Third day [morning session]

Practice to master numerical libraries

RIKEN CCS HPC Summer School
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Regarding software setup on OBCX

- Set Locale or LANG
  
  ```bash
  % export LANG=C
  ```

- For the third day sessions, you must load appropriate modules, such as impi, petsc and phdf, before make and run.
  
  ```bash
  % module load impi petsc phdf5
  ```

- When you find the message during make as `depend_c.inc:1: *** multiple target patterns. Stop.' please confirm modules, and load missing ones correctly. Erase `depend_c.inc', then do make again.

Currently Loaded Modulefiles:

1) impi/2019.5.281  3) phdf5/1.10.5
2) intel/2019.5.281  4) petsc/3.11.2
Most of source files and hints are on

/work/gt57/t57003/share/

PETSc/ → PETSc sample code, a job-script, and Makefile
mpi-petsc_merged_v6.tgz → tarball for CFD+mpi+PETSc
mpi-petsc_merged_v6/ → Karman Vortex CFD codes

● Please ‘cd’ your work directory, then ‘cp -R’ the directory (‘%’ is a prompt, and do not type). You will see

% cd /work/gt57/tXXXXX
% cp -R ../t57003/share ./third_day
% cd third_day
% ls

NN  PETSc  mpi-petsc_merged_v6  mpi-petsc_merged_v6.tgz
Numerical Library
What is it? For what?
Numerical Library

- Numerical Library is one of building blocks for ENSURING your advanced programming.
- It supports an API for very complex mathematical features, algorithm, schemes, also data handling…
- Solving sys. Eqs, FFT, Eigenvalue calculation, SVD, minimization, statistics, etc…
- There are reference codes.
- They might be examples of good (bad) programming.

* Form C := alpha*A*B + beta*C.
* DO 90 j = 1,n  
  IF (beta.EQ.zero) THEN  
    DO 50 i = 1,m  
      c(i,j) = zero  
    50 CONTINUE  
  ELSE IF (beta.NE.one) THEN  
    DO 60 i = 1,m  
      c(i,j) = beta*c(i,j)  
    60 CONTINUE  
  END IF  
  DO 80 l = 1,k  
    temp = alpha*b(l,j)  
    DO 70 i = 1,m  
      c(i,j) = c(i,j) + temp*a(i,l)  
    70 CONTINUE  
  80 CONTINUE  
90 CONTINUE

http://www.netlib.org/lapack/explore-html/d1/d54/group__double__blas__level3_gaeda3cbd99c8fb834a60a6412878226e1.html
Numerical Library

- Numerical Library is one of building blocks for **ENSURING** your advanced programming.
- It supports an API for very complex mathematical features, algorithm, schemes, also data handling...
  - Solving sys. Eqs, FFT, Eigenvalue calculation, SVD, minimization, statistics, etc...
  - **There are reference codes.**
    - They might be examples of good (bad) programming.
- **It provides us with better performance and finer accuracy than what you made.**
  - Commercial library: faster and more accurate but expensive
  - Open Source library: fast and free (sometimes faster than commercial library)
- You must check them before you run your application codes.

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Example

- **When you HOPE to SPEED UP your code bottlenecked in matmul, use (or link) an appropriate BLAS (Basic Linear Algebra Subprograms) library!**

- Standard API for linear algebra kernels (case of (C)BLAS).
  - **GEMM**: Matrix-matrix multiplication
    \[ C := \alpha AB + \beta C \]
  - **AXPY**: linear combination of 2 Vectors
  - **NRM2**: Norm of a vector, etc.

Reference codes are available from netlib@UTK.
http://www.netlib.org/BLAS/

- Commercial: Intel MKL, AMD ACML (free)
- Open Source: ATLAS(@UTK), GotoBLAS(@TACC), OpenBLAS for general purposed microprocessors
- nVIDIA CUBLAS, AMD clMATH, MAGMABLAS(@UTK), KBLAS(@KAUST), ASPEN.K2(@RIKEN): for GPGPU

```c
for(i=0; i<N; i++)
  for(j=0; j<N; j++) {
    t = 0.0;
    for(k=0; k<N; k++)
      t += a[i][k]*b[k][j];
    c[i][j] = t;
  }

ccblas_dgemm(CblasRowMajor,
              CblasNoTrans, CblasNoTrans,
              N, N, N,
              1.0, a, N, b, N, 0.0, c, N);
```

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When you HOPE to SPEED UP your code bottlenecked in matmul, use (or link) an appropriate BLAS (Basic Linear Algebra Subprograms) library!

- Standard API for linear algebra kernels (case of (C)BLAS).
- GEMM: Matrix-matrix multiplication
- AXPY: linear combination of 2 Vectors (aX+bY)
- NRM2: Norm of a vector, etc.

Reference codes are available from netlib@UTK.
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Example:

```
for(i=0; i<N; i++)
    for(j=0; j<N; j++) {
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            t += a[i][k]*b[k][j];
        c[i][j] = t;
    }
```

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Example

- When you HOPE to SPEED UP your code bottlenecked in matmul, use (or link) an appropriate BLAS (Basic Linear Algebra Subprograms) library!

