



Introduction to the PETSc library

Victor Eijkhout

`eijkhout@tacc.utexas.edu`

To set the stage

Developing parallel, nontrivial PDE solvers that deliver high performance is still difficult and requires months (or even years) of concentrated effort. PETSc is a toolkit that can ease these difficulties and reduce the development time, but it is not black-box PDE solver, nor a silver bullet.

Barry Smith

More specifically...

Portable Extendable Toolkit for Scientific Computations

- Scientific Computations: parallel linear algebra, in particular linear and nonlinear solvers
- Toolkit: Contains high level solvers, but also the low level tools to roll your own.
- Portable: Available on many platforms, basically anything that has MPI

Why use it? It's big, powerful, well supported.

What does PETSc target?

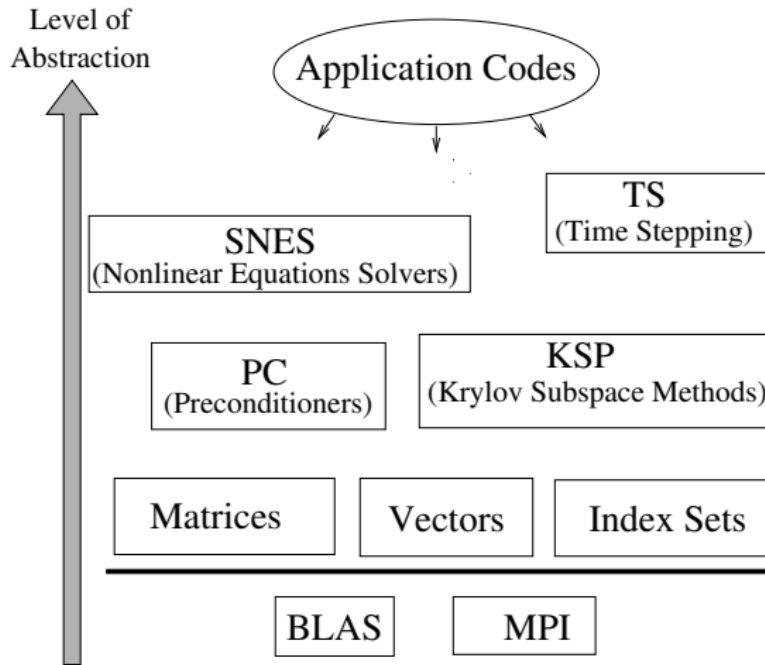
- Serial and Parallel
- Linear and nonlinear
- Finite difference and finite element
- Structured and unstructured

What is in PETSc?

- Linear system solvers (sparse/dense, iterative/direct)
- Nonlinear system solvers
- Optimization: TAO (used to be separate library)
- Tools for distributed matrices
- Support for profiling, debugging, graphical output

Documentation and help

- **Web page:** <http://www.mcs.anl.gov/petsc/>
- **PDF manual:** <http://www.mcs.anl.gov/petsc/petsc-current/docs/manual.pdf>
- Follow-up to this tutorial: eijkhout@tacc.utexas.edu
- PETSc on your local cluster: ask your local support
- General questions about PETSc: petsc-maint@mcs.anl.gov
- Example codes, found online, and in
 \$PETSC_DIR/src/mat/examples et cetera
- Sometimes consult include files, for instance
 \$PETSC_DIR/include/petscmat.h



Parallel Numerical Components of PETSc

Nonlinear Solvers	
Newton-based Methods	Other
Line Search	Trust Region

Time Steppers			
Euler	Backward Euler	Pseudo-Time Stepping	Other

Krylov Subspace Methods							
GMRES	CG	CGS	Bi-CG-Stab	TFQMR	Richardson	Chebychev	Other

Preconditioners						
Additive Schwarz	Block Jacobi	Jacobi	ILU	ICC	LU (sequential only)	Other

Matrices				
Compressed Sparse Row (AIJ)	Block Compressed Sparse Row (BAIJ)	Block Diagonal (BDiag)	Dense	Other

Vectors	Index Sets			
	Indices	Block Indices	Stride	Other

PETSc and parallelism

PETSc is layered on top of MPI

- MPI has basic tools: send elementary datatypes between processors
- PETSc has intermediate tools:
insert matrix element in arbitrary location,
do parallel matrix-vector product
- ⇒ you do not need to know much MPI when you use PETSc

Object oriented design

Petsc uses objects: vector, matrix, linear solver, nonlinear solver

Overloading:

```
MATMult(A,x,y); // y <- A x
```

same for sequential, parallel, dense, sparse

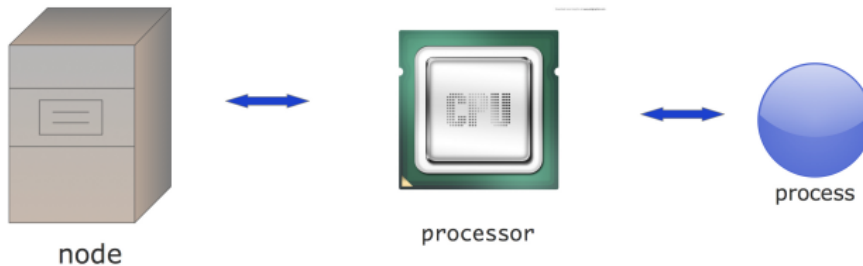
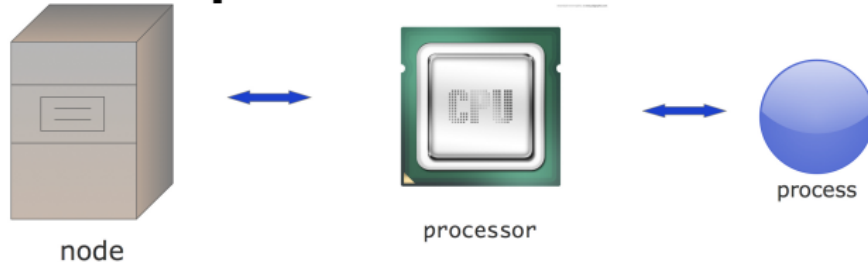
Data hiding

