

XBraid Tutorial

A flexible and scalable approach to parallel-in-time

Jacob Schroder and Rob Falgout

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Outline

1. **Introduction**
→ Tutorial software requirements and XBraid overview
2. **Simplest example of solving a scalar ODE with examples/ex-01**
→ Defining the `App` and `vector` structures, writing wrapper functions, running XBraid
3. **Explore more XBraid settings in examples/ex-01-expanded.c**
4. **Porting a user-code to XBraid with examples/ex-02**
→ Debugging the connection to XBraid
→ Intrusiveness versus efficiency
5. **A few application area highlights**

Appendix: Advanced XBraid features

- *Temporal adaptivity*
- *Shell-vectors and BDF-k*
- *Fortran90 Interface*
- *Residual and storage options*
- *Spatial coarsening*



To interact with the tutorial, you need

- This tutorial needs a working installation of XBraid 2.1 or higher
<http://llnl.gov/casc/xbraid/>
 - See the User's manual for instructions on how to install XBraid
 - See the "Publications" page for a copy of this tutorial
- XBraid v2.1 (or higher) *required*
- GCC compiler *required*
- MPI *recommended*
- Python 2.7 (or higher) with NumPy, Matplotlib *recommended*
- *hypre* installation for running ex-03 *optional*
<http://llnl.gov/casc/hypre>



To interact with the tutorial, you need

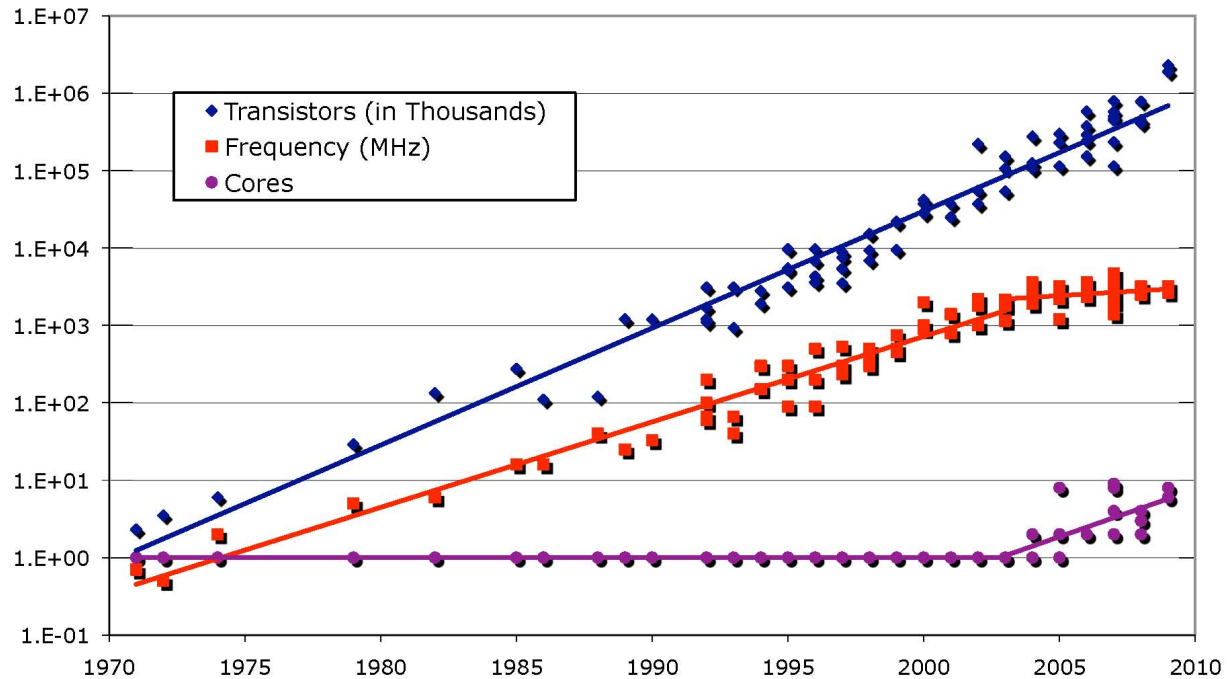
- Make sure you can run

```
$ cd xbraid
$ make
$ cd examples
$ make ex-01 ex-02
$ ./ex-01
Braid: || r_1 || = 2.845538e-02, conv factor = 1.00e+00, wall time = ...
Braid: || r_2 || = 8.621939e-04, conv factor = 3.03e-02, wall time = ...
Braid: || r_3 || = 0.000000e+00, conv factor = 0.00e+00, wall time = ...
...
$ ./ex-02
Braid: || r_0 || = 4.041694e+00, conv factor = 1.00e+00, wall time = ...
Braid: || r_1 || = 1.037471e-01, conv factor = 2.57e-02, wall time = ...
Braid: || r_2 || = 2.926906e-03, conv factor = 2.82e-02, wall time = ...
...
```



Traditional time integration will become a sequential bottleneck

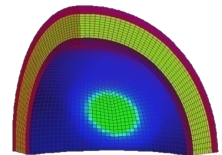
From Kathy Yelick's talk titled "Ten Ways to Waste a Parallel Computer."



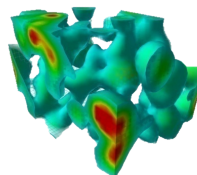
- Clock rates are no longer increasing – faster speed is now achieved through more concurrency
- Parallel time integration methods are needed (think exascale)!

Multigrid is well suited for exascale

- For many applications, the fastest and most scalable solvers are already multigrid methods



Elasticity / Plasticity



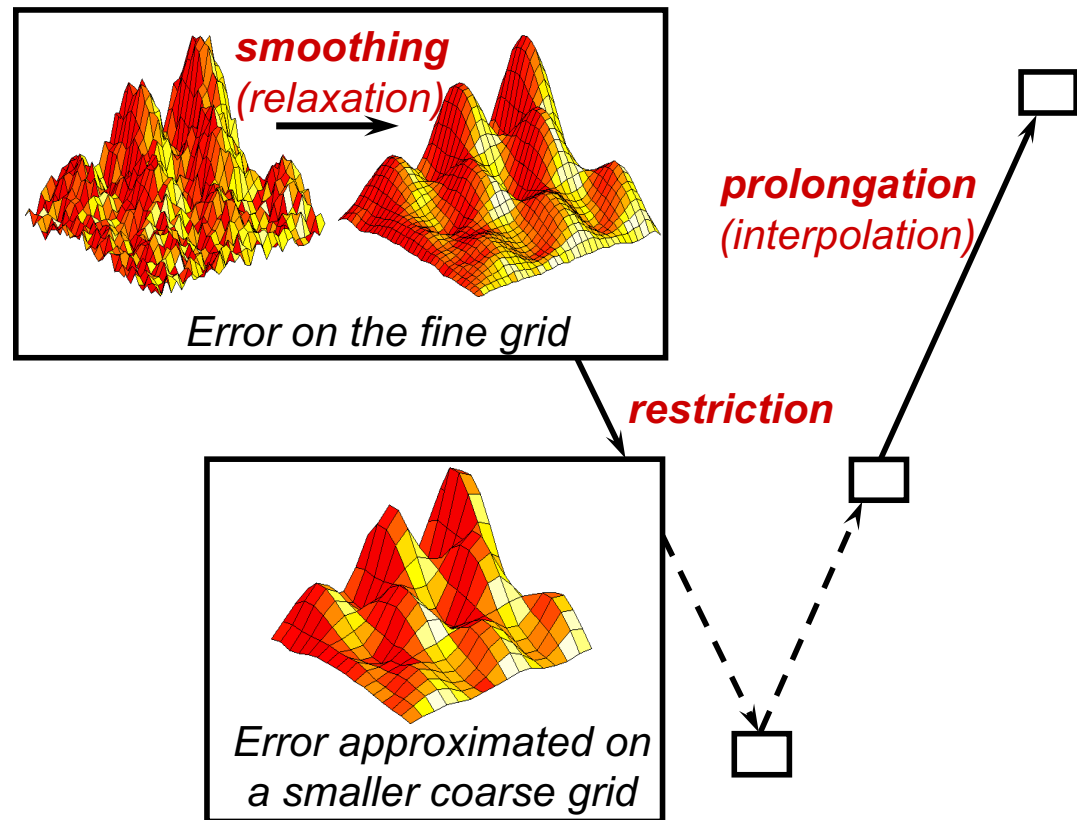
Quantum Chromodynamics

- Exascale solver algorithms will need to:
 - Exhibit extreme levels of parallelism (exascale → 1 billion cores)
Spatial multigrid has already scaled to over 1 million cores
 - Minimize data movement
Multigrid is $O(N)$ optimal
 - Exploit machine heterogeneity
If the user's problem can exploit heterogeneity, then so can multigrid
 - Be resilient to faults
Multigrid has already shown good resilience (being iterative and multilevel helps)
- Apply multigrid to the temporal dimension!

Our approach for parallel-in-time

- Apply the wealth of research on parallel spatial multigrid to multigrid in time
- This is where our team has extensive experience (*hypre* project)

The Multigrid V-cycle



Technical approach

- Consider the **general** one-step method

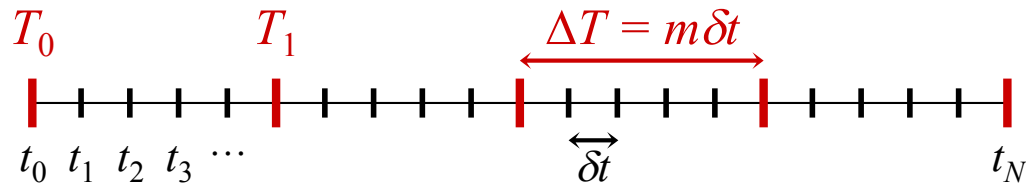
$$\mathbf{u}_i = \Phi_i(\mathbf{u}_{i-1}) + \mathbf{g}_i, \quad i = 1, 2, \dots, N$$

- In the linear setting (*for simplicity*), time marching \equiv forward solve
 - This is an $O(N)$ direct method, **but sequential**

$$A\mathbf{u} \equiv \begin{pmatrix} I & & & \\ -\Phi & I & & \\ & \ddots & \ddots & \\ & & -\Phi & I \end{pmatrix} \begin{pmatrix} \mathbf{u}_0 \\ \mathbf{u}_1 \\ \vdots \\ \mathbf{u}_N \end{pmatrix} = \begin{pmatrix} \mathbf{g}_0 \\ \mathbf{g}_1 \\ \vdots \\ \mathbf{g}_N \end{pmatrix} \equiv \mathbf{g}$$

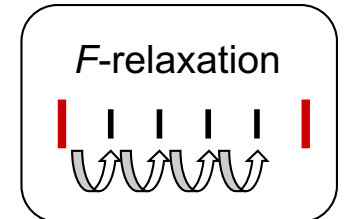
- We propose solving this system **iteratively** with a multigrid method
 - Extend multigrid reduction (MGR, 1979) to the time dimension
 - Coarsens only in time (non-intrusive)
 - $O(N)$, highly parallel

Technical approach



- *F*-point (fine grid only)
- *C*-point (coarse & fine grid)

- Relaxation is highly parallel
 - Alternates between *F*-points and *C*-points
 - *F*-point relaxation = integration over each coarse time interval
- Coarse system is a time rediscretization
 - Approximate impractical Φ^m with Φ_Δ a rediscretization with ΔT

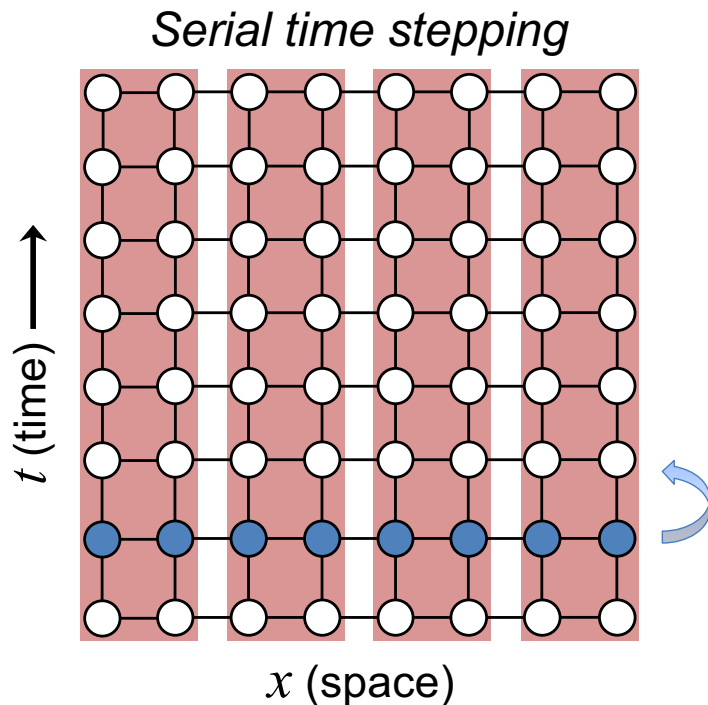


$$A_\Delta = \begin{pmatrix} I & & & \\ -\Phi^m & I & & \\ & \ddots & \ddots & \\ & & -\Phi^m & I \end{pmatrix} \Rightarrow A_\Delta = \begin{pmatrix} I & & & \\ -\Phi_\Delta & I & & \\ & \ddots & \ddots & \\ & & -\Phi_\Delta & I \end{pmatrix}$$