  - Standard API for linear algebra kernels.
    - GEMM : Matrix-matrix multiplication
      \[ C := \alpha AB + \beta C \]
    - AXPY: linear combination of 2 Vectors
      \[ y := \alpha x + y \]
    - NRM2: Norm of a vector, etc.
      \[ a = \|x\|_2 \]

  Reference codes are available from netlib@UTK.

- Commercial: Intel MKL, AMD ACML (free)
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- nVIDIA CUBLAS, AMD clMATH, MAGMABLAS(@UTK), KBLAS(@KAUST), ASPEN.K2(@RIKEN) : for GPGPU
Other cases

Suggestion: for more complex problems, use followings;

- LAPACK (http://www.netlib.org/lapack/)
- ScaLAPACK (http://www.netlib.org/scalapack/)
- Elemental (http://libelemental.org/)
- EigenExa (http://www.aics.riken.jp/labs/lpnctrtrt/EigenExa_e.html)
- ELPA (http://elpa.rzg.mpg.de/)
- PETSc (http://www.mcs.anl.gov/petsc/)
- Trillions (https://trilinos.org/)
- ARPACK (http://www.caam.rice.edu/software/ARPACK/)
- FFTW (http://www.fftw.org/)
- FFTE (http://www.ffte.jp/)
- 2decomp&FFT (http://www.2decomp.org/)
- MT, MTGP, dSFMT (http://www.math.sci.hiroshima-u.ac.jp/~mat/MT/SFMT/index.html)
- GMP big number librart(https://gmplib.org/)
- QD pack, MPACK, and so on
Review back to the CFD code
Core computational part in the CFD code

\[ \Delta \phi = d \]

\[ \frac{\phi^k_{i,j-1} - 2\phi^k_{i,j} + \phi^k_{i,j+1}}{\Delta y^2} + \frac{\phi^k_{i-1,j} - 2\phi^k_{i,j} + \phi^k_{i+1,j}}{\Delta x^2} - d^k_{i,j} = 0 \]

\[ A \phi = d \]

\[ A = \frac{1}{\Delta x^2} (I_{N_x-1} \otimes X_{N_y-1}) + \frac{1}{\Delta y^2} (X_{N_y-1} \otimes I_{N_x-1}) \]

\[ X_m = \begin{bmatrix} -2 & 1 & 1 \\ 1 & -2 & 1 \\ 1 & -2 & 1 \end{bmatrix}_{m \times m} \]

for(j=1; j<col_m_1; j++)
for(i=2; i<row_m_1 - 1; i++)
  *(at(phiT,i,j)) = const1 * ( (L(phi,i+1,j ) + L(phi,i-1,j )) * const2 +
                             (L(phi,i ,j+1) + L(phi,i ,j-1)) * const3 - L(d,i,j));
Solving Poisson’ eq

- You can write a Jacobi iteration code easily. BUT,
  - wasting your time (programming, execution time).
  - Poisson’s eq has a mathematical difficulty of convergence (sometimes diverges).
- Many algorithms are implemented as library codes.

You can choose Faster and reliable library routine to Solve Poisson’eq.

Today we try to use PETSc library

Krylov Subspace method GMRES, CG, GCR, etc.. with Preconditioners
Solving Poisson’ eq

- You can write a Jacobi iteration code easily. BUT,
  - wasting your time (programming, execution time).
  - Poisson’s eq has a mathematical difficulty of convergence (sometimes diverges).
- Many algorithms are implemented as library codes.
  - **Jacobi iteration**

\[ x^{n+1} = D^{-1}((A - D)x^n) - b \]

Convergence condition is very strict and the Jacobi method sometimes (or often?) diverges or stagnates. (diagonal dominant property is well known as a convergent condition).

Similar algorithms:
- Gauss-Seidel, SOR, SSOR, etc.

You can choose Faster and reliable library routine to Solve Poisson’eq.

Today we try to use PETSc library

Krylov Subspace method
- GMRES, CG, GCR, etc.
- with Preconditioners
Solving Poisson’ eq

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● Many algorithms are implemented as library codes.

• Jacobi iteration

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Convergence condition is very strict and the Jacobi method sometimes (or often?) diverges or stagnates. (diagonal dominant property is well known as a convergent condition).

Similar algorithms:
Gauss-Seidel, SOR, SSOR, etc.

• GMRES

\[ \text{argmin}_y \{ ||AVy - b|| | V \in \mathcal{K}^n_A(v_0) \} \]

Based on Krylov subspace iteration and Arnoldi procedure, it finds a local solution vector by using the Least square approximation within a spanned subspace.

Generally, GMRES is known as best method for a non-symmetric case.
Solving Poisson’ eq

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Krylov Subspace method GMRES, CG, GCR, etc.. with Preconditioners
General use of PETSc(SLEPC)

Official site on PETSc and SLEPC

Hands-on exercises
PETSc/TAO

- Developed by Argonne National Lab. USA
- Portable, Extensible Toolkit for Scientific Computation
- Toolkit for Advanced Optimization

https://www.mcs.anl.gov/petsc/

The current version of PETSc is 3.13; released March 29, 2020.

PETSc, pronounced PET-see (the S is silent), is a suite of data structures and routines for the scalable (parallel) solution of scientific applications modeled by partial differential equations. It supports MPI, and GPUs through CUDA or OpenCL, as well as hybrid MPI-GPU parallelism. PETSc (sometimes called PETSc/Tao) also contains the Tao optimization software library.

