

The Portable Extensible Toolkit for Scientific Computing

Matthew Knepley

Computational and Applied Mathematics
Rice University

PETSc Tutorial
CEMRACS Summer School
Marseille, FR Jul 18, 2016



Never believe *anything*,
unless you can run it.

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The PETSc Team



Matt Knepley



Barry Smith



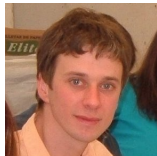
Satish Balay



Hong Zhang



Jed Brown



Lisandro Dalcin



Stefano Zampini

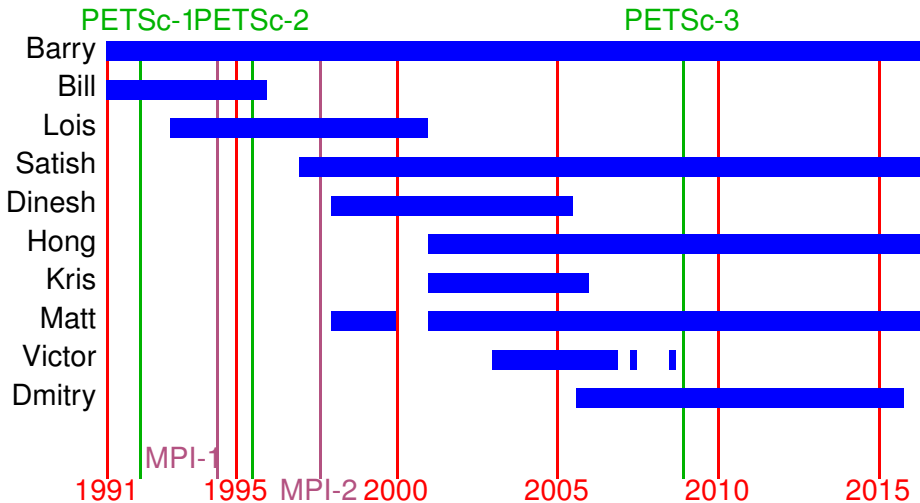


Mark Adams

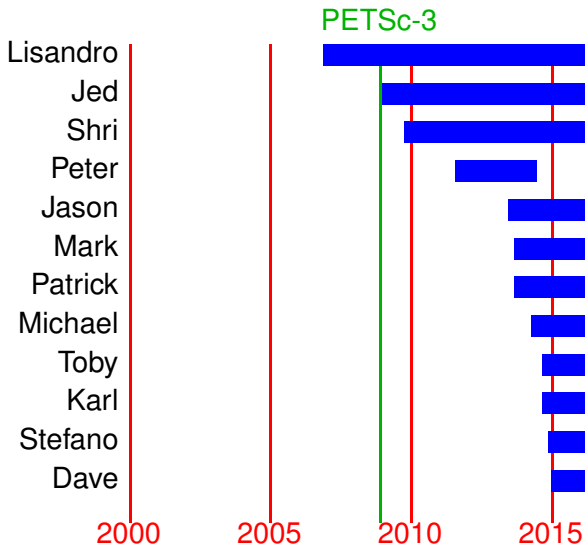


Toby Issac

Timeline (Old People)



Timeline (Young People)



What I Need From You

- Tell me if you do not understand
- Tell me if an example does not work
- Suggest better wording or **figures**
- Followup problems at petsc-maint@mcs.anl.gov

Ask Questions!!!

- Helps **me** understand what you are missing
- Helps **you** clarify misunderstandings
- Helps **others** with the same question

How We Can Help at the Tutorial

- Point out relevant documentation
- Quickly answer questions
- Help install
- Guide design of large scale codes
- Answer email at petsc-maint@mcs.anl.gov

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Outline

- 1 Getting Started with PETSc
 - Who uses PETSc?
 - Stuff for Windows
 - How can I get PETSc?
 - How do I Configure PETSc?
 - How do I Build PETSc?
 - How do I run an example?
 - How do I get more help?

2 PETSc Integration

3 DM

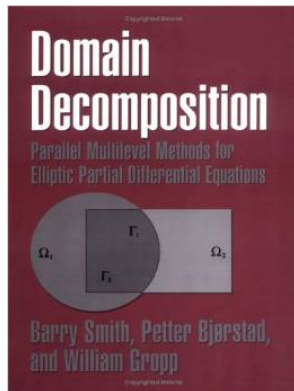
4 Advanced Solvers

How did PETSc Originate?

PETSc was developed as a Platform for Experimentation

We want to experiment with different

- Models
- Discretizations
- Solvers
- Algorithms
 - which blur these boundaries



The Role of PETSc

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 a black-box PDE solver, nor a **silver bullet**.*

— Barry Smith

Advice from Bill Gropp

You want to think about how you decompose your data structures, how you think about them globally. [...] If you were building a house, you'd start with a set of blueprints that give you a picture of what the whole house looks like. You wouldn't start with a bunch of tiles and say, "Well I'll put this tile down on the ground, and then I'll find a tile to go next to it." But all too many people try to build their parallel programs by creating the smallest possible tiles and then trying to have the structure of their code emerge from the chaos of all these little pieces. You have to have an organizing principle if you're going to survive making your code parallel.

(<http://www.rce-cast.com/Podcast/rce-28-mpich2.html>)

What is PETSc?

A freely available and supported research code for the parallel solution of nonlinear algebraic equations

Free

- Download from <http://www.mcs.anl.gov/petsc>
- Free for everyone, including industrial users

Supported

- Hyperlinked manual, examples, and manual pages for all routines
- Hundreds of tutorial-style examples
- Support via email: petsc-maint@mcs.anl.gov

Usable from C, C++, Fortran 77/90, Matlab, Julia, and Python

What is PETSc?

- Portable to any parallel system supporting MPI, including:
 - Tightly coupled systems
 - Cray XT6, BG/Q, NVIDIA Fermi, K Computer
 - Loosely coupled systems, such as networks of workstations
 - IBM, Mac, iPad/iPhone, PCs running Linux or Windows
- PETSc History
 - Begun September 1991
 - Over 60,000 downloads since 1995 (version 2)
 - Currently 400 per month
- PETSc Funding and Support
 - Department of Energy
 - ECP, AMR Program, SciDAC, MICS Program, INL Reactor Program
 - National Science Foundation
 - SI2, CIG, CISE, Multidisciplinary Challenge Program
 - Intel Parallel Computing Center

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Who Uses PETSc?

Computational Scientists

- Earth Science
 - **PyLith** (CIG)
 - **Underworld** (Monash)
 - **Magma Dynamics** (LDEO, Columbia, Oxford)
- Subsurface Flow and Porous Media
 - **STOMP** (DOE)
 - **PFLOTRAN** (DOE)

Who Uses PETSc?

Computational Scientists

- CFD
 - Firedrake
 - Fluidity
 - freeCFD
 - OpenFVM
- MicroMagnetics
 - MagPar
- Fusion
 - XGC
 - BOUT++
 - NIMROD

Who Uses PETSc?

Algorithm Developers

- Iterative methods
 - Deflated GMRES
 - LGMRES
 - QCG
 - SpecEst
- Preconditioning researchers
 - Prometheus (Adams)
 - ParPre (Eijkhout)
 - FETI-DP (Klawonn and Rheinbach)

Who Uses PETSc?

Algorithm Developers

- Finite Elements

- libMesh
- MOOSE
- PETSc-FEM
- Deal II
- OOFEM

- Other Solvers

- Fast Multipole Method (PetFMM)
- Radial Basis Function Interpolation (PetRBF)
- Eigensolvers (SLEPc)
- Optimization (TAO)

What Can We Handle?

- PETSc has run implicit problems with over **500 billion** unknowns
 - UNIC on BG/P and XT5
 - PFLOTRAN for flow in porous media
- PETSc has run on over **1,500,000** cores efficiently
 - Gordon Bell Prize Mantle Convection on IBM BG/Q Sequoia
- PETSc applications have run at 23% of peak (**600 Teraflops**)
 - Jed Brown on NERSC Edison
 - HPGMG code

What Can We Handle?

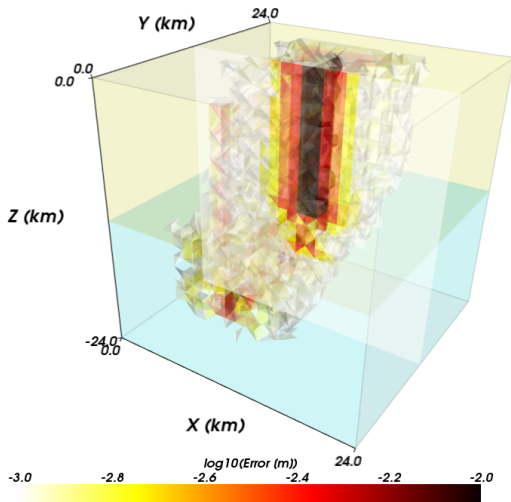
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PyLith

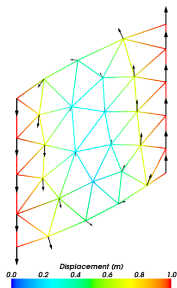
- Multiple problems
 - Dynamic rupture
 - Quasi-static relaxation
- Multiple models
 - Nonlinear visco-plastic
 - Finite deformation
 - Fault constitutive models
- Multiple meshes
 - 1D, 2D, 3D
 - Hex and tet meshes
- Parallel
 - PETSc solvers
 - DMPlex mesh management



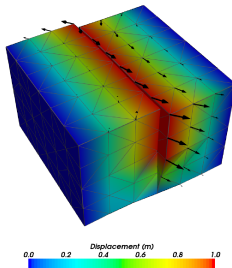
^aAagaard, Knepley, Williams

Multiple Mesh Types

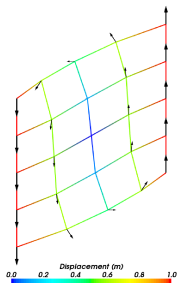
Triangular



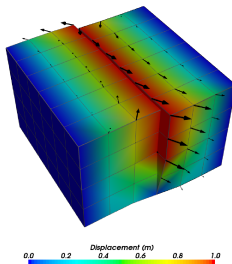
Tetrahedral



Rectangular

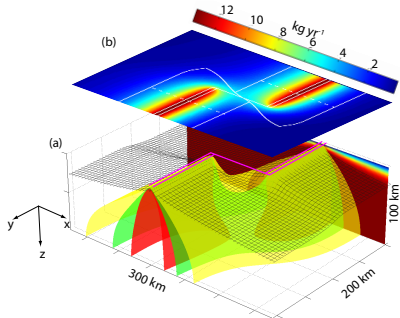
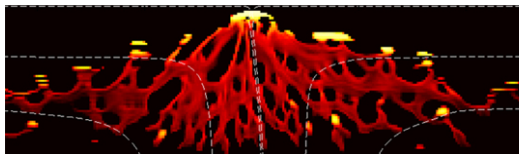


Hexahedral



Magma Dynamics

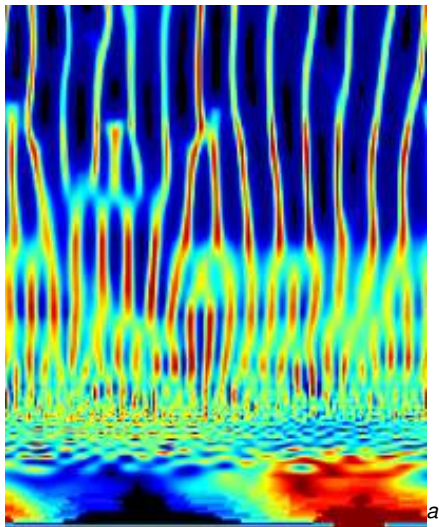
- Couples scales
 - Subduction
 - Magma Migration
- Physics
 - Incompressible fluid
 - Porous solid
 - Variable porosity
- Deforming matrix
 - Compaction pressure
- Code generation
 - FEniCS
- Multiphysics Preconditioning
 - PETSc FieldSplit



^aKatz

Magma Dynamics

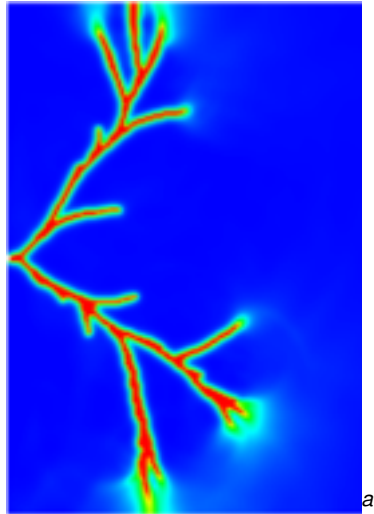
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^aKatz, Spiegelman

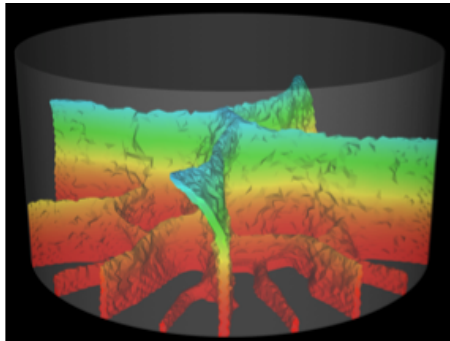
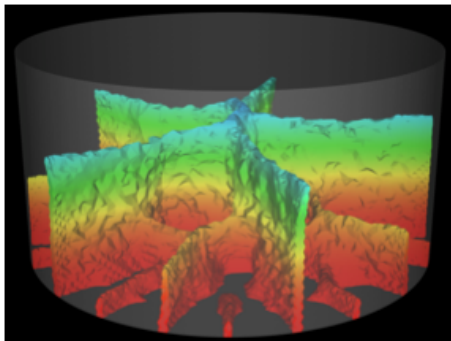
Fracture Mechanics

- Full variational formulation
 - Phase field
 - Linear or Quadratic penalty
- Uses TAO optimization
 - Necessary for linear penalty
 - Backtacking
- No prescribed cracks (**movie**)
 - Arbitrary crack geometry
 - Arbitrary intersections
- Multiple materials
 - Composite toughness



^aBourdin

Fracture Mechanics

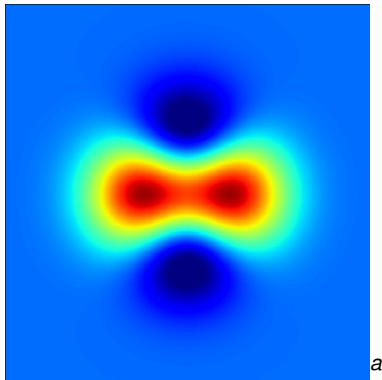


¹Bourdin

Vortex Method

$t = 000$

- Incompressible Flow
 - Gaussian vortex blobs
 - High Re
- PetFMM
 - 2D/3D domains
 - Automatic load balancing
 - Variety of kernels
 - Optimized with templates
- PetRBF
 - Variety of RBFs
 - Uses PETSc solvers
 - Scalable preconditioner
- Parallelism
 - MPI
 - GPU

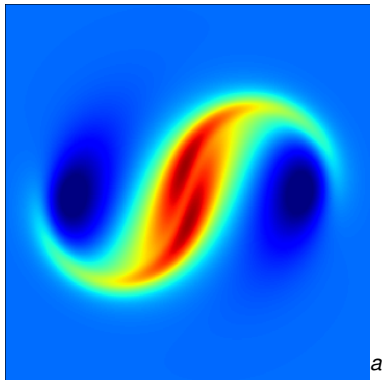


^aCruz, Yokota, Barba, Knepley

Vortex Method

$t = 100$

- Incompressible Flow
 - Gaussian vortex blobs
 - High Re
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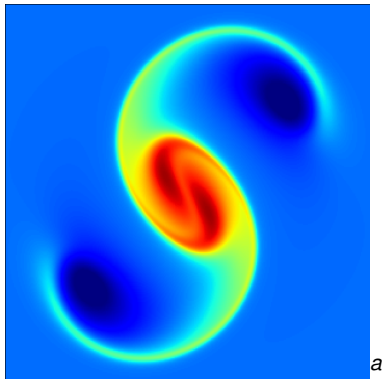


^aCruz, Yokota, Barba, Knepley

Vortex Method

$t = 200$

- Incompressible Flow
 - Gaussian vortex blobs
 - High Re
- PetFMM
 - 2D/3D domains
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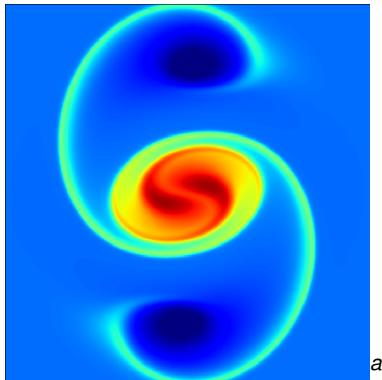


^aCruz, Yokota, Barba, Knepley

Vortex Method

$t = 300$

- Incompressible Flow
 - Gaussian vortex blobs
 - High Re
- PetFMM
 - 2D/3D domains
 - Automatic load balancing
 - Variety of kernels
 - Optimized with templates
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 - Variety of RBFs
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 - MPI
 - GPU

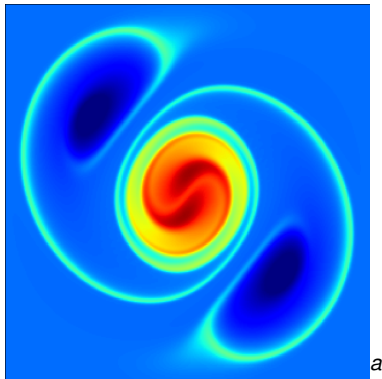


^aCruz, Yokota, Barba, Knepley

Vortex Method

$t = 400$

- Incompressible Flow
 - Gaussian vortex blobs
 - High Re
- PetFMM
 - 2D/3D domains
 - Automatic load balancing
 - Variety of kernels
 - Optimized with templates
- PetRBF
 - Variety of RBFs
 - Uses PETSc solvers
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 - GPU

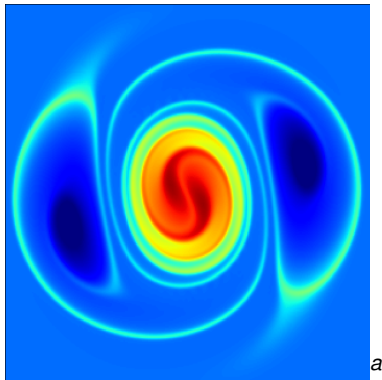


^aCruz, Yokota, Barba, Knepley

Vortex Method

$t = 500$

- Incompressible Flow
 - Gaussian vortex blobs
 - High Re
- PetFMM
 - 2D/3D domains
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 - Variety of kernels
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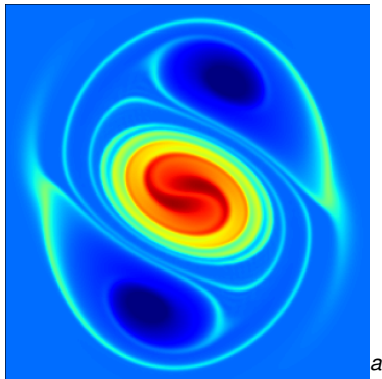


^aCruz, Yokota, Barba, Knepley

Vortex Method

$t = 600$

- Incompressible Flow
 - Gaussian vortex blobs
 - High Re
- PetFMM
 - 2D/3D domains
 - Automatic load balancing
 - Variety of kernels
 - Optimized with templates
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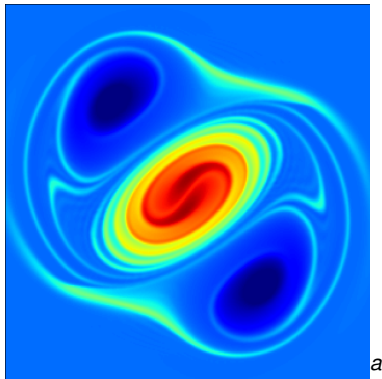


^aCruz, Yokota, Barba, Knepley

Vortex Method

$t = 700$

- Incompressible Flow
 - Gaussian vortex blobs
 - High Re
- PetFMM
 - 2D/3D domains
 - Automatic load balancing
 - Variety of kernels
 - Optimized with templates
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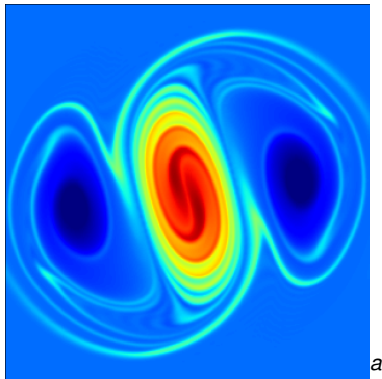
a

^aCruz, Yokota, Barba, Knepley

Vortex Method

$t = 800$

- Incompressible Flow
 - Gaussian vortex blobs
 - High Re
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 - 2D/3D domains
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 - Variety of kernels
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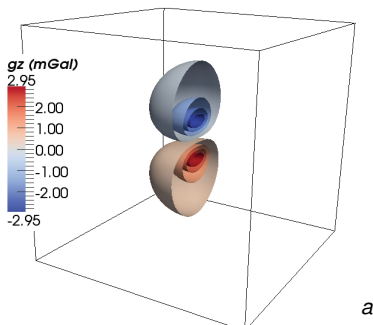