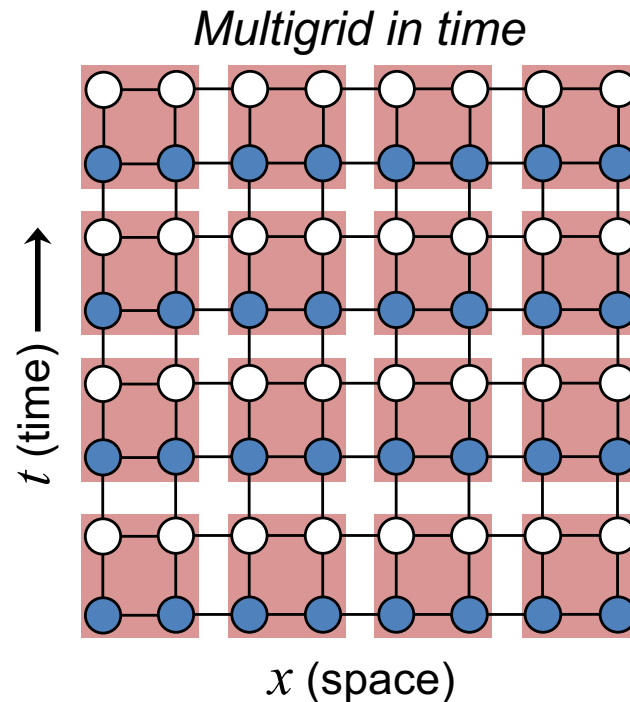
- Apply recursively for multilevel hierarchy

Parallel decomposition

- Our code **XBraid** is agnostic to spatial decomposition and only parallelizes in time



Negative: Parallelize in space only
Positive: Store only one time step



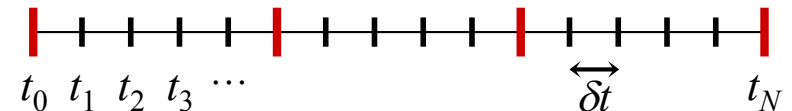
Positive: Parallelize in space and time
Negative: Store several time steps

Properties of the approach

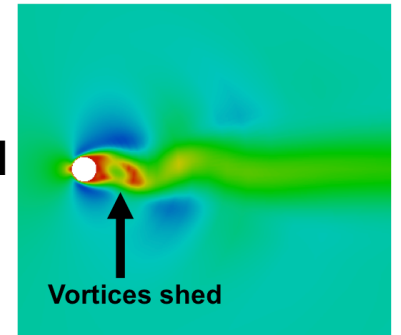
- Expose **concurrency** in the time dimension with multigrid
- Non-intrusive**, with unchanged time discretization
- Converges to **same solution** as sequential time stepping

$$\begin{pmatrix} I & & & & \\ -\Phi & I & & & \\ \ddots & \ddots & \ddots & & \\ & -\Phi & I & & \end{pmatrix}$$

- Only store **C-points** to minimize storage
- Optimal for variety of parabolic problems
 - Converges in ~10 iterations for any coarsening factor
 - Larger factors require larger (sequential) F-relaxation intervals

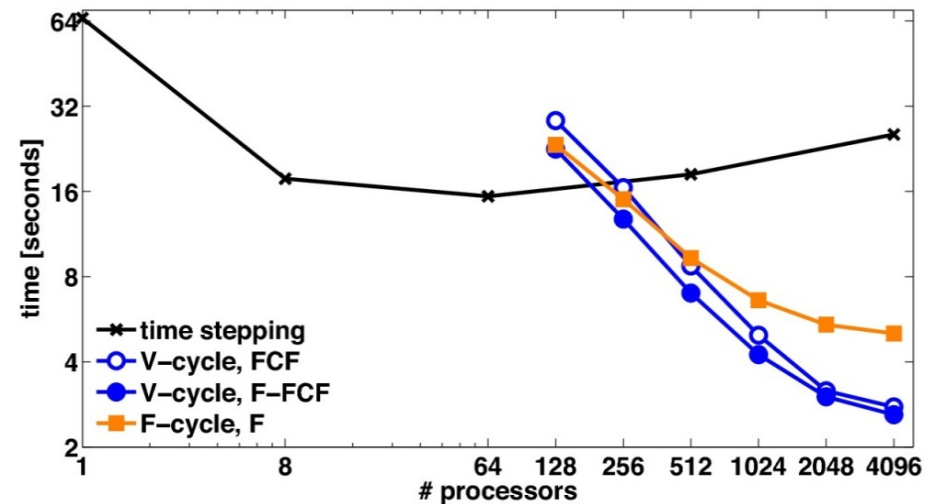


- Extends to **nonlinear** problems with FAS formulation
- In simple two-level setting, our method is **equivalent to parareal**
 - This is a popular method, typically not viewed as multigrid
- Many active research topics
 - Adaptivity in time, moving meshes and multistep methods all possible
 - Space-time coarsening possible (stability on coarse time-grids for explicit schemes)



Huge parallel speedups available, but in a new way

- Time stepping is already $O(N)$
- Useful only beyond a crossover
- Need 10-100x more parallelism just to break even



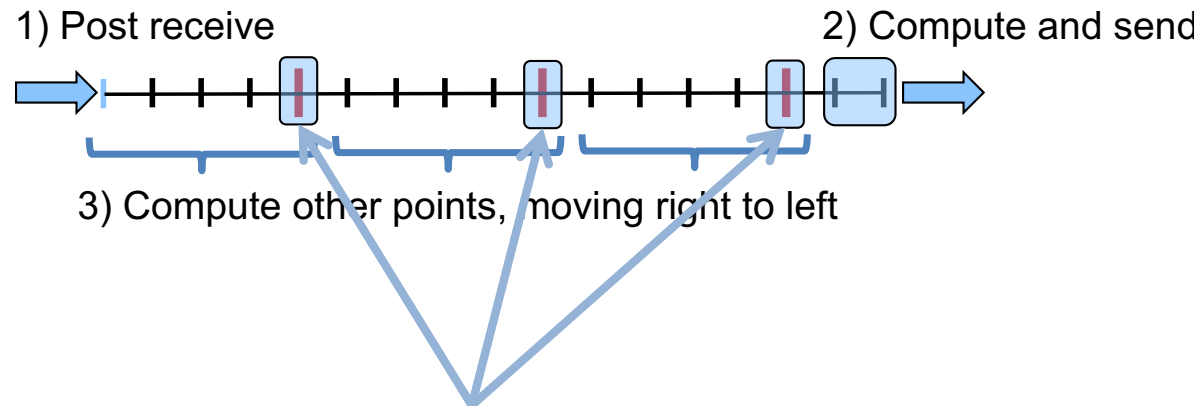
3D Heat Equation: $33^3 \times 4097$,
8 procs in space, **6x speedup**

- The more time steps, the more speedup potential
 - Applications that require lots of time steps will benefit first
 - Speedups (so far) up to 52x on 100K total cores

XBraid: open source, non-intrusive and flexible



- Overlap communication and computation
 - Consider relaxation over a processor's portion of the time interval
 - Start computation with right-most interval to overlap comm/comp



- Code stores only C -points to minimize storage
 - Ability to coarsen by large factors means fewer parallel resources
 - Memory multiplier per processor
 $O(\log N)$ with time coarsening, $O(1)$ with space-time coarsening

XBraid: open source, non-intrusive and flexible



- User defines two objects:
 - App and Vector
- User also writes several wrapper routines:
 - *Step, Init, Clone, Sum, SpatialNorm, Access, BufPack, BufUnpack, BufSize*
 - For optional spatial coarsening: Coarsen, Refine
- Example: Step(app, u, status)
 - Advances vector u from time t_{start} to t_{stop}
 - Returns a target refinement factor

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Appendix: Advanced XBraid features

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Simplest Example: Scalar ODE

- File: examples/ex-01.c Solves: $u_t = \lambda u$
- First, you must define your app and vector structures

This is your simulation application structure. Place any time-independent data here, which is needed to take a time step.

Here, we only need the MPI rank in the App structure (for later file output).

```
typedef struct _braid_App_struct{
    int          rank;
} my_App;

typedef struct _braid_Vector_struct{
    double value;
} my_Vector;
```

Simplest Example: Scalar ODE

- File: examples/ex-01.c Solves: $u_t = \lambda u$
- First, you must define your app and vector structures

This is your state vector structure. It holds any time-dependent information that should stay with a vector, e.g. mesh information and unknowns.

For this problem, the vector is one double.

```
typedef struct _braid_App_struct{
    int          rank;
} my_App;

[ ] typedef struct _braid_Vector_struct{
    double value;
} my_Vector;
```

Define the Step () function

- File: examples/ex-01.c

Solves: $u_t = \lambda u$

Step () evolves u from tstart to tstop

```
int my_Step(braid_App           app,
            braid_Vector        ustop,
            braid_Vector        fstop,
            braid_Vector        u,
            braid_StepStatus    status)
{
    double tstart;
    double tstop;
    braid_StepStatusGetTstartTstop(status, &tstart, &tstop);

    (u->value) = 1./(1. + tstop-tstart) * (u->value);

    return 0;
}
```

Define the Step() function

- File: examples/ex-01.c

Solves: $u_t = \lambda u$

The app structure is passed into every user-written function.

```
int my_Step(braid_App      app,
            braid_Vector  ustop,
            braid_Vector  fstop,
            braid_Vector  u,
            braid_StepStatus status)
{
    double tstart;
    double tstop;
    braid_StepStatusGetTstartTstop(status, &tstart, &tstop);

    (u->value) = 1./(1. + tstop-tstart) * (u->value);

    return 0;
}
```

Define the Step() function

- File: examples/ex-01.c

Solves: $u_t = \lambda u$

Vector at tstop from previous XBraid iteration (initial guess for implicit solvers)

```
int my_Step(braid_App           app,
            braid_Vector      ustop,
            braid_Vector      fstop,
            braid_Vector      u,
            braid_StepStatus  status)
{
    double tstart;
    double tstop;
    braid_StepStatusGetTstartTstop(status, &tstart, &tstop);

    (u->value) = 1./(1. + tstop-tstart) * (u->value);

    return 0;
}
```

Define the Step() function

- File: examples/ex-01.c

Solves: $u_t = \lambda u$

Vector at tstart

```
int my_Step(braid_App           app,
            braid_Vector      ustop,
            braid_Vector      fstop,
            braid_Vector      u,
            braid_StepStatus  status)

{
    double tstart;
    double tstop;
    braid_StepStatusGetTstartTstop(status, &tstart, &tstop);

    (u->value) = 1./(1. + tstop-tstart) * (u->value);

    return 0;
}
```



Define the Step() function

- File: examples/ex-01.c

Solves: $u_t = \lambda u$

Ignore by default. (XBraid forcing term, only needed if residual option is used)

```
int my_Step(braid_App           app,
            braid_Vector      ustop,
            braid_Vector      fstop,
            braid_Vector      u,
            braid_StepStatus  status)
{
    double tstart;
    double tstop;
    braid_StepStatusGetTstartTstop(status, &tstart, &tstop);

    (u->value) = 1./ (1. + tstop-tstart) * (u->value);

    return 0;
}
```

Define the Step() function

- File: examples/ex-01.c

Solves: $u_t = \lambda u$

Status structures can be queried for various information (level, iteration, etc...)

```
int my_Step(braid_App           app,
            braid_Vector      ustop,
            braid_Vector      fstop,
            braid_Vector      u,
            braid_StepStatus  status)
{
    double tstart;
    double tstop;
    braid_StepStatusGetTstartTstop(status, &tstart, &tstop);

    (u->value) = 1./(1. + tstop-tstart) * (u->value);

    return 0;
}
```

Define the Step() function

- File: examples/ex-01.c

Solves: $u_t = \lambda u$

For instance, to get tstart, tstop

```
int my_Step(braid_App           app,
            braid_Vector      ustop,
            braid_Vector      fstop,
            braid_Vector      u,
            braid_StepStatus  status)
{
    double tstart;
    double tstop;
    braid_StepStatusGetTstartTstop(status, &tstart, &tstop);

    (u->value) = 1./(1. + tstop-tstart) * (u->value);

    return 0;
}
```