(cf. PETSc/TAO homepage)
SLEPc

- Significant plugin modules of PETSc for EVP
- developed by Universitat Politecnica de Valencia, Spain

https://slepc.upv.es/

SLEPc is a software library for the solution of large scale sparse eigenvalue problems on parallel computers. It is an extension of PETSc and can be used for linear eigenvalue problems in either standard or generalized form, with real or complex arithmetic. It can also be used for computing a partial SVD of a large, sparse, rectangular matrix, and to solve nonlinear eigenvalue problems (polynomial or general). Additionally, SLEPc provides solvers for the computation of the action of a matrix function on a vector.

The current version of SLEPc is 3.13, released in March, 2020.
Access to a Sample code

https://slepc.upv.es/handson/handson1.html

Hyperlinked URL is not available. Please see ex1.c on the directory ‘PETSc’.

Other tutorial examples are on /work/gt57/t57003/slepc or /work/gt57/t57003/petsc
Compile and link

Makefile

ARCH=
SLEPC_DIR=/work/gt57/t57003/slepc/

CC=mpiicc
FC=mpiifort
BLAS=

CCFLAGS_PETSC=-std=c11 -qopenmp -Wall -O3 -g -I$(PETSC_DIR)/include
LDFLAGS_PETSC=-std=c11 -qopenmp -L$(PETSC_DIR)/lib -lpetsc -lhdf5_hl -L$(MKLROOT)/lib/intel64 -lmkl_rt
CCFLAGS_SLEPC=-std=c11 -qopenmp -Wall -O3 -g -I$(SLEPC_DIR)/include
LDFLAGS_SLEPC=-std=c11 -qopenmp -L$(PETSC_DIR)/lib -lpetsc -L$(SLEPC_DIR)/lib -lslepc -lhdf5_hl -L$(MKLROOT)/lib/intel64 -lmkl_rt

all: eps_exec
eps_exec: ex1.o
    $(CC) $(CCFLAGS_SLEPC) -o $@ $< $(LDFLAGS_SLEPC)
ex1.o: ex1.c
    $(CC) $(CCFLAGS_SLEPC) -c $<
clean:
    ¥rm *.o eps_exec

Use `make` command

→ → →

eps_exec is generated.
#!/bin/bash
#PJM -L rscgrp=lecture7
#PJM -L node=1
#PJM --mpi proc=4
#PJM -L elapse=0:10:00
#PJM -g gt57
#PJM -N slepc
#PJM -j

module load intel impi petsc phdf5

export SLEPC_DIR=/work/gt57/t57003/slepc/
export LD_LIBRARY_PATH=$SLEPC_DIR/lib:$LD_LIBRARY_PATH

mpiexec.hydra -n ${PJM_MPI_PROC} ./eps_exec -n 540 -m 180 -eps_type gd -eps_nev 4 -eps_monitor -eps_view
C version

1-D Laplacian Eigenproblem, n=100

Number of iterations of the method: 19
Solution method: krylovschur

Number of requested eigenvalues: 1
Stopping condition: tol=1e-08, maxit=100
Number of converged eigenpairs: 2

| k         | ||Ax-kx||/||kx|| |
|------------|------------------|
| 3.999033   | 4.02784e-09      |
| 3.996131   | 4.31174e-09      |

F90 version

1-D Laplacian Eigenproblem, n =100 (Fortran)

Number of iterations of the method: 19
Solution method: krylovschur
Number of requested eigenvalues: 1
Stopping condition: tol=1.0000E-08, maxit= 100
Number of converged eigenpairs: 2

| k         | ||Ax-kx||/||kx|| |
|------------|------------------|
| 3.9990E+00| 4.0278E-09       |
| 3.9961E+00| 4.3117E-09       |
Play with PETSc/SLEPC

- From the tutorial page,

```
$ ./eps_exec -n 400 -eps_nev 3 -eps_tol 1e-7

$ ./eps_exec -n 400 -eps_nev 3 -eps_ncv 24

$ ./eps_exec -n 100 -eps_nev 4 -eps_type lanczos
```

1-D Laplacian Eigenproblem, n=400

Number of iterations of the method: 100
Solution method: krylovshur

Number of requested eigenvalues: 3
Stopping condition: tol=1e-07, maxit=100
Number of converged eigenpairs: 1

| k          | ||Ax-kx|| / ||kx|| |
|------------|-------------------|
| 3.999939   | 9.48781e-08       |

1-D Laplacian Eigenproblem, n=100

Number of iterations of the method: 62
Solution method: lanczos

Number of requested eigenvalues: 4
Stopping condition: tol=1e-08, maxit=100
Number of converged eigenpairs: 4

| k          | ||Ax-kx|| / ||kx|| |
|------------|-------------------|
| 3.999939   | 9.48494e-09       |
| 3.99754    | 1.97435e-09       |
| 3.99448    | 9.15231e-09       |
| 3.99018    | 3.55339e-09       |

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Learn the sample code (Pseudo code)

SlepcInitialize( PETSC_NULL_CHARACTER, ierr )

MatCreate( PETSC_COMM_WORLD, A, ierr )
MatSetSizes( A, ..., n, n, ierr )
MatSetUp( A, ierr )

(.... Calculation of matrix elements and others .....)