To support this uniform interface, the implementation is hidden:

```
MatSetValue(A, i, j, v, INSERT_VALUES); // A[i, j] <- v
```

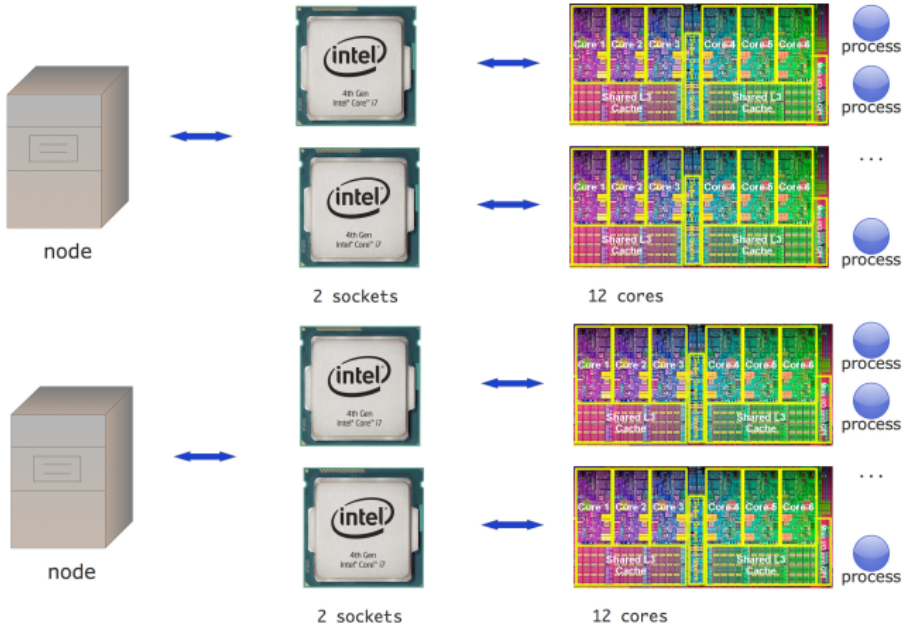
There are some direct access routines, but most of the time you don't need them.

Computers when MPI was designed



One processor and one process per node;
all communication goes through the network.

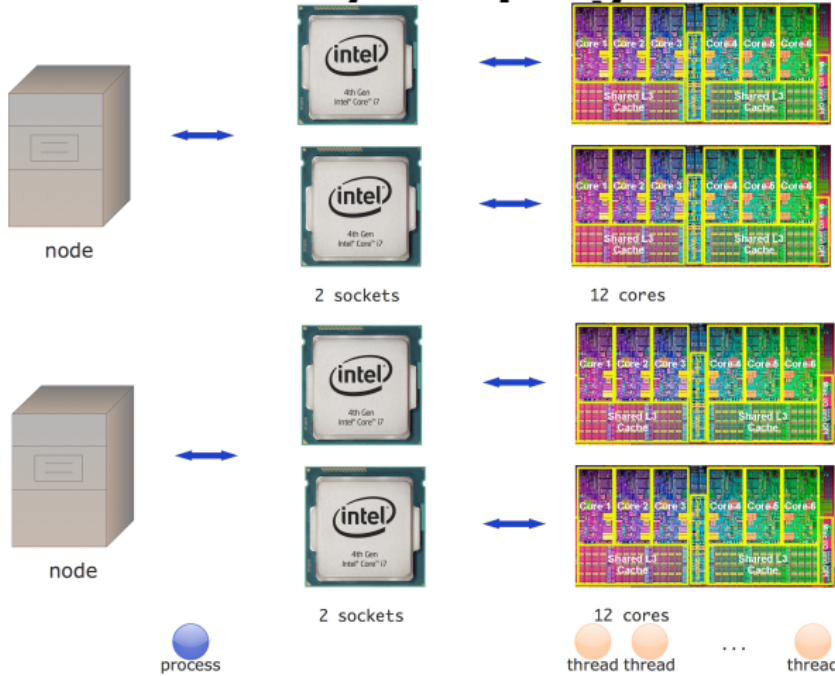
Pure MPI



A node has multiple sockets, each with multiple cores.

Pure MPI puts a process on each core: pretend shared memory doesn't exist.

Hybrid programming



Hybrid programming puts a process per node or per socket; further parallelism comes from threading.

Terminology

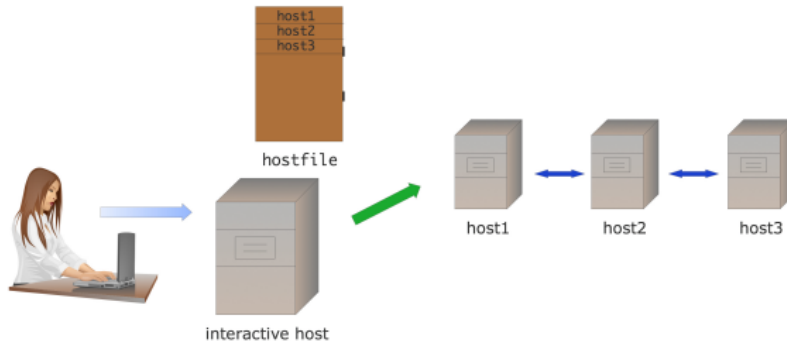
'Processor' is ambiguous: is that a chip or one independent instruction processing unit?