Define the Step() function

- File: examples/ex-01.c

Solves: $u_t = \lambda u$

Take backward Euler step

```
int my_Step(braid_App           app,
            braid_Vector        ustop,
            braid_Vector        fstop,
            braid_Vector        u,
            braid_StepStatus    status)
{
    double tstart;
    double tstop;
    braid_StepStatusGetTstartTstop(status, &tstart, &tstop);

    { (u->value) = 1./ (1. + tstop-tstart) * (u->value);

    return 0;
}
```

Define other wrapper functions

- File: examples/ex-01.c Solves: $u_t = \lambda u$
- Define functions: Init, Clone, Free, **Sum**, SpatialNorm, Access, BufPack, BufUnpack, BufSize

Again, we see the app structure being passed in

```
int my_Sum(braid_App      app,
            double        alpha,
            braid_Vector  x,
            double        beta,
            braid_Vector  y)
{
    (y->value) = alpha*(x->value) + beta*(y->value);
    return 0;
}
```

Define other wrapper functions

- File: examples/ex-01.c Solves: $u_t = \lambda u$
- Define functions: Init, Clone, Free, **Sum**, SpatialNorm, Access, BufPack, BufUnpack, BufSize

This function carries out a simple AXPY operation

```
int my_Sum(braid_App           app,
            double          alpha,
            braid_Vector   x,
            double          beta,
            braid_Vector   y)
{
    [y->value] = alpha*(x->value) + beta*(y->value);
    return 0;
}
```

Define other wrapper functions

- File: examples/ex-01.c Solves: $u_t = \lambda u$
- Define functions: Init, Clone, Free, Sum, SpatialNorm, **Access**, BufPack, BufUnpack, BufSize

This function is how the user accesses the solution

- **By default, it is called at the end of the simulation for every time point**
- **Using `braid_AccessSetLevel()` allows for more frequent access**

```
int my_Access(braid_App           app,
              braid_Vector        u,
              braid_AccessStatus astatus)
{
    int index;  char  filename[255]; FILE *file;

    braid_AccessStatusGetTIndex(astatus, &index);
    sprintf(filename, "%s.%04d.%03d", "ex-01.out", index, app->rank);
    file = fopen(filename, "w");
    fprintf(file, "%14e\n", (u->value));

    fflush(file); fclose(file); return 0;
}
```



Define other wrapper functions

- File: examples/ex-01.c Solves: $u_t = \lambda u$
- Define functions: Init, Clone, Free, Sum, SpatialNorm, **Access**, BufPack, BufUnpack, BufSize

Here, we just write a single solution value to individual files

```
int my_Access(braid_App           app,
               braid_Vector        u,
               braid_AccessStatus astatus)
{
    int index;   char  filename[255]; FILE *file;

    braid_AccessStatusGetTIndex(astatus, &index);
    sprintf(filename, "%s.%04d.%03d", "ex-01.out", index, app->rank);
    file = fopen(filename, "w");
    fprintf(file, "% .14e\n", (u->value));
    fflush(file); fclose(file); return 0;
}
```

Define other wrapper functions

- File: examples/ex-01.c Solves: $u_t = \lambda u$
- Define functions: Init, Clone, Free, Sum, SpatialNorm, Access, **BufPack**, BufUnpack, BufSize

The **Buf*** functions tell XBraid how to pack, unpack and size MPI Buffers

Define other wrapper functions

- File: examples/ex-01.c Solves: $u_t = \lambda u$
- Define functions: Init, Clone, Free, Sum, SpatialNorm, Access, **BufPack**, BufUnpack, BufSize

BufPack() flattens the vector **u** into **buffer**

```
int my_BufPack(braid_App app,
                braid_Vector u,
                void *buffer,
                braid_BufferStatus bstatus)
{
    double *dbuf = buffer;

    dbuf[0] = (u->value);
    braid_BufferStatusSetSize( bstatus, sizeof(double) );

    return 0;
}
```

Define other wrapper functions

- File: examples/ex-01.c Solves: $u_t = \lambda u$
- Define functions: Init, Clone, Free, Sum, SpatialNorm, Access, **BufPack**, BufUnpack, BufSize

Packing this buffer entails just setting a single double value

```
int my_BufPack(braid_App           app,
                braid_Vector        u,
                void                *buffer,
                braid_BufferStatus bstatus)
{
    double *dbuf = buffer;

    dbuf[0] = (u->value);
    braid_BufferStatusSetSize( bstatus, sizeof(double) );

    return 0;
}
```

Define other wrapper functions

- File: examples/ex-01.c Solves: $u_t = \lambda u$
- Define functions: Init, Clone, Free, Sum, SpatialNorm, Access, **BufPack**, BufUnpack, BufSize

This is an example of returning a value (the buffer size) with a status structure

```
int my_BufPack(braid_App           app,
                braid_Vector        u,
                void               *buffer,
                braid_BufferStatus bstatus)
{
    double *dbuf = buffer;

    dbuf[0] = (u->value);
    braid_BufferStatusSetSize( bstatus, sizeof(double) );

    return 0;
}
```

Initialize App and XBraid

- File: examples/ex-01.c Solves: $u_t = \lambda u$
- The next step is to setup XBraid in main()

```
int main()
{
    ...
    braid_Core    core;
    ntime   = 10;
    tstart  = 0.0; tstop   = 5.0;
    ...
    app = (my_App *) malloc(sizeof(my_App));
    (app->rank)    = rank;
    ...
    braid_Init(MPI_COMM_WORLD, MPI_COMM_WORLD, tstart, tstop,
               ntime, app, my_Step, my_Init, my_Clone,
               my_Free, my_Sum, my_SpatialNorm,
               my_Access, my_BufSize, my_BufPack,
               my_BufUnpack, &core);
}
```

Initialize App and XBraid

- File: examples/ex-01.c Solves: $u_t = \lambda u$
- The next step is to setup XBraid in `main()`

braid_Core is the core data structure, holding all of XBraid's internals

```
int main()
{
    ...
    braid_Core    core;
    ntime   = 10;
    tstart  = 0.0; tstop   = 5.0;
    ...
    app = (my_App *) malloc(sizeof(my_App));
    (app->rank)   = rank;
    ...
    braid_Init(MPI_COMM_WORLD, MPI_COMM_WORLD, tstart, tstop,
               ntime, app, my_Step, my_Init, my_Clone,
               my_Free, my_Sum, my_SpatialNorm,
               my_Access, my_BufSize, my_BufPack,
               my_BufUnpack, &core);
}
```

Initialize App and XBraid

- File: examples/ex-01.c Solves: $u_t = \lambda u$
- The next step is to setup XBraid in `main()`

Define your time domain

```
int main()
{
    ...
    braid_Core    core;
    ntime   = 10;
    tstart = 0.0; tstop   = 5.0;
    ...
    app = (my_App *) malloc(sizeof(my_App));
    (app->rank)   = rank;
    ...
    braid_Init(MPI_COMM_WORLD, MPI_COMM_WORLD, tstart, tstop,
               ntime, app, my_Step, my_Init, my_Clone,
               my_Free, my_Sum, my_SpatialNorm,
               my_Access, my_BufSize, my_BufPack,
               my_BufUnpack, &core);
}
```

Initialize App and XBraid

- File: examples/ex-01.c Solves: $u_t = \lambda u$
- The next step is to setup XBraid in `main()`

Initialize App structure

```
int main()
{
    ...
    braid_Core      core;
    ntime   = 10;
    tstart  = 0.0; tstop   = 5.0;
    ...
    app = (my_App *) malloc(sizeof(my_App));
    (app->rank)    = rank;
    ...
    braid_Init(MPI_COMM_WORLD, MPI_COMM_WORLD, tstart, tstop,
               ntime, app, my_Step, my_Init, my_Clone,
               my_Free, my_Sum, my_SpatialNorm,
               my_Access, my_BufSize, my_BufPack,
               my_BufUnpack, &core);
}
```

Initialize App and XBraid

- File: examples/ex-01.c Solves: $u_t = \lambda u$
- The next step is to setup XBraid in `main()`

Initialize `braid_Core`, passing in all user-written functions

```
int main()
{
    ...
    braid_Core    core;
    ntime   = 10;
    tstart  = 0.0; tstop   = 5.0;
    ...
    app = (my_App *) malloc(sizeof(my_App));
    (app->rank)   = rank;
    ...
    braid_Init(MPI_COMM_WORLD, MPI_COMM_WORLD, tstart, tstop,
               ntime, app, my_Step, my_Init, my_Clone,
               my_Free, my_Sum, my_SpatialNorm,
               my_Access, my_BufSize, my_BufPack,
               my_BufUnpack, &core);
}
```

Set XBraid options and run

- File: examples/ex-01.c Solves: $u_t = \lambda u$
- The next step is to setup XBraid in `main()`

Set all the XBraid options that you want

```
int main()
{
    ...
    braid_SetPrintLevel( core, 1);
    braid_SetMaxLevels(core, 2);
    braid_SetAbsTol(core, 1.0e-06);
    braid_SetCFactor(core, -1, 2);

    braid_Drive(core);

    braid_Destroy(core);
}
```

Set XBraid options and run

- File: examples/ex-01.c Solves: $u_t = \lambda u$
- The next step is to setup XBraid in main()

Run the simulation

```
int main()
{
    ...
    braid_SetPrintLevel( core, 1);
    braid_SetMaxLevels(core, 2);
    braid_SetAbsTol(core, 1.0e-06);
    braid_SetCFactor(core, -1, 2);

    [braid_Drive(core);

    braid_Destroy(core);
```



Set XBraid options and run

- File: examples/ex-01.c Solves: $u_t = \lambda u$
- The next step is to setup XBraid in main()

Clean up

```
int main()
{
    ...
    braid_SetPrintLevel( core, 1);
    braid_SetMaxLevels(core, 2);
    braid_SetAbsTol(core, 1.0e-06);
    braid_SetCFactor(core, -1, 2);

    braid_Drive(core);

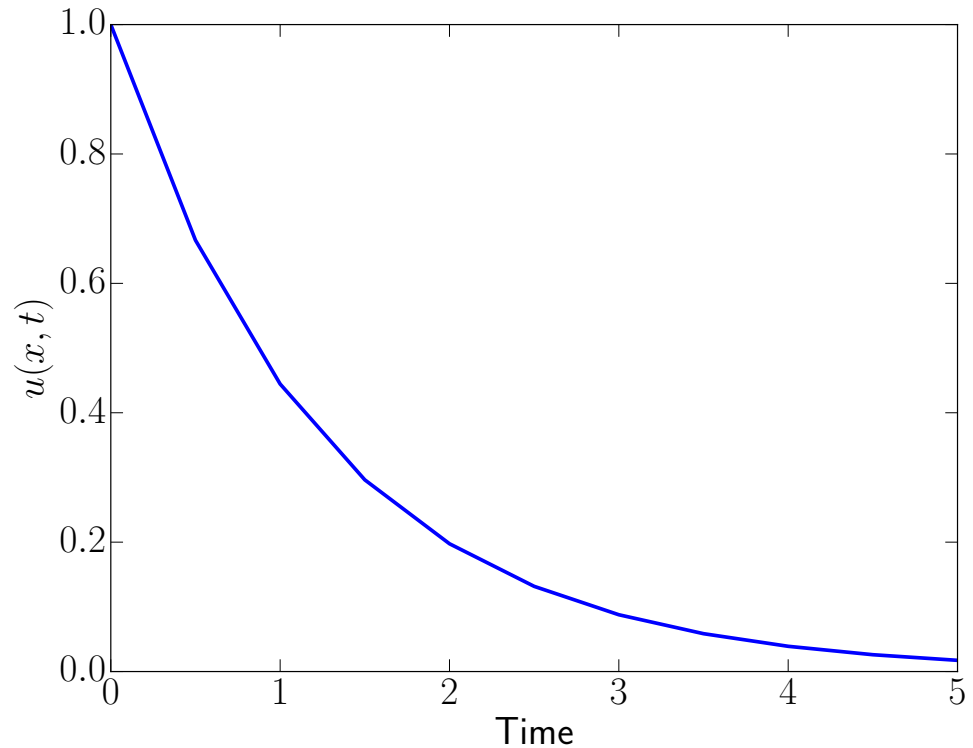
    braid_Destroy(core);
}
```

Output

- File: examples/ex-01.c
- Finally! We can run the example.

```
$ cd examples
$ make ex-01
$ ./ex-01
$ cat ex-01.out.00*
1.00000000000000e+00
6.66666666666667e-01
4.44444444444444e-01
2.96296296296296e-01
1.97530864197531e-01
1.31687242798354e-01
8.77914951989026e-02
5.85276634659351e-02
3.90184423106234e-02
2.60122948737489e-02
1.73415299158326e-02
```