ESPCreate( PETSC_COMM_WORLD, eps, ierr )
ESPSetOperators( eps, A, PETSC_NULL_OBJECT, ierr )
EPSSetProblemType( eps, EPS_HEP, ierr )

EPSSolve( eps, ierr )

EPSGetEigenPair( eps, ...... )

EPSDestroy( eps, ierr )

SlepcFinalize( ierr )

Modern style

1. Initialization
2. Create a Handle for data
3. Setup parameters
4. Create a Handle for solvers
5. Setup parameters
6. Kick the solver
7. Retrieve the results
8. Destroy handles
9. Finalize
How to setup a matrix? (in C)

- PETSc handles internal data format and interface data flexibly. Because of PETSc management mechanism, user does not to see actual state on memory. Matrix A is dealt with a handler variable, and matrix elements are accessed via a query API.

```c
// Simple matrix format
Mat A;
EPS eps;
EPSType tname;
PetscReal tol, error, *values;

MatCreate( PETSC_COMM_WORLD, &A )
MatSetSizes( A, PETSC_DECIDE, PETSC_DECIDE, M, N )
MatGetOwnershipRange(A, &Istart, &Iend )
MatSetValues( A, m, idxm, n, idxn, values, INSERT_VALUES|ADD_VALUES )

MatAssemblyBegin( A, MAT_FINAL_ASSEMBLY )
MatAssemblyEnd( A, MAT_FINAL_ASSEMBLY )
```
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For a mxn block matrix, array values are set.
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MatSetValue( A, m, idxm, n, idxn, values, INSERT_VALUES|ADD_VALUES )
MatAssemblyBegin( A, MAT_FINAL_ASSEMBLY )
MatAssemblyEnd( A, MAT_FINAL_ASSEMBLY )
```

Assemble the matrix data set on a distributed manner
How to setup a matrix? (in fortran90)

- PETSc handles internal data format and interface data flexibly. Because of PETSc management mechanism, user does not to see actual state on memory. Matrix A is dealt with a handler variable, and matrix elements are accessed via a query API.

```fortran
! Simple matrix format
Mat     A
EPS     eps
EPSType tname
PetscReal tol, error, values(:)

call MatCreate( PETSC_COMM_WORLD, A, ierr )
call MatSetSizes( A, PETSC_DECIDE, PETSC_DECIDE, M, N, ierr )

call MatGetOwnershipRange(A, Istart, Iend, ierr )
call MatSetValue( A, m, idxm, n, idxn, values, INSERT_VALUES|ADD_VALUES, ierr)

call MatAssemblyBegin( A, MAT_FINAL_ASSEMBLY, ierr)
call MatAssemblyEnd( A, MAT_FINAL_ASSEMBLY, ierr)
```
How to setup a matrix? (in fortran90)

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```fortran
! Simple matrix format
Mat    A
EPS    eps
EPSType tname
PetscReal tol, error, values(:)

call MatCreate( PETSC_COMM_WORLD, A, ierr )
call MatSetSizes( A, PETSC_DECIDE, PETSC_DECIDE, M, N, ierr )

call MatGetOwnershipRange(A, Istart, Iend, ierr )
call MatSetValues( A, m, idxm, n, idxn, values, INSERT_VALUES|ADD_VALUES, ierr)

call MatAssemblyBegin( A, MAT_FINAL_ASSEMBLY, ierr)
call MatAssemblyEnd( A, MAT_FINAL_ASSEMBLY, ierr)
```
How to setup a matrix? (in fortran90)

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```fortran
! Simple matrix format
Mat        A
EPS        eps
EPSType    tname
PetscReal  tol, error, values(:)

call MatCreate( PETSC_COMM_WORLD, A, ierr )
call MatSetSizes( A, PETSC_DECIDE, PETSC_DECIDE, M, N, ierr )
call MatGetOwnershipRange(A, Istart, Iend, ierr )
call MatSetValues(  A, m, idxm, n, idxn, values, INSERT_VALUES|ADD_VALUES, ierr)
call MatAssemblyBegin( A, MAT_FINAL_ASSEMBLY, ierr)
call MatAssemblyEnd( A, MAT_FINAL_ASSEMBLY, ierr)
```

Matrix size MxN
How to setup a matrix? (in fortran90)

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```fortran
! Simple matrix format
Mat A
EPS eps
EPSType tname
PetscReal tol, error, values(:)

call MatCreate( PETSC_COMM_WORLD, A, ierr )
call MatSetSizes( A, PETSC_DECIDE, PETSC_DECIDE, M, N, ierr )
call MatGetOwnershipRange(A, Istart, Iend, ierr )
call MatSetValues(  A, m, idxm, n, idxn, values, INSERT_VALUES|ADD_VALUES, ierr)
call MatAssemblyBegin( A, MAT_FINAL_ASSEMBLY, ierr)
call MatAssemblyEnd( A, MAT_FINAL_ASSEMBLY, ierr)
```

For a mxn block matrix, array values are set.
How to setup a matrix? (in fortran90)

