

# PETSc for Parallel FEM

**RIKEN CCS HPC Summer School**  
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# Sample files

Please access [/work/gt18/share/1D-PETSc](#)

```
$ cd /work/gt18/<YID>/
```

```
$ cp -r /work/gt18/share/1D-PETSc .
```

```
$ cd 1D-PETSc
```

```
$ ls
```

```
1d2-petsc.c    ex1.c    input.dat  run_1d2.sh
```

```
1d2-petscf.F90 ex1f.F90 Makefile  run_ex1.sh
```

# 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.

```

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

```

[http://www.netlib.org/lapack/explore-html/d1/d54/group\\_\\_double\\_\\_blas\\_\\_level3\\_gaeda3cbd99c8fb834a60a64128782226e1.html](http://www.netlib.org/lapack/explore-html/d1/d54/group__double__blas__level3_gaeda3cbd99c8fb834a60a64128782226e1.html)

# Numerical Library

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- **It supports an API for very complex mathematical features, algorithm, schemes, also data handling...**
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- **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.

# 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. ( )

```

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;
  }
  
```



```

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

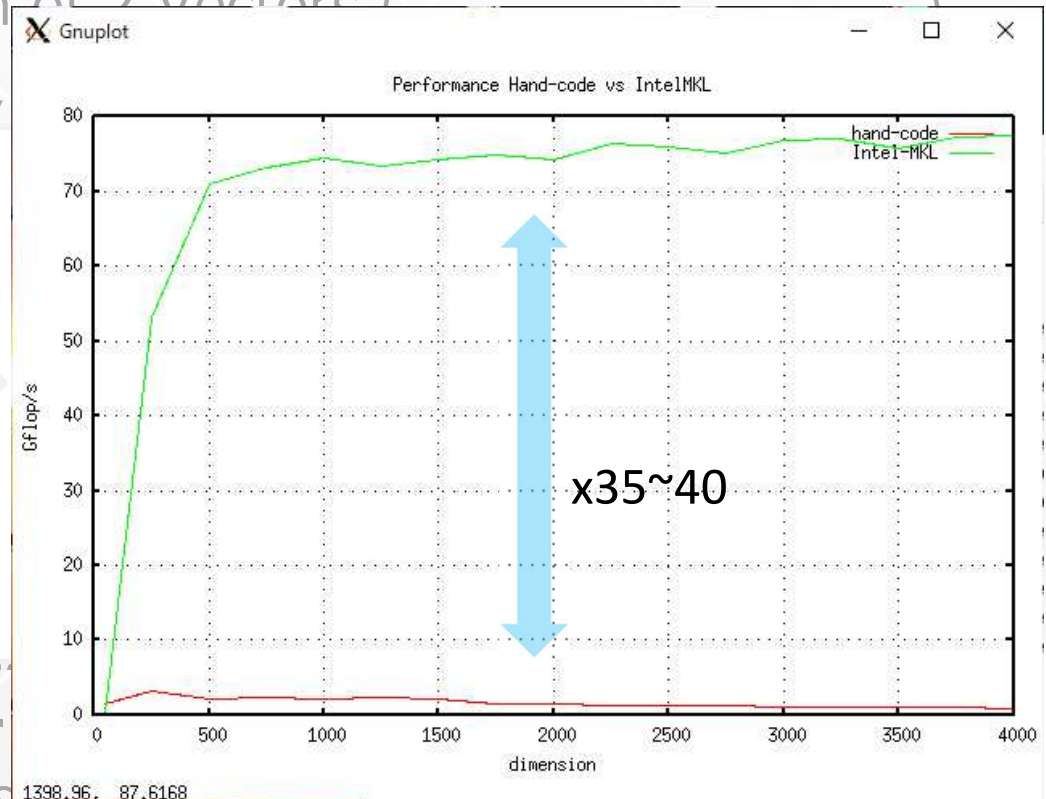
KBLAS(@KAUST), ASPEN.K2(@RIKEN) : for GPGPU

# Example

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



KBLAS(@KAUST), ASPEN.

# 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.

<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 cIMATH, 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/>) *Dense, General*
- Elemental (<http://libelemental.org/>)
- EigenExa ([http://www.aics.riken.jp/labs/lpnctr/EigenExa\\_e.html](http://www.aics.riken.jp/labs/lpnctr/EigenExa_e.html)) *Dense Eigenvalue*
- ELPA (<http://elpa.rzg.mpg.de/>)
- PETSc (<http://www.mcs.anl.gov/petsc/>) *Sparse, General*
- Trillions (<https://trilinos.org/>)
- ARPACK (<http://www.caam.rice.edu/software/ARPACK/>) *Sparse, Eigenvalue*
- FFTW (<http://www.fftw.org/>)
- FFTE (<http://www.ffte.jp/>) *FFT*
- 2decomp&FFT (<http://www.2decomp.org/>)
- MT, MTGP, dSFMT (<http://www.math.sci.hiroshima-u.ac.jp/~m-mat/MT/SFMT/index.html>) *Random number*