Solves: $u_t = \lambda u$



Outline

1. **Introduction**
→ Tutorial software requirements and XBraid overview
2. **Simplest example of solving a scalar ODE with examples/ex-01**
→ Defining the App and vector structures, writing wrapper functions, running XBraid
3. **Explore more XBraid settings in examples/ex-01-expanded.c**
4. **Porting a user-code to XBraid with examples/ex-02**
→ Debugging the connection to XBraid
→ Intrusiveness versus efficiency
5. **A few application area highlights**

Appendix: Advanced XBraid features

- *Temporal adaptivity*
- *Shell-vectors and BDF-k*
- *Fortran90 Interface*
- *Residual and storage options*
- *Spatial coarsening*



Moving to ex-01-expanded.c

- File: examples/ex-01-expanded.c Solves: $u_t = \lambda u$
- Adds more XBraid features and a command line interface to ex-01.c

Let's experiment with these options!

```
$ cd examples
$ make ex-01-expanded
$ ./ex-01-expanded -help

-ntime <ntime>      : set num time points
-ml  <max_levels>   : set max levels
-nu  <nrelax>        : set num F-C relaxations
-nu0 <nrelax>        : set num F-C relaxations on level 0
-tol <tol>           : set stopping tolerance
-cf   <cfactor>     : set coarsening factor
-mi  <max_iter>      : set max iterations
-fmg                         : use FMG cycling
-res                         : use my residual
-tg  <mydt>          : use user-specified time grid
                           1 - uniform time grid
                           2 - nonuniform time grid
```

Examine the standard XBraid output

- File: examples/ex-01-expanded.c Solves: $u_t = \lambda u$

Residual history is printed out, along with convergence factors and wall times

```
$ ./ex-01-expanded
Braid: Begin simulation, 10 time steps
Braid: || r_0 || not available, wall time = 1.81e-04
Braid: || r_1 || = 2.845538e-02, conv factor = 1.00e+00, wall time = ...
Braid: || r_2 || = 8.621939e-04, conv factor = 3.03e-02, wall time = ...
Braid: || r_3 || = 0.000000e+00, conv factor = 0.00e+00, wall time = ...

start time = 0.000000e+00
stop time  = 5.000000e+00
time steps = 10

use seq soln?          = 0
storage                = -1

stopping tolerance     = 1.000000e-06
use relative tol?      = 0
max iterations         = 100
iterations              = 4
residual norm          = 0.000000e+00
--> 2-norm TemporalNorm
```



Examine the standard XBraid output

- File: examples/ex-01-expanded.c Solves: $u_t = \lambda u$

Basic time domain information

```
$ ./ex-01-expanded
Braid: Begin simulation, 10 time steps
Braid: || r_0 || not available, wall time = 1.81e-04
Braid: || r_1 || = 2.845538e-02, conv factor = 1.00e+00, wall time = ...
Braid: || r_2 || = 8.621939e-04, conv factor = 3.03e-02, wall time = ...
Braid: || r_3 || = 0.000000e+00, conv factor = 0.00e+00, wall time = ...

[ start time = 0.000000e+00
  stop time  = 5.000000e+00
  time steps = 10

  use seq soln?          = 0
  storage                = -1

  stopping tolerance     = 1.000000e-06
  use relative tol?      = 0
  max iterations         = 100
  iterations              = 4
  residual norm           = 0.000000e+00
                           --> 2-norm TemporalNorm
```



Examine the standard XBraid output

- File: examples/ex-01-expanded.c Solves: $u_t = \lambda u$

Advanced options

```
$ ./ex-01-expanded
Braid: Begin simulation, 10 time steps
Braid: || r_0 || not available, wall time = 1.81e-04
Braid: || r_1 || = 2.845538e-02, conv factor = 1.00e+00, wall time = ...
Braid: || r_2 || = 8.621939e-04, conv factor = 3.03e-02, wall time = ...
Braid: || r_3 || = 0.000000e+00, conv factor = 0.00e+00, wall time = ...

start time = 0.000000e+00
stop time  = 5.000000e+00
time steps = 10

use seq soln?          = 0
storage                = -1

stopping tolerance     = 1.000000e-06
use relative tol?      = 0
max iterations         = 100
iterations              = 4
residual norm          = 0.000000e+00
--> 2-norm TemporalNorm
```



Examine the standard XBraid output

- File: examples/ex-01-expanded.c Solves: $u_t = \lambda u$

Describe the XBraid options set for this run

```
$ ./ex-01-expanded
Braid: Begin simulation, 10 time steps
Braid: || r_0 || not available, wall time = ...
Braid: || r_1 || = 2.845538e-02, conv factor = 1.00e+00, wall time = ...
Braid: || r_2 || = 8.621939e-04, conv factor = 3.03e-02, wall time = ...
Braid: || r_3 || = 0.000000e+00, conv factor = 0.00e+00, wall time = ...

start time = 0.000000e+00
stop time  = 5.000000e+00
time steps = 10

use seq soln?          = 0
storage                = -1

[stopping tolerance    = 1.000000e-06
use relative tol?     = 0
max iterations        = 100
iterations             = 4
residual norm          = 0.000000e+00
                         --> 2-norm TemporalNorm]
```

Examine the standard XBraid output

- File: examples/ex-01-expanded.c Solves: $u_t = \lambda u$

Describe the XBraid options set for this run

```
$ ./ex-01-expanded
Braid: Begin simulation, 10 time steps
...
use fmg?          = 0
access_level      = 1
print_level       = 1

max number of levels = 2
min coarse         = 2
number of levels   = 2
skip down cycle    = 1
number of refinements = 0

level  time-pts  cfactor  nrelax
  0        10        2        1
  1         5

wall time = ...
```



Examine the standard XBraid output

- File: examples/ex-01-expanded.c Solves: $u_t = \lambda u$

Describes the levels in the XBraid hierarchy

```
$ ./ex-01-expanded
Braid: Begin simulation, 10 time steps
...
use fmg?          = 0
access_level      = 1
print_level       = 1

max number of levels = 2
min coarse         = 2
number of levels   = 2
skip down cycle    = 1
number of refinements = 0

[ level    time-pts    cfactor    nrelax
  0        10          2           1
  1        5           

wall time = ...
```



Increase number of time points

- File: examples/ex-01-expanded.c Solves: $u_t = \lambda u$

Now, compare the effects of increasing the time domain size

```
$ ./ex-01-expanded -ntime 16
Braid: Begin simulation, 16 time steps
Braid: || r_0 || not available, wall time = ...
Braid: || r_1 || = 2.851025e-02, conv factor = 1.00e+00, wall time = ...
Braid: || r_2 || = 1.040035e-03, conv factor = 3.65e-02, wall time = ...
Braid: || r_3 || = 3.530338e-05, conv factor = 3.39e-02, wall time = ...
Braid: || r_4 || = 3.716892e-07, conv factor = 1.05e-02, wall time = ...
...
$ ./ex-01-expanded -ntime 128
Braid: Begin simulation, 128 time steps
Braid: || r_0 || not available, wall time = ...
Braid: || r_1 || = 2.851112e-02, conv factor = 1.00e+00, wall time = ...
Braid: || r_2 || = 1.049429e-03, conv factor = 3.68e-02, wall time = ...
Braid: || r_3 || = 4.437913e-05, conv factor = 4.23e-02, wall time = ...
Braid: || r_4 || = 1.990483e-06, conv factor = 4.49e-02, wall time = ...
Braid: || r_5 || = 9.174722e-08, conv factor = 4.61e-02, wall time = ...
...
```



FCF-relaxation

- File: examples/ex-01-expanded.c Solves: $u_t = \lambda u$

Observe how changing the number of FCF-relaxations improves convergence

```
$ ./ex-01-expanded -ntime 128 -nu 0
Braid: Begin simulation, 128 time steps
Braid: || r_0 || not available, wall time = ...
Braid: || r_1 || = 6.415003e-02, conv factor = 1.00e+00, wall time = ...
Braid: || r_2 || = 5.312734e-03, conv factor = 8.28e-02, wall time = ...
Braid: || r_3 || = 5.055060e-04, conv factor = 9.51e-02, wall time = ...
Braid: || r_4 || = 5.101391e-05, conv factor = 1.01e-01, wall time = ...
Braid: || r_5 || = 5.290607e-06, conv factor = 1.04e-01, wall time = ...
Braid: || r_6 || = 5.570496e-07, conv factor = 1.05e-01, wall time = ...

$ ./ex-01-expanded -ntime 128 -nu 3
Braid: Begin simulation, 128 time steps
Braid: || r_0 || not available, wall time = ...
Braid: || r_1 || = 5.631827e-03, conv factor = 1.00e+00, wall time = ...
Braid: || r_2 || = 4.094709e-05, conv factor = 7.27e-03, wall time = ...
Braid: || r_3 || = 3.420453e-07, conv factor = 8.35e-03, wall time = ...
```



Halting tolerance and max-iterations

- File: examples/ex-01-expanded.c Solves: $u_t = \lambda u$

Observe how changing the tolerance and max-iter (-mi) parameters affect XBraid

```
$ ./ex-01-expanded -ntime 128 -tol 1e-3
...
iterations          = 4
...

$ ./ex-01-expanded -ntime 128 -tol 1e-12
...
iterations          = 10
...

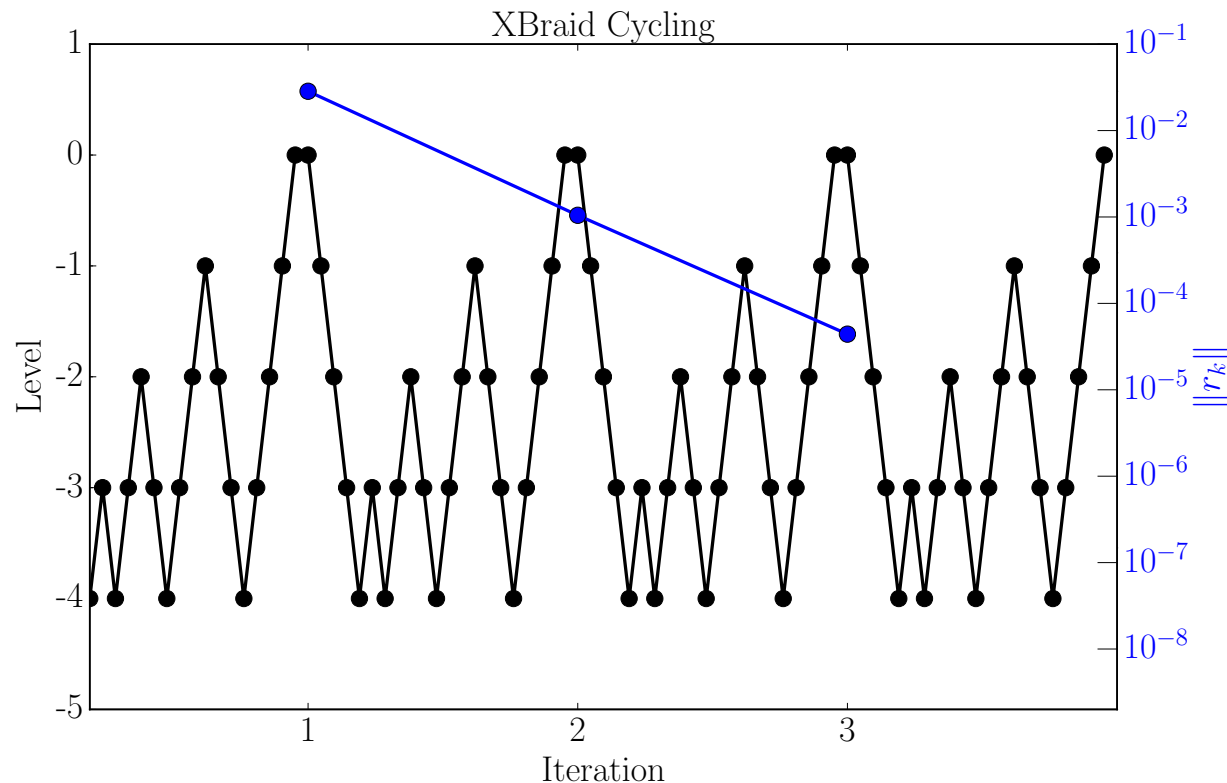
$ ./ex-01-expanded -ntime 128 -tol 1e-12 -mi 3
...
iterations          = 3
...
```

Full multigrid cycles (FMG)