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```fortran
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Mat A
EPS eps
EPSType tname
PetscReal tol, error, values(:)

call MatCreate( PETSC_COMM_WORLD, A, ierr )
call MatSetSizes( A, PETSC_DECIDE, PETSC_DECIDE, M, N, ierr )
call MatGetOwnershipRange(A, Istart, Iend, ierr )
call MatSetValues( A, m, idxm, n, idxn, values, INSERT_VALUES|ADD_VALUES, ierr)
call MatAssemblyBegin( A, MAT_FINAL_ASSEMBLY, ierr)
call MatAssemblyEnd( A, MAT_FINAL_ASSEMBLY, ierr)
```

Assemble the matrix data set on a distributed manner.

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Play back the CFD code and link it with PETSc
Core computational part in the CFD code

● Solving a Poisson equation

\[ \Delta \phi = d \]

● Discretization:

\[
\begin{align*}
\phi_{i-1,j}^{k+1} & - 2\phi_{i,j}^{k+1} + \phi_{i+1,j}^{k +} \\
\phi_{i,j-1}^{k+1} & - 2\phi_{i,j}^{k+1} + \phi_{i,j+1}^{k +} \\
\Delta x^2 & \\
\Delta y^2 &
\end{align*}
\]

\[
\Delta \phi_{i,j} = 0
\]

● Finally, Core iteration: it is a well-know iteration

for(j=1; j<col_m_1; j++)
for(i=2; i<row_m_1 - 1; i++)
*(at(phiT,i,j)) = const1 * ( (L(phi,i+1,j ) + L(phi,i-1,j )) * const2 + (L(phi,i ,j+1) + L(phi,i ,j-1)) * const3 - L(d,i,j));
Use a PETSc KSP solver

- **What I did were,**
  - Download a sample source code
  - Modify main.cpp and cfd.cpp
  - Key points are i) initialization of PETSc environment and matrix generation, ii) calling a solver routine.

PETSc supports DMDA (distributed multidimensional array format) for vector data and a stencil mechanism for matrix representation.

- I Prepared 3 call-back functions, and 1 utility function;
  - extern PetscErrorCode ComputeStencil (KSP, Mat, Mat, void *);
  - extern PetscErrorCode ComputeRHS (KSP, Vec, void *);
  - extern PetscErrorCode SetInitialGuess (KSP, Vec, void *);
  - extern PetscErrorCode GetComputedResults (KSP, void *);

- Then, associated them with a DM handler.
  - KSPSolve() organized RHS and MatVec, etc.
Use a PETSc KSP solver

What I did were,

- Download a sample source code
- Modify main.cpp and cfd.cpp
- Key points are i) initialization of PETSc environment and matrix generation, ii) calling a solver routine.

**PETSc supports DMDA (distributed multidimensional array format)** for vector data and a stencil mechanism for matrix representation.

- I Prepared 3 call-back functions, and 1 utility function;
  - extern PetscErrorCode ComputeStencil (KSP, Mat, Mat, void *);
  - extern PetscErrorCode ComputeRHS (KSP, Vec, void *);
  - extern PetscErrorCode SetInitialGuess (KSP, Vec, void *);
  - extern PetscErrorCode GetComputedResults (KSP, void *);

- Then, associated them with a DM handler.
  - KSPSolve() organized RHS and MatVec, etc.
Use a PETSc KSP solver

- **What I did were,**
  - Download a sample source code
  - Modify main.cpp and cfd.cpp
  - Key points are i) initialization of PETSc environment and matrix generation, ii) calling a solver routine.

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   - Translate above relation into a Matrix-vector product.

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* Key functions *

1. MatSetStencil( Mat m, int dim, int dims[], int starts[], int dof)
   It sets the grid information for setting values into a matrix

2. KSPSetComputeOperators(KSP ksp, (*func)(), void *ctx)
   It set a Stencil callback routine to contexts such as DM/user-defined handler

   - Retrieve the result computed by using PETS.
ComputeStencil

ComputeStencil (KSP ksp, Mat A, Mat Aac, void *ctx)
{
    DM da;
    int i, j, M, N, xm, ym, xs, ys;
    grid2D *g = (grid2D *) ctx;

    PetscFunctionBeginUser;

    KSPGetDM (ksp, &da);
    DMDAGetInfo (da, 0, &M, &N, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0);
    DMDAGetCorners (da, &xs, &ys, 0, &xm, &ym, 0);

    const PetscScalar const2 = 1.0 / DX2;
    const PetscScalar const3 = 1.0 / DY2;
    const PetscScalar inv_const1 = 2 * (const2 + const3);

    for (j = ys; j < ys + ym; j++) {
        for (i = xs; i < xs + xm; i++) {
            PetscScalar v[5];
            MatStencil global_index, local_indices[5];

            global_index.i = i;
            global_index.j = j;

            ComputeStencil • Declaration of a work-array for the 5-point stencil
            • Obtain DA and DMDA infos
            • Get the global domain info, corresponding to the local process

            N
            M
            (xs, ys)
            xm
            ym
ComputeStencil

if (i <= 1 || j <= 0 || i >= M - 2 || j >= N - 1) {

/* Regular Boundary : not changed */

v[0] = 1;
local_indices[0].i = i;
local_indices[0].j = j;
MatSetValuesStencil (Aac, 1, &global_index, 1,
local_indices, v, INSERT_VALUES);
}
else {

int is_object = 0;

int ctr_obj;
for( ctr_obj=0; ctr_obj<g->objs.n; ctr_obj++) {

/* Boundary of the object */

const int obj_x = (int) g->objs.obj[ctr_obj].x;
const int obj_y = (int) g->objs.obj[ctr_obj].y;
const int obj_w = (int) g->objs.obj[ctr_obj].w;
const int obj_h = (int) g->objs.obj[ctr_obj].h;

• Compute the object region in global grid.
• (obj_x, obj_y) : Center point
• (obj_w, obj_h): Shape, width/height

Boundary grid points