^a

^aCruz, Yokota, Barba, Knepley

Gravity Anomaly Modeling

- Potential Solution
 - Kernel of inverse problem
 - Needs optimal algorithm
- Implementations
 - Direct Summation
 - FEM
 - FMM
- Parallelism
 - MPI
 - 4000+ cores
 - All methods scalable

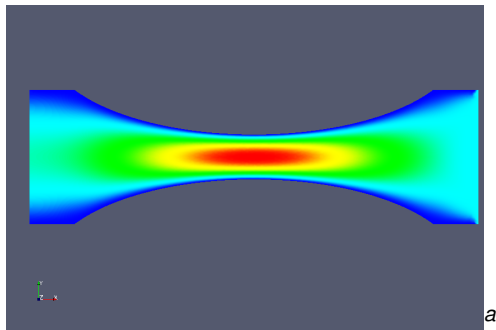


^aMay, Knepley

FEniCS-Apps

Rheagen

- Rheologies
 - Maxwell
 - Grade 2
 - Oldroyd-B
- Stabilization
 - DG
 - SUPG
 - EVSS
 - DEVSS
 - Macroelement
- Automation
 - FIAT (elements)
 - FFC (weak forms)

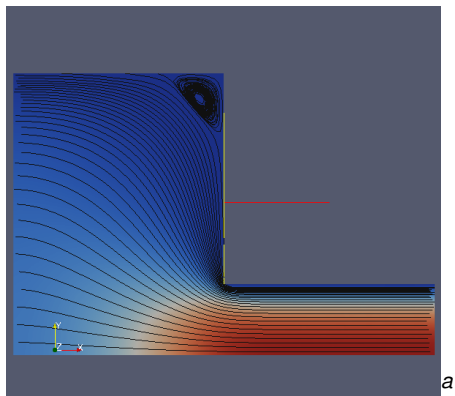


^aTerrel

FEniCS-Apps

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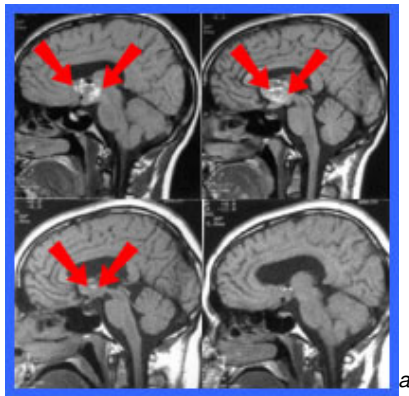
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^aTerrel

Real-time Surgery

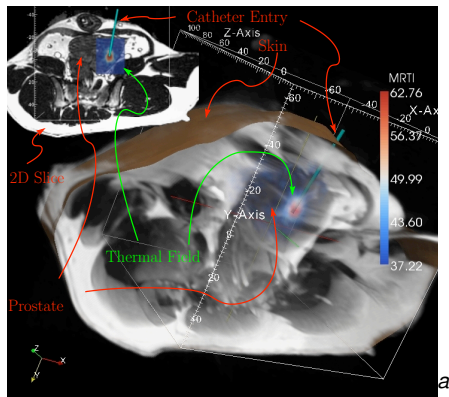
- Brain Surgery
 - Elastic deformation
 - Overlaid on MRI
 - Guides surgeon
- Laser Thermal Therapy
 - PDE constrained optimization
 - Per-patient calibration
 - Thermal inverse problem



^aWarfield, Ferrant, et.al.

Real-time Surgery

- Brain Surgery
 - Elastic deformation
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^aFuentes, Oden, et.al.

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Questions for Windows Users

- Have you installed cygwin?
 - Need python, make, and build-utils packages
- Will you use the GNU compilers?
 - If not, remove `link.exe`
 - If MS, check compilers from `cmd` window and use `win32fe`
- Which MPI will you use?
 - You can use `--with-mpi=0`
 - If MS, need to install MPICH2
 - If GNU, can use `--download-mpich`

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Downloading PETSc

- The latest tarball is on the PETSc site:
<http://www.mcs.anl.gov/petsc/download>
- There is a **Debian package** (`aptitude install petsc-dev`)
- There is a **Git development repository**

Cloning PETSc

- The full development repository is open to the public
 - <https://bitbucket.org/petsc/petsc/>
- Why is this better?
 - You can clone to any release (or any specific ChangeSet)
 - You can easily rollback changes (or releases)
 - You can get fixes from us the same day
 - You can easily submit changes using a pull request
- All releases are just tags:
 - [Source at tag v3.7.2](#)

Unpacking PETSc

- Just clone development repository

- `git clone http://bitbucket.org/petsc/petsc.git`
- `git checkout -rv3.7.2`

or

- Unpack the tarball

- `tar xzf petsc.tar.gz`

Exercise 1

Download and Unpack PETSc!

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Configuring PETSc

- Set `$PETSC_DIR` to the installation root directory
- Run the configuration utility
 - `$PETSC_DIR/configure`
 - `$PETSC_DIR/configure --help`
 - `$PETSC_DIR/configure --download-mpich`
 - `$PETSC_DIR/configure --prefix=/usr`
- There are many examples on the installation page
- Config files in `$PETSC_DIR/$PETSC_ARCH/lib/petsc/conf`
 - Config header in `$PETSC_DIR/$PETSC_ARCH/include`
 - `$PETSC_ARCH` has a default if not specified

Configuring PETSc

- You can easily reconfigure with the same options
 - `./$PETSC_ARCH/lib/petsc/conf/reconfigure-$PETSC_ARCH.py`
- Can maintain several different configurations
 - `./configure -PETSC_ARCH=linux-fast --with-debugging=0`
- All configuration information is in the logfile
 - `./$PETSC_ARCH/lib/petsc/conf/configure.log`
 - **ALWAYS** send this file with bug reports

Configuring PETSc for FEM

`$PETSC_DIR/configure`

`–download-triangle –download-ctetgen –download-p4est`

`–download-chaco –download-metis –download-parmetis`

`–download-pragmatic`

`–download-hdf5 –download-netcdf –download-exodusii`

`–download-gmp –download-mpfr`

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Configuring PETSc for Accelerators

`$PETSC_DIR/configure`

`-with-cuda`

`-with-cudac='nvcc -m64' -with-cuda-arch=sm_10`

`-with-cuda-only`

`-with-opengl`

`-with-opengl-include=/System/Library/Frameworks/OpenCL.framework/Headers/`

`-with-opengl-lib=/System/Library/Frameworks/OpenCL.framework/OpenCL`

`-with-precision=single`

Configuring PETSc for Accelerators

```
$PETSC_DIR/configure
```

```
–with-cuda
```

```
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`-with-opengl`

`-with-opengl-include=/System/Library/Frameworks/OpenCL.framework/Headers/`

`-with-opengl-lib=/System/Library/Frameworks/OpenCL.framework/OpenCL`

`-with-precision=single`

Configuring PETSc for Accelerators

`$PETSC_DIR/configure`

`-with-cuda`

`-with-cudac='nvcc -m64' -with-cuda-arch=sm_10`

`-with-cuda-only`

`-with-opengl`

`-with-opengl-include=/System/Library/Frameworks/OpenCL.framework/Headers/`

`-with-opengl-lib=/System/Library/Frameworks/OpenCL.framework/OpenCL`

`-with-precision=single`

Configuring PETSc for Accelerators

`$PETSC_DIR/configure`

`–with-cuda`

`–with-cudac='nvcc -m64' –with-cuda-arch=sm_10`

`–with-cuda-only`

`–with-ocl`

`–with-ocl-include=/System/Library/Frameworks/OpenCL.framework/Headers/`

`–with-ocl-lib=/System/Library/Frameworks/OpenCL.framework/OpenCL`

`–with-precision=single`

Configuring PETSc for Accelerators

`$PETSC_DIR/configure`

`-with-cuda`

`-with-cudac='nvcc -m64' -with-cuda-arch=sm_10`

`-with-cuda-only`

`-with-ocl`

`-with-ocl-include=/System/Library/Frameworks/OpenCL.framework/Headers/`

`-with-ocl-lib=/System/Library/Frameworks/OpenCL.framework/OpenCL`

`-with-precision=single`

Automatic Downloads

- Starting in 2.2.1, some packages are automatically
 - Downloaded
 - Configured and Built (in `$PETSC_DIR/externalpackages`)
 - Installed with PETSc
- Currently works for
 - petsc4py, mpi4py
 - PETSc documentation utilities (Sowing, c2html)
 - BLAS, LAPACK, Elemental, ScaLAPACK
 - MPICH, OpenMPI
 - ParMetis, Chaco, Jostle, Party, Scotch, Zoltan
 - SuiteSparse, MUMPS, SuperLU, SuperLU_Dist, PaStiX, Pardiso
 - HYPRE, ML
 - BLOPEX, FFTW, STRUMPACK, SPAI, CUSP, Sundials
 - Triangle, TetGen, p4est, Pragmatic
 - HDF5, NetCDF, ExodusII
 - AfterImage, gifLib, libjpeg, opengl
 - GMP, MPFR
 - ConcurrencyKit, hwloc

Exercise 2

Configure your downloaded PETSc.

Outline

- 1 Getting Started with PETSc
 - Who uses PETSc?
 - Stuff for Windows
 - How can I get PETSc?
 - How do I Configure PETSc?
 - **How do I Build PETSc?**
 - How do I run an example?
 - How do I get more help?

Building PETSc

- There is now One True Way to build PETSc:
 - `make`
 - `make install` if you configured with `--prefix`
 - Check build when done with `make test`
- Can build multiple configurations
 - `PETSC_ARCH=linux-fast make`
 - Libraries are in `$PETSC_DIR/$PETSC_ARCH/lib/`
- Complete log for each build is in logfile
 - `./$PETSC_ARCH/lib/petsc/conf/make.log`
 - **ALWAYS** send this with bug reports

Exercise 3

Build your configured PETSc.

Exercise 4

Reconfigure PETSc to use ParMetis.

- 1 `linux-debug/lib/petsc/conf/reconfigure-linux-debug.py`
 - `--PETSC_ARCH=linux-parmetis`
 - `--download-metis --download-parmetis`
- 2 `PETSC_ARCH=linux-parmetis make`
- 3 `PETSC_ARCH=linux-parmetis make test`

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Running PETSc

- Try running PETSc examples first
 - `cd $PETSC_DIR/src/snes/examples/tutorials`
- Build examples using make targets
 - `make ex5`
- Run examples using the make target
 - `make runex5`
- Can also run using MPI directly
 - `mpirun ./ex5 -snes_max_it 5`
 - `mpiexec ./ex5 -snes_monitor`

Running PETSc with Python

- Can run any PETSc example

- `./config/builder2.py check ./src/snes/examples/tutorials/ex5.c`

- Checks against test output

- Ignores if no output is present

- Can specify multiple files

- `builder2.py check [./src/snes/examples/tutorials/ex5.c,code.c]`

- Can also run using MPI directly

- Use `--retain` to keep executable

- `mpiexec ./PETSC_ARCH/lib/lib-ex5/ex5 -snes_monitor`

Using MPI

- The **M**essage **P**assing **I**nterface is:
 - a library for parallel communication
 - a system for launching parallel jobs (mpirun/mpiexec)
 - a community standard
- Launching jobs is easy
 - `mpiexec -n 4 ./ex5`
- You should never have to make MPI calls when using PETSc
 - Almost never

MPI Concepts

- Communicator
 - A context (or scope) for parallel communication (“Who can I talk to”)
 - There are two defaults:
 - yourself (PETSC_COMM_SELF),
 - and everyone launched (PETSC_COMM_WORLD)
 - Can create new communicators by splitting existing ones
 - Every PETSc object has a communicator
 - Set PETSC_COMM_WORLD to put all of PETSc in a subcomm
- Point-to-point communication
 - Happens between two processes (like in `MatMult()`)
- Reduction or scan operations
 - Happens among all processes (like in `VecDot()`)

Common Viewing Options

- Gives a text representation
 - `-vec_view`
- Generally views subobjects too
 - `-snes_view`
- Can visualize some objects
 - `-mat_view draw::`
- Alternative formats
 - `-vec_view binary:sol.bin::`, `-vec_view ::matlab`, `-vec_view socket`
- Sometimes provides extra information
 - `-mat_view ::ascii_info`, `-mat_view ::ascii_info_detailed`
- Use `-help` to see all options

Common Monitoring Options

- Display the residual
 - `-ksp_monitor`, **graphically** `-ksp_monitor_draw`
- Can disable dynamically
 - `-ksp_monitors_cancel`
- Does not display subsolvers
 - `-snes_monitor`
- Can use the true residual
 - `-ksp_monitor_true_residual`
- Can display different subobjects
 - `-snes_monitor_residual`, `-snes_monitor_solution`,
`-snes_monitor_solution_update`
 - `-snes_monitor_range`
 - `-ksp_gmres_krylov_monitor`
- Can display the spectrum
 - `-ksp_monitor_singular_value`

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 - How do I Build PETSc?
 - How do I run an example?
 - How do I get more help?

Getting More Help

- <http://www.mcs.anl.gov/petsc>
- Hyperlinked documentation
 - Manual
 - Manual pages for every method
 - HTML of all example code (linked to manual pages)
- FAQ
- Full support at petsc-maint@mcs.anl.gov
- High profile users
 - David Keyes
 - Marc Spiegelman
 - Richard Katz
 - Brad Aagaard
 - Aron Ahmadi
 - Patrick Farrell
 - Gerard Gorman

Outline

1 Getting Started with PETSc

2 PETSc Integration

- Initial Operations
- Vector Algebra
- Matrix Algebra
- Algebraic Solvers
- Debugging PETSc
- Profiling PETSc
- Serial Performance

3 DM

4 Advanced Solvers

Outline

- 2 **PETSc Integration**
 - **Initial Operations**
 - Vector Algebra
 - Matrix Algebra
 - Algebraic Solvers
 - Debugging PETSc
 - Profiling PETSc
 - Serial Performance

Application Integration

- Be willing to experiment with algorithms
 - No optimality without interplay between physics and algorithmics
- Adopt flexible, extensible programming
 - Algorithms and data structures not hardwired
- Be willing to play with the real code
 - Toy models are rarely helpful
- If possible, profile before integration
 - Automatic in PETSc

PETSc Integration

PETSc is a set a library interfaces

- We do not seize `main()`
- We do not control output
- We propagate errors from underlying packages
- We present the same interfaces in:
 - C
 - C++
 - F77
 - F90
 - Python

See Gropp in [SIAM, OO Methods for Interop SciEng, '99](#)

Integration Stages

- **Version Control**
 - It is impossible to overemphasize
 - We use **Git**
- **Initialization**
 - Linking to PETSc
- **Profiling**
 - Profile **before** changing
 - Also incorporate command line processing
- **Linear Algebra**
 - First PETSc data structures
- **Solvers**
 - Very easy after linear algebra is integrated

Initialization

- Call `PetscInitialize ()`
 - Setup static data and services
 - Setup MPI if it is not already
- Call `PetscFinalize ()`
 - Calculates logging summary
 - Shutdown and release resources
- Checks compile and link

Profiling

- Use `-log_view` for a performance profile
 - Event timing
 - Event flops
 - Memory usage
 - MPI messages

This used to be `-log_summary`

- Call `PetscLogStagePush()` and `PetscLogStagePop()`
 - User can add new stages
- Call `PetscLogEventBegin()` and `PetscLogEventEnd()`
 - User can add new events

Command Line Processing

- Check for an option
 - PetscOptionsHasName()
- Retrieve a value
 - PetscOptionsGetInt(), PetscOptionsGetIntArray()
- Set a value
 - PetscOptionsSetValue()
- Check for unused options
 - `-options_left`
- Clear, alias, reject, etc.
- Modern form uses
 - PetscOptionsBegin(), PetscOptionsEnd()
 - PetscOptionsInt(), PetscOptionsReal()
 - Integrates with `-help`

Outline

- 2 PETSc Integration
 - Initial Operations
 - **Vector Algebra**
 - Matrix Algebra
 - Algebraic Solvers
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 - Profiling PETSc
 - Serial Performance

Vector Algebra

What are PETSc vectors?

- Fundamental objects representing
 - solutions
 - right-hand sides
 - coefficients
- Each process locally owns a subvector of contiguous global data

Vector Algebra

How do I create vectors?

- `VecCreate(MPI_Comm, Vec*)`
- `VecSetSizes(Vec, PetscIntn, PetscInt N)`
- `VecSetType(Vec, VecType typeName)`
- `VecSetFromOptions(Vec)`
 - Can set the type at runtime

Vector Algebra

A PETSc Vec

- Supports all vector space operations
 - `VecDot()`, `VecNorm()`, `VecScale()`
- Has a direct interface to the values
 - `VecGetArray()`, `VecGetArrayF90()`
- Has unusual operations
 - `VecSqrtAbs()`, `VecStrideGather()`
- Communicates automatically during assembly
- Has customizable communication (`PetscSF`, `VecScatter`)

Parallel Assembly

Vectors and Matrices

- Processes may set an arbitrary entry
 - Must use proper interface
- Entries need not be generated locally
 - Local meaning the process on which they are stored
- PETSc automatically moves data if necessary
 - Happens during the assembly phase

Vector Assembly

- A three step process
 - Each process sets or adds values
 - Begin communication to send values to the correct process
 - Complete the communication

```
VecSetValues(Vec v, PetscInt n, PetscInt rows[],  
            PetscScalar values[], InsertMode mode)
```

- Mode is either INSERT_VALUES or ADD_VALUES
- Two phases allow overlap of communication and computation
 - VecAssemblyBegin(v)
 - VecAssemblyEnd(v)

One Way to Set the Elements of a Vector

```
VecGetSize(x, &N);
MPI_Comm_rank(PETSC_COMM_WORLD, &rank);
if (rank == 0) {
    val = 0.0;
    for(i = 0; i < N; ++i) {
        VecSetValues(x, 1, &i, &val, INSERT_VALUES);
        val += 10.0;
    }
}
/* These routines ensure that the data is
   distributed to the other processes */
VecAssemblyBegin(x);
VecAssemblyEnd(x);
```

A Better Way to Set the Elements of a Vector

```
VecGetOwnershipRange(x, &low, &high);
val = low*10.0;
for(i = low; i < high; ++i) {
    VecSetValues(x, 1, &i, &val, INSERT_VALUES);
    val += 10.0;
}
/* No data will be communicated here */
VecAssemblyBegin(x);
VecAssemblyEnd(x);
```

Selected Vector Operations

Function Name	Operation
VecAXPY(Vec y, PetscScalar a, Vec x)	$y = y + a * x$
VecAYPX(Vec y, PetscScalar a, Vec x)	$y = x + a * y$
VecWAYPX(Vec w, PetscScalar a, Vec x, Vec y)	$w = y + a * x$
VecScale(Vec x, PetscScalar a)	$x = a * x$
VecCopy(Vec y, Vec x)	$y = x$
VecPointwiseMult(Vec w, Vec x, Vec y)	$w_i = x_i * y_i$
VecMax(Vec x, PetscInt *idx, PetscScalar *r)	$r = \max r_i$
VecShift(Vec x, PetscScalar r)	$x_i = x_i + r$
VecAbs(Vec x)	$x_i = x_i $
VecNorm(Vec x, NormType type, PetscReal *r)	$r = x $

Working With Local Vectors

It is sometimes more efficient to directly access local storage of a `Vec`.

- PETSc allows you to access the local storage with
 - `VecGetArray(Vec, double *[])`
- You must return the array to PETSc when you finish
 - `VecRestoreArray(Vec, double *[])`
- Allows PETSc to handle data structure conversions
 - Commonly, these routines are fast and do not involve a copy

VecGetArray in C

```
Vec          v;  
PetscScalar *array;  
PetscInt    n, i;  
  
VecGetArray(v, &array);  
VecGetLocalSize(v, &n);  
PetscSynchronizedPrintf(PETSC_COMM_WORLD,  
    "First element of local array is %f\n", array[0]);  
PetscSynchronizedFlush(PETSC_COMM_WORLD);  
for(i = 0; i < n; ++i) {  
    array[i] += (PetscScalar) rank;  
}  
VecRestoreArray(v, &array);
```

VecGetArray in F77

```
#include "finclude/petsc.h"

Vec          v;
PetscScalar  array(1)
PetscOffset  offset
PetscInt     n, i
PetscErrorCode ierr

call VecGetArray(v, array, offset, ierr)
call VecGetLocalSize(v, n, ierr)
do i=1,n
  array(i+offset) = array(i+offset) + rank
end do
call VecRestoreArray(v, array, offset, ierr)
```

VecGetArray in F90

```
#include "finclude/petsc.h90"