- GMP big number librant(<https://gmplib.org/>)
- QD pack, MPACK, and so on *Multi-precision number*

# General use of PETSc

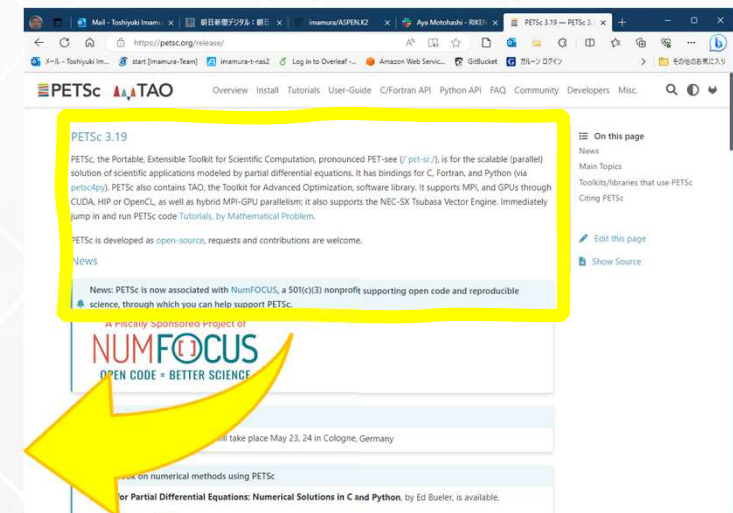
# PETSc/TAO

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

<https://petsc.org/>

PETSc, the Portable, Extensible Toolkit for Scientific Computation, pronounced PET-see (/ˈpet-si:/), is for the scalable (parallel) solution of scientific applications modeled by partial differential equations. It has bindings for C, Fortran, and Python (via `petsc4py`). PETSc also contains TAO, the Toolkit for Advanced Optimization, software library. It supports MPI, and GPUs through CUDA, HIP or OpenCL, as well as hybrid MPI-GPU parallelism; it also supports the NEC-SX Tsubasa Vector Engine. Immediately jump in and run PETSc code.

(*cf. PETSc/TAO homepage*)



# Simple example (ex1.c)

$$A = \begin{pmatrix} -2 & 1 & & & \\ 1 & -2 & & & \\ & & \ddots & & \\ & & & 1 & \\ & & & 1 & -2 \end{pmatrix}, b = \begin{pmatrix} 1 \\ 0 \\ \vdots \\ 0 \\ 1 \end{pmatrix}$$

- Solve the linear system  $Ax = b$ .
- PETSc handles the internal data format and interface data; due to PETSc's management mechanism, the user never sees the actual state in memory. Matrix A is handled by handler variables, and matrix elements are accessed by the query API.

# Preparation

- **Module load is necessary to access Odyssey and use some software packages**
- **Here, at least basic packages for [odyssey] is significant.**

```
% module purge
% module list
No Modulefiles Currently Loaded.
% module load odyssey
Loading odyssey
  Loading requirement: fjmpi/1.2.37 fj/1.2.37
% module list
Currently Loaded Modulefiles:
1) fjmpi/1.2.37  2) fj/1.2.37(default)  3) odyssey
```

## Simple example (C/ex1.c)

```
Vec      x, b, u;    /* approx solution, RHS, exact solution */
Mat      A;         /* linear system matrix */
KSP      ksp;       /* linear solver context */
PC       pc;        /* preconditioner context */
PetscReal norm;     /* norm of solution error */
```

```
VecCreate(PETSC_COMM_WORLD,&x);    //Define x
VecSetSizes(x,PETSC_DECIDE,n); // Set size of x
VecDuplicate(x,&b);
VecDuplicate(x,&u);
```