For the inlet/outlet flow, both left/right wall is bounded (i.e. i=0,1 and M-2,M-1)
ComputeStencil

const int sta_i = (int) (obj_x - obj_w / 2);
const int end_i = (int) (sta_i + obj_w);
const int sta_j = (int) (obj_y - obj_h / 2);
const int end_j = (int) (sta_j + obj_h);

if (i >= sta_i && i <= end_i && j >= sta_j && j <= end_j) {
    is_object = 1;

    int num, numi, numj;
    num = numi = numj = 0;

    if (i == sta_i) {
        v[num] = -DY;
        local_indices[num].i = i - 1;
        local_indices[num].j = j;
        num++;
        numi++;
    }
    else if (i == end_i) {
        v[num] = -DY;
        local_indices[num].i = i + 1;
        }
local_indices[num].j = j;
num++;
umi++;
}
else {
    if (j == sta_j) {
        v[num] = -DX;
        local_indices[num].i = i;
        local_indices[num].j = j - 1;
        num++;
        numj++;
    }
    else if (j == end_j) {
        v[num] = -DX;
        local_indices[num].i = i;
        local_indices[num].j = j + 1;
        num++;
        numj++;
    }
}

• Check whether on the object wall, if so, setup special boundary condition according to the numerical scheme.
ComputeStencil

if (num > 0) {       // surface of the object
    v[num] = (numj * DX + numi * DY);
}
else {               // object internal
    v[0] = 1;
}
local_indices[num].i = i;
local_indices[num].j = j;
num++;
MatSetValuesStencil (Aac, 1,
    &global_index,
    num, local_indices, v, INSERT_VALUES);

• Set up the coefficient of the central point in the 5-point stencil.
• If the point is object internal, output is invariant to the input.
ComputeStencil

if (!is_object) {

    v[0] = const3;
    local_indices[0].i = i;
    local_indices[0].j = j - 1;
    v[1] = const2;
    local_indices[1].i = i - 1;
    local_indices[1].j = j;
    v[2] = -inv_const1;
    local_indices[2].i = i;
    local_indices[2].j = j;
    v[3] = const2;
    local_indices[3].i = i + 1;
    local_indices[3].j = j;
    v[4] = const3;
    local_indices[4].i = i;
    local_indices[4].j = j + 1;
    MatSetValuesStencil (Aac, 1,
        &global_index, 5,
        local_indices, v, INSERT_VALUES);
}

• Set up the coefficient of the central point in the 5-point stencil on the Fluid region.
• Do not forget to finalize the assembling the matrix data elements.
Use a PETSc KSP solver

1. **ComputeStencil**
   - Jacobi iteration is written in a 5-point Stencil fashion.
   - Any stencil codes are

2. **ComputeRHS**
   - Update the vector which appears in Right hand side

3. **SetInitialGuess**
   - Set up an initial guess for Krylov iterative solver

4. **GetComputedResult.**
   - Retrieve the result computed by using PETS.

* Key functions *

1. **DMDAVecGetArray( DM da, Vec vec, void *array)**
   - It returns a pointer (an underlying vector) with global-index

2. **KSPSetComputeRHS (KSP ksp, (*func)(), void *ctx)**
   - It sets an RHS callback routine and a user-defined context
ComputeRHS

ComputeRHS (KSP ksp, Vec b, void *ctx)
{
    DM da;
    int i, j, M, N, xm, ym, xs, ys;
    grid2D *g = (grid2D *) ctx;

    PetscFunctionBeginUser;

    array2D *phi = &(g->phi);
    array2D *d = &(g->d);
    array2D *u = &(g->u);
    array2D *v = &(g->v);

    KSPGetDM (ksp, &da);
    DMDAGetInfo (da, 0, &M, &N, 0, 0, 0, 0, 0, 0, 0, 0, 0);
    DMDAGetCorners (da, &xs, &ys, 0, &xm, &ym, 0);
    PetscScalar **array;
    DMDAVecGetArray (da, b, &array);

    • Ctx is a user-defined context handle, which contains any info to calculate matrix and vectors.
    • Here, the pointer to the grid2D structured data set was assigned.

    • DMDAVecGetArray() returns a multiple dimension array that shares data with the underlying vector and is indexed using the global dimensions. (see. PETSc DMDA manual)
#pragma omp parallel for private(i)
for (j = ys; j < ys + ym; j++) {
    for (i = xs; i < xs + xm; i++) {
        double tmp = 0.0;
        if (i <= 1 || j <= 1 || i >= M - 2 || j >= N - 2) {
            /* Boundary wall 2-point layer */
            tmp = L(phi, i-xs, j-ys);
        } else {
            int is_object = 0;

            int ctr_obj;
            for( ctr_obj=0; ctr_obj<g->objs.n; ctr_obj++ ) {
                const int obj_x = (int) g->objs.obj[ctr_obj].x;
                const int obj_y = (int) g->objs.obj[ctr_obj].y;
                const int obj_w = (int) g->objs.obj[ctr_obj].w;
                const int obj_h = (int) g->objs.obj[ctr_obj].h;

                ComputeRHS • Set the ctr_obj-th object information
            }
        }
    }
}
const int sta_i = (int) (obj_x - obj_w / 2);
const int end_i = (int) (sta_i + obj_w);
const int sta_j = (int) (obj_y - obj_h / 2);
const int end_j = (int) (sta_j + obj_h);

// Boundary of the object
if (i >= sta_i && i <= end_i && j >= sta_j && j <= end_j) {
    is_object = 1;
}

// in case surface of the object
int num = 0;
if (i == sta_i) { num++;
    tmp += DY * (2 * NU / DX) * L(u, i-xs - 1, j-ys);
}
else if (i == end_i) { num++;
    tmp += DY * (2 * NU / DX) * L(u, i-xs + 1, j-ys);
}
else {
    if (j == sta_j) { num++;
        tmp += DX * (2 * NU / DY) * L(v, i-xs, j-ys - 1);
    }

• Check Region of the object

• Check Boundary of the object
ComputeRHS