Vec          v;
PetscScalar  pointer :: array(:)
PetscInt     n, i
PetscErrorCode ierr

call VecGetArrayF90(v, array, ierr)
call VecGetLocalSize(v, n, ierr)
do i=1,n
  array(i) = array(i) + rank
end do
call VecRestoreArrayF90(v, array, ierr)
```

VecGetArray in Python

```
with v as a:  
    for i in range(len(a)):  
        a[i] = 5.0*i
```

DMDAVecGetArray in C

```
DM          da;
Vec         v;
DMDALocalInfo *info;
PetscScalar **array;

DMDAVecGetArray(da, v, &array);
for(j = info->ys; j < info->ys+info->ym; ++j) {
  for(i = info->xs; i < info->xs+info->xm; ++i) {
    u      = x[j][i];
    uxx    = (2.0*u - x[j][i-1] - x[j][i+1])*hydhx;
    uyy    = (2.0*u - x[j-1][i] - x[j+1][i])*hxdhy;
    f[j][i] = uxx + uyy;
  }
}
DMDAVecRestoreArray(da, v, &array);
```

DMDAVecGetArray in F90

```

DM          da
Vec         v
PetscScalar, pointer :: array (:, :)

call DMDAGetCorners(ada, xs, ys, PETSC_NULL_INTEGER,
                  xm, ym, PETSC_NULL_INTEGER, ierr)
call DMDAVecGetArrayF90(da, v, array, ierr);
do i = xs, xs+xm
  do j = ys, ys+ym
    u      = x(i, j)
    uxx    = (2.0*u - x(i-1, j) - x(i+1, j))*hydhx;
    uyy    = (2.0*u - x(i, j-1) - x(i, j+1))*hxdhy;
    f(i, j) = uxx + uyy;
  enddo
enddo
call DMDAVecRestoreArrayF90(da, v, array, ierr);

```

Outline

2 PETSc Integration

- Initial Operations
- Vector Algebra
- **Matrix Algebra**
- Algebraic Solvers
- Debugging PETSc
- Profiling PETSc
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Matrix Algebra

What are PETSc matrices?

- Fundamental objects for storing stiffness matrices and Jacobians
- Each process locally owns a contiguous set of rows
- Supports many data types
 - AIJ, Block AIJ, Symmetric AIJ, Block Matrix, etc.
- Supports structures for many packages
 - MUMPS, Spooles, SuperLU, UMFPack, DSCPack

How do I create matrices?

- `MatCreate(MPI_Comm, Mat*)`
- `MatSetSizes(Mat, PetscInt m, PetscInt n, M, N)`
- `MatSetType(Mat, MatType typeName)`
- `MatSetFromOptions(Mat)`
 - Can set the type at runtime
- `MatSeqAIJPreallocation(Mat, PetscInt nz, const PetscInt nnz[])`
- `MatXAIJPreallocation(Mat, bs, dnz[], onz[], dnzu[], onzu[])`
- `MatSetValues(Mat, m, rows[], n, cols [], values [], InsertMode)`
 - **MUST** be used, but does automatic communication

Matrix Polymorphism

The PETSc `Mat` has a single user interface,

- Matrix assembly
 - `MatSetValues()`
 - `MatGetLocalSubMatrix()`
- Matrix-vector multiplication
 - `MatMult()`
- Matrix viewing
 - `MatView()`

but multiple underlying implementations.

- AIJ, Block AIJ, Symmetric Block AIJ,
- Dense
- Matrix-Free
- etc.

A matrix is defined by its **interface**, not by its **data structure**.

Matrix Assembly

- A three step process
 - Each process sets or adds values
 - Begin communication to send values to the correct process
 - Complete the communication
- `MatSetValues(Matm, m, rows[], n, cols [], values [], mode)`
 - mode is either `INSERT_VALUES` or `ADD_VALUES`
 - Logically dense block of values
- Two phase assembly allows overlap of communication and computation
 - `MatAssemblyBegin(Matm, MatAssemblyType type)`
 - `MatAssemblyEnd(Matm, MatAssemblyType type)`
 - type is either `MAT_FLUSH_ASSEMBLY` or `MAT_FINAL_ASSEMBLY`

One Way to Set the Elements of a Matrix

Simple 3-point stencil for 1D Laplacian

```
v[0] = -1.0; v[1] = 2.0; v[2] = -1.0;
if (rank == 0) {
  for(row = 0; row < N; row++) {
    cols[0] = row-1; cols[1] = row; cols[2] = row+1;
    if (row == 0) {
      MatSetValues(A,1,&row,2,&cols[1],&v[1],INSERT_VALUES);
    } else if (row == N-1) {
      MatSetValues(A,1,&row,2,cols,v,INSERT_VALUES);
    } else {
      MatSetValues(A,1,&row,3,cols,v,INSERT_VALUES);
    }
  }
}
MatAssemblyBegin(A, MAT_FINAL_ASSEMBLY);
MatAssemblyEnd(A, MAT_FINAL_ASSEMBLY);
```

Parallel Sparse Matrix Layout



A Better Way to Set the Elements of a Matrix

Simple 3-point stencil for 1D Laplacian

```
v[0] = -1.0; v[1] = 2.0; v[2] = -1.0;
MatGetOwnershipRange(A,&start,&end);
for(row = start; row < end; row++) {
  cols[0] = row-1; cols[1] = row; cols[2] = row+1;
  if (row == 0) {
    MatSetValues(A,1,&row,2,&cols[1],&v[1],INSERT_VALUES);
  } else if (row == N-1) {
    MatSetValues(A,1,&row,2,cols,v,INSERT_VALUES);
  } else {
    MatSetValues(A,1,&row,3,cols,v,INSERT_VALUES);
  }
}
MatAssemblyBegin(A, MAT_FINAL_ASSEMBLY);
MatAssemblyEnd(A, MAT_FINAL_ASSEMBLY);
```

Why Are PETSc Matrices That Way?

- No one data structure is appropriate for all problems
 - Blocked and diagonal formats provide performance benefits
 - PETSc has many formats
 - Makes it easy to add new data structures
- Assembly is difficult enough without worrying about partitioning
 - PETSc provides parallel assembly routines
 - High performance still requires making most operations local
 - However, programs can be incrementally developed.
 - [MatPartitioning](#) and [MatOrdering](#) can help
 - Its better to partition and reorder the underlying grid
- Matrix decomposition in contiguous chunks is simple
 - Makes interoperation with other codes easier
 - For other ordering, PETSc provides “Application Orderings” (AO)

Outline

2 PETSc Integration

- Initial Operations
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- Matrix Algebra
- **Algebraic Solvers**
- Debugging PETSc
- Profiling PETSc
- Serial Performance

Experimentation is Essential!

Proof is not currently enough to examine solvers

- N. M. Nachtigal, S. C. Reddy, and L. N. Trefethen, *How fast are nonsymmetric matrix iterations?*, SIAM J. Matrix Anal. Appl., **13**, pp.778–795, 1992.
- Anne Greenbaum, Vlastimil Ptak, and Zdenek Strakos, *Any Nonincreasing Convergence Curve is Possible for GMRES*, SIAM J. Matrix Anal. Appl., **17** (3), pp.465–469, 1996.

Linear Solvers

Krylov Methods

- Using PETSc linear algebra, just add:
 - `KSPSetOperators(KSPksp, MatA, MatM, MatStructure flag)`
 - `KSPSolve(KSPksp, Vecb, Vecx)`
- Can access subobjects
 - `KSPGetPC(KSPksp, PC*pc)`
- Preconditioners must obey PETSc interface
 - Basically just the KSP interface
- Can change solver dynamically from the command line
 - `-ksp_type bicgstab`

Nonlinear Solvers

- Using PETSc linear algebra, just add:
 - `SNESSetFunction(SNESsnes, Vecr, residualFunc, void *ctx)`
 - `SNESSetJacobian(SNESsnes, MatA, MatM, jacFunc, void *ctx)`
 - `SNESsolve(SNESsnes, Vecb, Vecx)`
- Can access subobjects
 - `SNESGetKSP(SNESsnes, KSP*ksp)`
- Can customize subobjects from the cmd line
 - Set the subdomain preconditioner to ILU with `-sub_pc_type ilu`

Basic Solver Usage

Use `SNESSetFromOptions()` so that everything is set dynamically

- Set the type
 - Use `-snes_type` (or take the default)
- Set the preconditioner
 - Use `-npc_snes_type` (or take the default)
- Override the tolerances
 - Use `-snes_rtol` and `-snes_atol`
- View the solver to make sure you have the one you expect
 - Use `-snes_view`
- For debugging, monitor the residual decrease
 - Use `-snes_monitor`
 - Use `-ksp_monitor` to see the underlying linear solver

3rd Party Solvers in PETSc

Complete table of solvers

- Sequential LU
 - ESSL (IBM)
 - SuperLU (Sherry Li, LBNL)
 - Suitesparse (Tim Davis, U. of Florida)
 - LUSOL (MINOS, Michael Saunders, Stanford)
 - PILUT (Hypre, David Hysom, LLNL)
- Parallel LU
 - Elemental/Clique (Jack Poulson, Google)
 - MUMPS (Patrick Amestoy, IRIT)
 - SuperLU_Dist (Jim Demmel and Sherry Li, LBNL)
 - Pardiso (MKL, Intel)
 - STRUMPACK (Pieter Ghysels, LBNL)
- Parallel Cholesky
 - Elemental (Jack Poulson, Google)
 - DSCPACK (Padma Raghavan, Penn. State)
 - MUMPS (Patrick Amestoy, Toulouse)

3rd Party Preconditioners in PETSc

Complete table of solvers

- Parallel Algebraic Multigrid
 - GAMG (Mark Adams, LBNL)
 - BoomerAMG (Hypre, LLNL)
 - ML (Trilinos, Ray Tuminaro and Jonathan Hu, SNL)
- Parallel BDDC (Stefano Zampini, KAUST)
- Parallel ILU, PaStiX (Faverge Mathieu, INRIA)
- Parallel Redistribution (Dave May, Oxford)
- Parallel Sparse Approximate Inverse
 - Parasails (Hypre, Edmund Chow, LLNL)
 - SPAI 3.0 (Marcus Grote and Barnard, NYU)

User Solve

```
MPI_Comm comm;  
SNES snes;  
DM dm;  
Vec u;
```

```
SNESCreate(comm, &snest);  
SNESSetDM(snest, dm);  
SNESSetFromOptions(snest);  
DMCreateGlobalVector(dm, &u);  
SNESolve(snest, NULL, u);
```

Solver use in SNES ex62

Solver code does not change for different algorithms:

```
SNES          snes;
DM            dm;
Vec           u;
PetscErrorCode ierr;

ierr = SNESCreate(PETSC_COMM_WORLD, &snes);CHKERRQ(ierr);
ierr = SNESSetDM(snes, dm);CHKERRQ(ierr);
/* Specify residual computation */
ierr = SNESSetFromOptions(snes);CHKERRQ(ierr); /* Configure solver */
ierr = DMCreateGlobalVector(dm, &u);CHKERRQ(ierr);
ierr = SNESolve(snes, PETSC_NULL, u);CHKERRQ(ierr);
```

- **Never recompile!** all configuration is dynamic
- **DM** controls data layout and communication
- Type of nested solvers can be changed at runtime

Solver use in SNES ex62

I will omit error checking and declarations:

```
SNESCreate(PETSC_COMM_WORLD, &snes);
SNESSetDM(snes, dm);
/* Specify residual computation */
SNESSetFromOptions(snes); /* Configure solver */
DMCreateGlobalVector(dm, &u);
SNESolve(snes, PETSC_NULL, u);
```

Solver use in SNES ex62

The configuration API can also be used:

```
SNESCreate(PETSC_COMM_WORLD, &snes);
SNESSetDM(snes, dm);
/* Specify residual computation */
SNESNGMRESRestartType(snes, SNES_NGMRES_RESTART_PERIODIC);
SNESSetFromOptions(snes);
DMCreateGlobalVector(dm, &u);
SNESolve(snes, PETSC_NULL, u);
```

- Ignored when not applicable (no ugly check)
- Type safety of arguments is retained
- No downcasting

Solver use in SNES ex62

Adding a prefix namespaces command line options:

```
SNESCreate(PETSC_COMM_WORLD, &snes);
SNESSetDM(snes, dm);
/* Specify residual computation */
SNESSetOptionsPrefix(snes, "stokes_");
SNESSetFromOptions(snes);
DMCreateGlobalVector(dm, &u);
SNESolve(snes, PETSC_NULL, u);
```

`-stokes_snes_type qn` changes the solver type,
whereas `-snes_type qn` does not

Solver use in SNES ex62

User provides a function to compute the residual:

```
SNESCreate(PETSC_COMM_WORLD, &snes);  
SNESSetDM(snes, dm);  
DMCreateGlobalVector(dm, &r);  
SNESSetFunction(snes, r, FormFunction, &user);  
SNESSetFromOptions(snes);  
DMCreateGlobalVector(dm, &u);  
SNESolve(snes, PETSC_NULL, u);
```

$$r = F(u)$$

- User handles parallel communication
- User handles domain geometry and discretization

Solver use in SNES ex62

DM allows the user to compute only on a local patch:

```
SNESCreate(PETSC_COMM_WORLD, &snes);  
SNESSetDM(snes, dm);  
SNESSetFromOptions(snes);  
DMCreateGlobalVector(dm, &u);  
SNESolve(snes, PETSC_NULL, u);
```

```
DMSNESSetLocalFunction(dm, FormFunctionLocal);
```

- Code looks serial to the user
- PETSc handles global residual assembly
- Also works for unstructured meshes

Solver use in SNES ex62

Optionally, the user can also provide a Jacobian:

```
SNESCreate(PETSC_COMM_WORLD, &snes);  
SNESSetDM(snes, dm);  
SNESSetFromOptions(snes);  
DMCreateGlobalVector(dm, &u);  
SNESolve(snes, PETSC_NULL, u);
```

```
DMSNESSetLocalFunction(dm, FormFunctionLocal);  
DMSNESSetLocalJacobian(dm, FormJacobianLocal);
```

SNES ex62 allows both

- finite difference (JFNK), and
- FEM action

versions of the Jacobian.

Solver use in SNES ex62

The `DM` also handles storage:

```
CreateMesh(PETSC_COMM_WORLD, &user, &dm);  
DMCreateLocalVector(dm, &lu);  
DMCreateGlobalVector(dm, &u);  
DMCreateMatrix(dm, &J);
```

- `DM` can create local and global vectors
- Matrices are correctly preallocated
- Easy supported for discretization

Outline

2 PETSc Integration

- Initial Operations
- Vector Algebra
- Matrix Algebra
- Algebraic Solvers
- **Debugging PETSc**
- Profiling PETSc
- Serial Performance

Correctness Debugging

- Automatic generation of tracebacks
- Detecting memory corruption and leaks
- Optional user-defined error handlers

Interacting with the Debugger

- Launch the debugger

- `-start_in_debugger [gdb,dbx,noxterm]`
- `-on_error_attach_debugger [gdb,dbx,noxterm]`

- Attach the debugger only to some parallel processes

- `-debugger_nodes 0,1`

- Set the display (often necessary on a cluster)

- `-display khan.mcs.anl.gov:0.0`

Debugging Tips

- Put a breakpoint in `PetscError()` to catch errors as they occur
- PETSc tracks memory overwrites at both ends of arrays
 - The `CHKMEMQ` macro causes a check of all allocated memory
 - Track memory overwrites by bracketing them with `CHKMEMQ`
- PETSc checks for leaked memory
 - Use `PetscMalloc()` and `PetscFree()` for all allocation
 - Print unfreed memory on `PetscFinalize()` with `-malloc_dump`
- Simply the best tool today is **valgrind**
 - It checks memory access, cache performance, memory usage, etc.
 - <http://www.valgrind.org>
 - Need `--trace-children=yes` when running under MPI

Exercise 7

Use the debugger to find a SEGV Locate a memory overwrite using CHKMEMQ.

- Get the example

- `hg clone -r1 http://petsc.cs.iit.edu/petsc/SimpleTutorial`

- Build the example `make`

- Run it and watch the fireworks

- `mpiexec -n 2 ./bin/ex5 -use_coords`

- Run it under the debugger and correct the error

- `mpiexec -n 2 ./bin/ex5 -use_coords -start_in_debugger`

- `hg update -r2`

- Build it and run again smoothly

Outline

2 PETSc Integration

- Initial Operations
- Vector Algebra
- Matrix Algebra
- Algebraic Solvers
- Debugging PETSc
- **Profiling PETSc**
- Serial Performance

Performance Debugging

- PETSc has integrated profiling
 - Option `-log_view` prints a report on `PetscFinalize()`
- PETSc allows user-defined events
 - Events report time, calls, flops, communication, etc.
 - Memory usage is tracked by object
- Profiling is separated into stages
 - Event statistics are aggregated by stage

Using Stages and Events

- Use `PetscLogStageRegister()` to create a new stage
 - Stages are identifier by an integer handle
- Use `PetscLogStagePush/Pop()` to manage stages
 - Stages may be nested, but will not aggregate in a nested fashion
- Use `PetscLogEventRegister()` to create a new stage
 - Events also have an associated class
- Use `PetscLogEventBegin/End()` to manage events
 - Events may also be nested and will aggregate in a nested fashion
 - Can use `PetscLogFlops()` to log user flops

Adding A Logging Stage

C

```
int stageNum ;

PetscLogStageRegister (&stageNum , "name" );
PetscLogStagePush (stageNum );

/* Code to Monitor */

PetscLogStagePop ();
```

Adding A Logging Stage

Python

```
with PETSc.LogStage('Fluid Stage') as fluidStage:  
    # All operations will be aggregated in fluidStage  
    fluid.solve()
```

Adding A Logging Event

C

```
static int USER_EVENT;
```

```
PetscLogEventRegister(&USER_EVENT, "name", CLS_ID);  
PetscLogEventBegin(USER_EVENT,0,0,0,0);
```

```
/* Code to Monitor */
```

```
PetscLogFlops(user_event_flops);  
PetscLogEventEnd(USER_EVENT,0,0,0,0);
```

Adding A Logging Event

Python

```
with PETSc.logEvent('Reconstruction') as recEvent:  
    # All operations are timed in recEvent  
    reconstruct(sol)  
    # Flops are logged to recEvent  
    PETSc.Log.logFlops(user_event_flops)
```

Adding A Logging Class

```
static int CLASS_ID;
```

```
PetscLogClassRegister(&CLASS_ID, "name");
```

- Class ID identifies a class uniquely
- Must initialize before creating any objects of this type

Matrix Memory Preallocation

- PETSc sparse matrices are dynamic data structures
 - can add additional nonzeros freely
- Dynamically adding many nonzeros
 - requires additional memory allocations
 - requires copies
 - can kill performance
- Memory preallocation provides
 - the freedom of dynamic data structures
 - good performance
- Easiest solution is to replicate the assembly code
 - Remove computation, but preserve the indexing code
 - Store set of columns for each row
- Call preallocation routines for all datatypes
 - `MatSeqAIJSetPreallocation()`
 - `MatMPIAIJSetPreallocation()`
 - Only the relevant data will be used

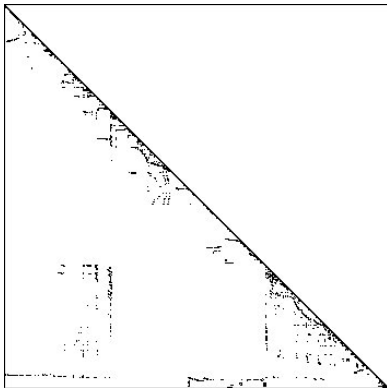
Matrix Memory Preallocation

Sequential Sparse Matrices

`MatSeqAIJPreallocation(MatA, int nz, int nnz[])`

`nz`: expected number of nonzeros in any row

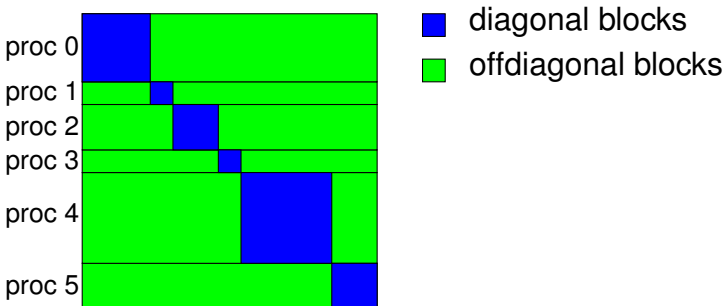
`nnz(i)`: expected number of nonzeros in row `i`



Matrix Memory Preallocation

ParallelSparseMatrix

- Each process locally owns a submatrix of contiguous global rows
- Each submatrix consists of diagonal and off-diagonal parts



- `MatGetOwnershipRange(MatA,int *start,int *end)`

`start`: first locally owned row of global matrix

`end-1`: last locally owned row of global matrix

Matrix Memory Preallocation

Parallel Sparse Matrices

`MatMPIAIJPreallocation(MatA, int dnz, int dnnz[], int onz, int onnz[])`

`dnz`: expected number of nonzeros in any row in the diagonal block

`dnnz(i)`: expected number of nonzeros in row i in the diagonal block

`onz`: expected number of nonzeros in any row in the offdiagonal portion

`onnz(i)`: expected number of nonzeros in row i in the offdiagonal portion

Matrix Memory Preallocation

Verifying Preallocation

- Use runtime option `-info`

- Output:

