-mumps=1'
  ]
if __name__ == '__main__':
  import sys, os
  sys.path.insert(0,os.path.abspath('config'))
  import configure
  configure.petsc_configure(configure_options)
```

2. Make "gfortran.py" executable

```
chmod 755 gfortran.py
```

3. Setup environment variables for PETSc

```
export PETSC_DIR=/opt/PETSc/petsc
export PETSC_ARCH=gfortran
```

4. Configure

```
./gfortran.py
```

1.3 Compile

1. Compile

make all

2 Compile of task/mtxp module

2.1 Setup make.mtxp file

1. Goto mtxp directory

```
cd task/mtxp
```

2. Create a setup file

```
cp make.mtxp.org make.mtxp
```

- 3. Edit the setup file "make.mtxp"
 - \bullet If MPI is not available, remove comment mark "#" on lines 4–10
 - If MPI is available but PETs not, remove comment mark "#" on lines 13-19
 - If MPI and PETSc are available, remove comment mark "#" on lines 24, 32,33, 35–40

2.2 Modify make header file

1. Go to task directory

```
cd ..
```

- 2. Edit make.header to use lapack and blas libraries for fortran
 - Near the beginning of the file, remove comment and ajust the path

```
LAPACK = lapack.f
LIBLA=-L /opt/PETSc/petsc/gfortran/lib -lflapack -lfblas
```

• In the following, add comment marks

```
#LAPACK = nolapack.f
#LIBLA =
MODLA95 =
```

Keep MODLA95 as it is, since lapack95 is not used.

• Go back to mtxp directory

```
cd mtxp
```

2.3 Compile

1. Compile

```
make clean
```

2.4 Test mtxp

- 1. Test programs solving 1d, 2D and 3D diffusion equation are generated
 - testbnd: Direct band matrix solver (non-parallel)
 - testpcg: Iterative band matrix solver (non-parallel)
 - testdmumps: Direct band matrix solver (parallel using MUMPS)
 - testkdsp: Iterative band matrix solver (parallel using PETSc)

2. Input parameters

- idimen: number of dimension (i or 2 or 3), 0 for quit
- isiz: number of mesh point in one dimension
- isource : source position is all dimensions
- itype: tyoe of initial guess for PETSc 0..5 (default=0)
- m1: type of solver (methodKSP) of PETSc 0..13 (default=4)
- m2: type of preconditioner (methodPC) 0..12 (default=5)
- tolerance: tolerance in iterative method
- 3. Example input For parallel processing

```
mpirun -np 4 ./testdmumps
# INPUT: idimen,isiz,isource,itype,m1,m2,tolerance,idebug=
1,11,6,0,4,5,1.D-7
3/
0/
```

3 Compile of task/fp and related modules

3.1 Update Makefile

1. Change directory

```
cd ../fp
```

- 2. Edit Makefile
 - To use serial band matrix solver, remove comment mark "#" on lines 4-5.
 - To use serial iterative solver, remove comment mark "#" on lines 6-7.
 - To use parallel direct solver MUMPS, remove comment mark "#" on lines 8-9.
 - \bullet To use parallel direct solver library PETSc, remove comment mark "#" on lines 10-11. .

3.2 Compile

1. Compile related modules and task/fp files

make

4 Install on Ubuntu

4.1 Install of required modules

```
sudo apt-get install gfortran-8
sudo apt-get install gcc-8
sudo apt-get install g++-8

sudo apt-get install emacs
sudo apt-get install git
sudo apt-get install xorg-dev
```

```
sudo apt-get install valgrind
sudo apt-get install cmake
sudo apt-get install python
    Install of MPICH
  1. Download of mpich-3.3.1.tar.gz
                                     (See 1.1)
  2. Expand at /soft/mpich
                              (See 1.1)
                                     (See 1.2)
  3. Configure by executing "./run"
         run:
         CC=gcc-8 CFLAGS=''-m64'' CXX=g++-8 CXXFLAGS=''-m64'' FC=gfortran-8
         FFLAGS=''-m64'' ./configure --prefix=/usr/local/mpich331-gfortran-gcc8
         --enable-cxx --enable-fast --enable-romio --disable-shared
  4. Compile and install
                           (See 1.3)
4.3 Install of PETSc
  1. Download of petsc-3.11.3.tar.gz
                                     (See 2.1)
  2. Expand at /opt/PETSc/
                                (See 2.1)
                                  (See 2.2)
  3. Setup environment variables
         export PETSC_DIR=/opt/PETSc/petsc-3.11.3
         export PETSC_ARCH=gfortran
  4. Configure by gfortran.py
                               (See 2.2)
  5. Execute fortran.py
         gortran.py:
         #!/usr/bin/env python
         # Build PETSc, with gfortran
         configure_options = [
           '--with-mpi=1',
           '--with-mpi-dir=/usr/local/mpich331-gfortran-gcc8',
           '--with-shared-libraries=0',
           '--with-cxx-dialect=C++11',
           '--download-mpich=0',
           '--download-hypre=0',
           '--download-fblaslapack=1',
           '--download-spooles=1',
           '--download-superlu=1',
           '--download-metis=1',
           '--download-parmetis=1',
```

'--download-superlu_dist=1',

'--download-blacs=1',

```
'--download-scalapack=1',
'--download-mumps=1'
]

if __name__ == '__main__':
    import sys,os
    sys.path.insert(0,os.path.abspath('config'))
    import configure
    configure.petsc_configure(configure_options)
```