- File: examples/ex-01-expanded.c Solves: $u_t = \lambda u$

Now, use the fmfg parameter and plot braid.out.cycle (file generated at runtime)

```
$ ./ex-01-expanded -ntime 32 -ml 15 -mi 4 -fmfg  
$ python ../user_utils/cycleplot.py
```



This functionality can
be used to adaptively
refine in time
(nested iteration)

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Appendix: Advanced XBraid features

- *Temporal adaptivity*
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- *Spatial coarsening*



How to convert a user-code

- File: examples/ex-02*

Solves: $u_t = -u_{xx}$

ex-02-serial.c

```
/* Set up simulation */
t= 0.0; tstop= 2*PI; ...

/* Initialize u(t=0) */
get_solution(values, ...);

/* Loop over all time values */
for(step=1; step < ntime; step++){
    t = t + deltaT;
    take_step(values, t, ...);
}

/* Process result */
compute_error_norm(values, ...);
save_solution(fname, values, ...);
```

```
$ex-02-serial -ntime 64 -nspatial 17
```

ex-02-lib.c

```
/* Common functions with XBraid */

/* Initialization routine */
void get_solution(...)

/* Helpers for take_step */
void solve_tridiag(...)
void matvec_tridiag(...)
void compute_stencil(...)

/* Core time-stepping routine */
void take_step(...)

/* Output Functions */
double compute_error_norm(...)
void save_solution(...)

/* XBraid specific spatial
interpolation/coarsening */
void interpolate_1D(...)
void coarsen_1D(...)
```

ex-02.c

```
XBraid
Driver
...
```



How to convert a user-code

- File: examples/ex-02*

Solves: $u_t = -u_{xx}$

ex-02-serial.c

```
/* Set up simulation */
t= 0.0; tstop= 2*PI; ...

/* Initialize u(t=0) */
get_solution(values, ...);

/* Loop over all time values */
for(step=1; step < ntime; step++){
    t = t + deltaT;
    take_step(values, t, ...);
}

/* Process result */
compute_error_norm(values, ...);
save_solution(fname, values, ...);
```

\$ex-02-serial -ntime 64 -nspatial 17

ex-02-lib.c

```
/* Common functions with XBraid */

/* Initialization routine */
void get_solution(...)

/* Helpers for take_step */
void solve_tridiag(...)
void matvec_tridiag(...)
void compute_stencil(...)

/* Core time-stepping routine */
void take_step(...)

/* Output Functions */
double compute_error_norm(...)
void save_solution(...)

/* XBraid specific spatial
interpolation/coarsening */
void interpolate_1D(...)
void coarsen_1D(...)
```

ex-02.c

XBraid
Driver
...



How to convert a user-code

- File: examples/ex-02*

Solves: $u_t = -u_{xx}$

ex-02-serial.c

```
/* Set up simulation */
t= 0.0; tstop= 2*PI; ...

/* Initialize u(t=0) */
get_solution(values, ...);

/* Loop over all time values */
for(step=1; step < ntime; step++){
    t = t + deltaT;
    take_step(values, t, ...);
}

/* Process result */
compute_error_norm(values, ...);
save_solution(fname, values, ...);
```

```
$ex-02-serial -ntime 64 -nspatial 17
```

ex-02-lib.c

```
/* Common functions with XBraid */

/* Initialization routine */
void get_solution(...)

/* Helpers for take_step */
void solve_tridiag(...)
void matvec_tridiag(...)
void compute_stencil(...)

/* Core time-stepping routine */
void take_step(...)

/* Output Functions */
double compute_error_norm(...)
void save_solution(...)

/* XBraid specific spatial
interpolation/coarsening */
void interpolate_1D(...)
void coarsen_1D(...)
```

ex-02.c

```
XBraid
Driver
...
```



How to convert a user-code

- File: examples/ex-02*

Solves: $u_t = -u_{xx}$

ex-serial.c

Serial
Driver
...

```
/* Common functions with XBraid */

/* Initialization routine */
void get_solution(...)

/* Helpers for take_step */
void solve_tridiag(...)
void matvec_tridiag(...)
void compute_stencil(...)

/* Core time-stepping routine */
void take_step(...)

/* Output Functions */
double compute_error_norm(...)
void save_solution(...)

/* XBraid specific spatial
interpolation/coarsening */
void interpolate_1D(...)
void coarsen_1D(...)
```

App structure holds time-independent data for stepping

How to convert a user-code

- File: examples/ex-02*

Solves: $u_t = -u_{xx}$

ex-serial.c

Serial
Driver
...

```
/* Common functions with XBraid */

/* Initialization routine */
void get_solution(...)

/* Helpers for take_step */
void solve_tridiag(...)
void matvec_tridiag(...)
void compute_stencil(...)

/* Core time-stepping routine */
void take_step(...)

/* Output Functions */
double compute_error_norm(...)
void save_solution(...)

/* XBraid specific spatial
interpolation/coarsening */
void interpolate_1D(...)
void coarsen_1D(...)
```

Vector holds time-dependent data for stepping

ex-02.c

```
typedef struct _braid_App_struct
MPI_Comm    comm;
double      matrix[3];
...

typedef struct _braid_Vector_struct
int         size;
double *values;

int my_Step(u, ...)
take_step(u->values, ...);

int my_Access(u, ...)
compute_error_norm(u->values, ...);
save_solution(fname, u->values, ...);

int my_Init(u, ...)
get_solution(u->values, ...);

main()
braid_Core core; app = (my_App *) ...
braid_Init(..., core);
braid_Drive(core);
```

How to convert a user-code

- File: examples/ex-02*

Solves: $u_t = -u_{xx}$

ex-serial.c

Serial
Driver
...

```
/* Common functions with XBraid */

/* Initialization routine */
void get_solution(...)

/* Helpers for take_step */
void solve_tridiag(...)
void matvec_tridiag(...)
void compute_stencil(...)

/* Core time-stepping routine */
void take_step(...)

/* Output Functions */
double compute_error_norm(...)
void save_solution(...)

/* XBraid specific spatial
interpolation/coarsening */
void interpolate_1D(...)
void coarsen_1D(...)
```

Various wrapper functions re-use library routines

ex-02.c

```
typedef struct _braid_App_struct
MPI_Comm    comm;
double      matrix[3];
...

typedef struct _braid_Vector_struct
int         size;
double *values;

int my_Step(u, ...)
take_step(u->values, ...);

int my_Access(u, ...)
compute_error_norm(u->values, ...);
save_solution(fname, u->values, ...);

int my_Init(u, ...)
get_solution(u->values, ...);

main()
braid_Core core; app = (my_App *) ...
braid_Init(..., core);
braid_Drive(core);
```

How to convert a user-code

- File: examples/ex-02*

Solves: $u_t = -u_{xx}$

ex-serial.c

Serial
Driver
...

```
/* Common functions with XBraid */

/* Initialization routine */
void get_solution(...)

/* Helpers for take_step */
void solve_tridiag(...)
void matvec_tridiag(...)
void compute_stencil(...)

/* Core time-stepping routine */
void take_step(...)

/* Output Functions */
double compute_error_norm(...)
void save_solution(...)

/* XBraid specific spatial
interpolation/coarsening */
void interpolate_1D(...)
void coarsen_1D(...)
```

Actually running XBraid is easy!

ex-02.c

```
typedef struct _braid_App_struct
MPI_Comm    comm;
double      matrix[3];
...

typedef struct _braid_Vector_struct
int         size;
double *values;

int my_Step(u, ...)
take_step(u->values, ...);

int my_Access(u, ...)
compute_error_norm(u->values, ...);
save_solution(fname, u->values, ...);

int my_Init(u, ...)
get_solution(u->values, ...);

main()
braid_Core core; app = (my_App *) ...
braid_Init(..., core);
braid_Drive(core);
```

How to convert a user-code

- File: examples/ex-02*

Solves: $u_t = -u_{xx}$

ex-serial.c

Serial
Driver
...

```
/* Common functions with XBraid */

/* Initialization routine */
void get_solution(...)

/* Helpers for take_step */
void solve_tridiag(...)
void matvec_tridiag(...)
void compute_stencil(...)

/* Core time-stepping routine */
void take_step(...)

/* Output Functions */
double compute_error_norm(...)
void save_solution(...)

/* XBraid specific spatial
interpolation/coarsening */
void interpolate_1D(...)
void coarsen_1D(...)
```

These final functions allow for spatial coarsening

ex-02.c

```
typedef struct _braid_App_struct
MPI_Comm    comm;
double      matrix[3];
...

typedef struct _braid_Vector_struct
int         size;
double *values;

int my_Step(u, ...)
take_step(u->values, ...);

int my_Access(u, ...)
compute_error_norm(u->values, ...);
save_solution(fname, u->values, ...);

int my_Init(u, ...)
get_solution(u->values, ...);

main()
braid_Core core; app = (my_App *) ...
braid_Init(..., core);
braid_Drive(core);
```

How to convert a user-code

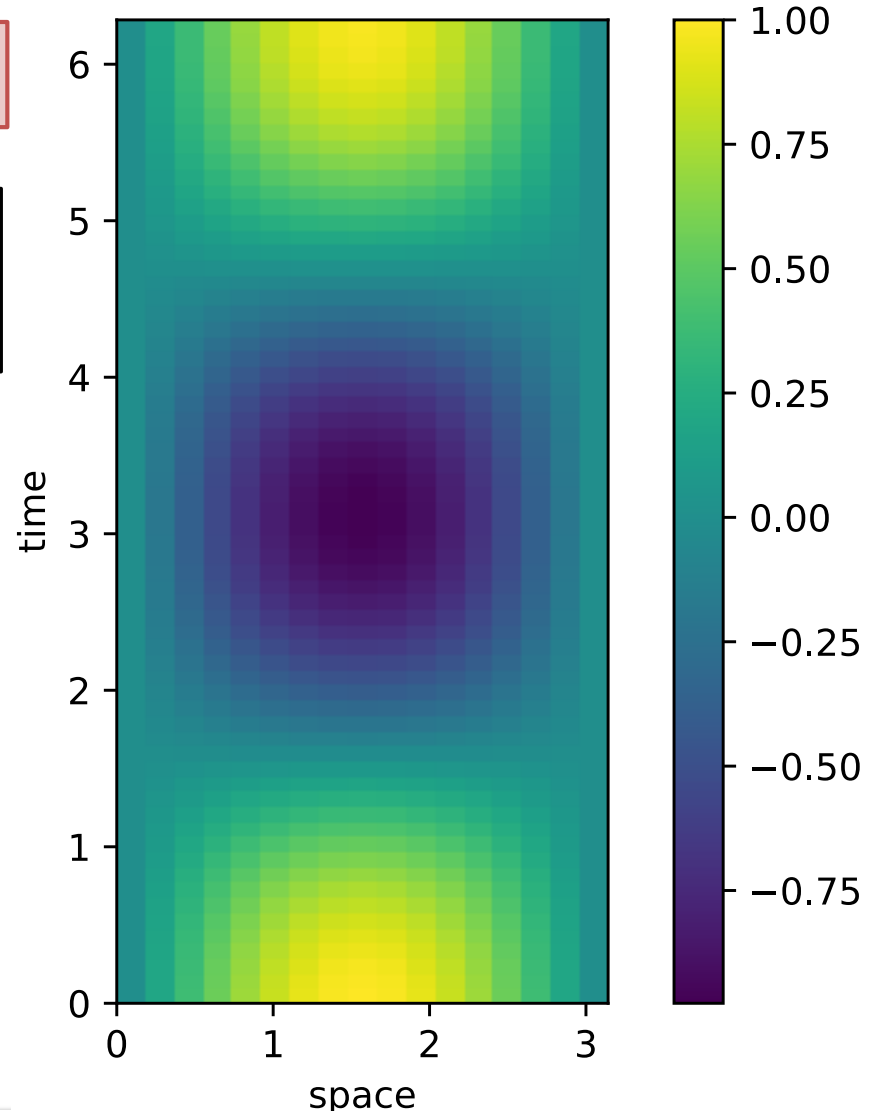
- File: examples/ex-02*

Solves: $u_t = -u_{xx}$

```
$ ./ex-02 -ntime 64 -nspac 17  
$ python viz-ex-02.py
```

The solution for ex-02 is

$$u(t, x) = \sin(x) \cos(t)$$



How to debug your new code

- File: examples/ex-02.c

Solves: $u_t = -u_{xx}$

There is a test function for each wrapper, e.g., `braid_TestInit()`

```
$ ./ex-02 -wrapper_tests  
...  
Finished braid_TestAll: no fails detected
```

Set `max-levels=1`. The answer should exactly match sequential time stepping.

```
$ ./ex-02 -ntime 64 -nspaced 17 -ml 1  
$ python viz-ex-02.py  
(In reality, you'd want to check the agreement to 15 or 16 decimals)
```