```
[proc #] Matrix size:  %d X %d; storage space:  
%d unneeded, %d used
```

```
[proc #] Number of mallocs during MatSetValues( )  
is %d
```

```
[merlin] mpirun ex2 -log_info  
[0]MatAssemblyEnd_SeqAIJ:Matrix size: 56 X 56; storage space:  
[0] 310 unneeded, 250 used  
[0]MatAssemblyEnd_SeqAIJ:Number of mallocs during MatSetValues() is 0  
[0]MatAssemblyEnd_SeqAIJ:Most nonzeros in any row is 5  
[0]Mat_AIJ_CheckInode: Found 56 nodes out of 56 rows. Not using Inode routine  
[0]Mat_AIJ_CheckInode: Found 56 nodes out of 56 rows. Not using Inode routine  
Norm of error 0.000156044 iterations 6  
[0]PetscFinalize:PETSc successfully ended!
```


Exercise 8

Return to Exercise 7 and add more profiling.

- Update to the next revision
 - `hg update -r3`
- Build, run, and look at the profiling report
 - `make ex5`
 - `./bin/ex5 -use_coords -log_summary`
- Add a new stage for setup
- Add a new event for `FormInitialGuess()` and log the flops
- Build it again and look at the profiling report

Outline

- 2 PETSc Integration
 - Initial Operations
 - Vector Algebra
 - Matrix Algebra
 - Algebraic Solvers
 - Debugging PETSc
 - Profiling PETSc
 - **Serial Performance**

Importance of Computational Modeling

Without a model,
performance measurements are meaningless!

Before a code is written, we should have a model of

- computation
- memory usage
- communication
- bandwidth
- achievable concurrency

This allows us to

- **verify** the implementation
- **predict** scaling behavior

Complexity Analysis

The key performance indicator, which we will call the *balance factor* β , is the ratio of **flops** executed to **bytes** transferred.

- We will designate the unit $\frac{\text{flop}}{\text{byte}}$ as the *Keyes*
- Using the peak flop rate r_{peak} , we can get the required bandwidth B_{req} for an algorithm

$$B_{\text{req}} = \frac{r_{\text{peak}}}{\beta} \quad (1)$$

- Using the peak bandwidth B_{peak} , we can get the maximum flop rate r_{max} for an algorithm

$$r_{\text{max}} = \beta B_{\text{peak}} \quad (2)$$

Performance Caveats

- The peak flop rate r_{peak} on modern CPUs is attained through the usage of a SIMD multiply-accumulate instruction on special 128-bit registers.
- SIMD MAC operates in the form of 4 simultaneous operations (2 adds and 2 multiplies):

$$c_1 = c_1 + a_1 * b_1 \quad (3)$$

$$c_2 = c_2 + a_2 * b_2 \quad (4)$$

You will miss peak by the corresponding number of operations you are missing. In the worst case, you are reduced to 25% efficiency if your algorithm performs naive summation or products.

- Memory alignment is also crucial when using SSE, the instructions used to load and store from the 128-bit registers throw very costly alignment exceptions when the data is not stored in memory on 16 byte (128 bit) boundaries.

Analysis of BLAS `axpy()`

$$\vec{y} \leftarrow \alpha \vec{x} + \vec{y}$$

For vectors of length N and b -byte numbers, we have

- Computation
 - $2N$ flops
- Memory Access
 - $(3N + 1)b$ bytes

Thus, our balance factor $\beta = \frac{2N}{(3N+1)b} \approx \frac{2}{3b}$ Keyes

Analysis of BLAS `axpy()`

$$\vec{y} \leftarrow \alpha \vec{x} + \vec{y}$$

For Matt's Laptop,

- $r_{\text{peak}} = 1700 \text{ MF/s}$
implies that
- $B_{\text{req}} = 2550b \text{ MB/s}$
 - Much greater than B_{peak}
- $B_{\text{peak}} = 1122 \text{ MB/s}$
implies that
- $r_{\text{max}} = \frac{748}{b} \text{ MF/s}$
 - 5.5% of r_{peak}

STREAM Benchmark

Simple benchmark program measuring **sustainable** memory bandwidth

- Prototypical operation is Triad (WAXPY): $\mathbf{w} = \mathbf{y} + \alpha\mathbf{x}$
- Measures the memory bandwidth bottleneck (much below peak)
- Datasets outstrip cache

Machine	Peak (MF/s)	Triad (MB/s)	MF/MW	Eq. MF/s
Matt's Laptop	1700	1122.4	12.1	93.5 (5.5%)
Intel Core2 Quad	38400	5312.0	57.8	442.7 (1.2%)
Tesla 1060C	984000	102000.0*	77.2	8500.0 (0.8%)

Table: Bandwidth limited machine performance

<http://www.cs.virginia.edu/stream/>

Analysis of Sparse Matvec (SpMV)

Assumptions

- No cache misses
- No waits on memory references

Notation

m Number of matrix rows

nz Number of nonzero matrix elements

V Number of vectors to multiply

We can look at bandwidth needed for peak performance

$$\left(8 + \frac{2}{V}\right) \frac{m}{nz} + \frac{6}{V} \text{ byte/flop} \quad (5)$$

or achievable performance given a bandwidth BW

$$\frac{Vnz}{(8V + 2)m + 6nz} BW \text{ Mflop/s} \quad (6)$$

Towards Realistic Performance Bounds for Implicit CFD Codes, Gropp, Kaushik, Keyes, and Smith.

Improving Serial Performance

For a single matvec with 3D FD Poisson, Matt's laptop can achieve at most

$$\frac{1}{(8 + 2)\frac{1}{7} + 6} \text{ bytes/flop} (1122.4 \text{ MB/s}) = \mathbf{151} \text{ MFlops/s}, \quad (7)$$

which is a dismal **8.8%** of peak.

Can improve performance by

- Blocking
- Multiple vectors

but operation issue limitations take over.

Improving Serial Performance

For a single matvec with 3D FD Poisson, Matt's laptop can achieve at most

$$\frac{1}{(8 + 2)^{\frac{1}{7}} + 6} \text{ bytes/flop} (1122.4 \text{ MB/s}) = 151 \text{ MFlops/s}, \quad (7)$$

which is a dismal 8.8% of peak.

Better approaches:

- Unassembled operator application (Spectral elements, FMM)
 - N data, N^2 computation
- Nonlinear evaluation (Picard, FAS, Exact Polynomial Solvers)
 - N data, N^k computation

Performance Tradeoffs

We must balance storage, bandwidth, and cycles

- Assembled Operator Action
 - Trades cycles and storage for bandwidth in application
- Unassembled Operator Action
 - Trades bandwidth and storage for cycles in application
 - For high orders, storage is impossible
 - Can make use of FERari decomposition to save calculation
 - Could store element matrices to save cycles
- Partial assembly gives even finer control over tradeoffs
 - Also allows introduction of parallel costs (load balance, ...)

Homework 2

Consider the Gram-Schmidt Orthogonalization process. Starting with a set of vectors $\{v_i\}$, create a set of orthonormal vectors $\{n_i\}$.

$$n_1 = \frac{v_1}{\|v_1\|} \quad (8)$$

$$n_2 = \frac{w_2}{\|w_2\|} \text{ where } w_2 = v_2 - (n_1 \cdot v_2)n_1 \quad (9)$$

$$n_k = \frac{w_k}{\|w_k\|} \text{ where } w_k = v_k - \sum_{j < k} (n_j \cdot v_k)n_j \quad (10)$$

What is

- 1 the balance factor β for this algorithm?
- 2 the bandwidth required to run at peak (B_{req}) on your computer?
- 3 the maximum achievable flop rate (r_{max}) on your computer?

Extra Credit: Can this algorithm be improved?

Homework 3

Run SNES ex5 for a variety of solver and preconditioner combinations. Plot the total number of linear iterations against the problem size.

Outline

- 1 Getting Started with PETSc
- 2 PETSc Integration
- 3 DM**
 - Structured Meshes (DMDA)
 - Unstructured Meshes (DMPLex/DMForest)
- 4 Advanced Solvers
- 5 More Stuff

DM Interface

• Allocation

- `DMCreateGlobalVector(DM, Vec *)`
- `DMCreateLocalVector(DM, Vec *)`
- `DMCreateMatrix(DM, MatType, Mat *)`

• Mapping

- `DMGlobalToLocalBegin/End(DM, Vec, InsertMode, Vec)`
- `DMLocalToGlobalBegin/End(DM, Vec, InsertMode, Vec)`
- `DMGetLocalToGlobalMapping(DM, IS *)`

DM Interface

● Geometry

- `DMGetCoordinateDM(DM, DM *)`
- `DMGetCoordinates(DM, Vec *)`
- `DMGetCoordinatesLocal(DM, Vec *)`

● Layout

- `DMGetDefaultSection(DM, PetscSection *)`
- `DMGetDefaultGlobalSection(DM, PetscSection *)`
- `DMGetDefaultSF(DM, PetscSF *)`

DM Interface

• Hierarchy

- `DMRefine(DM, MPI_Comm, DM *)`
- `DMCoarsen(DM, MPI_Comm, DM *)`
- `DMGetSubDM(DM, MPI_Comm, DM *)`

• Intergrid transfer

- `DMGetInterpolation(DM, DM, Mat *, Vec *)`
- `DMGetAggregates(DM, DM, Mat *)`
- `DMGetInjection(DM, DM, VecScatter *)`

Multigrid Paradigm

The **DM** interface uses the *local* callback functions to

- assemble global functions/operators from local pieces
- assemble functions/operators on coarse grids

Then **PCMG** organizes

- control flow for the multilevel solve, and
- projection and smoothing operators at each level.

Outline

3 DM

- Structured Meshes (DMDA)
- Unstructured Meshes (DMPLex/DMForest)

What is a DMDA?

DMDA is a topology interface on structured grids

- Handles parallel data layout
- Handles local and global indices
 - `DMDAGetGlobalIndices()` **and** `DMDAGetAO()`
- Provides local and global vectors
 - `DMGetGlobalVector()` **and** `DMGetLocalVector()`
- Handles ghost values coherence
 - `DMGlobalToLocalBegin/End()` **and** `DMLocalToGlobalBegin/End()`

Residual Evaluation

The **DM** interface is based upon *local* callback functions

- `FormFunctionLocal()`
- `FormJacobianLocal()`

Callbacks are registered using

- `SNESSetDM()`, `TSSetDM()`
- `DMSNESSetFunctionLocal()`, `DMTSSetJacobianLocal()`

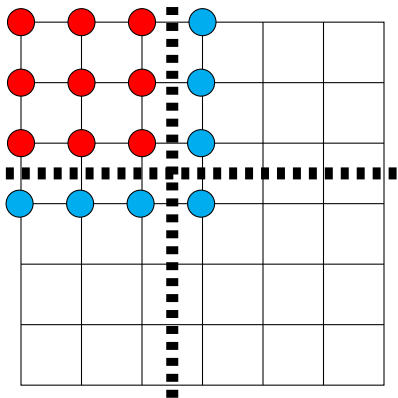
When PETSc needs to evaluate the nonlinear residual **F(x)**,

- Each process evaluates the local residual
- PETSc assembles the global residual automatically
 - Uses `DMLocalToGlobal()` method

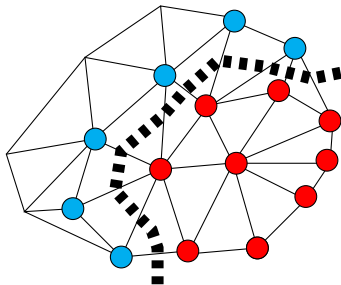
Ghost Values

To evaluate a local function $f(x)$, each process requires

- its local portion of the vector x
- its **ghost values**, bordering portions of x owned by neighboring processes



- Local Node
- Ghost Node



DMDA Global Numberings

Proc 2			Proc 3	
25	26	27	28	29
20	21	22	23	24
15	16	17	18	19
10	11	12	13	14
5	6	7	8	9
0	1	2	3	4
Proc 0			Proc 1	

Natural numbering

Proc 2			Proc 3	
21	22	23	28	29
18	19	20	26	27
15	16	17	24	25
6	7	8	13	14
3	4	5	11	12
0	1	2	9	10
Proc 0			Proc 1	

PETSc numbering

DMDA Global vs. Local Numbering

- **Global:** Each vertex has a unique id belongs on a unique process
- **Local:** Numbering includes vertices from neighboring processes
 - These are called **ghost** vertices

Proc 2			Proc 3	
X	X	X	X	X
X	X	X	X	X
12	13	14	15	X
8	9	10	11	X
4	5	6	7	X
0	1	2	3	X
Proc 0			Proc 1	

Local numbering

Proc 2			Proc 3	
21	22	23	28	29
18	19	20	26	27
15	16	17	24	25
6	7	8	13	14
3	4	5	11	12
0	1	2	9	10
Proc 0			Proc 1	

Global numbering

DMDA Local Function

User provided function calculates the nonlinear residual (in 2D)

```
(* If)(DMDALocalInfo *info, PetscScalar**x, PetscScalar**r, void *ctx)
```

`info`: All layout and numbering information

`x`: The current solution (a multidimensional array)

`r`: The residual

`ctx`: The user context passed to `DMDASNESSetFunctionLocal()`

The local DMDA function is activated by calling

```
DMDASNESSetFunctionLocal(dm, INSERT_VALUES, lfunc, &ctx)
```

Bratu Residual Evaluation

$$\Delta u + \lambda e^u = 0$$

```

ResLocal(DMDALocalInfo *info, PetscScalar **x, PetscScalar **f, void *ctx)
for(j = info->ys; j < info->ys+info->ym; ++j) {
  for(i = info->xs; i < info->xs+info->xm; ++i) {
    u = x[j][i];
    if (i==0 || j==0 || i == M || j == N) {
      f[j][i] = 2.0*(hydhx+hxhdy)*u; continue;
    }
    u_xx    = (2.0*u - x[j][i-1] - x[j][i+1])*hydhx;
    u_yy    = (2.0*u - x[j-1][i] - x[j+1][i])*hxhdy;
    f[j][i] = u_xx + u_yy - hx*hy*lambda*exp(u);
  }}

```

[\\$PETSC_DIR/src/snes/examples/tutorials/ex5.c](#)

DMDA Local Jacobian

User provided function calculates the Jacobian (in 2D)

```
(* ljac )(DMDALocalInfo *info, PetscScalar**x, Mat J, void *ctx)
```

`info`: All layout and numbering information

`x`: The current solution

`J`: The Jacobian

`ctx`: The user context passed to `DASetLocalJacobian()`

The local DMDA function is activated by calling

```
DMDASNESSetJacobianLocal(dm, ljac, &ctx)
```

Bratu Jacobian Evaluation

```

JacLocal(DMDALocalInfo *info, PetscScalar **x, Mat jac, void *ctx) {
for(j = info->ys; j < info->ys + info->ym; j++) {
    for(i = info->xs; i < info->xs + info->xm; i++) {
        row.j = j; row.i = i;
        if (i == 0 || j == 0 || i == mx-1 || j == my-1) {
            v[0] = 1.0;
            MatSetValuesStencil(jac, 1, &row, 1, &row, v, INSERT_VALUES);
        } else {
            v[0] = -(hx/hy); col[0].j = j-1; col[0].i = i;
            v[1] = -(hy/hx); col[1].j = j; col[1].i = i-1;
            v[2] = 2.0*(hy/hx+hx/hy)
                - hx*hy*lambda*PetscExpScalar(x[j][i]);
            v[3] = -(hy/hx); col[3].j = j; col[3].i = i+1;
            v[4] = -(hx/hy); col[4].j = j+1; col[4].i = i;
            MatSetValuesStencil(jac, 1, &row, 5, col, v, INSERT_VALUES);
        }
    }
}
}
}
}

```

[\\$PETSC_DIR/src/snes/examples/tutorials/ex5.c](#)

DMDA Vectors

- The **DMDA** object contains only layout (topology) information
 - All field data is contained in PETSc **Vecs**
- Global vectors are parallel
 - Each process stores a unique local portion
 - `DMCreateGlobalVector(DM da, Vec *gvec)`
- Local vectors are sequential (and usually temporary)
 - Each process stores its local portion plus ghost values
 - `DMCreateLocalVector(DM da, Vec *lvec)`
 - includes ghost and boundary values!

Updating Ghosts

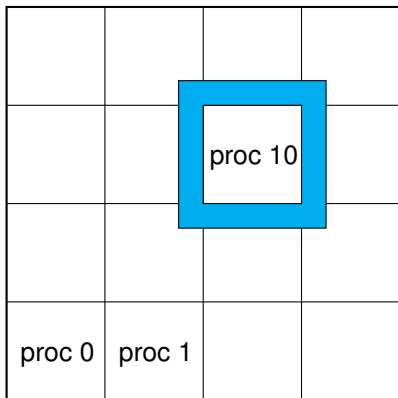
Two-step process enables overlapping computation and communication

- `DMGlobalToLocalBegin(da, gvec, mode, lvec)`
 - `gvec` provides the data
 - `mode` is either `INSERT_VALUES` or `ADD_VALUES`
 - `lvec` holds the local and ghost values
- `DMGlobalToLocalEnd(da, gvec, mode, lvec)`
 - Finishes the communication

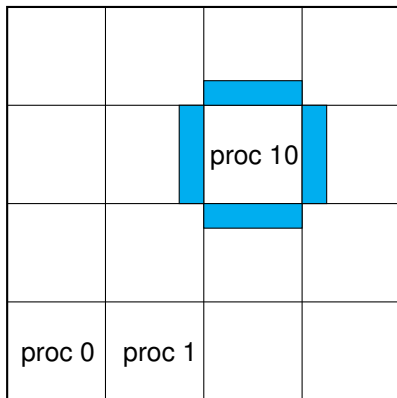
The process can be reversed with `DALocalToGlobalBegin/End()`.

DMDA Stencils

Both the **box** stencil and **star** stencil are available.



Box Stencil



Star Stencil

Setting Values on Regular Grids

PETSc provides

```
MatSetValuesStencil(Mat A, m, MatStencil idxm[], n, MatStencil idxn[],  
                  PetscScalar values[], InsertMode mode)
```

- Each row or column is actually a **MatStencil**
 - This specifies grid coordinates and a component if necessary
 - Can imagine for unstructured grids, they are *vertices*
- The values are the same logically dense block in row/col

Creating a DMDA

`DMDACreate2d(comm, bdX, bdY, type, M, N, m, n, dof, s, lm[], ln[], DMDA *da)`

`bd`: Specifies boundary behavior

- `DM_BOUNDARY_NONE`, `DM_BOUNDARY_GHOSTED`, or `DM_BOUNDARY_PERIODIC`

`type`: Specifies stencil

- `DMDA_STENCIL_BOX` or `DMDA_STENCIL_STAR`

`M/N`: Number of grid points in x/y-direction

`m/n`: Number of processes in x/y-direction

`dof`: Degrees of freedom per node

`s`: The stencil width

`lm/n`: Alternative array of local sizes

- Use `NULL` for the default

Viewing the DA

We use **SNES ex5**

- `ex5 -dm_view`
 - Shows both the DA and coordinate DA:
- `ex5 -dm_view draw -draw_pause -1`
- `ex5 -da_grid_x 10 -da_grid_y 10 -dm_view draw -draw_pause -1`
- `${PETSC_ARCH}/bin/mpiexec -n 4 ex5 -da_grid_x 10 -da_grid_y 10 -dm_view draw -draw_pause -1`
 - Shows PETSc numbering

DA Operators

- Evaluate only the local portion
 - No nice local array form without copies
- Use `MatSetValuesStencil()` to convert (i,j,k) to indices

Also use **SNES ex48**