```
MatCreate(PETSC_COMM_WORLD,&A); // Define A
MatSetSizes(A,PETSC_DECIDE,PETSC_DECIDE,n,n); // Specify size of A
MatSetFromOptions(A); // Reflects -mat_type option (default is AIJ format)
MatSetUp(A);
```

## Simple example (C/ex1.c)

```
value[0] = -1.0; value[1] = 2.0; value[2] = -1.0;
for (i=1; i<n-1; i++) {
    col[0] = i-1; col[1] = i; col[2] = i+1;
    MatSetValues(A,1,&i,3,col,value,INSERT_VALUES);
}
i = n - 1; col[0] = n - 2; col[1] = n - 1;
MatSetValues(A,1,&i,2,col,value,INSERT_VALUES);
i = 0; col[0] = 0; col[1] = 1; value[0] = 2.0; value[1] = -1.0;
MatSetValues(A,1,&i,2,col,value,INSERT_VALUES);
MatAssemblyBegin(A,MAT_FINAL_ASSEMBLY);
MatAssemblyEnd(A,MAT_FINAL_ASSEMBLY);
```

# Simple example (F/ex1f.F90)

```
call MatCreate(PETSC_COMM_WORLD,A,ierr)
call MatSetSizes(A,PETSC_DECIDE,PETSC_DECIDE,n,n,ierr)
call MatSetFromOptions(A,ierr)
call MatSetUp(A,ierr)

value(1) = -1.0
value(2) = 2.0
value(3) = -1.0
do 50 i=1,n-2
  col(1) = i-1
  col(2) = i
  col(3) = i+1
  call MatSetValues(A,i1,i,i3,col,value,INSERT_VALUES,ierr)
50 continue
i = n - 1
col(1) = n - 2
col(2) = n - 1
call MatSetValues(A,i1,i,i2,col,value,INSERT_VALUES,ierr)
i = 0
col(1) = 0
col(2) = 1
value(1) = 2.0
value(2) = -1.0
call MatSetValues(A,i1,i,i2,col,value,INSERT_VALUES,ierr)
call MatAssemblyBegin(A,MAT_FINAL_ASSEMBLY,ierr)
call MatAssemblyEnd(A,MAT_FINAL_ASSEMBLY,ierr)
```



# Computed result

```
KSP Object: 1 MPI processes
  type: gmres
    restart=30, using Classical (unmodified) Gram-Schmidt Orthogonalization with no
iterative refinement
    happy breakdown tolerance 1e-30
    maximum iterations=10000, initial guess is zero
    tolerances: relative=1e-05, absolute=1e-50, divergence=10000.
    left preconditioning
    using PRECONDITIONED norm type for convergence test
PC Object: 1 MPI processes
  type: jacobi
  type DIAGONAL
  linear system matrix = precondition matrix:
Mat Object: 1 MPI processes
  type: seqaij
  rows=100, cols=100
  total: nonzeros=298, allocated nonzeros=500
  total number of mallocs used during MatSetValues calls=0
  not using l-node routines
Norm of error 0.0114852, Iterations 318
```

# Computed result

```
mpiexec ./ex1 -n 100 -ksp_type cg -pc_type none
```



```
KSP Object: 1 MPI processes  
type: cg  
maximum iterations=10000, initial guess is zero  
tolerances: relative=1e-05, absolute=1e-50, divergence=10000.  
left preconditioning  
using PRECONDITIONED norm type for convergence test  
PC Object: 1 MPI processes  
type: none  
linear system matrix = precondition matrix:  
Mat Object: 1 MPI processes  
type: seqaij  
rows=100, cols=100  
total: nonzeros=298, allocated nonzeros=500  
total number of mallocs used during MatSetValues calls=0  
not using l-node routines  
Norm of error 1.85656e-14, Iterations 50
```

# Hands-on time

Please access `/work/gt18/share/1D-PETSc`

```
$ cd /work/gt18/<YID>/
```

```
$ cp -r /work/gt18/share/1D-PETSc .
```

```
$ cd 1D-PETSc
```

```
$ ls
```

```
1d2-petsc.c    ex1.c    input.dat  run_1d2.sh
```

```
1d2-petscf.F90 ex1f.F90 Makefile  run_ex1.sh
```

# Compile and link

## Makefile