Continue with `max-levels=1`, but switch to multiple processors in time. Check that the answer again exactly matches sequential time stepping.

```
$ mpirun -np 2 ex-02 -ntime 64 -nspaced 17 -ml 1  
$ python viz-ex-02.py  
(In reality, you'd want to check the agreement to 15 or 16 decimals)
```



How to debug your new code

- File: examples/ex-02.c

Solves: $u_t = -u_{xx}$

Check that XBraid is a fixed point method

Set `max-levels=2`, `tol=0.0`, `max-iter=3`, and initialize XBraid with the sequential solution

```
$ ./ex-02 -ntime 64 -nspaced 17 -ml 2 -tol 0.0 -mi 3 -use_seq
Braid: || r_0 || = 0.000000e+00, conv factor = 1.00e+00, wall time = ...
Braid: || r_1 || = 0.000000e+00, conv factor = nan, wall time = ...
Braid: || r_2 || = 0.000000e+00, conv factor = nan, wall time = ...
Braid: || r_3 || = 0.000000e+00, conv factor = nan, wall time = ...
Braid: || r_4 || = 0.000000e+00, conv factor = nan, wall time = ...
```



How to debug your new code

- File: examples/ex-02.c Solves: $u_t = -u_{xx}$

Turn on debug-level printing and check that the exact solution is propagating

With FCF-relaxation, the exact solution propagates forward 2 C-points each iter

```
$ ./ex-02 -ntime 8 -nspace 17 -mi 3 -print_level 2
Braid: time step:      0, rnorm: 0.00e+00
Braid: time step:      2, rnorm: 0.00e+00
Braid: time step:      4, rnorm: 6.86e-01
Braid: time step:      6, rnorm: 1.10e+00
Braid: time step:      8, rnorm: 2.04e-02
Braid: || r_0 || = 1.292837e+00, conv factor = 1.00e+00, wall time = ...
Braid: time step:      0, rnorm: 0.00e+00
Braid: time step:      2, rnorm: 0.00e+00
Braid: time step:      4, rnorm: 0.00e+00
Braid: time step:      6, rnorm: 0.00e+00
Braid: time step:      8, rnorm: 1.62e-02
...
...
```

Then, run some larger, multilevel tests of XBraid, checking that the sequential and time-parallel versions agree to within the halting tolerance



Intrusiveness versus efficiency

- The more intrusive XBraid is allowed to be, the more efficient it is
 - **Residual option:** computing the residual with a naive implementation of XBraid is as expensive in FLOPs as sequential time stepping. Writing this extra function allows you to avoid this for implicit schemes.
 - This function also allows relaxation to be significantly less expensive
 - **Adaptivity:** constructing the correct adaptive space-time grid is active research
 - For instance a development branch is currently using threshold refinement across the temporal communicator to choose time intervals to refine
 - **Storage:** requires a little extra coding, i.e., a new initial guess for implicit scheme
 - **Level-dependent time-stepper:** how to change `Step()` on coarse-levels is problem dependent, but almost always yields big benefits, e.g., vary the tolerance for taking a (non)linear implicit step
 - **Spatial coarsening:** this can affect convergence, but is required for an $O(N)$ method in both time and space
 - **Stephanie Friedhoff's talk** covers this in more detail, e.g., results from taking a naive XBraid implementation and moving to an STMG (space-time MG) method

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Appendix: Advanced XBraid features

- *Temporal adaptivity*
- *Shell-vectors and BDF-k*
- *Fortran90 Interface*
- *Residual and storage options*
- *Spatial coarsening*



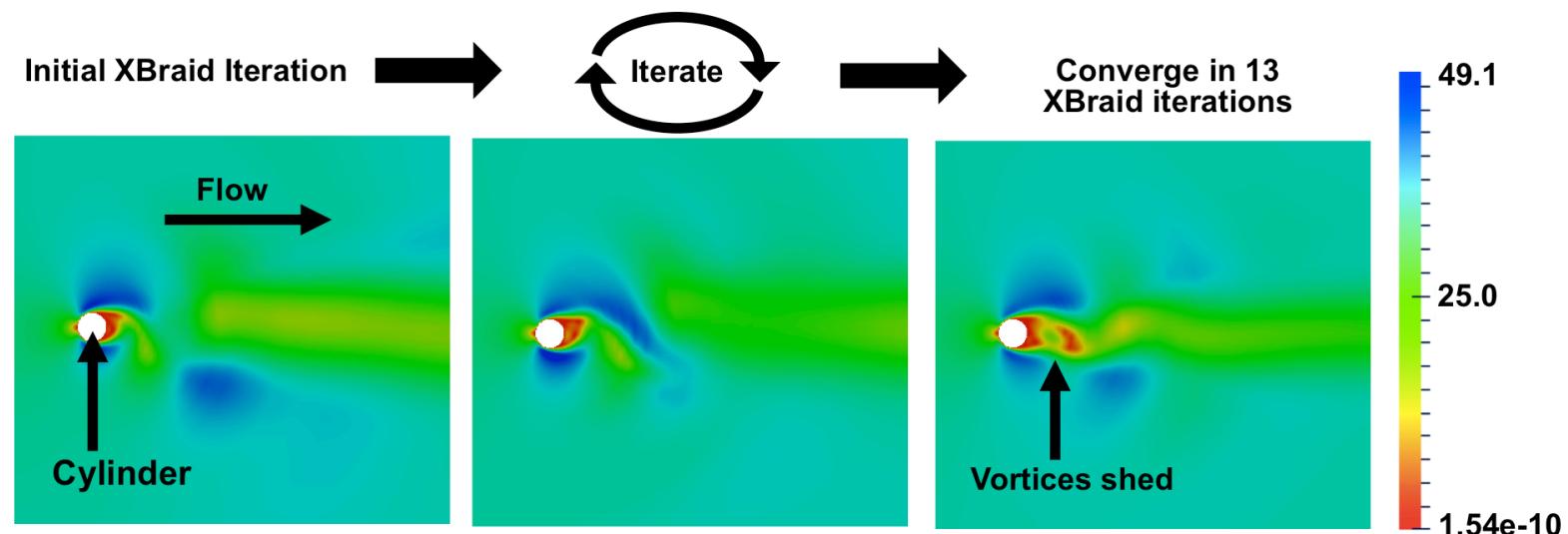
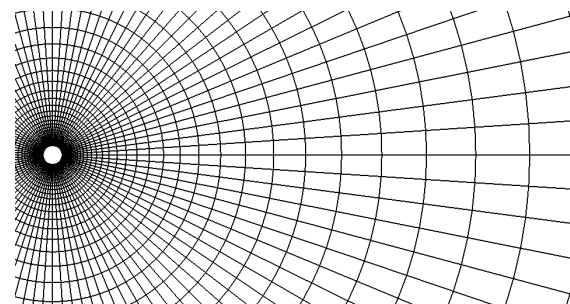
Experiments coupling our code XBraid with various application research codes

- Navier-Stokes (compressible and incompressible)
 - Strand2D, CarT3D, LifeV (Trilinos-based)
- Heat equation (including moving mesh example)
 - MFEM, hypre
- Nonlinear diffusion, the p -Laplacian
 - MFEM
- Power-grid simulations
 - GridDyn
- Explicit time-stepping coupled with space-time coarsening
 - Heat equation
 - Advection plus artificial dissipation
 - MFEM, hypre



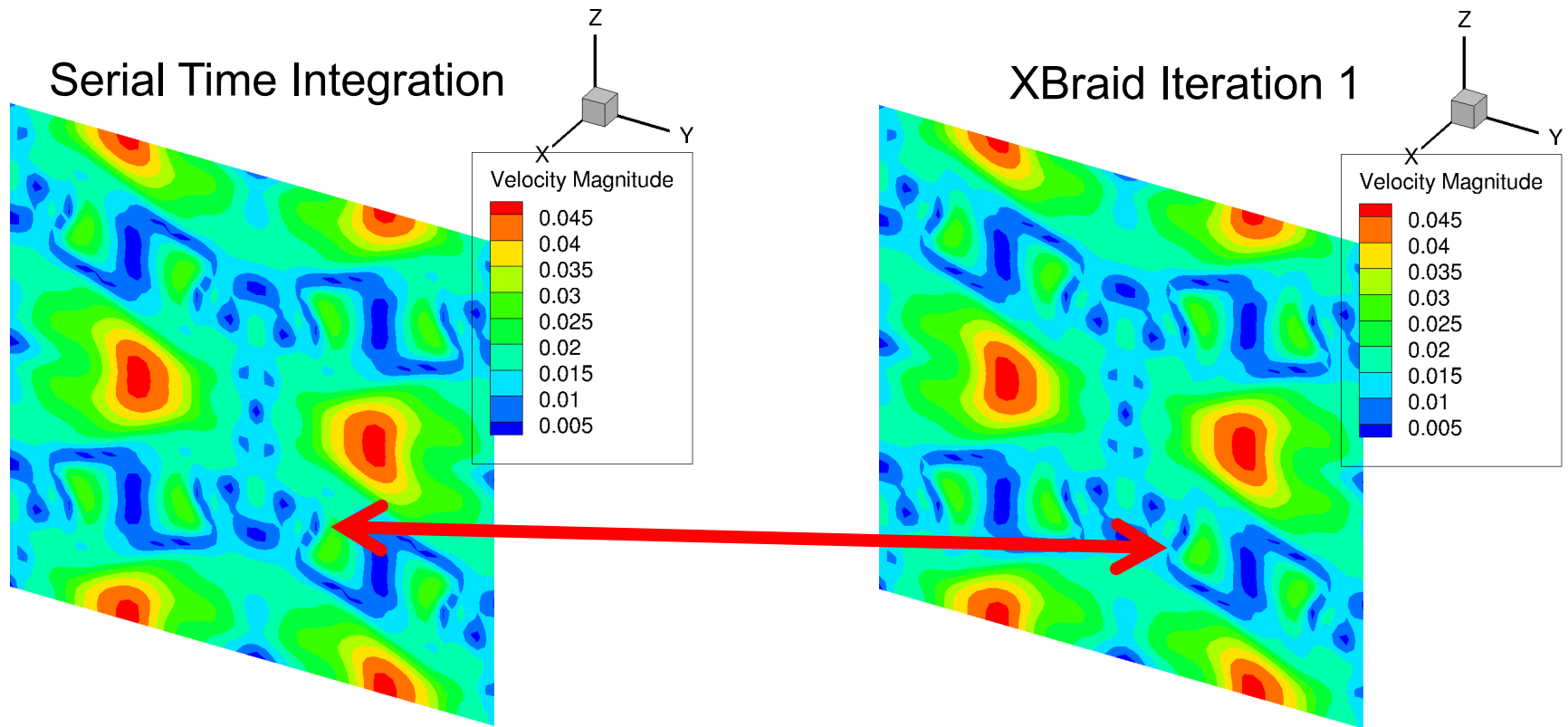
Compressible Navier-Stokes (nonlinear) – speedups to 7.5x with typical MG scaling

- Coupled XBraid with existing code Strand2D (DoD project)
 - ~500 lines of XBraid wrapper code plus minor changes to Strand2D
 - ~3 weeks with minimal outside help
- Plots of velocity magnitude at time step 5120