- ```
mpiexec -n 2
./ex5 -da_grid_x 10 -da_grid_y 10 -mat_view draw -draw_pause -1
```
- ```
mpiexec -n 3
./ex48 -mat_view draw -draw_pause 1 -da_refine 3 -mat_type ai
```

Viewing FD Operator Actions

We cannot currently visualize the 3D results,

- `make runbratu EXTRA_ARGS="-run_test_-vec_view_draw_-draw_pause_-1"`
- `make runbratu`
`EXTRA_ARGS="-run_test_-da_grid_x_10_-da_grid_y_10_-vec_view_draw_-dra`
- `make runbratu EXTRA_ARGS="-run_test_-dim_3_-vec_view"`

but can check the ASCII output if necessary.

Debugging Assembly

On two processes, I get a **SEGV!**

So we try running with:

- `make NP=2 debugbratu`
`EXTRA_ARGS="-run_test_-vec_view_draw_-draw_pause_-1"`
- Spawns one debugger window per process
- SEGV on access to ghost coordinates
- Fix by using a local ghosted vector
 - Update to fixed version
- Notice
 - we already use ghosted assembly (completion) for FEM
 - FD does not need ghosted assembly

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Structured Functions

- Functions takes values at the DA vertices
- Used as approximations to functions on the continuous domain
 - Values are really coefficients of linear basis
- User only constructs the local portion
- `mpiexec -n 2 ./ex5 -run test -vec_view_draw -draw_pause -1`

Outline

3 DM

- Structured Meshes (DMDA)
- Unstructured Meshes (DMPLex/DMForest)

Problem

Traditional PDE codes cannot:

- Compare different discretizations
 - Different orders, finite elements
 - finite volume vs. finite element
- Compare different mesh types
 - Simplicial, hexahedral, polyhedral
- Run 1D, 2D, and 3D problems
- Enable an optimal solver
 - Fields, auxiliary operators

Problem

Traditional Mesh/Solver Interface is Too **General**:

- Solver not told about discretization data, e.g. fields
- Cannot take advantage of problem structure
 - blocking
 - saddle point structure
- Cannot use auxiliary data
 - Eigen-estimates
 - null spaces

Problem

Traditional Mesh/Solver Interface is Too Specific:

- Assembly code specialized to each discretization
 - dimension
 - cell shape
 - approximation space
- Explicit references to element type
 - `getVertices(faceID)`, `getAdjacency(edgeID, VERTEX)`,
`getAdjacency(edgeID, dim = 0)`
- No interface for transitive closure
 - Awkward nested loops to handle different dimensions

Mesh Representation

We represent each mesh as a **Hasse Diagram**:

- Can represent any CW complex
- Can be implemented as a Directed Acyclic Graph
- Reduces mesh information to a single *covering* relation
- Can discover dimension, since meshes are ranked posets

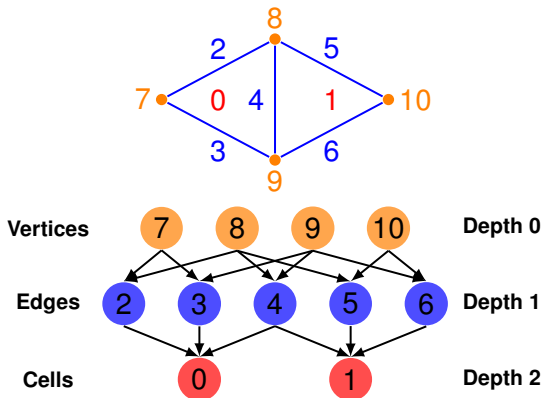
We use an abstract **topological** interface to organize traversals for:

- discretization integrals
- solver size determination
- computing communication patterns

Mesh geometry is treated as just another mesh function.

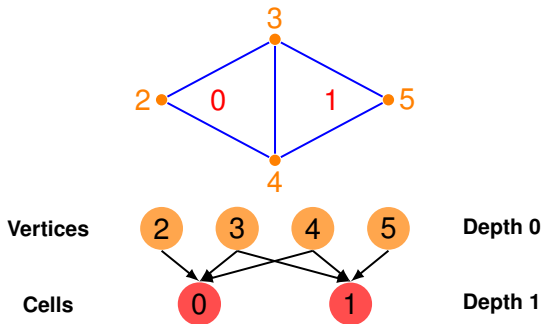
Sample Meshes

Interpolated triangular mesh



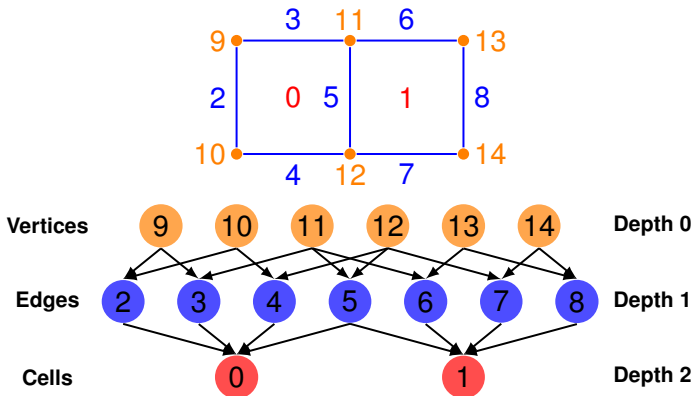
Sample Meshes

Optimized triangular mesh



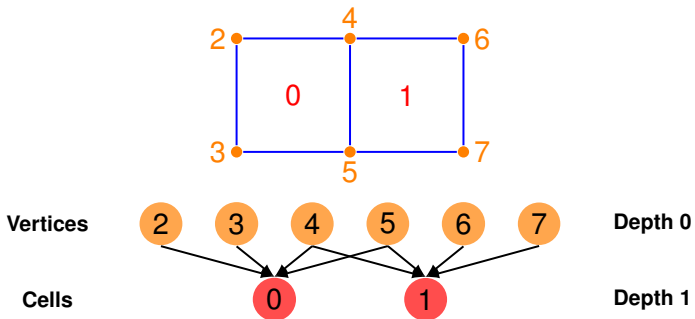
Sample Meshes

Interpolated quadrilateral mesh



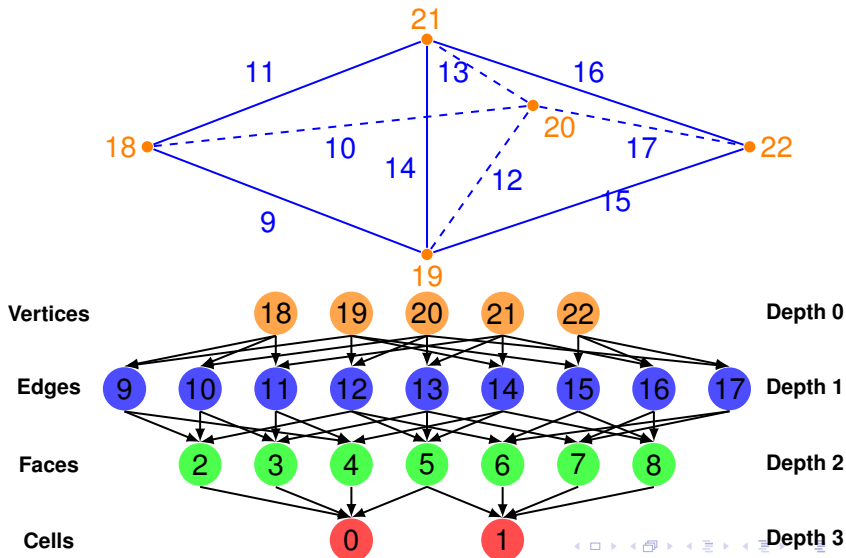
Sample Meshes

Optimized quadrilateral mesh



Sample Meshes

Interpolated tetrahedral mesh



Mesh Interface

By focusing on the key topological relations, the interface can be both concise and quite general

- Single relation
- Dual is obtained by reversing arrows
- Can associate functions with DAG points
 - Dual operation gives the support of the function

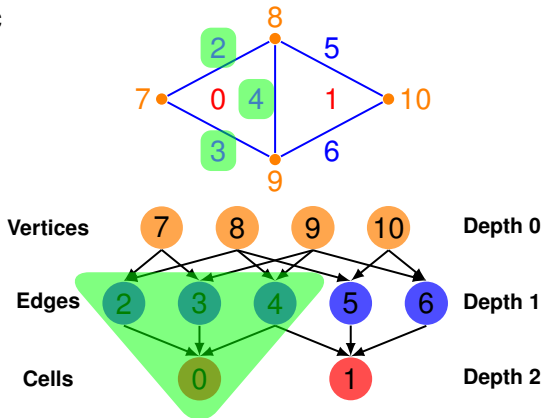
Mesh Algorithms for PDE with Sieve I: Mesh Distribution, Knepley, Karpeev, Sci. Prog., 2009.

Basic Operations

Cone

We begin with the basic covering relation,

$$\text{cone}(0) = \{2, 3, 4\}$$

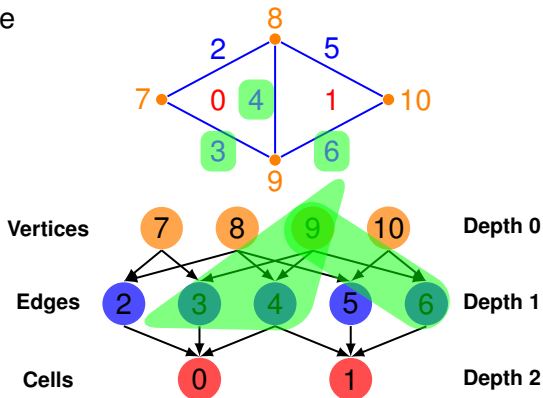


Basic Operations

Support

reverse arrows to get the
dual operation,

$$\text{support}(9) = \{3, 4, 6\}$$

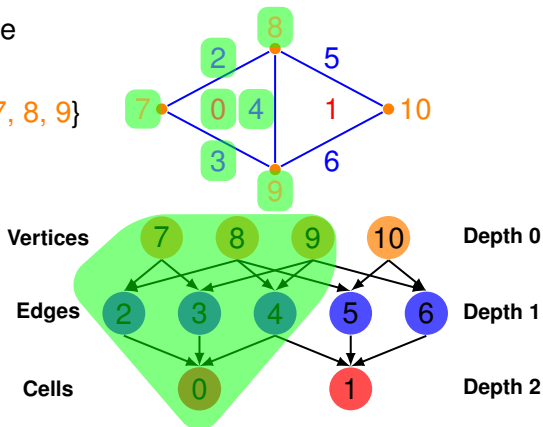


Basic Operations

Closure

add the transitive closure
of the relation,

$$\text{closure}(0) = \{0, 2, 3, 4, 7, 8, 9\}$$

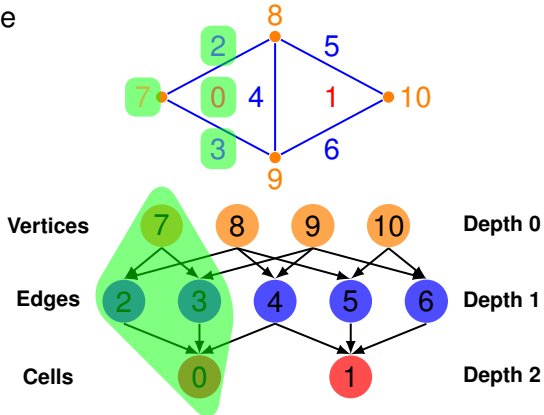


Basic Operations

Star

and the transitive closure
of the dual,

$$\text{star}(7) = \{7, 2, 3, 0\}$$

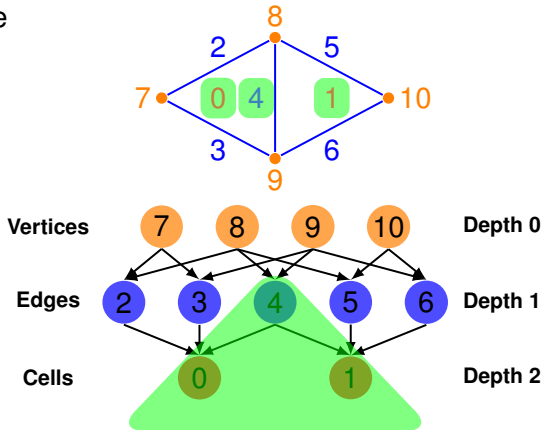


Basic Operations

Meet

and augment with lattice operations.

$$\text{meet}(0, 1) = \{4\}$$

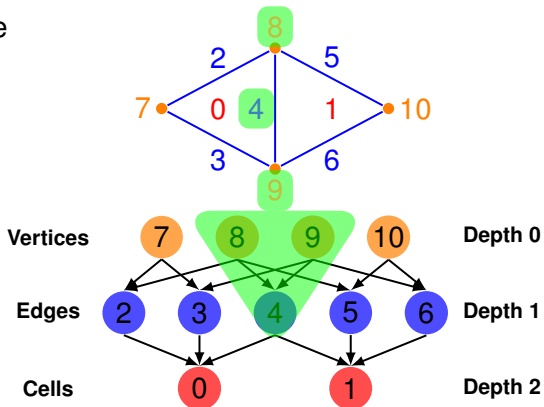


Basic Operations

Join

and augment with lattice operations.

$$\text{join}(8, 9) = \{4\}$$



DMplex

What is It?

DMplex stands for a DM modeling a CW Complex

- Handles any kind of mesh
 - Simplicial
 - Hex
 - Hybrid
 - Non-manifold
- Small interface
 - Simple to input a mesh using the API
- Accepts mesh generator input
 - ExodusII, Triangle, TetGen, LaGriT, Cubit

DMPlex

How Do I Use It?

The operations used in SNES ex62 get and set values from a **Vec**, organized by the **DM** and **PetscSection**

```
DMPlexVecGetClosure(  
    DM dm, PetscSection section, Vec v, PetscInt point,  
    PetscInt *csize, const PetscScalar *values[])
```

- Element vector on cell
- Coordinates on cell vertices

Used in `FormFunctionLocal()`,

```
for(c = cStart; c < cEnd; ++c) {  
    const PetscScalar *x;  
  
    DMPlexVecGetClosure(dm, PETSC_NULL, X, c, PETSC_NULL, &x);  
    for(PetscInt i = 0; i < cellDof; ++i) {  
        u[c+cellDof+i] = x[i];  
    }  
    DMPlexVecRestoreClosure(dm, PETSC_NULL, X, c, PETSC_NULL, &x);  
}
```

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        u[c*cellDof+i] = x[i];
    }
    DMPlexVecRestoreClosure(dm, PETSC_NULL, X, c, PETSC_NULL, &x);
}
```

DMPlex

How Do I Use It?

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```
DMPlexVecSetClosure(  
    DM dm, PetscSection section, Vec v, PetscInt point,  
    const PetscScalar values[], InsertMode mode)  
DMPlexMatSetClosure(  
    DM dm, PetscSection section, PetscSection globalSection, Mat A, PetscInt  
    PetscScalar values[], InsertMode mode)
```

- Element vector and matrix on cell

Used in `FormJacobianLocal()`,

```
for(c = cStart; c < cEnd; ++c) {  
    DMPlexMatSetClosure(dm, PETSC_NULL, PETSC_NULL, JacP, c,  
                       &elemMat[c*cellDof*cellDof], ADD_VALUES);  
}
```


DMPlex

How Do I Use It?

The operations used in SNES ex62 get and set values from a **Vec**, organized by the **DM** and **PetscSection**

```
DMPlexVecSetClosure(
    DM dm, PetscSection section, Vec v, PetscInt point,
    const PetscScalar values[], InsertMode mode)
DMPlexMatSetClosure(
    DM dm, PetscSection section, PetscSection globalSection, Mat A, PetscInt
    PetscScalar values[], InsertMode mode)
```

- Element vector and matrix on cell

Used in `FormJacobianLocal()`,

```
for(c = cStart; c < cEnd; ++c) {
    DMPlexMatSetClosure(dm, PETSC_NULL, PETSC_NULL, JacP, c,
        &elemMat[c*cellDof*cellDof], ADD_VALUES);
}
```

DMPlex

How Do I Use It?

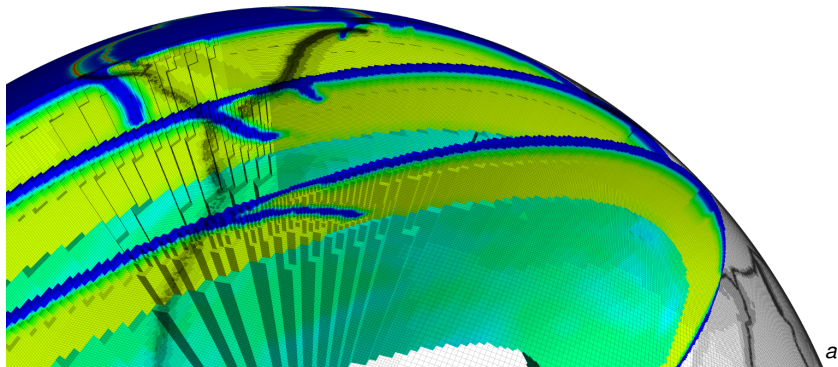
The functions above are built upon

```
DMPlexGetTransitiveClosure(  
  DM dm, PetscInt p, PetscBool useCone,  
  PetscInt *numPoints, PetscInt *points[])
```

- Returns points *and* orientations
- Iterate over points to stack up the data in the array

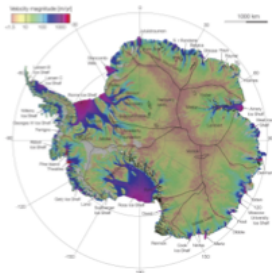
Adaptive Mesh Refinement

- DM interface with **p4est** package from Burstedde and Isaac
- PETSc solvers can be used seamlessly
- 2015 Gordon Bell Winner for Mantle Convection Simulation
- Inversion for basal traction on the full Antarctic ice sheet

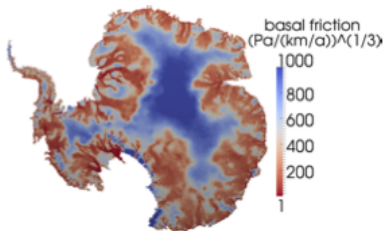


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Observed surface flow velocity (Rignot et al., 2011)



Antarctic ice sheet inversion for the basal friction parameter field using InSAR surface velocity measurements

a

^aIsaac

What does a DM do?

- Problem Definition
 - Discretization/Dof mapping (**PetscSection**)
 - Residual calculation
- Decomposition
 - Partitioning, `DMCreateSubDM()`
 - **Vec** and **Mat** creation
 - Global \iff Local mapping
- Hierarchy
 - `DMCoarsen()` and `DMRefine()`
 - `DMInterpolate()` and `DMRestrict()`
 - Hooks for resolution-dependent data

PetscSection

What Is It?

Similar to **PetscLayout**, maps point \rightarrow (size, offset)

- Processes are replaced by **points**
 - Also what we might use for multicore **PetscLayout**
- Boundary conditions are just another **PetscSection**
 - Map points to number of constrained dofs
 - Offsets into integer array of constrained local dofs
- Fields are just another **PetscSection**
 - Map points to number of field dofs
 - Offsets into array with all fields
- Usable by all **DM** subclasses
 - Structured grids with **DMDA**
 - Unstructured grids with **DMPLex**

PetscSection

Why Use It?

PETSc Solvers only understand Integers

Decouples Mesh From Discretization

- Mesh does not need to know how dofs are generated, just how many are attached to each point.
- It does not matter whether you use FD, FVM, FEM, etc.

Decouples Mesh from Solver

- Solver gets the data layout and partitioning from **Vec** and **Mat**, nothing else from the mesh.
- Solver gets restriction/interpolation matrices from **DM**.

Decouples Discretization from Solver

- Solver only gets the field division, nothing else from discretization.

PetscSection

How Do I Build One?

High Level Interface

```
DMPLexCreateSection(
    DM dm, PetscInt dim, PetscInt numFields,
    PetscInt numComp[], PetscInt numDof[],
    PetscInt numBC, PetscInt bcField[], IS bcPoints[],
    PetscSection *section);
```

Discretization	Dof/Dimension
$P_1 - P_0$	[3 0 0 0 0 0 0 1]
$Q_2 - Q_1$	[3 3 3 3 1 0 0 0]
$Q_2 - P_1^{\text{disc}}$	[3 3 3 3 0 0 0 3]

PetscSection

How Do I Build One?

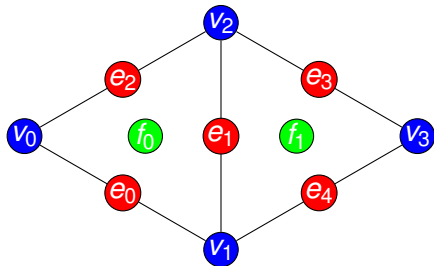
Low Level Interface