```
#
# The minimal example of Makefile for compiling PETSc
executables
# By Toshiyuki Imamura (imamura.toshiyuki@riken.jp)
#
PETSC_DIR = /work/gt18/share/petsc/3.15.0
PETSC_LIB = $(PETSC_DIR)/lib
PETSC_INC = $(PETSC_DIR)/include
HYPRE_DIR = /work/gt18/share/hypre/2.20.0
HYPRE_LIB = $(HYPRE_DIR)/lib
HYPRE_INC = $(HYPRE_DIR)/include

CC      = mpifccpx -Nclang -Kopenmp
FC      = mpifrtpx -Nclang -Kopenmp
INCFLAGS = -I$(PETSC_INC) -I$(HYPRE_INC)
LDLAGS  = -Wl,-rpath,$(PETSC_LIB) -L$(PETSC_LIB) -
lpetsc
LDLAGS := $(LDLAGS) -Wl,-rpath,$(HYPRE_LIB) -
L$(HYPRE_LIB) -IHYPRE
LDLAGS := $(LDLAGS) -lfjprofmpif -lfjprofmpi -lfjcr -
SCALAPACK -SSL2BLAMP

all: $(programs)

.SUFFIXES: .o .c .F90
.c.o:
    $(CC) -c $< -I./ -I$(PETSC_INC)
.F90.o:
    $(FC) -c $< -cpp -I./ -I$(PETSC_INC)

1d2-petsc: 1d2-petsc.o
    $(CC) -o 1d2-petsc 1d2-petsc.o $(INCFLAGS)
$(LDLAGS)
1d2-petscf: 1d2-petscf.o
    $(FC) -o 1d2-petscf 1d2-petscf.o $(INCFLAGS)
$(LDLAGS)

ex1: ex1.o
    $(CC) -o ex1 ex1.o $(INCFLAGS) $(LDLAGS)
ex1f: ex1f.o
    $(FC) -o ex1f ex1f.o $(INCFLAGS) $(LDLAGS)

clean:
    rm -rf *.o *~ $(programs) *bak

programs=1d2-petsc ex1 1d2-petscf ex1f
```

# Job submission

```
#!/bin/bash
```

```
#PJM -L node=1  
#PJM -L rscgrp=lecture-o  
#PJM -L elapse=00:10:00  
#PJM -g gt18  
#PJM --mpi proc=8
```

```
module load odyssey
```

```
# C
```

```
mpiexec -np 1 ./1d2-petsc  
#mpiexec -np 8 ./1d2-petsc -ksp_type cg -pc_type none  
#mpiexec -np 8 ./1d2-petsc -ksp_type cg -pc_type bjacobi  
#mpiexec -np 8 ./1d2-petsc -ksp_type cg -pc_type asm  
#mpiexec -np 8 ./1d2-petsc -ksp_type gmres -pc_type none  
#mpiexec -np 8 ./1d2-petsc -ksp_type gmres -pc_type bjacobi  
#mpiexec -np 8 ./1d2-petsc -ksp_type gmres -pc_type asm
```

```
# Fortran
```

```
#mpiexec -np 1 ./1d2-petscf
```

# 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.**

```
Vec Rhs_PETSC;
VecCreate(PETSC_COMM_WORLD, &Rhs_PETSC);
VecSetSizes(Rhs_PETSC, N, PETSC_DECIDE);
VecSetFromOptions(Rhs_PETSC);
Vec x_PETSC;
```

Create a vector handler

```
Mat AMat_PETSC;
MatCreate(PETSC_COMM_WORLD, &AMat_PETSC);
MatSetType(AMat_PETSC, MATMPIAIJ);
MatSetSizes(AMat_PETSC, PETSC_DECIDE, N, Ng, PETSC_DECIDE);
MatSetUp(AMat_PETSC);
```

Create a matrix handler

# 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.**

```
Vec Rhs_PETSC;
VecCreate(PETSC_COMM_WORLD, &Rhs_PETSC);
VecSetSizes(Rhs_PETSC, N, PETSC_DECIDE);
VecSetFromOptions(Rhs_PETSC);
Vec x_PETSC;
```

```
Mat AMat_PETSC;
MatCreate(PETSC_COMM_WORLD, &AMat_PETSC);
MatSetType(AMat_PETSC, MATMPIAIJ);
MatSetSizes(AMat_PETSC, PETSC_DECIDE, N, Ng, PETSC_DECIDE);
MatSetUp(AMat_PETSC);
```



Define the matrix type

# How to setup a matrix? (in C)

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```

Vec Rhs_PETSC;
VecCreate(PETSC_COMM_WORLD, &Rhs_PETSC);
VecSetSizes(Rhs_PETSC, N, PETSC_DECIDE);
VecSetFromOptions(Rhs_PETSC);
Vec x_PETSC;
  