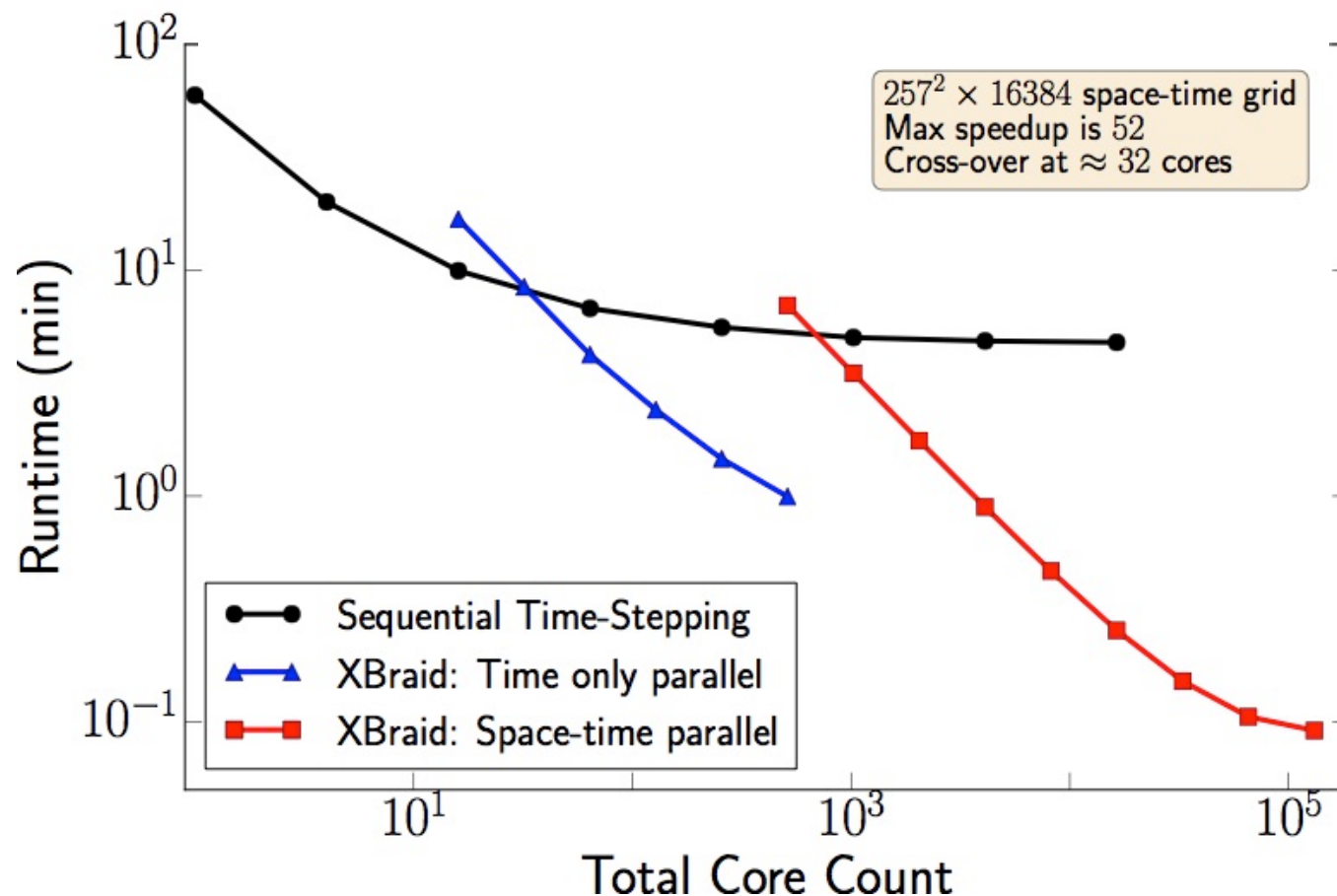
Compressible Navier-Stokes with Cart3D – convergence is very fast, ~5 iterations

- Taylor-Green problem: turbulent decay of vortex, $Re=1600$
 - Higher-order spatial discretization on $58^3 \times 20,000$ Cartesian grid
- Plot velocity magnitude at $x=0$ cross-section



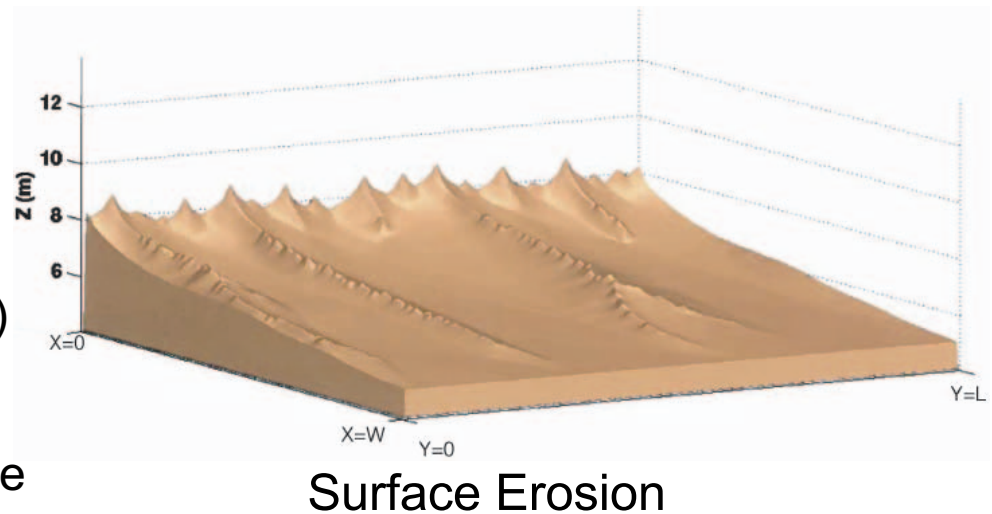
Strong scaling for heat equation

- XBraid uses V-cycles and FCF-relaxation
- Excellent strong scaling, until parallelism is exhausted



The p -Laplacian: nonlinear diffusion

- Solve $u_t = \nabla \cdot (|\nabla u|^{p-2} \nabla u)$
- 2D linear finite elements
 - 16K x 20K space-time problem
 - Backward Euler (Newton's method)
- Current results
 - Crossover at ~40 processors in time
 - Speedup of 18x at 130K cores
- Important parameters for performance
 - Full storage and space-time coarsening
 - Adjusting the Newton tolerance for the early iterations



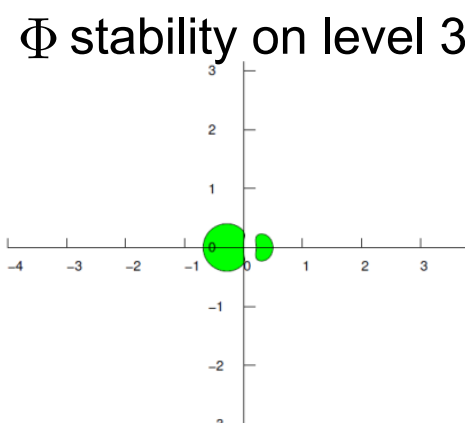
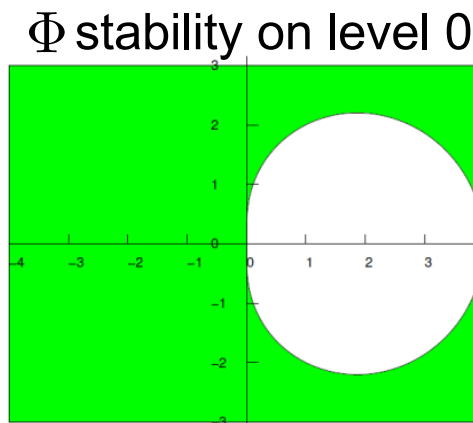
+++ Image courtesy of Birnir, Rowlett. "Mathematical Models for Erosion and the optimal Transportation of Sediment. Int. J. Nonlinear Sci. Numer. Simul. 2013

Initial speedups for power-grid

- Simulate 4 generators (30 unknowns) for 30s with 30K time steps

	Sequential	128 cores	256 cores	512 cores	1024 cores
Implicit RK4	227s	144s	113s	102s	105s
BDF-4	12.4s	13.6s	9.46s	7.73s	7.30s

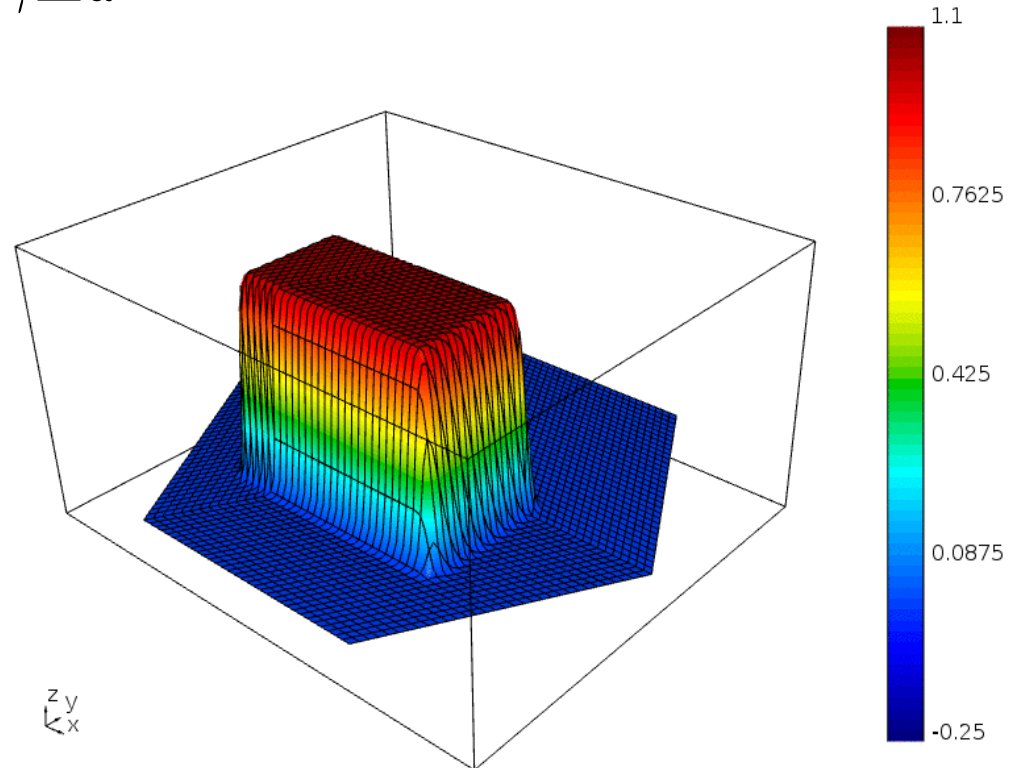
- XBraid is designed for one-step methods, so we make BDF- k “one-step” by grouping k time-steps together
 - Creates non-uniform time-step sizes on coarse grids and stability issues for Φ



Solution: reduce the BDF order on coarse levels

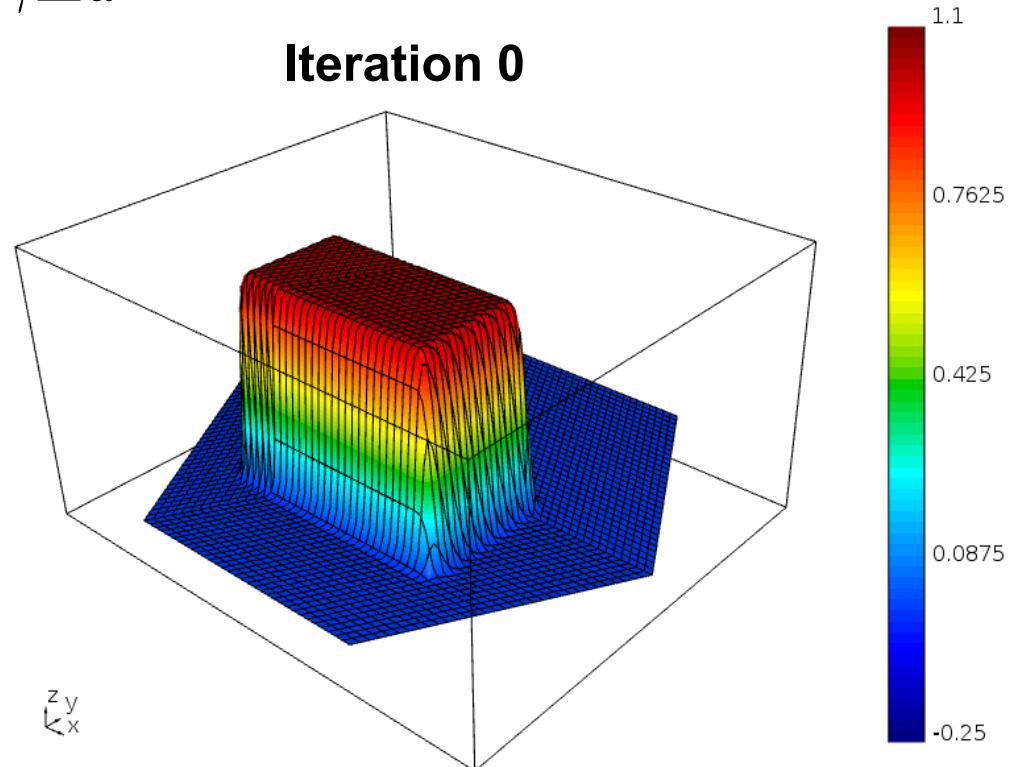
Explicit methods with MFEM

- 2D advection $u_t = \mathbf{b}(\mathbf{x}) \cdot \nabla u + \gamma \Delta u$
 - Stability determined by convection (convection dominated)
 - Diffusion term 0.001
- **Sequential Time Stepping**
 - Sharp profile is transported over 1100 time steps
 - 3rd order explicit method