```
PetscSectionCreate(PETSC_COMM_WORLD, &s);
PetscSectionSetNumFields(s, 2);
PetscSectionSetFieldComponents(s, 0, 3);
PetscSectionSetFieldComponents(s, 1, 1);
PetscSectionSetChart(s, cStart, vEnd);
for(PetscInt v = vStart; v < vEnd; ++v) {
    PetscSectionSetDof(s, v, 3);
    PetscSectionSetFieldDof(s, v, 0, 3);
}
for(PetscInt c = cStart; c < cEnd; ++c) {
    PetscSectionSetDof(s, c, 1);
    PetscSectionSetFieldDof(s, c, 1, 1);
}
PetscSectionSetUp(s);
```

DMComplex

SNES ex62

$P_2 - P_1$ Stokes Example

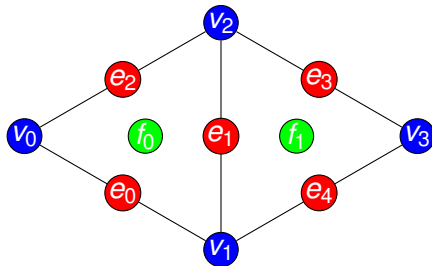


Naively, we have

$$cl(\text{cell}) = [f_{e_0} e_1 e_2 v_0 v_1 v_2 \quad]$$

$$x(\text{cell}) = [u_{e_0} v_{e_0} u_{e_1} v_{e_1} u_{e_2} v_{e_2} \\ u_{v_0} v_{v_0} p_{v_0} u_{v_1} v_{v_1} p_{v_1} u_{v_2} v_{v_2} p_{v_2} \quad]$$

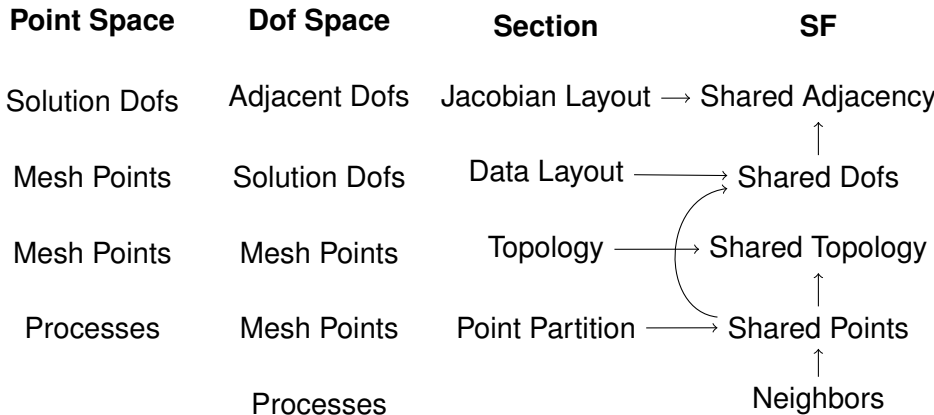
$P_2 - P_1$ Stokes Example



We reorder so that fields are contiguous

$$x'(\text{cell}) = \begin{bmatrix} u_{e_0} & v_{e_0} & u_{e_1} & v_{e_1} & u_{e_2} & v_{e_2} \\ u_{v_0} & v_{v_0} & u_{v_1} & v_{v_1} & u_{v_2} & v_{v_2} \\ p_{v_0} & p_{v_1} & p_{v_2} \end{bmatrix}$$

Section Hoisting



Residual Evaluation

I developed a single residual evaluation routine independent of spatial dimension, cell geometry, and finite element:

$$F(\vec{u}) = 0$$

Dim

1

2

3

6[†]

Cell Types

Simplex

Tensor Product

Polyhedral

Prism

Discretizations

Lagrange FEM

H(div) FEM*

H(curl) FEM*

DG FEM *[‡]

[†] Peter Brune, ANL

* FEniCS Project

[‡] Blaise Bourdin, LSU

We have also implemented a polyhedral FVM.

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FEM Integration Model

Proposed by Jed Brown

We consider weak forms dependent only on fields and gradients,

$$\int_{\Omega} \phi \cdot \mathbf{f}_0(u, \nabla u) + \nabla \phi : \vec{\mathbf{f}}_1(u, \nabla u) = 0. \quad (11)$$

Discretizing we have

$$\sum_e \mathcal{E}_e^T \left[B^T W^q \mathbf{f}_0(u^q, \nabla u^q) + \sum_k D_k^T W^q \vec{\mathbf{f}}_1^k(u^q, \nabla u^q) \right] = 0 \quad (12)$$

- f_n pointwise physics functions
- u^q field at a quad point
- W^q diagonal matrix of quad weights
- B, D basis function matrices which reduce over quad points
- \mathcal{E} assembly operator

Batch Integration

```
DMPlexComputeResidualFEM(dm, X, F, user)
{
  VecSet(F, 0.0);
  <Put boundary conditions into local input vector>
  <Extract coefficients and geometry for batch>
  <Integrate batch of elements>
  <Insert batch of element vectors into global vector>
}
```

Batch Integration

Set boundary conditions

```
DMPlexComputeResidualFEM(dm, X, F, user)
{
  VecSet(F, 0.0);
  DMPlexProjectFunctionLocal(dm, numComponents,
    bcFuncs, INSERT_BC_VALUES, X);
  <Extract coefficients and geometry for batch>
  <Integrate batch of elements>
  <Insert batch of element vectors into global vector>
}
```

Batch Integration

Extract coefficients and geometry

```
DMPlexComputeResidualFEM(dm, X, F, user)
{
  VecSet(F, 0.0);
  <Put boundary conditions into local input vector>
  DMPlexGetHeightStratum(dm, 0, &cStart, &cEnd);
  for (c = cStart; c < cEnd; ++c) {
    DMPlexComputeCellGeometry(dm, c, &v0[c*dim],
      &J[c*dim*dim], &invJ[c*dim*dim], &detJ[c]);
    DMPlexVecGetClosure(dm, NULL, X, c, NULL, &x);
    for (i = 0; i < cellDof; ++i) u[c*cellDof+i] = x[i];
    DMPlexVecRestoreClosure(dm, NULL, X, c, NULL, &x);
  }
  <Integrate batch of elements>
  <Insert batch of element vectors into global vector>
}
```

Batch Integration

Integrate element batch

```
DMPlexComputeResidualFEM(dm, X, F, user)
{
  VecSet(F, 0.0);
  <Put boundary conditions into local input vector>
  <Extract coefficients and geometry for batch>
  for (field = 0; field < numFields; ++field) {
    (*mesh->integrateResidualFEM)(Ne, numFields, field,
      quad, u,
      v0, J, invJ, detJ,
      f0, f1, elemVec);
    (*mesh->integrateResidualFEM)(Nr, ...);
  }
  <Insert batch of element vectors into global vector>
}
```

Batch Integration

Insert element vectors

```
DMPlexComputeResidualFEM(dm, X, F, user)
{
  VecSet(F, 0.0);
  <Put boundary conditions into local input vector>
  <Extract coefficients and geometry for batch>
  <Integrate batch of elements>
  for (c = cStart; c < cEnd; ++c) {
    DMPlexVecSetClosure(dm, NULL, F, c,
      &elemVec[c*cellDof], ADD_VALUES);
  }
}
```

Element Integration

```
FEMIntegrateResidualBatch(Ne, numFields, field,
    quad[], coefficients[],
    v0s[], jacobians[], jacobianInv[], jacobianDet[],
    f0_func, f1_func)
{
    <Loop over batch of elements (e)>
        <Loop over quadrature points (q)>
            <Make x_q>
            <Make u_q and gradU_q>
            <Call f_0 and f_1>
        <Loop over element vector entries (f, fc)>
            <Add contributions from f_0 and f_1>
}
```


Element Integration

Calculate x_q

```
FEMIntegrateResidualBatch(...)
{
  <Loop over batch of elements (e)>
    <Loop over quadrature points (q)>
      for (d = 0; d < dim; ++d) {
        x[d] = v0[d];
        for (d2 = 0; d2 < dim; ++d2) {
          x[d] += J[d*dim+d2]*(quadPoints[q*dim+d2]+1);
        }
      }
    <Make x_q>
    <Make u_q and gradU_q>
    <Call f_0 and f_1>
    <Loop over element vector entries (f, fc)>
      <Add contributions from f_0 and f_1>
}
```

Element Integration

Calculate u_q and ∇u_q

```
FEMIntegrateResidualBatch(...)
```

```
{
```

```
  <Loop over batch of elements (e)>
```

```
    <Loop over quadrature points (q)>
```

```
      <Make x_q>
```

```
      for (f = 0; f < numFields; ++f) {
```

```
        for (b = 0; b < Nb; ++b) {
```

```
          for (comp = 0; comp < Ncomp; ++comp) {
```

```
            u[comp] += coefficients[cidx]*basis[q+cidx];
```

```
            for (d = 0; d < dim; ++d) {
```

```
              <Transform derivative to real space>
```

```
              gradU[comp*dim+d] +=
```

```
                coefficients[cidx]*realSpaceDer[d];
```

```
            }
```

```
          }
```

```
        }
```

```
      }
```

```
      <Call f_0 and f_1>
```

```
    <Loop over element vector entries (f, fc)>
```

Element Integration

Calculate u_q and ∇u_q

```

FEMIntegrateResidualBatch(...)
{
    <Loop over batch of elements (e)>
        <Loop over quadrature points (q)>
            <Make x_q>
            for (f = 0; f < numFields; ++f) {
                for (b = 0; b < Nb; ++b) {
                    for (comp = 0; comp < Ncomp; ++comp) {
                        u[comp] += coefficients[ciIdx]*basis[q+ciIdx];
                        for (d = 0; d < dim; ++d) {
                            realSpaceDer[d] = 0.0;
                            for (g = 0; g < dim; ++g) {
                                realSpaceDer[d] +=
                                    invJ[g*dim+d]*basisDer[(q+ciIdx)*dim+g];
                            }
                            gradU[comp*dim+d] +=
                                coefficients[ciIdx]*realSpaceDer[d];
                        }
                    }
                }
            }
}

```

Element Integration

Call f_0 and f_1

```
FEMIntegrateResidualBatch(...)
{
  <Loop over batch of elements (e)>
    <Loop over quadrature points (q)>
      <Make x_q>
      <Make u_q and gradU_q>
      f0_func(u, gradU, x, &f0[q*Ncomp]);
      for (i = 0; i < Ncomp; ++i) {
        f0[q*Ncomp+i] *= detJ*quadWeights[q];
      }
      f1_func(u, gradU, x, &f1[q*Ncomp*dim]);
      for (i = 0; i < Ncomp*dim; ++i) {
        f1[q*Ncomp*dim+i] *= detJ*quadWeights[q];
      }
    <Loop over element vector entries (f, fc)>
    <Add contributions from f_0 and f_1>
  }
}
```

Element Integration

Update element vector

```
FEMIntegrateResidualBatch(...)
{
  <Loop over batch of elements (e)>
    <Loop over quadrature points (q)>
      <Make x_q>
      <Make u_q and gradU_q>
      <Call f_0 and f_1>
    <Loop over element vector entries (f, fc)>
      for (q = 0; q < Nq; ++q) {
        elemVec[ciidx] += basis[q+ciidx]*f0[q+comp];
        for (d = 0; d < dim; ++d) {
          <Transform derivative to real space>
          elemVec[ciidx] +=
            realSpaceDer[d]*f1[(q+comp)*dim+d];
        }
      }
}
```

Outline

1 Getting Started with PETSc

2 PETSc Integration

3 DM

4 Advanced Solvers

- Fieldsplit
- Multigrid
- Timestepping

5 More Stuff

Outline

4 Advanced Solvers

- Fieldsplit
- Multigrid
- Timestepping

FieldSplit Preconditioner

- Analysis

- Use **ISes** to define **fields**
- Decouples **PC** from problem definition

- Synthesis

- Additive, Multiplicative, Schur
- Commutes with Multigrid

FieldSplit Customization

• Analysis

- `-pc_fieldsplit_<split num>_fields 2,1,5`
- `-pc_fieldsplit_detect_saddle_point`

• Synthesis

- `-pc_fieldsplit_type <additive, multiplicative, schur>`
- `-pc_fieldsplit_diag_use_amat`
`-pc_fieldsplit_off_diag_use_amat`

Use diagonal blocks of operator to build PC

• Schur complements

- `-pc_fieldsplit_schur_precondition <user, all, full, self, selfp>`

How to build preconditioner for S



`-pc_fieldsplit_schur_factorization_type <diag, lower, upper, full>`

Which off-diagonal parts of the block factorization to use

Solver Configuration: No New Code

ex62: P_2/P_1 Stokes Problem on Unstructured Mesh

$$\begin{pmatrix} A & B \\ B^T & 0 \end{pmatrix}$$

Solver Configuration: No New Code

ex62: P_2/P_1 Stokes Problem on Unstructured Mesh

Block-Jacobi (Exact), Cohouet & Chabard, *IJNMF*, 1988.

```
-ksp_type gmres -pc_type fieldsplit -pc_fieldsplit_type additive  
-fieldsplit_velocity_ksp_type preonly -fieldsplit_velocity_pc_type lu  
-fieldsplit_pressure_ksp_type preonly -fieldsplit_pressure_pc_type jacobi
```

$$\begin{pmatrix} A & 0 \\ 0 & I \end{pmatrix}$$

Solver Configuration: No New Code

ex62: P_2/P_1 Stokes Problem on Unstructured Mesh

Block-Jacobi (Inexact), Cohouet & Chabard, *IJNMF*, 1988.

```
-ksp_type fgmres -pc_type fieldsplit -pc_fieldsplit_type additive  
-fieldsplit_velocity_ksp_type preonly -fieldsplit_velocity_pc_type gamg  
-fieldsplit_pressure_ksp_type preonly -fieldsplit_pressure_pc_type jacobi
```

$$\begin{pmatrix} \hat{A} & 0 \\ 0 & I \end{pmatrix}$$

Solver Configuration: No New Code

ex62: P_2/P_1 Stokes Problem on Unstructured Mesh

Gauss-Seidel (Inexact), Elman, DTIC, 1994.

```
-ksp_type fgmres -pc_type fieldsplit -pc_fieldsplit_type multiplicative  
-fieldsplit_velocity_ksp_type preonly -fieldsplit_velocity_pc_type gamg  
-fieldsplit_pressure_ksp_type preonly -fieldsplit_pressure_pc_type jacobi
```

$$\begin{pmatrix} \hat{A} & B \\ 0 & I \end{pmatrix}$$

Solver Configuration: No New Code

ex62: P_2/P_1 Stokes Problem on Unstructured Mesh

Gauss-Seidel (Inexact), Elman, DTIC, 1994.

```
-ksp_type fgmres -pc_type fieldsplit -pc_fieldsplit_type multiplicative  
-pc_fieldsplit_0_fields 1 -pc_fieldsplit_1_fields 0  
-fieldsplit_velocity_ksp_type preonly -fieldsplit_velocity_pc_type gamg  
-fieldsplit_pressure_ksp_type preonly -fieldsplit_pressure_pc_type jacobi
```

$$\begin{pmatrix} I & B^T \\ 0 & \hat{A} \end{pmatrix}$$

Solver Configuration: No New Code

ex62: P_2/P_1 Stokes Problem on Unstructured Mesh

Diagonal Schur Complement, Olshanskii, et.al., [Numer. Math.](#), 2006.

```
-ksp_type fgmres -pc_type fieldsplit -pc_fieldsplit_type schur  
-pc_fieldsplit_schur_factorization_type diag  
-fieldsplit_velocity_ksp_type preonly -fieldsplit_velocity_pc_type gamg  
-fieldsplit_pressure_ksp_type minres -fieldsplit_pressure_pc_type none
```

$$\begin{pmatrix} \hat{A} & 0 \\ 0 & -\hat{S} \end{pmatrix}$$

Solver Configuration: No New Code

ex62: P_2/P_1 Stokes Problem on Unstructured Mesh

Lower Schur Complement, May and Moresi, PEPI, 2008.

```
-ksp_type fgmres -pc_type fieldsplit -pc_fieldsplit_type schur  
-pc_fieldsplit_schur_factorization_type lower  
-fieldsplit_velocity_ksp_type preonly -fieldsplit_velocity_pc_type gamg  
-fieldsplit_pressure_ksp_type minres -fieldsplit_pressure_pc_type none
```

$$\begin{pmatrix} \hat{A} & 0 \\ B^T & \hat{S} \end{pmatrix}$$

Solver Configuration: No New Code

ex62: P_2/P_1 Stokes Problem on Unstructured Mesh

Upper Schur Complement, May and Moresi, PEPI, 2008.

```
-ksp_type fgmres -pc_type fieldsplit -pc_fieldsplit_type schur  
-pc_fieldsplit_schur_factorization_type upper  
-fieldsplit_velocity_ksp_type preonly -fieldsplit_velocity_pc_type gamg  
-fieldsplit_pressure_ksp_type minres -fieldsplit_pressure_pc_type none
```

$$\begin{pmatrix} \hat{A} & B \\ & \hat{S} \end{pmatrix}$$

Solver Configuration: No New Code

ex62: P_2/P_1 Stokes Problem on Unstructured Mesh

Uzawa Iteration, Uzawa, 1958

```
-ksp_type fgmres -pc_type fieldsplit -pc_fieldsplit_type schur  
-pc_fieldsplit_schur_factorization_type upper  
-fieldsplit_velocity_ksp_type preonly -fieldsplit_velocity_pc_type lu  
-fieldsplit_pressure_ksp_type richardson  
-fieldsplit_pressure_ksp_max_its 1
```

$$\begin{pmatrix} A & B \\ & \hat{S} \end{pmatrix}$$

Solver Configuration: No New Code

ex62: P_2/P_1 Stokes Problem on Unstructured Mesh

Full Schur Complement, Schur, 1905.

```
-ksp_type fgmres -pc_type fieldsplit -pc_fieldsplit_type schur
-pc_fieldsplit_schur_factorization_type full
-fieldsplit_velocity_ksp_type preonly -fieldsplit_velocity_pc_type lu
-fieldsplit_pressure_ksp_rtol 1e-10 -fieldsplit_pressure_pc_type jacobi
```

$$\begin{pmatrix} I & 0 \\ B^T A^{-1} & I \end{pmatrix} \begin{pmatrix} A & 0 \\ 0 & S \end{pmatrix} \begin{pmatrix} I & A^{-1} B \\ 0 & I \end{pmatrix}$$

Solver Configuration: No New Code

ex62: P_2/P_1 Stokes Problem on Unstructured Mesh

SIMPLE, Patankar and Spalding, *IJHMT*, 1972.

```
-ksp_type fgmres -pc_type fieldsplit -pc_fieldsplit_type schur
-pc_fieldsplit_schur_factorization_type full
-fieldsplit_velocity_ksp_type preonly -fieldsplit_velocity_pc_type lu
-fieldsplit_pressure_ksp_rtol 1e-10 -fieldsplit_pressure_pc_type jacobi
-fieldsplit_pressure_inner_ksp_type preonly
-fieldsplit_pressure_inner_pc_type jacobi
-fieldsplit_pressure_upper_ksp_type preonly
-fieldsplit_pressure_upper_pc_type jacobi
```

$$\begin{pmatrix} I & 0 \\ B^T A^{-1} & I \end{pmatrix} \begin{pmatrix} A & 0 \\ 0 & B^T D_A^{-1} B \end{pmatrix} \begin{pmatrix} I & D_A^{-1} B \\ 0 & I \end{pmatrix}$$

Solver Configuration: No New Code

ex62: P_2/P_1 Stokes Problem on Unstructured Mesh

Least-Squares Commutator, Kay, Loghin and Wathen, **SISC**, 2002.

```
-ksp_type fgmres -pc_type fieldsplit -pc_fieldsplit_type schur
-pc_fieldsplit_schur_factorization_type full
-pc_fieldsplit_schur_precondition self
-fieldsplit_velocity_ksp_type gmres -fieldsplit_velocity_pc_type lu
-fieldsplit_pressure_ksp_rtol 1e-5 -fieldsplit_pressure_pc_type lsc
```

$$\begin{pmatrix} I & 0 \\ B^T A^{-1} & I \end{pmatrix} \begin{pmatrix} A & 0 \\ 0 & \hat{S}_{\text{LSC}} \end{pmatrix} \begin{pmatrix} I & A^{-1} B \\ 0 & I \end{pmatrix}$$

Solver Configuration: No New Code

ex31: P_2/P_1 Stokes Problem with Temperature on Unstructured Mesh

Additive Schwarz + Full Schur Complement, Elman, Howle, Shadid, Shuttleworth, and Tuminaro, **SISC**, 2006.

```
-ksp_type fgmres -pc_type fieldsplit -pc_fieldsplit_type additive
-pc_fieldsplit_0_fields 0,1 -pc_fieldsplit_1_fields 2
-fieldsplit_0_ksp_type fgmres -fieldsplit_0_pc_type fieldsplit
-fieldsplit_0_pc_fieldsplit_type schur
-fieldsplit_0_pc_fieldsplit_schur_factorization_type full
  -fieldsplit_0_fieldsplit_velocity_ksp_type preonly
  -fieldsplit_0_fieldsplit_velocity_pc_type lu
  -fieldsplit_0_fieldsplit_pressure_ksp_rtol 1e-10
  -fieldsplit_0_fieldsplit_pressure_pc_type jacobi
-fieldsplit_temperature_ksp_type preonly
-fieldsplit_temperature_pc_type lu
```

$$\begin{pmatrix} \begin{pmatrix} I & 0 \\ B^T A^{-1} & I \end{pmatrix} & \begin{pmatrix} \hat{A} & 0 \\ 0 & \hat{S} \end{pmatrix} & \begin{pmatrix} I & A^{-1} B \\ 0 & I \end{pmatrix} & 0 \\ 0 & & & L_T \end{pmatrix}$$

Solver Configuration: No New Code

ex31: P_2/P_1 Stokes Problem with Temperature on Unstructured Mesh

Upper Schur Comp. + Full Schur Comp. + Least-Squares Comm.