```
else {
    if ( j == sta_j ) { num++;
        tmp += DX * (2 * NU / DY) * L (v, i-xs, j-ys - 1);
    }
    else if ( j == end_j ) { num++;
        tmp += DX * (2 * NU / DY) * L (v, i-xs, j-ys + 1);
    }
}
// in case inside of the object
    if ( num == 0 ) { tmp = -150.; }
}
}
if ( ! is_object ) {
    tmp = L (d, i-xs, j-ys);
}
array[j][i] = tmp;
}
```
ComputeRHS

DMDAVecRestoreArray (da, b, &array);
VecAssemblyBegin (b);
VecAssemblyEnd (b);

PetscFunctionReturn (0);

• Regular points in the Fluid domain

• Restores a multiple dimension array obtained with DMDAVecGetArray().

• Do not forget to finalize the assembling the vector data elements.
Use a PETSc KSP solver

1. **ComputeStencil**
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     \[
     u_{i,j}^{\text{new}} := au_{i-1,j} + bu_{i,j-1} + cu_{i,j} + du_{i+1,j} + eu_{i,j+1} + f_{i,j}
     \]
   - Translate above relation into a Matrix-vector product.

2. **ComputeRHS**
   - Update the vector which appears in Right hand side

3. **SetInitialGuess**
   - Set up an initial guess for Krylov iterative solver

4. **GetComputedResult.**
   - Retrieve the result computed by using PETS.
SetInitialGuess

SetInitialGuess (KSP ksp, Vec b, void *ctx)
{
    DM da;
    int i, j, xm, ym, xs, ys;
    grid2D *g = (grid2D *) ctx;
    PetscFunctionBeginUser;
    KSPGetDM (ksp, &da);
    DMDAGetCorners (da, &xs, &ys, 0, &xm, &ym, 0);
    double **array;
    DMDAVecGetArray (da, b, &array);
    array2D *phi = &(g->phi);
    #pragma omp parallel for private(i)
    for (j = ys; j < ys+ym; j++) {
        for (i = xs; i < xs+xm; i++) {
            array[j][i] = (L (phi, i-xs-1, j-ys) + L (phi, i-xs, j-ys-1) +
                           L (phi, i-xs+1, j-ys) + L (phi, i-xs, j-ys+1))/4;
        }
    }
    DMDAVecRestoreArray (da, b, &array);

    VecAssemblyBegin (b);
    VecAssemblyEnd (b);
    PetscFunctionReturn (0);
}

- For the initial guess for KSP, just use a mean value of four neighboring points. (similar to the Jacobi solver)
- Do not forget to finalize the assembling the vector data elements.
Use a PETSc KSP solver

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   \[ u_{i,j}^{\text{new}} := a u_{i-1,j} + b u_{i,j-1} + c u_{i,j} + d u_{i+1,j} + e u_{i,j+1} + f_{i,j} \]
   
   - Translate above relation into a Matrix-vector product.

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GetComputedResult

GetComputedResults (KSP ksp, void *ctx)
{
    DM da;
    int i, j, xm, ym, xs, ys;
    grid2D *g = (grid2D *) ctx;
    PetscFunctionBeginUser;

    KSPGetDM (ksp, &da);
    DMDAGetCorners (da, &xs, &ys, 0, &xm, &ym, 0);
    Vec b;
    KSPGetSolution (ksp, &b);
    double **array;
    DMDAVecGetArray (da, b, &array);

    #pragma omp parallel for private(i)
    for (j = ys; j < ys+ym; j++) {
        for (i = xs; i < xs+xm; i++) {
            *(at (phi, i-xs, j-ys)) = array[j][i];
        }
    }
    PetscFunctionReturn (0);
}

- Obtain the corresponding data, while DMDA holds Halo (overlap region).
- Also, needs to reshape the data structure.
Use a PETSc KSP solver

1. **ComputeStencil**
   - Jacobi iteration is written in a 5-point Stencil fashion. Any stencil codes are
     \[
     u_{i,j}^{\text{new}} := a u_{i-1,j} + b u_{i,j-1} + c u_{i,j} + d u_{i+1,j} + e u_{i,j+1} + f_{i,j}
     \]
   - Translate above relation into a Matrix-vector product.

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(PETSc) KSP solver and CFD code

Once KSP starts up, user can access the data within PETSc rule, or the backdoor via the application context Ctx pointer.

PoissonSolver

PETSc: KSP solver

Setup the Matrix on a stencil
Setup the RHS vector, and initial guess
KSP main iteration
Obtain the result vector

CFD main code

Application context

Callback handler functions

ComputeRHS()
SetInitialGuess()
ComputeStencil()
GetComputedResult()
Integrated or registration part for the call-back functions

KSP ksp;  // global variable, sorry it is not a good manner
.
void grid2D_initialize( … )
{
    ...
    KSPCreate (PETSC_COMM_WORLD, &ksp);
    DM da;
    DMDACreate2d (PETSC_COMM_WORLD,
        DM_BOUNDARY_NONE, DM_BOUNDARY_NONE,