```



Vector size

```

Mat AMat_PETSC;
MatCreate(PETSC_COMM_WORLD, &AMat_PETSC);
MatSetType(AMat_PETSC, MATMPIAIJ);
MatSetSizes(AMat_PETSC, PETSC_DECIDE, N, Ng, PETSC_DECIDE);
MatSetUp(AMat_PETSC);
  
```



Matrix size



# 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.**

```

for(i=0;i<N;i++){
  int Row = Base + i;
  MatSetValue(AMat_PETSC, Row, Row, Diag[i], INSERT_VALUES);
  for(j=Index[i];j<Index[i+1];j++){
    int Col = Base + Item[j];
    if(Item[j] >= N){
      Col = Base + N;
      if(MyRank>0 && Item[j] == N){ Col = Base - 1; }}
    MatSetValue(AMat_PETSC, Row, Col, AMat[j], INSERT_VALUES);
  }
}

```

Set matrix value to  
PETSc

```

MatAssemblyBegin( A, MAT_FINAL_ASSEMBLY )
MatAssemblyEnd( A, MAT_FINAL_ASSEMBLY )

```

Assemble the matrix  
data set on a  
distributed manner

# Play with PETSc

- Change the KSP or PC types

```
mpirun -np 8 ./1d2-petsc -ksp_type cg -pc_type none
```

Converged Reason = DIVERGED\_ITS  
2000 iters, RESID= 4.599971e+04  
5.002301e-02 9.699515e-01 sec.

```
mpirun -np 8 ./1d2-petsc -ksp_type cg -pc_type bjacobi
```

Converged Reason = CONVERGED\_ATOL  
164 iters, RESID= 1.759163e-09  
2.389900e-02 2.276256e-01 sec.

```
mpirun -np 8 ./1d2-petsc -ksp_type cg -pc_type asm
```

Converged Reason = CONVERGED\_ATOL  
21 iters, RESID= 1.420679e-09  
2.384981e-02 4.237503e-02 sec.

Converged Reason = DIVERGED\_ITS  
2000 iters, RESID= 2.184571e+02  
2.386021e-02 1.957389e+00 sec.

```
mpirun -np 8 ./1d2-petsc -ksp_type gmres -pc_type none
```

```
mpirun -np 8 ./1d2-petsc -ksp_type gmres -pc_type bjacobi
```

Converged Reason = DIVERGED\_BREAKDOWN  
30 iters, RESID= 1.985900e+02  
2.430236e-02 6.757502e-02 sec.

```
mpirun -np 8 ./1d2-petsc -ksp_type gmres -pc_type asm
```

Converged Reason = CONVERGED\_ATOL  
137 iters, RESID= 2.002330e-10  
2.392390e-02 2.671833e-01 sec.

# Sample files

Please access [/work/gt18/share/1D-PETSc](#)

```
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```

```
$ cp -r /work/gt18/share/1D-PETSc .
```

```
$ cd 1D-PETSc
```

```
$ ls
```

```
1d2-petsc.c    ex1.c    input.dat  run_1d2.sh
```

```
1d2-petscf.F90 ex1f.F90 Makefile  run_ex1.sh
```

- **KSP tpes**

Solver	KSPType	Options Database Name
Richardson	KSPRICHARDSON	richardson
Chebychev	KSPCHEBYCHEV	chebychev
Conjugate Gradient	KSPCG	cg
BiConjugate Gradient	KSPBICG	bicg
Generalized Minimal Residual	KSPGMRES	gmres
BiCGSTAB	KSPBCGS	bcgs
Conjugate Gradient Squared	KSPCGS	cgs

- **There are other types.**
- **See user manual for details**

- **PC types**

SPrecondition	PCType	Options Database Name
Jacobi	PCJACOBI	jacobi
Block Jacob	PCBJACOBI	bjacobi
SOR (and SSOR)	PCSOR	sor
Incomplete Cholesky	PCICC	icc
Incomplete LU	PCILU	ilu
Additive Schwarz	PCASM	asm
No preconditioning	PCNONE	none

- **There are other types.**
- **See user manual for details**
- **Some combinations cannot be calculated.**