- Socket: the processor chip
- Processor: we don't use that word
- Core: one instruction-stream processing unit
- Process: preferred terminology in talking about MPI.

Cluster setup

Typical cluster:

- Login nodes, where you ssh into; usually shared with 100 (or so) other people. You don't run your parallel program there!
- Compute nodes: where your job is run. They are often exclusive to you: no other users getting in the way of your program.



Hostfile: the description of where your job runs. Usually generated by a job scheduler such as SLURM.

Do I need a supercomputer?

- With `mpiexec` and such, you start a bunch of processes that execute your MPI program.
- Does that mean that you need a cluster or a big multicore?
- No! You can start a large number of MPI processes, even on your laptop. The OS will use ‘time slicing’.
- Of course it will not be very efficient...

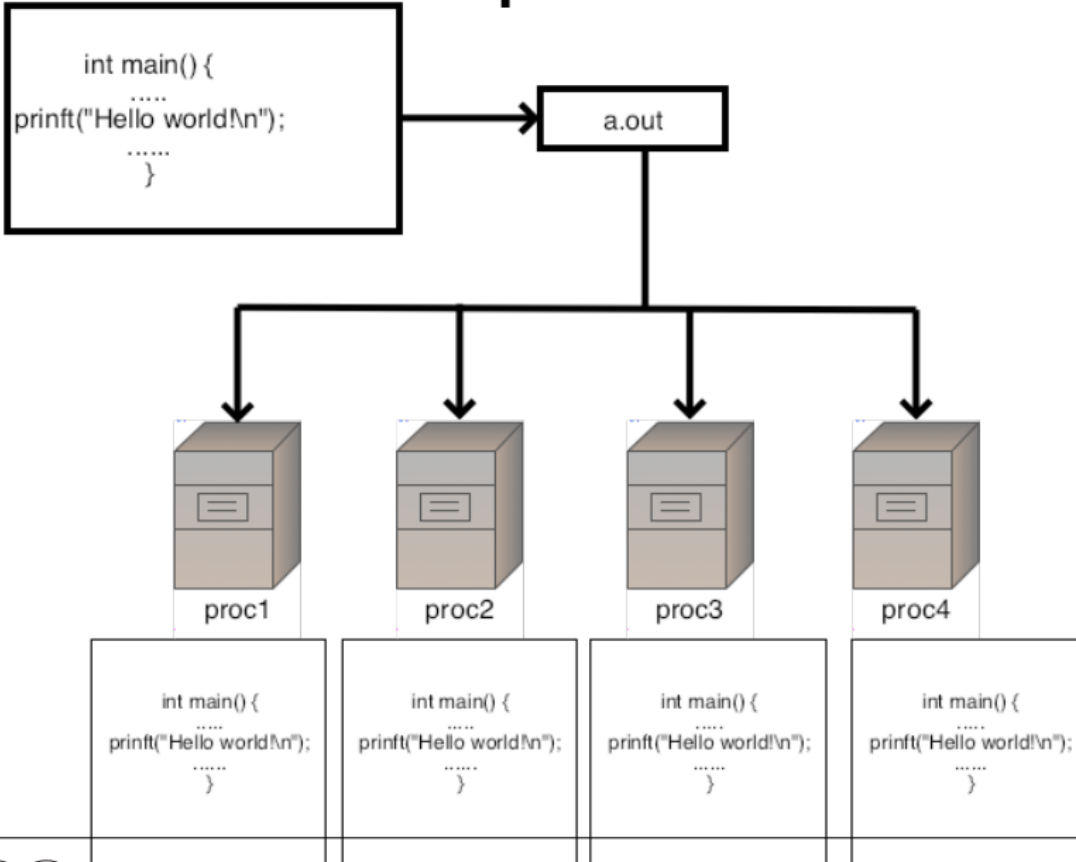
SPMD

The basic model of MPI is
'Single Program Multiple Data':
each process is an instance of the same program.

Symmetry: There is no 'master process', all processes are equal, start and end at the same time.

Communication calls do not see the cluster structure:
data sending/receiving is the same for all neighbours.

In a picture



In a picture

Four processes on two nodes (`idev -N 2 -n 4`)

```
Program:  
number <- MPI_Comm_rank  
name <- MPI_Get_processor_name
```

```
Program:  
number <- MPI_Comm_rank  
0  
name <- MPI_Get_processor_name  
c111.tacc.utexas.edu
```

```
Program:  
number <- MPI_Comm_rank  
1  
name <- MPI_Get_processor_name  
c111.tacc.utexas.edu
```

```
Program:  
number <- MPI_Comm_rank  
2  
name <- MPI_Get_processor_name  
c222.tacc.utexas.edu
```

```
Program:  
number <- MPI_Comm_rank  
3  
name <- MPI_Get_processor_name  
c222.tacc.utexas.edu
```

c111.tacc.utexas.edu

c222.tacc.utexas.edu

Sequential semantics

A PETSc program almost looks like sequential:

```
|| MatMult(A, x, y);      //  $y \leftarrow Ax$ 
|| VecCopy(y, res);       //  $r \leftarrow y$ 
|| VecAXPY(res, -1., b); //  $r \leftarrow r - b$ 
```

but note all objects are distributed.

Minimal MPI knowledge

With PETSc you barely need to know MPI.

Here is just about the only thing:

```
int MPI_Comm_rank( MPI_Comm comm, int *procno )
int MPI_Comm_size( MPI_Comm comm, int *nprocs )
```

For now, the communicator will be MPI_COMM_WORLD.

Running at TACC

Get the course files:

```
cp ~train00/petsc-course.tgz .
tar fxz petsc-course.tgz
cd petsc_course_dir
```

Running at TACC

First:

```
module load petsc/3.12-debug
# later: module load petsc/3.12
```

Compute node for 2 hours, on 2 nodes with 12 cores total:

```
idev -t 2:0:0 -N 2 -n 12
```

If you do this during an official TACC training, there will probably be a reservation in place.