        DMDA_STENCIL_STAR, row, col, Px, Py, 1, 1,
        NULL, NULL, &da);
    DMSsetUp (da);
    KSPSetDM (ksp, (DM) da);
    DMDSetApplicationContext (da, (void *) g);
    KSPSetComputeInitialGuess (ksp, SetInitialGuess, (void *) g);
    KSPSetComputeRHS (ksp, ComputeRHS, (void *) g);
    KSPSetComputeOperators (ksp, ComputeStencil, (void *) g);

    ...
}

• These are super-important functions!
Let’s Run the CFD code on OBCX

You already copy mpi-petsc_merged_v6.tgz from /work/gt57/t57003/share/

% tar zxvf mpi-petsc_merged_v6.tgz
% cd mpi-petsc_merged_v6
% make

Examples of job scripts are stored in scripts/.
If you want to run larger jobs, modify NX and NY larger (aspect must be 3:1), and make DT defined in cfd.h smaller.
Do, make clean, then again, make.

NOTE: We are sharing a queuing system. So, please specify the CPU time limit less than 5mins, and do not submit a lot of jobs at the same time!

https://www.cc.u-tokyo.ac.jp/supercomputer/images/Oakbridge-CX.jpg
Example of Problem Settings (cfd.h)

● For a large test
#define NX (360)
#define NY (120)
#define DT (5e-5)
#define NU (0.01)
#define END_TIMESTEP (10000)
#define SAVE_INTERVAL (200)

● For a huge case
#define NX (540)
#define NY (180)
#define DT (2.5e-5)
#define NU (0.01)
#define END_TIMESTEP (20000)
#define SAVE_INTERVAL (400)

● For a Challenging case
#define NX (2160)
#define NY (720)
#define DT (2.5e-6)
#define NU (0.01)
#define END_TIMESTEP (4000*50)
#define SAVE_INTERVAL (4000)

Challenging case needs 8nodes and 15min elapsed time

25, September 2020 RIKEN CCS HPC Summer School
Large test

1. Edit cfd.h (available _LARGE_ and disable others)
   
   #define TEST_CASE _LARGE_

2. Compile by ‘make’.
3. Prepare files required to a job
   - Make a work directory ‘run_large’ and change directory to it.
     
     % mkdir run_large
     % cd run_large

   - Then, copy ../script/run_large.sh and the executable ../solver_fractional on the directory.
     
     % cp ../script/run_large.sh .
     % cp ../solver_fractional .
Large test

4. Submit a job script
   
   `% pjsub run_large.sh`

5. Find computed results on the directory.
   ● Data files are stored on the directory, namely,
   ● `AVEse_****.dat`

6. Copy `../view-x.py`
   ● Follow the next pages
Visualize a result

- Use a python script

Now you are on the work directory.

% python view-new.py
view-x.py

- If you run X-window server on your local PC,
  - Use view-x.py as it is.

- If your network connection is bad or not using X server on your local PC,
  - Please edit two lines in view-x.py as

    ```python
    #plt.show()
    anim.save("plot-z-4.mp4", fps=5)
    ```

You will get plot-z-4.mp4 after python run, then download it on your local PC.

Viewer on your PC depends on your environment. Probably just double-clicking the file will play the animation.
Hands-on time

Please access `/work/gt57/t57003/share/` if have not yet copied files.

PETSc/ → PETSc sample code, a job-script, and Makefile

`mpi-petsc_merged_v6.tgz` → Karman Vortex CFD codes
Practices

1. Change the shape of obstacle or put some obstacles in a fluid region
   1. One square block is already put in the original CFD domain.
   2. Please modify cfd.h, and main(), consistently.
   3. Ghost grid points inside or at the boundary of the obstacle objects must be introduced.
   4. Visualize as the same way to display the original.
   5. Confirm Karman vortices.

[Cautions]
The objects must have no intersection are.
If the objects are smaller than grid width or almost touches the wall, the CFL condition worsens, then CFD code might diverge. Carefully setup the boundary, and tweak delta-t as a smaller value.
2. **Use other iterative algorithms**
   1. Current PETSc solver uses a default KSP method
   2. You can select other algorithms by a command-line or add PetscErrorCode KSPSetType(KSP ksp, KSPTYPE type)
   3. Please refer to the list of KSPTYPE
      [https://www.mcs.anl.gov/petsc/petsc-current/docs/manualpages/KSP/KSPTYPE.html#KSPTYPE](https://www.mcs.anl.gov/petsc/petsc-current/docs/manualpages/KSP/KSPTYPE.html#KSPTYPE)
   4. Confirm the convergent histories.