```
-ksp_type fgmres -pc_type fieldsplit -pc_fieldsplit_type schur
-pc_fieldsplit_0_fields 0,1 -pc_fieldsplit_1_fields 2
-pc_fieldsplit_schur_factorization_type upper
-fieldsplit_0_ksp_type fgmres -fieldsplit_0_pc_type fieldsplit
-fieldsplit_0_pc_fieldsplit_type schur
-fieldsplit_0_pc_fieldsplit_schur_factorization_type full
-fieldsplit_0_fieldsplit_velocity_ksp_type preonly
-fieldsplit_0_fieldsplit_velocity_pc_type lu
-fieldsplit_0_fieldsplit_pressure_ksp_rtol 1e-10
-fieldsplit_0_fieldsplit_pressure_pc_type jacobi
-fieldsplit_temperature_ksp_type gmres
-fieldsplit_temperature_pc_type lsc
```

$$\begin{pmatrix} \begin{pmatrix} I & 0 \\ B^T A^{-1} & I \end{pmatrix} & \begin{pmatrix} \hat{A} & 0 \\ 0 & \hat{S} \end{pmatrix} & \begin{pmatrix} I & A^{-1} B \\ 0 & I \end{pmatrix} & G \\ & 0 & & \hat{S}_{LSC} \end{pmatrix}$$

SNES ex62

Preconditioning

Jacobi

ex62

```
-run_type full -bc_type dirichlet -show_solution 0  
-refinement_limit 0.00625 -interpolate 1  
-vel_petscspace_order 2 -pres_petscspace_order 1  
-snes_monitor_short -snes_converged_reason  
  -snes_view  
-ksp_gmres_restart 100 -ksp_rtol 1.0e-9  
  -ksp_monitor_short  
-pc_type jacobi
```


SNES ex62

Preconditioning

Block diagonal

ex62

```
-run_type full -bc_type dirichlet -show_solution 0
-refinement_limit 0.00625 -interpolate 1
-vel_petscspace_order 2 -pres_petscspace_order 1
-snes_monitor_short -snes_converged_reason
  -snes_view
-ksp_type fgmres -ksp_gmres_restart 100
  -ksp_rtol 1.0e-9 -ksp_monitor_short
-pc_type fieldsplit -pc_fieldsplit_type additive
-fieldsplit_velocity_pc_type lu
-fieldsplit_pressure_pc_type jacobi
```

SNES ex62

Preconditioning

Block triangular

ex62

```
-run_type full -bc_type dirichlet -show_solution 0
-refinement_limit 0.00625 -interpolate 1
-vel_petscspace_order 2 -pres_petscspace_order 1
-snes_monitor_short -snes_converged_reason
  -snes_view
-ksp_type fgmres -ksp_gmres_restart 100
  -ksp_rtol 1.0e-9 -ksp_monitor_short
-pc_type fieldsplit -pc_fieldsplit_type multiplicative
-fieldsplit_velocity_pc_type lu
-fieldsplit_pressure_pc_type jacobi
```

SNES ex62

Preconditioning

Diagonal Schur complement

ex62

```
-run_type full -bc_type dirichlet -show_solution 0
-refinement_limit 0.00625 -interpolate 1
-vel_petscspace_order 2 -pres_petscspace_order 1
-snes_monitor_short -snes_converged_reason
  -snes_view
-ksp_type fgmres -ksp_gmres_restart 100
  -ksp_rtol 1.0e-9 -ksp_monitor_short
-pc_type fieldsplit -pc_fieldsplit_type schur
  -pc_fieldsplit_schur_factorization_type diag
-fieldsplit_velocity_ksp_type gmres
  -fieldsplit_velocity_pc_type lu
-fieldsplit_pressure_ksp_rtol 1e-10
  -fieldsplit_pressure_pc_type jacobi
```

SNES ex62

Preconditioning

Upper triangular Schur complement

ex62

```
-run_type full -bc_type dirichlet -show_solution 0
-refinement_limit 0.00625 -interpolate 1
-vel_petscspace_order 2 -pres_petscspace_order 1
-snes_monitor_short -snes_converged_reason
  -snes_view
-ksp_type fgmres -ksp_gmres_restart 100
  -ksp_rtol 1.0e-9 -ksp_monitor_short
-pc_type fieldsplit -pc_fieldsplit_type schur
  -pc_fieldsplit_schur_factorization_type upper
-fieldsplit_velocity_ksp_type gmres
  -fieldsplit_velocity_pc_type lu
-fieldsplit_pressure_ksp_rtol 1e-10
  -fieldsplit_pressure_pc_type jacobi
```

SNES ex62

Preconditioning

Lower triangular Schur complement

ex62

```
-run_type full -bc_type dirichlet -show_solution 0
-refinement_limit 0.00625 -interpolate 1
-vel_petscspace_order 2 -pres_petscspace_order 1
-snes_monitor_short -snes_converged_reason
  -snes_view
-ksp_type fgmres -ksp_gmres_restart 100
  -ksp_rtol 1.0e-9 -ksp_monitor_short
-pc_type fieldsplit -pc_fieldsplit_type schur
  -pc_fieldsplit_schur_factorization_type lower
-fieldsplit_velocity_ksp_type gmres
  -fieldsplit_velocity_pc_type lu
-fieldsplit_pressure_ksp_rtol 1e-10
  -fieldsplit_pressure_pc_type jacobi
```

SNES ex62

Preconditioning

Full Schur complement

ex62

```
-run_type full -bc_type dirichlet -show_solution 0
-refinement_limit 0.00625 -interpolate 1
-vel_petscspace_order 2 -pres_petscspace_order 1
-snes_monitor_short -snes_converged_reason
  -snes_view
-ksp_type fgmres -ksp_gmres_restart 100
  -ksp_rtol 1.0e-9 -ksp_monitor_short
-pc_type fieldsplit -pc_fieldsplit_type schur
  -pc_fieldsplit_schur_factorization_type full
-fieldsplit_velocity_ksp_type gmres
  -fieldsplit_velocity_pc_type lu
-fieldsplit_pressure_ksp_rtol 1e-10
  -fieldsplit_pressure_pc_type jacobi
```

Programming with Options

ex55: Allen-Cahn problem in 2D

- constant mobility
- triangular elements

Geometric multigrid method for saddle point variational inequalities:

```
./ex55 -ksp_type fgmres -pc_type mg -mg_levels_ksp_type fgmres  
-mg_levels_pc_type fieldsplit -mg_levels_pc_fieldsplit_detect_saddle_point  
-mg_levels_pc_fieldsplit_type schur -da_grid_x 65 -da_grid_y 65  
-mg_levels_pc_fieldsplit_factorization_type full  
-mg_levels_pc_fieldsplit_schur_precondition user  
-mg_levels_fieldsplit_1_ksp_type gmres -mg_coarse_ksp_type preonly  
-mg_levels_fieldsplit_1_pc_type none -mg_coarse_pc_type svd  
-mg_levels_fieldsplit_0_ksp_type preonly  
-mg_levels_fieldsplit_0_pc_type sor -pc_mg_levels 5  
-mg_levels_fieldsplit_0_pc_sor_forward -pc_mg_galerkin  
-snes_vi_monitor -ksp_monitor_true_residual -snes_atol 1.e-11  
-mg_levels_ksp_monitor -mg_levels_fieldsplit_ksp_monitor  
-mg_levels_ksp_max_it 2 -mg_levels_fieldsplit_ksp_max_it 5
```

Programming with Options

ex55: Allen-Cahn problem in 2D

Run flexible GMRES with 5 levels of multigrid as the preconditioner

```
./ex55 -ksp_type fgmres -pc_type mg -pc_mg_levels 5  
-da_grid_x 65 -da_grid_y 65
```

Use the Galerkin process to compute the coarse grid operators

```
-pc_mg_galerkin
```

Use SVD as the coarse grid saddle point solver

```
-mg_coarse_ksp_type preonly -mg_coarse_pc_type svd
```


Programming with Options

ex55: Allen-Cahn problem in 2D

Run flexible GMRES with 5 levels of multigrid as the preconditioner

```
./ex55 -ksp_type fgmres -pc_type mg -pc_mg_levels 5  
-da_grid_x 65 -da_grid_y 65
```

Use the Galerkin process to compute the coarse grid operators

```
-pc_mg_galerkin
```

Use SVD as the coarse grid saddle point solver

```
-mg_coarse_ksp_type preonly -mg_coarse_pc_type svd
```

Programming with Options

ex55: Allen-Cahn problem in 2D

Run flexible GMRES with 5 levels of multigrid as the preconditioner

```
./ex55 -ksp_type fgmres -pc_type mg -pc_mg_levels 5  
-da_grid_x 65 -da_grid_y 65
```

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```
-pc_mg_galerkin
```

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```
-mg_coarse_ksp_type preonly -mg_coarse_pc_type svd
```

Programming with Options

ex55: Allen-Cahn problem in 2D

Run flexible GMRES with 5 levels of multigrid as the preconditioner

```
./ex55 -ksp_type fgmres -pc_type mg -pc_mg_levels 5  
-da_grid_x 65 -da_grid_y 65
```

Use the Galerkin process to compute the coarse grid operators

```
-pc_mg_galerkin
```

Use SVD as the coarse grid saddle point solver

```
-mg_coarse_ksp_type preonly -mg_coarse_pc_type svd
```

Programming with Options

ex55: Allen-Cahn problem in 2D

Smoother: Flexible GMRES (2 iterates) with a Schur complement PC

```
-mg_levels_ksp_type fgmres -mg_levels_pc_fieldsplit_detect_saddle_point  
-mg_levels_ksp_max_it 2 -mg_levels_pc_type fieldsplit  
-mg_levels_pc_fieldsplit_type schur  
-mg_levels_pc_fieldsplit_factorization_type full  
-mg_levels_pc_fieldsplit_schur_precondition diag
```

Schur complement solver: GMRES (5 iterates) with no preconditioner

```
-mg_levels_fieldsplit_1_ksp_type gmres  
-mg_levels_fieldsplit_1_pc_type none -mg_levels_fieldsplit_ksp_max_it 5
```

Schur complement action: Use only the lower diagonal part of A00

```
-mg_levels_fieldsplit_0_ksp_type preonly  
-mg_levels_fieldsplit_0_pc_type sor  
-mg_levels_fieldsplit_0_pc_sor_forward
```

Programming with Options

ex55: Allen-Cahn problem in 2D

Smoother: Flexible GMRES (2 iterates) with a Schur complement PC

```
-mg_levels_ksp_type fgmres -mg_levels_pc_fieldsplit_detect_saddle_point  
-mg_levels_ksp_max_it 2 -mg_levels_pc_type fieldsplit  
-mg_levels_pc_fieldsplit_type schur  
-mg_levels_pc_fieldsplit_factorization_type full  
-mg_levels_pc_fieldsplit_schur_precondition diag
```

Schur complement solver: GMRES (5 iterates) with no preconditioner

```
-mg_levels_fieldsplit_1_ksp_type gmres  
-mg_levels_fieldsplit_1_pc_type none -mg_levels_fieldsplit_ksp_max_it 5
```

Schur complement action: Use only the lower diagonal part of A00

```
-mg_levels_fieldsplit_0_ksp_type preonly  
-mg_levels_fieldsplit_0_pc_type sor  
-mg_levels_fieldsplit_0_pc_sor_forward
```

Programming with Options

ex55: Allen-Cahn problem in 2D

Smoother: Flexible GMRES (2 iterates) with a Schur complement PC

```
-mg_levels_ksp_type fgmres -mg_levels_pc_fieldsplit_detect_saddle_point  
-mg_levels_ksp_max_it 2 -mg_levels_pc_type fieldsplit  
-mg_levels_pc_fieldsplit_type schur  
-mg_levels_pc_fieldsplit_factorization_type full  
-mg_levels_pc_fieldsplit_schur_precondition diag
```

Schur complement solver: GMRES (5 iterates) with no preconditioner

```
-mg_levels_fieldsplit_1_ksp_type gmres  
-mg_levels_fieldsplit_1_pc_type none -mg_levels_fieldsplit_ksp_max_it 5
```

Schur complement action: Use only the lower diagonal part of A00

```
-mg_levels_fieldsplit_0_ksp_type preonly  
-mg_levels_fieldsplit_0_pc_type sor  
-mg_levels_fieldsplit_0_pc_sor_forward
```

Programming with Options

ex55: Allen-Cahn problem in 2D

Smoother: Flexible GMRES (2 iterates) with a Schur complement PC

```
-mg_levels_ksp_type fgmres -mg_levels_pc_fieldsplit_detect_saddle_point  
-mg_levels_ksp_max_it 2 -mg_levels_pc_type fieldsplit  
-mg_levels_pc_fieldsplit_type schur  
-mg_levels_pc_fieldsplit_factorization_type full  
-mg_levels_pc_fieldsplit_schur_precondition diag
```

Schur complement solver: GMRES (5 iterates) with no preconditioner

```
-mg_levels_fieldsplit_1_ksp_type gmres  
-mg_levels_fieldsplit_1_pc_type none -mg_levels_fieldsplit_ksp_max_it 5
```

Schur complement action: Use only the lower diagonal part of A00

```
-mg_levels_fieldsplit_0_ksp_type preonly  
-mg_levels_fieldsplit_0_pc_type sor  
-mg_levels_fieldsplit_0_pc_sor_forward
```

Null spaces

For a single matrix, use

```
MatSetNullSpace(J, nullSpace);
```

to alter the **KSP**, and

```
MatSetNearNullSpace(J, nearNullSpace);
```

to set the coarse basis for AMG.

But this will not work for dynamically created operators.

Null spaces

For a single matrix, use

```
MatSetNullSpace(J, nullSpace);
```

to alter the **KSP**, and

```
MatSetNearNullSpace(J, nearNullSpace);
```

to set the coarse basis for AMG.

But this will not work for dynamically created operators.

Null spaces

Field Split

Can attach a nullspace to the **IS** that creates a split,

```
PetscObjectCompose( pressureIS , "nullspace" ,  
                    (PetscObject) nullSpacePres );
```

If the **DM** makes the **IS**, use

```
PetscObject pressure ;  
  
DMGetField( dm , 1 , &pressure );  
PetscObjectCompose( pressure , "nullspace" ,  
                    (PetscObject) nullSpacePres );
```

Outline

4 Advanced Solvers

- Fieldsplit
- **Multigrid**
- Timestepping

Why not use AMG?

- Of course we will try AMG
 - GAMG, `-pc_type gamg`
 - ML, `-download-ml, -pc_type ml`
 - BoomerAMG, `-download-hypre, -pc_type hypre`
`-pc_hypre_type boomeramg`
- Problems with
 - vector character
 - anisotropy
 - scalability of setup time

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- Problems with
 - vector character
 - anisotropy
 - scalability of setup time

Multigrid with DM

Allows multigrid with some simple command line options

- `-pc_type mg, -pc_mg_levels`
- `-pc_mg_type, -pc_mg_cycle_type, -pc_mg_galerkin`
- `-mg_levels_1_ksp_type, -mg_levels_1_pc_type`
- `-mg_coarse_ksp_type, -mg_coarse_pc_type`
- `-da_refine, -ksp_view`

Interface also works with GAMG and 3rd party packages like ML

A 2D Problem

Problem has:

- 1,640,961 unknowns (on the fine level)
- 8,199,681 nonzeros

	Options	Explanation
./ex5	-da_grid_x 21 -da_grid_y 21	Original grid is 21x21
	-ksp_rtol 1.0e-9	Solver tolerance
	-da_refine 6	6 levels of refinement
	-pc_type mg	4 levels of multigrid
	-pc_mg_levels 4	
	-snes_monitor -snes_view	Describe solver

A 3D Problem

Problem has:

- 1,689,600 unknowns (on the fine level)
- 89,395,200 nonzeros

	Options	Explanation
<code>./ex48</code>	<code>-M 5 -N 5</code>	Coarse problem size
	<code>-da_refine 5</code>	5 levels of refinement
	<code>-ksp_rtol 1.0e-9</code>	Solver tolerance
	<code>-thi_mat_type baij</code>	Needs SOR
	<code>-pc_type mg</code>	4 levels of multigrid
	<code>-pc_mg_levels 4</code>	
	<code>-snes_monitor -snes_view</code>	Describe solver

Full Multigrid

The Full Multigrid algorithm (FMG)

- V-cycle at each level,
- then interpolate to the next finer grid
- Can solve to discretization error with a *single* iteration

Full Multigrid Work

$$\begin{aligned}C_{FMG} &= \left(1 + \frac{1}{2^d} + \frac{1}{2^{2d}} + \dots\right) C_V \\&= \sum_{n=0}^{\infty} \frac{1}{2^{nd}} C_V \\&= \frac{2^d}{2^d - 1} C_V \\&= \left(\frac{2^d}{2^d - 1}\right)^2 C_{\text{twolevel}}.\end{aligned}$$

1D FMG is $2 \times C_V$, 3D FMG is $\frac{8}{7} \times C_V$

Full Multigrid Work

$$\begin{aligned}C_{FMG} &= \left(1 + \frac{1}{2^d} + \frac{1}{2^{2d}} + \dots\right) C_V \\&= \sum_{n=0}^{\infty} \frac{1}{2^{nd}} C_V \\&= \frac{2^d}{2^d - 1} C_V \\&= \left(\frac{2^d}{2^d - 1}\right)^2 C_{\text{twolevel}}.\end{aligned}$$

1D FMG is $2 \times C_V$, 3D FMG is $\frac{8}{7} \times C_V$

Full Multigrid Accuracy

Suppose we have an order α method,

$$\|x - x_h\| < Ch^\alpha$$

FD and P_1 both have $\alpha = 2$

Full Multigrid Accuracy

E_d Discretization Error

E_a Algebraic Error

Choose iterative tolerance so that

$$E_a = rE_d \quad r < 1$$

and

$$E \leq E_d + E_a = (1 + r)Ch^\alpha$$

Full Multigrid Accuracy

Suppose

- Finish V-cycle for $2h$ grid,
- Use as coarse correction for h grid
- Perform final V-cycle for h grid
- Need V-cycle error reduction factor η to get r reduction in E_a

Full Multigrid Accuracy

$$\eta E_a < rCh^\alpha$$

$$\eta(E - E_d) < rCh^\alpha$$

$$\eta((1+r)C(2h)^\alpha - Ch^\alpha) < rCh^\alpha$$

$$\eta((1+r)2^\alpha - 1) < r$$

$$\eta < \frac{1}{2^\alpha + \frac{2^\alpha - 1}{r}}.$$

If $\alpha = 2$ and $r = \frac{1}{2}$, then $\eta < \frac{1}{10}$.

Full Multigrid Experiment

V-cycle

```
./ex5 -mms 1 -par 0.0 -da_refine 3 -snes_type newtonls -snes_max_it 1  
-ksp_rtol 1e-10 -pc_type mg -snes_monitor_short -ksp_monitor_short
```

gives

```
0 SNES Function norm 0.0287773  
0 KSP Residual norm 0.793727  
1 KSP Residual norm 0.00047526  
2 KSP Residual norm 4.18007e-06  
3 KSP Residual norm 1.1668e-07  
4 KSP Residual norm 3.25952e-09  
5 KSP Residual norm 7.274e-11  
1 SNES Function norm 2.251e-10  
N: 625 error 12 1.21529e-13 inf 9.53484e-12
```

Full Multigrid Experiment

V-cycle

```
./ex5 -mms 1 -par 0.0 -da_refine 3 -snes_type newtonls -snes_max_it 1  
-ksp_rtol 1e-10 -pc_type mg -snes_monitor_short -ksp_monitor_short
```

and it changes little if we refine six more times

```
0 SNES Function norm 0.000455131  
  0 KSP Residual norm 50.6842  
  1 KSP Residual norm 0.00618427  
  2 KSP Residual norm 9.87833e-07  
  3 KSP Residual norm 2.99517e-09  
1 SNES Function norm 2.83358e-09  
N: 2362369 error l2 1.28677e-15 inf 7.68693e-12
```

Full Multigrid Experiment

FMG

```
./ex5 -mms 1 -par 0.0 -da_refine 3 -snes_type newtonls -snes_max_it 1  
-ksp_rtol 1e-10 -pc_type mg -snes_monitor_short -ksp_monitor_short  
-pc_mg_type full
```

We do not seem to see the convergence acceleration

```
0 SNES Function norm 0.0287773  
  0 KSP Residual norm 0.799687  
  1 KSP Residual norm 6.95292e-05  
  2 KSP Residual norm 1.50836e-06  
  3 KSP Residual norm 2.62524e-08  
  4 KSP Residual norm 6.184e-10  
  5 KSP Residual norm 1.275e-11  
1 SNES Function norm 3.757e-11  
N: 625 error 12 2.1428e-14 inf 1.80611e-12
```

Full Multigrid Experiment

FMG

```
./ex5 -mms 1 -par 0.0 -da_refine 3 -snes_type newtonls -snes_max_it 1  
-ksp_rtol 1e-10 -pc_type mg -snes_monitor_short -ksp_monitor_short  
-pc_mg_type full
```

although its a little more apparent as we refine,