How to make exercises

- Directory: exercises-3.10-c or f or (maybe) p
- Type `make exercisename` to compile it
- Python: setup once per session

`module load python`

No compilation needed. Run:

`ibrun python yourprogram`

Include files

C:

```
#include "petsc.h"
int main(int argc, char **argv)
```

Fortran:

```
program basic
#include <petsc/finclude/petsc.h>
use petsc
implicit none
```

Python:

```
from petsc4py import PETSc
```

Variable declarations, C

```
KSP           solver;  
Mat           A;  
Vec           x,y;  
PetscInt      n = 20;  
PetscScalar   v;  
PetscReal     nrm;
```

Note Scalar vs Real

Variable declarations, F

```
KSP      :: solver
Mat      :: A
Vec      :: x,y
PetscInt :: j(3)
PetscScalar :: mv
PetscReal :: nrm
```

Much like in C; uses cpp

Library setup, C

```
ierr = PetscInitialize(&argc,&argv,0,0); CHKERRQ(ierr);
// all the petsc work
ierr = PetscFinalize(); CHKERRQ(ierr);
```

Can replace MPI_Init

General: Every routine has an error return. Catch that value!

Library setup, F

```
call PetscInitialize(PETSC_NULL_CHARACTER, ierr)
    CHKERRQ(ierr)
// all the petsc work
call PetscFinalize(ierr); CHKERRQ(ierr)
```

Error code is now final parameter. This holds for every PETSc routine.

- CHKERRA in main program
- CKKERRQ in subprograms

Exercise 1 (hello)

Look up the function `PetscPrintf` and print a message
'This program runs on 27 processors'
from process zero.

- Start with the template code `hello.c/hello.F`
- Compile with `make hello`
- Part two: use `PetscSynchronizedPrintf`

About routine prototypes: C/C++

Prototype:

```
PetscErrorCode VecCreate(MPI_Comm comm, Vec *v);
```

Use:

```
PetscErrorCode ierr;
MPI_Comm comm = MPI_COMM_WORLD;
Vec v;
ierr = VecCreate( comm, &vec ); CHKERRQ(ierr).
```

(always good idea to catch that error code)

About routine prototypes: Fortran

Prototype

```
MPI_Comm_size(comm, size, ierror)
Type(MPI_Comm), INTENT(IN) :: comm
INTEGER, INTENT(OUT) :: size
INTEGER, OPTIONAL, INTENT(OUT) :: ierror
```

Use:

```
Type(MPI_Comm) :: comm = MPI_COMM_WORLD
integer :: size
CALL MPI_Comm_size( comm, size, ierr )
```

- Final parameter always error parameter. Do not forget!
- Most MPI_... types are INTEGER.

About routine prototypes: Python

Prototype:

```
# object method  
MPI.Comm.Send(self, buf, int dest, int tag=0)  
# class method  
MPI.Request.Waitall(type cls, requests, statuses=None)
```

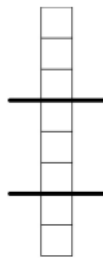
Use:

```
from mpi4py import MPI  
comm = MPI.COMM_WORLD  
comm.Send(sendbuf, dest=other)  
MPI.Request.Waitall(requests)
```

Parallel layout up to PETSc

```
VecSetSizes(Vec v, int m, int M);
```

Local size can be specified as PETSC_DECIDE.



```
VecSetSizes(V,PETSC_DECIDE,8)
```

```
VecSetSizes(V,PETSC_DECIDE,8)
```

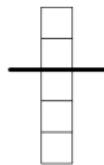
```
VecSetSizes(V,PETSC_DECIDE,8)
```

Parallel layout specified

Local or global size in

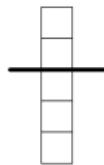
```
VecSetSizes(Vec v, int m, int M);
```

Global size can be specified as PETSC_DECIDE.



VecSetSizes(V,2,5)

VecSetSizes(V,3,5)



VecSetSizes(V,2,PETSC_DECIDE)

VecSetSizes(V,3,PETSC_DECIDE)

Setting values

Set vector to constant value:

```
VecSet(Vec x,PetscScalar value);
```

Set individual elements (global indexing!):

```
VecSetValue(Vec x,int row,PetscScalar value, InsertMode mode);
VecSetValues(Vec x,int n,int *rows,PetscScalar *values,
    InsertMode mode); // INSERT_VALUES or ADD_VALUES
```

```
i = 1; v = 3.14;
VecSetValue(x,i,v,INSERT_VALUES);
ii[0] = 1; ii[1] = 2; vv[0] = 2.7; vv[1] = 3.1;
VecSetValues(x,2,ii,vv,INSERT_VALUES);
```

```
call VecSetValue(x,i,v,INSERT_VALUES,ierr,e)
ii(1) = 1; ii(2) = 2; vv(1) = 2.7; vv(2) = 3.1
call VecSetValues(x,2,ii,vv,INSERT_VALUES,ierr,e)
```

Setting values

No restrictions on parallelism;
after setting, move values to appropriate processor:

```
VecAssemblyBegin(Vec x);  
VecAssemblyEnd(Vec x);
```

Basic operations

```
VecAXPY(Vec y,PetscScalar a,Vec x); /* y <- y + a x */
VecAYPX(Vec y,PetscScalar a,Vec x); /* y <- a y + x */
VecScale(Vec x, PetscScalar a);
VecDot(Vec x, Vec y, PetscScalar *r); /* several variants */
VecMDot(Vec x,int n,Vec y[],PetscScalar *r);
VecNorm(Vec x,NormType type, double *r);
VecSum(Vec x, PetscScalar *r);
VecCopy(Vec x, Vec y);
VecSwap(Vec x, Vec y);
VecPointwiseMult(Vec w,Vec x,Vec y);
VecPointwiseDivide(Vec w,Vec x,Vec y);
VecMAXPY(Vec y,int n, PetscScalar *a, Vec x[]);
VecMax(Vec x, int *idx, double *r);
VecMin(Vec x, int *idx, double *r);
VecAbs(Vec x);
VecReciprocal(Vec x);
VecShift(Vec x,PetscScalar s);
```

Exercise 2 (vec)

Use the routines `VecDot`, `VecScale` and `VecNorm` to compute the inner product of vectors x , y , scale the vector x , and check its norm:

$$\begin{aligned} p &\leftarrow x^t y \\ x &\leftarrow x/p \\ n &\leftarrow \|x\|_2 \end{aligned}$$

Matrix creation

The usual create/destroy calls:

```
MatCreate(MPI_Comm comm, Mat *A)  
MatDestroy(Mat *A)
```

Several more aspects to creation:

```
MatSetType(A, MATSEQAIJ) /* or MATMPIAIJ or MATAIJ */  
MatSetSizes(Mat A, int m, int n, int M, int N)  
MatSeqAIJSetPreallocation /* more about this later*/  
(Mat B, PetscInt nz, const PetscInt nnz[])
```

Local or global size can be PETSC_DECIDE (as in the vector case)

If you already have a CRS matrix

```
PetscErrorCode MatCreateSeqAIJWithArrays  
  (MPI_Comm comm, PetscInt m, PetscInt n,  
   PetscInt* i, PetscInt* j, PetscScalar *a, Mat *mat)
```

(also from triplets)

Do not use this unless you interface to a legacy code. And even then...

Matrix Preallocation

- PETSc matrix creation is very flexible:
- No preset sparsity pattern
- any processor can set any element
⇒ potential for lots of malloc calls
- tell PETSc the matrix' sparsity structure
(do construction loop twice: once counting, once making)
- Re-allocating is expensive:

`MatSetOption(A, MAT_NEW_NONZERO_LOCATIONS, PETSC_FALSE);`

(is default) Otherwise:

[1]PETSC ERROR: Argument out of range

[1]PETSC ERROR: New nonzero at (0,1) caused a malloc

Sequential matrix structure

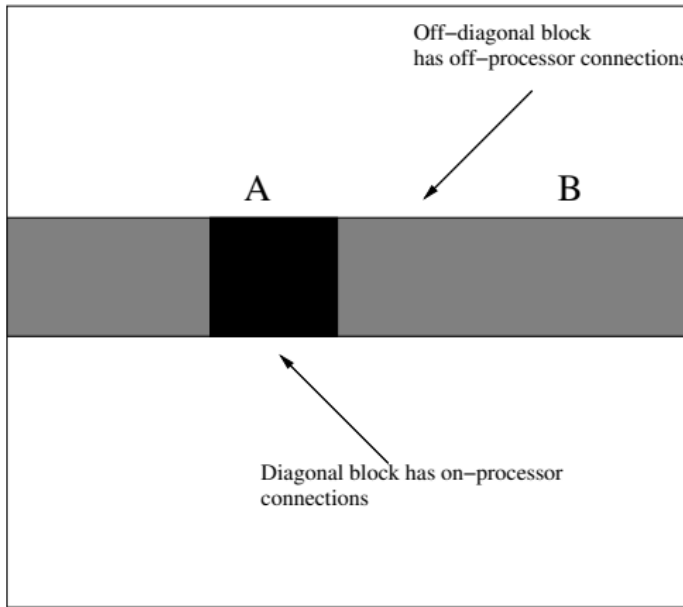
`MatSeqAIJSetPreallocation`

`(Mat B, PetscInt nz, const PetscInt nnz[])`

- `nz` number of nonzeros per row
(or slight overestimate)
- `nnz` array of row lengths (or overestimate)
- considerable savings over dynamic allocation!

In Fortran use `PETSC_NULL_INTEGER` if not specifying `nnz` array

Parallel matrix structure



(why does it do this?)

- $y \leftarrow Ax_A + Bx_B$
- x_B needs to be communicated; Ax_A can be computed in the meantime
- Algorithm
 - Initiate asynchronous sends/receives for x_B
 - compute Ax_A
 - make sure x_B is in
 - compute Bx_B
- so by splitting matrix storage into A, B part, code for the sequential case can be reused.
- This is one of the few places where PETSc's design is visible to the user.

Parallel matrix structure description

- m, n local size; M, N global. Note: If the matrix is square, specify m, n equal, even though distribution by block rows
- d_{nz} : number of nonzeros per row in diagonal part
- o_{nz} : number of nonzeros per row in off-diagonal part
- d_{nnz} : array of numbers of nonzeros per row in diagonal part
- o_{nnz} : array of numbers of nonzeros per row in off-diagonal part

```
MatMPIAIJSetPreallocation  
(Mat B,  
 PetscInt d_nz,const PetscInt d_nnz[],  
 PetscInt o_nz,const PetscInt o_nnz[])
```

In Fortran use PETSC_NULL_INTEGER if not specifying arrays

Querying parallel structure

Matrix partitioned by block rows:

```
MatGetSize(Mat mat,PetscInt *M,PetscInt* N);  
MatGetLocalSize(Mat mat,PetscInt *m,PetscInt* n);  
MatGetOwnershipRange(Mat A,int *first row,int *last row);
```

In query functions, unneeded components can be specified as
PETSC_NULL.