```
0 SNES Function norm 0.000455131  
  0 KSP Residual norm 51.2  
  1 KSP Residual norm 2.92416e-06  
  2 KSP Residual norm 3.76404e-09  
1 SNES Function norm 8.50096e-09  
N: 2362369 error 12 1.70304e-15 inf 6.22476e-11
```

Full Multigrid Experiment

Script

```
#!/usr/bin/env python
import argparse
import subprocess
import numpy as np

parser = argparse.ArgumentParser(
    description = 'CAAM 519 FMG',
    epilog = 'For more information, visit http://www.mcs.anl.gov/petsc',
    formatter_class = argparse.ArgumentDefaultsHelpFormatter)
parser.add_argument('--kmax', type=int, default=5,
                    help='The number of doublings to test')
parser.add_argument('--save', action='store_true', default=False,
                    help='Save the figures')
args = parser.parse_args()

sizesA = []
sizesB = []
errorA = []
errorB = []
```

Full Multigrid Experiment

Script

```
for k in range(args.kmax):
    options = ['-snes_type', 'newtonls', '-snes_max_it', '1', '-da_refine',
              '-par', '0.0', '-ksp_atol', '1e-1', '-mms', '1',
              '-pc_type', 'mg', '-pc_mg_type', 'multiplicative',
              '-mg_levels_ksp_max_it', '5']
    cmd = './ex5 '+' '.join(options)
    out = subprocess.check_output(['./ex5']+options).split(' ')
    # This is l_2, out[6] is l_infty
    sizesA.append(int(out[1]))
    errorA.append(float(out[4]))
for k in range(args.kmax):
    options = ['-snes_type', 'newtonls', '-snes_max_it', '1', '-da_refine',
              '-par', '0.0', '-ksp_atol', '1e-1', '-mms', '1',
              '-pc_type', 'mg', '-pc_mg_type', 'full',
              '-mg_levels_ksp_max_it', '5']
    cmd = './ex5 '+' '.join(options)
    out = subprocess.check_output(['./ex5']+options).split(' ')
    # This is l_2, out[6] is l_infty
    sizesB.append(int(out[1]))
    errorB.append(float(out[4]))
```

Full Multigrid Experiment

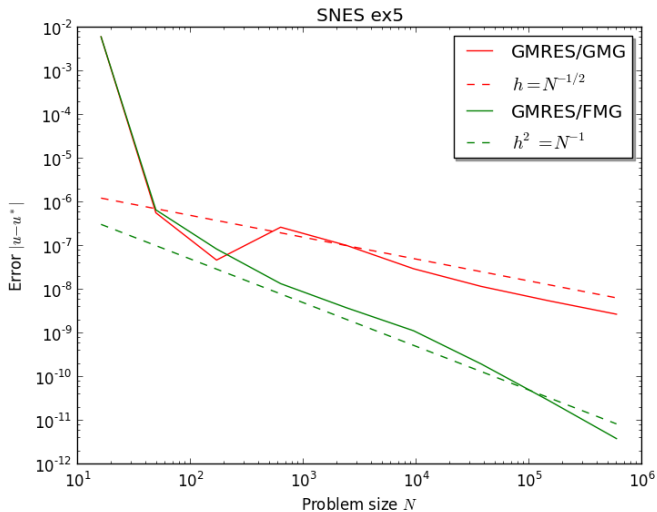
Script

```
SA = np.array(sizesA)
SB = np.array(sizesB)

from pylab import legend, plot, loglog, show, title, xlabel, ylabel, savefig
loglog(SA, errorA, 'r', SA, 5e-6 * SA ** -0.5, 'r--',
        SB, errorB, 'g', SB, 5e-6 * SB ** -1., 'g--')
title('SNES ex5')
xlabel('Problem size $N$')
ylabel('Error $\|u - u^*\|_{\infty}$')
legend(['GMRES/GMG', '$h = N^{-1/2}$', 'GMRES/FMG', '$h^2 = N^{-1}$'],
        'upper right', shadow = True)
if args.save:
    savefig('fmg.png')
else:
    show()
```

Full Multigrid Experiment

Comparison



Outline

4 Advanced Solvers

- Fieldsplit
- Multigrid
- Timestepping

What about TS?

Time Integration in PETSc Has Made a Quantum Leap

and is now a premier library in the field,

thanks to **Jed**, **Emil**, and **Peter** \implies

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Some TS methods

TSSSPRK104 10-stage, fourth order, low-storage, optimal explicit SSP Runge-Kutta $c_{\text{eff}} = 0.6$ (Ketcheson 2008)

TSARKIMEX2E second order, one explicit and two implicit stages, L -stable, optimal (Constantinescu)

TSARKIMEX3 (and 4 and 5), L -stable (Kennedy and Carpenter, 2003)

TSROSWRA3PW three stage, third order, for index-1 PDAE, A -stable, $R(\infty) = 0.73$, second order strongly A -stable embedded method (Rang and Angermann, 2005)

TSROSWRA34PW2 four stage, third order, L -stable, for index 1 PDAE, second order strongly A -stable embedded method (Rang and Angermann, 2005)

TSROSWLLSSP3P4S2C four stage, third order, L -stable implicit, SSP explicit, L -stable embedded method (Constantinescu)

IMEX time integration in PETSc

Additive Runge-Kutta IMEX methods

$$G(t, x, \dot{x}) = F(t, x)$$

$$J_{\alpha} = \alpha G_{\dot{x}} + G_x$$

User provides:

- `FormRHSFunction(ts, t, x, F, void *ctx)`
- `FormIFunction(ts, t, x, xdot, G, void *ctx)`
- `FormIJacobian(ts, t, x, xdot, alpha, J, J_p, mstr, void *ctx)`
- Single step interface so user can have own time loop
- Choice of explicit method, e.g. SSP
- L-stable DIRK for stiff part G
- Orders 2 through 5, embedded error estimates
- Dense output, hot starts for Newton
- More accurate methods if G is linear, also Rosenbrock-W
- Can use preconditioner from classical “semi-implicit” methods
- Extensible adaptive controllers, can change order within a family
- Easy to register new methods: `TSARKIMEXRegister()`

Stiff linear advection-reaction test problem

Equations

TS ex22.c

$$u_t + a_1 u_x = -k_1 u + k_2 v + s_1$$

$$v_t + a_2 v_x = k_1 u - k_2 v + s_2$$

Upstream boundary condition:

$$u(0, t) = 1 - \sin(12t)^4$$

Stiff linear advection-reaction test problem

Equations

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$$u_t + a_1 u_x = -k_1 u + k_2 v + s_1$$

$$v_t + a_2 v_x = k_1 u - k_2 v + s_2$$

```

FormIFunction(TS ts, PetscReal t, Vec X, Vec Xdot, Vec F, void *ptr) {
    TSGetDM(ts, &da);
    DMGetLocalInfo(da, &info);
    DMGetVecArray(da, X, &x);
    DMGetVecArray(da, Xdot, &xdot);
    DMGetVecArray(da, F, &f);
    /* Compute function over the locally owned part of the grid */
    for (i = info.xs; i < info.xs+info.xm; ++i) {
        f[i][0] = xdot[i][0] + k[0]*x[i][0] - k[1]*x[i][1] - s[0];
        f[i][1] = xdot[i][1] - k[0]*x[i][0] + k[1]*x[i][1] - s[1];
    }
    DMRestoreVecArray(da, X, &x);
    DMRestoreVecArray(da, Xdot, &xdot);
    DMRestoreVecArray(da, F, &f);
}

```


Stiff linear advection-reaction test problem

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$$u_t + a_1 u_x = -k_1 u + k_2 v + s_1$$

$$v_t + a_2 v_x = k_1 u - k_2 v + s_2$$

```

FormIJacobian(TS ts, PetscReal t, Vec X, Vec Xdot, PetscReal a, Mat *J,
              Mat *Jpre, MatStructure *str, void *ptr) {
  for (i = info.xs; i < info.xs+info.xm; ++i) {
    PetscScalar v[2][2];
    v[0][0] = a + k[0]; v[0][1] = -k[1];
    v[1][0] = -k[0]; v[1][1] = a+k[1];
    MatSetValuesBlocked(*Jpre, 1, &i, 1, &i, &v[0][0], INSERT_VALUES);
  }
  MatAssemblyBegin(*Jpre, MAT_FINAL_ASSEMBLY);
  MatAssemblyEnd(*Jpre, MAT_FINAL_ASSEMBLY);
  if (*J != *Jpre) {
    MatAssemblyBegin(*J, MAT_FINAL_ASSEMBLY);
    MatAssemblyEnd(*J, MAT_FINAL_ASSEMBLY);
  }
}

```

Stiff linear advection-reaction test problem

Equations

TS ex22.c

$$u_t + a_1 u_x = -k_1 u + k_2 v + s_1$$

$$v_t + a_2 v_x = k_1 u - k_2 v + s_2$$

```

FormRHSFunction(TS ts, PetscReal t, Vec X, Vec F, void *ptr) {
    PetscReal u0t[2] = {1. - PetscPowScalar(sin(12*t), 4.), 0};
    DMGetLocalVector(da, &Xloc);
    DMGlobalToLocalBegin(da, X, INSERT_VALUES, Xloc);
    DMGlobalToLocalEnd(da, X, INSERT_VALUES, Xloc);
    for (i = info.xs; i < info.xs+info.xm; ++i) {
        /* CALCULATE RESIDUAL f[i][j] */
    }
}

```

Stiff linear advection-reaction test problem

Equations

TS ex22.c

$$u_t + a_1 u_x = -k_1 u + k_2 v + s_1$$

$$v_t + a_2 v_x = k_1 u - k_2 v + s_2$$

```

for (i = info.xs; i < info.xs+info.xm; ++i) {
for (j = 0; j < 2; ++j) {
    const PetscReal a = a[j]/hx;
    if (i == 0)          f[i][j] =
        a*(1/3*u0t[j] + 1/2*x[i][j] - x[i+1][j] + 1/6*x[i+2][j]);
    else if (i == 1)    f[i][j] =
        a*(-1/12*u0t[j] + 2/3*x[i-1][j] - 2/3*x[i+1][j] + 1/12*x[i+2][j]);
    else if (i == info.mx-2) f[i][j] =
        a*(-1/6*x[i-2][j] + x[i-1][j] - 1/2*x[i][j] - 1/3*x[i+1][j]);
    else if (i == info.mx-1) f[i][j] =
        a*(-x[i][j] + x[i-1][j]);
    else                f[i][j] =
        a*(-1/12*x[i-2][j] + 2/3*x[i-1][j] - 2/3*x[i+1][j] + 1/12*x[i+2][j]);
}
}

```

Stiff linear advection-reaction test problem

Parameters

TS ex22.c

$$a_1 = 1,$$

$$a_2 = 0,$$

$$k_1 = 10^6,$$

$$k_2 = 2k_1,$$

$$s_1 = 0,$$

$$s_2 = 1$$

Stiff linear advection-reaction test problem

Initial conditions

TS ex22.c

$$u(x, 0) = 1 + s_2 x$$

$$v(x, 0) = \frac{k_0}{k_1} u(x, 0) + \frac{s_1}{k_1}$$

```

PetscErrorCode FormInitialSolution(TS ts, Vec X, void *ctx) {
    TSGetDM(ts, &da);
    DMGetLocalInfo(da, &info);
    DMGetVecArray(da, X, &x);
    /* Compute function over the locally owned part of the grid */
    for (i = info.xs; i < info.xs+info.xm; ++i) {
        PetscReal r = (i+1)*hx;
        PetscReal ik = user->k[1] != 0.0 ? 1.0/user->k[1] : 1.0;
        x[i][0] = 1 + user->s[1]*r;
        x[i][1] = user->k[0]*ik*x[i][0] + user->s[1]*ik;
    }
    DMRestoreArray(da, X, &x);
}

```

Stiff linear advection-reaction test problem

Initial conditions

TS ex22.c

$$u(x, 0) = 1 + s_2 x$$

$$v(x, 0) = \frac{k_0}{k_1} u(x, 0) + \frac{s_1}{k_1}$$

```

PetscErrorCode FormInitialSolution(TS ts, Vec X, void *ctx) {
    TSGetDM(ts, &da);
    DMGetLocalInfo(da, &info);
    DMGetVecArray(da, X, &x);
    /* Compute function over the locally owned part of the grid */
    for (i = info.xs; i < info.xs+info.xm; ++i) {
        PetscReal r = (i+1)*hx;
        PetscReal ik = user->k[1] != 0.0 ? 1.0/user->k[1] : 1.0;
        x[i][0] = 1 + user->s[1]*r;
        x[i][1] = user->k[0]*ik*x[i][0] + user->s[1]*ik;
    }
    DMRestoreVecArray(da, X, &x);
}

```

Stiff linear advection-reaction test problem

Examples

TS ex22.c

- `./ex22 -da_grid_x 200 -ts_monitor_draw_solution -ts_arkimex_type 4 -ts_adapt_type none`
- `./ex22 -da_grid_x 200 -ts_monitor_draw_solution -ts_type ros w -ts_dt 1e-3 -ts_adapt_type none`
- `./ex22 -da_grid_x 200 -ts_monitor_draw_solution -ts_type ros w -ts_ros w_type sandu3 -ts_dt 5e-3 -ts_adapt_type none`
- `./ex22 -da_grid_x 200 -ts_monitor_draw_solution -ts_type ros w -ts_ros w_type ra34pw2 -ts_adapt_monitor`

1D Brusselator reaction-diffusion

Equations

TS ex25.c

$$u_t - \alpha u_{xx} = A - (B + 1)u + u^2 v$$

$$v_t - \alpha v_{xx} = Bu - u^2 v$$

Boundary conditions:

$$u(0, t) = u(1, t) = 1$$

$$v(0, t) = v(1, t) = 3$$

1D Brusselator reaction-diffusion

Equations

TS ex25.c

$$u_t - \alpha u_{xx} = A - (B + 1)u + u^2v$$

$$v_t - \alpha v_{xx} = Bu - u^2v$$

```

FormIFunction(TS ts, PetscReal t, Vec X, Vec Xdot, Vec F, void *ptr) {
  DMGlobalToLocalBegin(da, X, INSERT_VALUES, Xloc);
  DMGlobalToLocalEnd(da, X, INSERT_VALUES, Xloc);
  for (i = info.xs; i < info.xs+info.xm; ++i) {
    if (i == 0) {
      f[i].u = hx * (x[i].u - uleft);
      f[i].v = hx * (x[i].v - vleft);
    } else if (i == info.mx-1) {
      f[i].u = hx * (x[i].u - uright);
      f[i].v = hx * (x[i].v - vright);
    } else {
      f[i].u = hx * xdot[i].u - alpha * (x[i-1].u - 2.*x[i].u + x[i+1].u)
      f[i].v = hx * xdot[i].v - alpha * (x[i-1].v - 2.*x[i].v + x[i+1].v)
    }
  }
}

```

1D Brusselator reaction-diffusion

Equations

TS ex25.c

$$u_t - \alpha u_{xx} = A - (B + 1)u + u^2v$$

$$v_t - \alpha v_{xx} = Bu - u^2v$$

```

FormIJacobian(TS ts, PetscReal t, Vec X, Vec Xdot, PetscReal a, Mat *J,
              Mat *Jpre, MatStructure *str, void *ptr) {
  for (i = info.xs; i < info.xs+info.xm; ++i) {
    if (i == 0 || i == info.mx-1) {
      const PetscInt   row = i, col = i;
      const PetscScalar vals[2][2] = {{hx, 0}, {0, hx}};
      MatSetValuesBlocked(*Jpre, 1, &row, 1, &col, &vals[0][0], INSERT_VALUES);
    } else {
      const PetscInt   row = i, col[] = {i-1, i, i+1};
      const PetscScalar dL = -alpha/hx, dC = 2*alpha/hx, dR = -alpha/hx;
      const PetscScalar v[2][3][2] = {{{dL, 0}, {a*hx+dC, 0}, {dR, 0}},
                                       {{0, dL}, {0, a*hx+dC}, {0, dR}}};
      MatSetValuesBlocked(*Jpre, 1, &row, 3, col, &v[0][0][0], INSERT_VALUES);
    }
  }
}

```

1D Brusselator reaction-diffusion

Equations

TS ex25.c

$$U_t - \alpha U_{xx} = A - (B + 1)u + u^2 v$$

$$V_t - \alpha V_{xx} = Bu - u^2 v$$

```

FormRHSFunction(TS ts, PetscReal t, Vec X, Vec F, void *ptr) {
    TSGetDM(ts, &da);
    MDMAGetLocalInfo(da, &info);
    MDMAVecGetArray(da, X, &x);
    MDMAVecGetArray(da, F, &f);
    for (i = info.xs; i < info.xs+info.xm; ++i) {
        PetscScalar u = x[i].u, v = x[i].v;
        f[i].u = hx*(A - (B+1)*u + u*u*v);
        f[i].v = hx*(B*u - u*u*v);
    }
    MDMAVecRestoreArray(da, X, &x);
    MDMAVecRestoreArray(da, F, &f);
}

```

1D Brusselator reaction-diffusion

Parameters

TS ex25.c

$$A = 1,$$

$$B = 3,$$

$$\alpha = 1/50$$

1D Brusselator reaction-diffusion

Initial conditions

TS ex25.c

$$u(x, 0) = 1 + \sin(2\pi x)$$

$$v(x, 0) = 3$$

```
PetscErrorCode FormInitialSolution(TS ts, Vec X, void *ctx) {
    TSGetDM(ts, &da);
    DMGetLocalInfo(da, &info);
    DMGetVecArray(da, X, &x);
    /* Compute function over the locally owned part of the grid */
    for (i = info.xs; i < info.xs+info.xm; ++i) {
        PetscReal xi = i*hx;
        x[i].u = uleft*(1-xi) + uright*xi + sin(2*PETSC_PI*xi);
        x[i].v = vleft*(1-xi) + vright*xi;
    }
    DMGetVecArray(da, X, &x);
}
```

1D Brusselator reaction-diffusion

Initial conditions

TS ex25.c

$$u(x, 0) = 1 + \sin(2\pi x)$$

$$v(x, 0) = 3$$

```
PetscErrorCode FormInitialSolution(TS ts, Vec X, void *ctx) {
    TSGetDM(ts, &da);
    DMGetLocalInfo(da, &info);
    DMGetVecArray(da, X, &x);
    /* Compute function over the locally owned part of the grid */
    for (i = info.xs; i < info.xs+info.xm; ++i) {
        PetscReal xi = i*hx;
        x[i].u = uleft*(1-xi) + uright*xi + sin(2*PETSC_PI*xi);
        x[i].v = vleft*(1-xi) + vright*xi;
    }
    DMRestoreVecArray(da, X, &x);
}
```

1D Brusselator reaction-diffusion

Examples

TS ex25.c

- `./ex25 -da_grid_x 20 -ts_monitor_draw_solution -ts_type ros`
`-ts_dt 5e-2 -ts_adapt_type none`
- `./ex25 -da_grid_x 20 -ts_monitor_draw_solution -ts_type ros`
`-ts_ros_type 2p -ts_dt 5e-2`
- `./ex25 -da_grid_x 20 -ts_monitor_draw_solution -ts_type ros`
`-ts_ros_type 2p -ts_dt 5e-2 -ts_adapt_type none`

Second Order TVD Finite Volume Method

Example

TS ex11.c

```
MESH_DIR = $PETSC_DIR/share/petsc/datafiles/meshes
```

- `./ex11 -f $MESH_DIR/sevenside.exo`
- `./ex11 -f $MESH_DIR/sevenside-quad-15.exo`
- `./ex11 -f $MESH_DIR/sevenside.exo -ts_type ros`

Second Order TVD Finite Volume Method

Example

TS ex11.c

```
./ex11 -f -grid_size 2,1 -grid_bounds -1,1.,0.,1 -grid_skew_60  
-physics euler -bc_wall 1,2,3,4 -eu_type iv_shock -ufv_cfl 10  
-eu_alpha 60. -eu_gamma 1.4 -eu_amach 2.02 -eu_rho2 3.  
-dm_type p4est -dm_forest_partition_overlap 1  
-dm_forest_maximum_refinement 6 -dm_forest_minimum_refinement 2  
-dm_forest_initial_refinement 2  
-petscfv_type leastsquares -petscfv_compute_gradients 0  
-petsclimiter_type minmod  
-ts_final_time 1 -ts_ssp_type rks2 -ts_ssp_nstages 10  
-ufv_use_amr -ufv_refine_tol 0.5 -ufv_coarsen_tol 1.e-2  
-ufv_vtk_interval 1 -monitor density,energy  
-ufv_view_initial_refinement
```

Outline

1 Getting Started with PETSc

2 PETSc Integration

3 DM

4 Advanced Solvers

5 More Stuff

Things I Will Not Talk About

- Communication abstractions
 - `PetscSF` and `VecScatter`
- Finite elements and finite volumes
 - `PetscFE` and `PetscFV`
- Non-nested geometric multigrid
 - Uses `DMplex` and `Pragmatic`
- Timestepping for second order systems and stiff systems
 - `TSALPHA2` and `TSBDF`