Fortran: PETSC_NULL_INTEGER

Setting values

Set one value:

```
MatSetValue(Mat A,  
           PetscInt i,PetscInt j,PetscScalar va,InsertMode mode)
```

where insert mode is INSERT_VALUES, ADD_VALUES

Set block of values:

```
MatSetValues(Mat A,int m,const int idxm[],  
            int n,const int idxn[],const PetscScalar values[],  
            InsertMode mode)
```

(v is row-oriented)

Special case of the general case:

```
MatSetValues (A, 1, &i, 1, &j, &v, INSERT_VALUES); // C  
MatSetValues (A, 1, i, 1, j, v, INSERT_VALUES, e); ! F
```

Assembling the matrix

Setting elements is independent of parallelism; move elements to proper processor:

```
MatAssemblyBegin(Mat A, MAT_FINAL_ASSEMBLY);  
MatAssemblyEnd(Mat A, MAT_FINAL_ASSEMBLY);
```

Cannot mix inserting/adding values: need to do assembly in between
with MAT_FLUSH_ASSEMBLY

Exercise 3 (matvec)

Pretend that you do not know how the matrix is created. Use `MatGetOwnershipRange` or `MatGetLocalSize` to create a vector with the same distribution, and then compute $y \leftarrow Ax$.

(Part of the code has been disabled with `#if 0`. We will get to that next.)

Matrix operations

Main operations are matrix-vector:

MatMult (Mat A, Vec in, Vec out);

MatMultAdd

MatMultTranspose

MatMultTransposeAdd

Simple operations on matrices:

MatNorm

MatScale

MatDiagonalScale

Some matrix-matrix operations

```
MatMatMult (Mat, Mat, MatReuse, PetscReal, Mat*) ;
```

```
MatPtAP (Mat, Mat, MatReuse, PetscReal, Mat*) ;
```

```
MatMatMultTranspose (Mat, Mat, MatReuse, PetscReal, Mat*) ;
```

```
MatAXPY (Mat, PetscScalar, Mat, MatStructure) ;
```

What are iterative solvers?

Solving a linear system $Ax = b$ with Gaussian elimination can take lots of time/memory.

Alternative: iterative solvers use successive approximations of the solution:

- Convergence not always guaranteed
- Possibly much faster / less memory
- Basic operation: $y \leftarrow Ax$ executed once per iteration
- Also needed: preconditioner $B \approx A^{-1}$

Topics

- All linear solvers in PETSc are iterative, even the direct ones
- Preconditioners
- Fargoin control through commandline options
- Tolerances, convergence and divergence reason
- Custom monitors and convergence tests

Iterative solver basics

```
KSPCreate(comm,&solver); KSPDestroy(solver);
// set Amat and Pmat
KSPSetOperators(solver,A,B); // usually: A,A
// solve
KSPSolve(solver,rhs,sol);
```

Optional: KSPSetup(solver)

Reuse Amat **or** Pmat: KSPGetOperators and
PetscObjectReference.

Solver type

```
KSPSetType(solver, KSPGMRES);
```

KSP can be controlled from the commandline:

```
KSPSetFromOptions(solver);  
/* right before KSPSolve or KSPSetUp */
```

then options `-ksp....` are parsed.

- `type`: `-ksp_type gmres -ksp_gmres_restart 20`
- `-ksp_view`

Convergence

Iterative solvers can fail

- Solve call itself gives no feedback: solution may be completely wrong
- KSPGetConvergedReason(solver, &reason) :
positive is convergence, negative divergence
KSPConvergedReasons[reason] is string
- KSPGetIterationNumber(solver, &nits) : after how many iterations did the method stop?

```
KSPSolve(solver,B,X);
KSPGetConvergedReason(solver,&reason);
if (reason<0) {
    PetscPrintf(comm,
        "Failure to converge after %d iterations; reason %s\n",
        its,KSPConvergedReasons[reason]);
} else {
    KSPGetIterationNumber(solver,&its);
    PetscPrintf(comm,
        "Convergence in %d iterations.\n",its);
}
```

Preconditioners

System $Ax = b$ is transformed:

$$M^{-1}A = M^{-1}b$$

- M is constructed once, applied in every iteration
- If $M = A$: convergence in one iteration
- Tradeoff: M expensive to construct \Rightarrow low number of iterations; construction can sometimes be amortized.
- Other tradeoff: M more expensive to apply and only modest decrease in number of iterations
- Symmetry: A, M symmetric $\not\Rightarrow M^{-1}A$ symmetric, however can be symmetrized by change of inner product
- Can be tricky to make both parallel and efficient

PC basics

- PC usually created as part of KSP: separate create and destroy calls exist, but are (almost) never needed

```
KSP solver; PC precon;  
KSPCreate(comm, &solver);  
KSPGetPC(solver, &precon);  
PCSetType(precon, PCJACOBI);
```

- Many choices, some with options: PCJACOBI, PCILU (only sequential), PCASM, PCBjacobi, PCMG, et cetera
- Controllable through commandline options:
-pc_type ilu -pc_factor_levels 3

Exercise 4 (ksp)

File `ksp.c` / `ksp.F90` contains the solution of a (possibly nonsymmetric) linear system.

Compile the code and run it. Now experiment with commandline options. Make notes on your choices and their outcomes.

- The code has two custom commandline switch:
 - `-n 123` set the domain size to 123 and therefore the matrix size to 123^2 .
 - `-unsymmetry 456` adds a convection-like term to the matrix, making it unsymmetric. The numerical value is the actual element size that is set in the matrix.
- What is the default solver in the code? Run with `-ksp_view`
- Print out the matrix for a small size with `-mat_view`.
- Now out different solvers for different matrix sizes and amounts of unsymmetry. See the instructions in the code.

Monitors and convergence tests

```
KSPSetTolerances(solver,rtol,atol,dtol,maxit);
```

Monitors can be set in code, but simple cases:

- `-ksp_monitor`
- `-ksp_monitor_true_residual`

More KSP topics

- Custom monitors and convergence tests
- Method-specific options (especially GMRES)
- Null spaces

Fieldsplit preconditioners

If a problem contains multiple physics, separate preconditioning can make sense

Matrix block storage: MatCreateNest

$$\begin{pmatrix} A_{00} & A_{01} & A_{02} \\ A_{10} & A_{11} & A_{12} \\ A_{20} & A_{21} & A_{22} \end{pmatrix}$$

However, it makes more sense to interleave these fields

Easy case: all fields are the same size

```
PCSetType(prec, PCFIELDSPLIT);  
PCFieldSplitSetBlockSize(prec, 3);  
PCFieldSplitSetType(prec, PC_COMPOSITE_ADDITIVE);
```

Subpreconditioners can be specified in code, but easier with options:

```
PetscOptionsSetValue  
("-fieldsplit_0_pc_type", "lu");  
PetscOptionsSetValue  
("-fieldsplit_0_pc_factor_mat_solver_package", "mumps");
```

Fields can be named instead of numbered.

Non-strided, arbitrary fields: PCFieldSplitSetIS()

Stokes equation can be detected:

-pc_fieldsplit_detect_saddle_point

Combining fields multiplicatively: solve

$$\begin{pmatrix} I & \\ A_{10}A_{00}^{-1} & I \end{pmatrix} \begin{pmatrix} A_{00} & A_{01} \\ & A_{11} \end{pmatrix}$$

If there are just two fields, they can be combined by Schur complement

$$\begin{pmatrix} I & \\ A_{10}A_{00}^{-1} & I \end{pmatrix} \begin{pmatrix} A_{00} & A_{01} \\ & A_{11} - A_{10}A_{00}^{-1}A_{01} \end{pmatrix}$$

Fieldsplit example

```
KSPGetPC(solver,&prec);
PCSetType(prec,PCFIELDSPLIT);
PCFieldSplitSetBlockSize(prec,2);
PCFieldSplitSetType(prec,PC_COMPOSITE_ADDITIVE);
PetscOptionsSetValue
  ("-fieldsplit_0_pc_type","lu");
PetscOptionsSetValue
  ("-fieldsplit_0_pc_factor_mat_solver_package","mumps");
PetscOptionsSetValue
  ("-fieldsplit_1_pc_type","lu");
PetscOptionsSetValue
  ("-fieldsplit_1_pc_factor_mat_solver_package","mumps");
```

Global preconditioners: MG

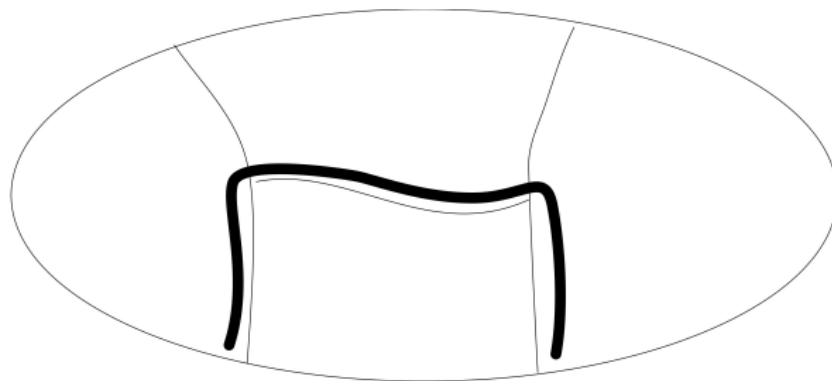
```
PCSetType (PC pc, PCMG) ;  
PCMGSetsLevels (pc, int levels, MPI Comm *comms) ;  
PCMGSetsType (PC pc, PCMGTyoe mode) ;  
PCMGSetsCycleType (PC pc, PCMGCycleType ctype) ;  
PCMGSetsNumberSmoothUp (PC pc, int m) ;  
PCMGSetsNumberSmoothDown (PC pc, int n) ;  
PCMGGetsCoarseSolve (PC pc, KSP *ksp) ;  
PCMGSetsInterpolation (PC pc, int level, Mat P) ; and  
PCMGSetsRestriction (PC pc, int level, Mat R) ;  
PCMGSetsResidual (PC pc, int level, PetscErrorCode  
(*residual) (Mat, Vec, Vec, Vec), Mat mat) ;
```

Global preconditioners: Hypre

- Hypre is a package like PETSc
- selling point: fancy preconditioners
- Install with `--with-hypre=yes --download-hypre=yes`
- then use `-pc_type hypre -pc_hypre_type`
`parasails/boomeramg/euclid/pilut`

Irregular data movement

Example: collect distributed boundary onto a single processor (this happens in the matrix-vector product)



Problem: figuring out communication is hard, actual communication is cheap

VecScatter

Preprocessing: determine mapping between input vector and output:

```
VecScatterCreate(Vec, IS, Vec, IS, VecScatter*)
// also Destroy
```

Application to specific vectors:

```
VecScatterBegin(VecScatter,
    Vec, Vec, InsertMode mode, ScatterMode direction)
VecScatterEnd  (VecScatter,
    Vec, Vec, InsertMode mode, ScatterMode direction)
```

IS: index set

Index Set is a set of indices

```
ISCreateGeneral(comm,n,indices,PETSC_COPY_VALUES,&is);
    /* indices can now be freed */
ISCreateStride (comm,n,first,step,&is);
ISCreateBlock  (comm,bs,n,indices,&is);

ISDestroy(is);
```

Use MPI_COMM_SELF most of the time.

Various manipulations: ISSum, ISDifference,
ISInvertPermutations **et cetera.**

Also ISGetIndices / ISRestoreIndices / ISGetSize

Example: simulate allgather

```
/* create the distributed vector with one element per processor */
ierr = VecCreate(MPI_COMM_WORLD,&global);
ierr = VecSetType(global,VECMPI);
ierr = VecSetSizes(global,1,PETSC_DECIDE);

/* create the local copy */
ierr = VecCreate(MPI_COMM_SELF,&local);
ierr = VecSetType(local,VECSEQ);
ierr = VecSetSizes(local,ntids,ntids);
```

```
IS indices;
ierr = ISCreateStride(MPI_COMM_SELF,ntids,0,1, &indices);
/* create a scatter from the source indices to target */
ierr = VecScatterCreate
(global,indices,local,indices,&scatter);
/* index set is no longer needed */
ierr = ISDestroy(&indices);
```

Example: even and odd indices

```
if (mytid==0) {  
    ierr = ISCreateStride  
        (MPI_COMM_SELF, (N+1)/2, 0, 2, &gindices); // even  
} else {  
    ierr = ISCreateStride  
        (MPI_COMM_SELF, N/2, 1, 2, &gindices); // odd  
}  
ierr = ISGetSize(gindices, &localsize);  
/* create a vector with the requisite local size */  
ierr = VecCreate(MPI_COMM_WORLD, &in);  
ierr = VecSetType(in, VECMPI);  
ierr = VecSetSizes(in, localsize, N);
```

Create local target vectors:

```
/* create IS objects for the target locations */
ierr = ISCreateStride
      (MPI_COMM_SELF,localsize,0,1,&lindices);
/* output vector is local */
ierr = VecCreate(MPI_COMM_SELF,&out);
ierr = VecSetType(out,VECSEQ);
ierr = VecSetSizes(out,localsize,localsize);
```

Create and apply scatter:

```
/* create a scatter from the source indices to target */
ierr = VecScatterCreate(in,gindices,out,lindices,&scatter);
/* index sets are no longer needed */
ierr = ISDestroy(&gindices);
ierr = ISDestroy(&gindices);

ierr = VecScatterBegin
(scatter,in,out,INSERT_VALUES,SCATTER_FORWARD);
ierr = VecScatterEnd
(scatter,in,out,INSERT_VALUES,SCATTER_FORWARD);
VecView(out,0);
```

```
Vector Object: 1 MPI processes
```

```
  type: seq
```

```
 0  
 2  
 4  
 6  
 8  
10  
12  
14  
16  
18  
20
```

```
Vector Object: 1 MPI processes
```

```
  type: seq
```

```
 1  
 3  
 5  
 7  
 9  
11  
13  
15  
17  
19  
21
```

Same, but distributed output vector

```
/* create IS objects for the target locations */
ierr = VecGetOwnershipRange(in,&myfirst,PETSC_NULL);
ierr = ISCreateStride
      (MPI_COMM_SELF,localsize,myfirst,1,&lindices);
/* output vectors will have the same size & layout as input
ierr = VecDuplicate(in,&out);
```

Basic profiling

- `-log_summary` flop counts and timings of all PETSc events
- `-info` all sorts of information, in particular

```
%% mpiexec yourprogram -info | grep malloc
[0] MatAssemblyEnd_SeqAIJ():
    Number of mallocs during MatSetValues() is 0
```

- `-log_trace` start and end of all events: good for hanging code

Log summary: overall

	Max	Max/Min	Avg	Total
Time (sec):	5.493e-01	1.00006	5.493e-01	
Objects:	2.900e+01	1.00000	2.900e+01	
Flops:	1.373e+07	1.00000	1.373e+07	2.746e+07
Flops/sec:	2.499e+07	1.00006	2.499e+07	4.998e+07
Memory:	1.936e+06	1.00000		3.871e+06
MPI Messages:	1.040e+02	1.00000	1.040e+02	2.080e+02
MPI Msg Lengths:	4.772e+05	1.00000	4.588e+03	9.544e+05
MPI Reductions:	1.450e+02	1.00000		

Log summary: details

	Max	Ratio	Max	Ratio	Max	Ratio	Avg	len	%T	%F	%M	%L	%R	%T	%F	%M	%L	%R	Mflop/s
MatMult	100	1.0	3.4934e-02	1.0	1.28e+08	1.0	8.0e+02	6	32	96	17	0	6	32	96	17	0	255	
MatSolve	101	1.0	2.9381e-02	1.0	1.53e+08	1.0	0.0e+00	5	33	0	0	0	5	33	0	0	0	305	
MatLUFactorNum	1	1.0	2.0621e-03	1.0	2.18e+07	1.0	0.0e+00	0	0	0	0	0	0	0	0	0	0	43	
MatAssemblyBegin	1	1.0	2.8350e-03	1.1	0.00e+00	0.0	1.3e+05	0	0	3	83	1	0	0	3	83	1	0	
MatAssemblyEnd	1	1.0	8.8258e-03	1.0	0.00e+00	0.0	4.0e+02	2	0	1	0	3	2	0	1	0	3	0	
VecDot	101	1.0	8.3244e-03	1.2	1.43e+08	1.2	0.0e+00	1	7	0	0	35	1	7	0	0	35	243	
KSPSetup	2	1.0	1.9123e-02	1.0	0.00e+00	0.0	0.0e+00	3	0	0	0	2	3	0	0	0	2	0	
KSPSolve	1	1.0	1.4158e-01	1.0	9.70e+07	1.0	8.0e+02	26100	96	17	92	26100	96	17	92	26100	194		

Debugging

- Use of `CHKERRQ` and `SETERRQ` for catching and generating error
- Use of `PetscMalloc` and `PetscFree` to catch memory problems;
`CHKMEMQ` for instantaneous memory test (debug mode only)
- Better than `PetscMalloc`: `PetscMalloc1` aligned to
`PETSC_MEMALIGN`