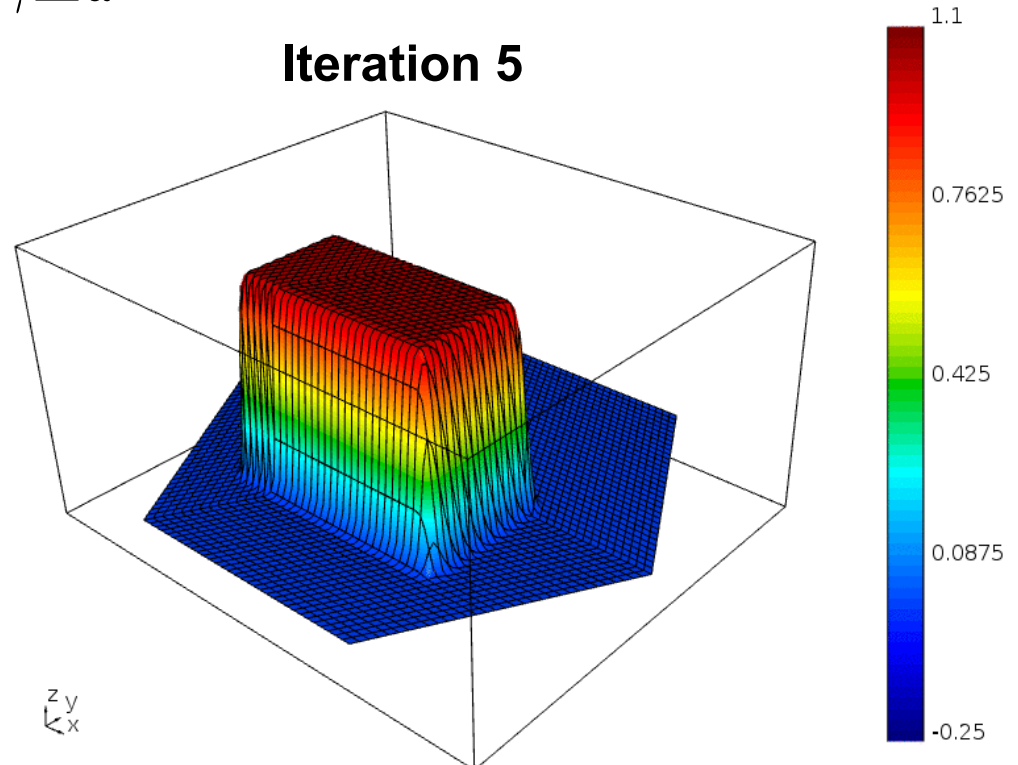
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 - Diffusion term 0.001
- **Parallel-in-time solution**
 - Sharp profile is transported over 1100 time steps
 - 3rd order explicit method
 - 3-level XBraid hierarchy



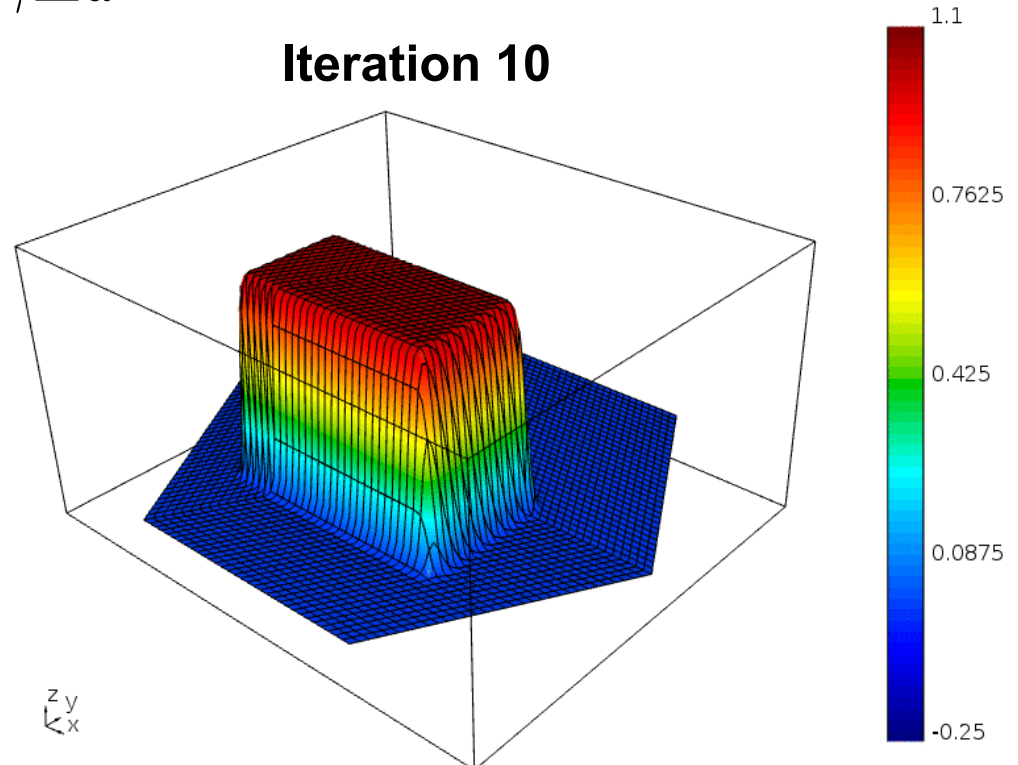
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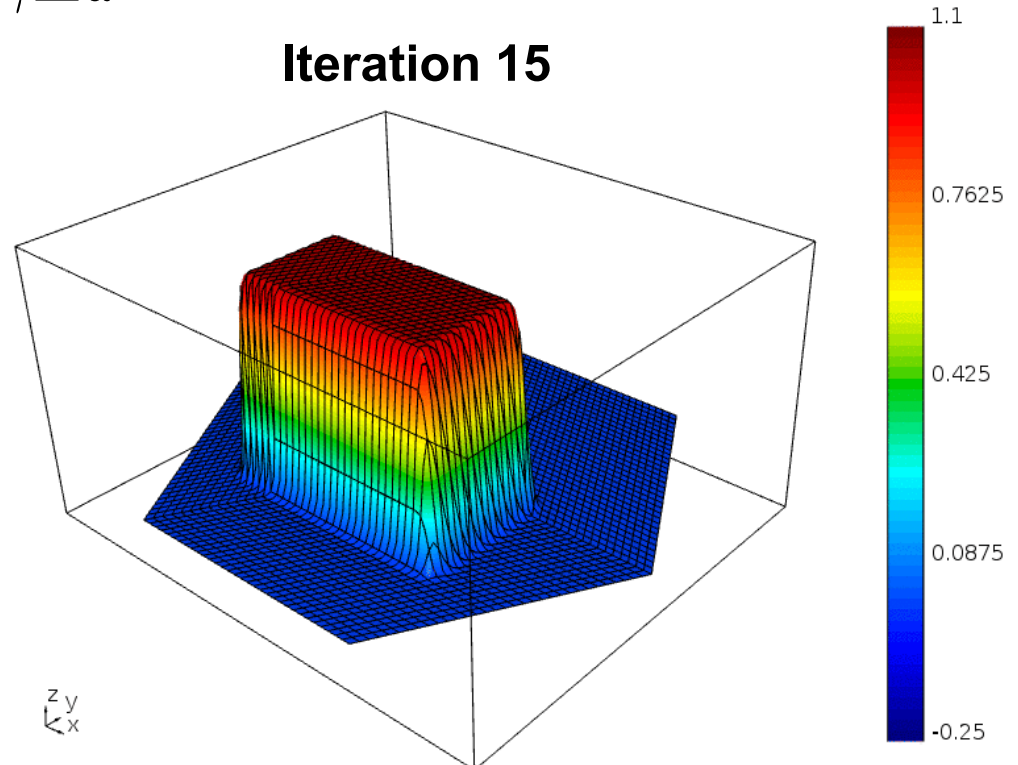
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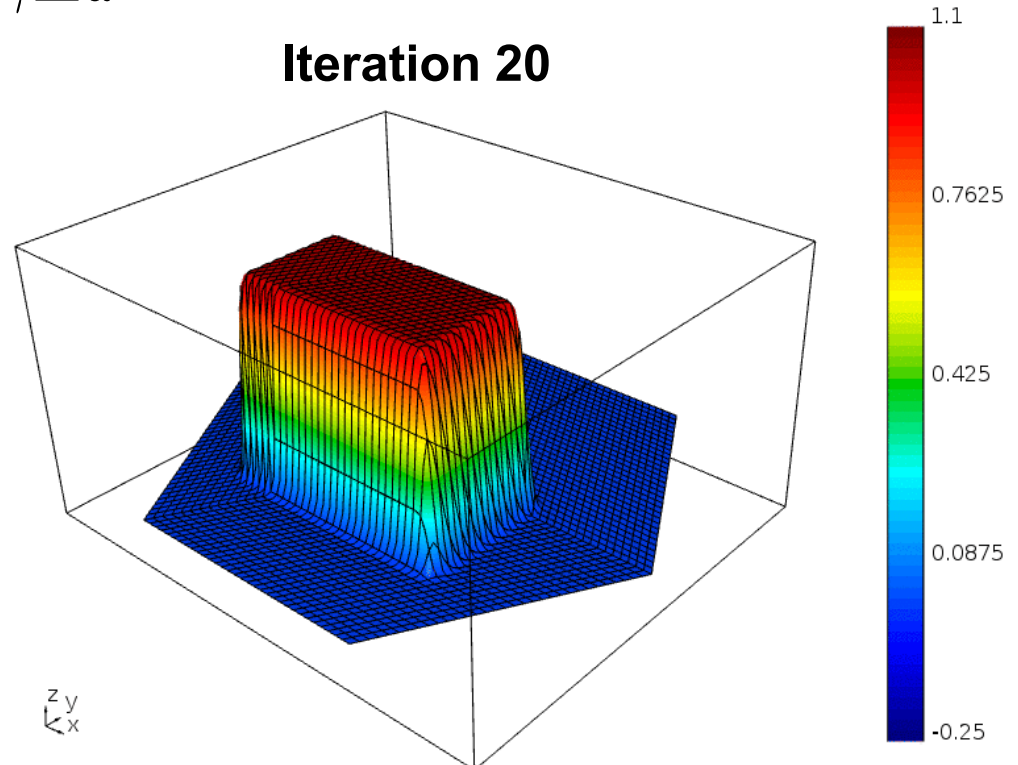
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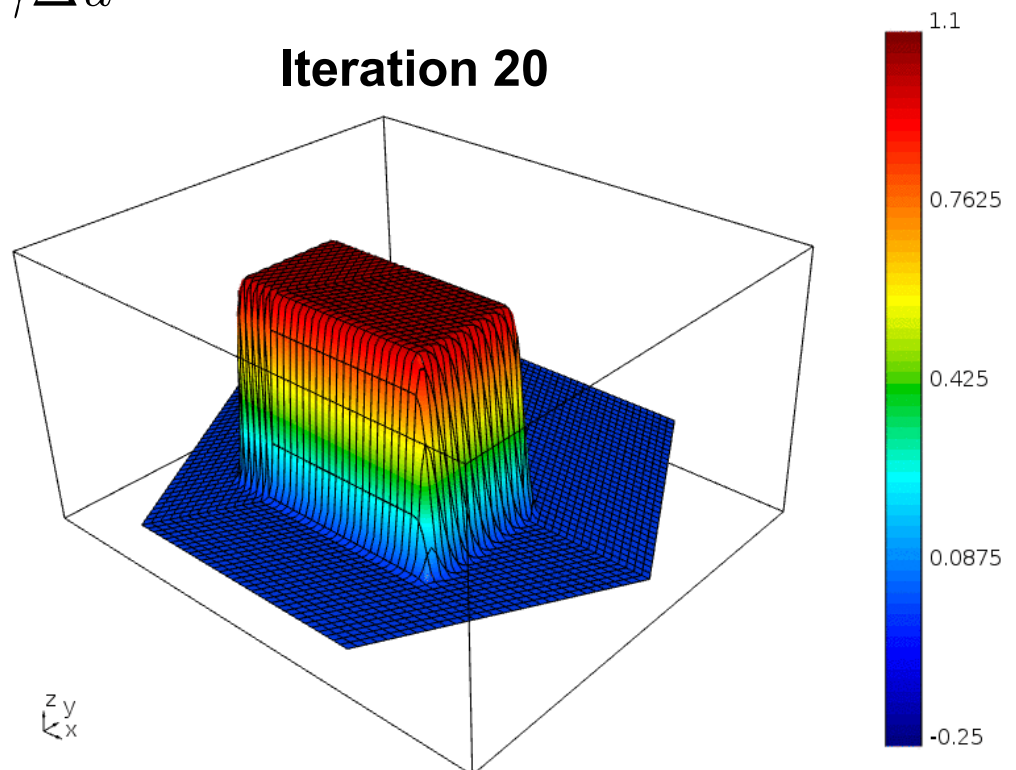
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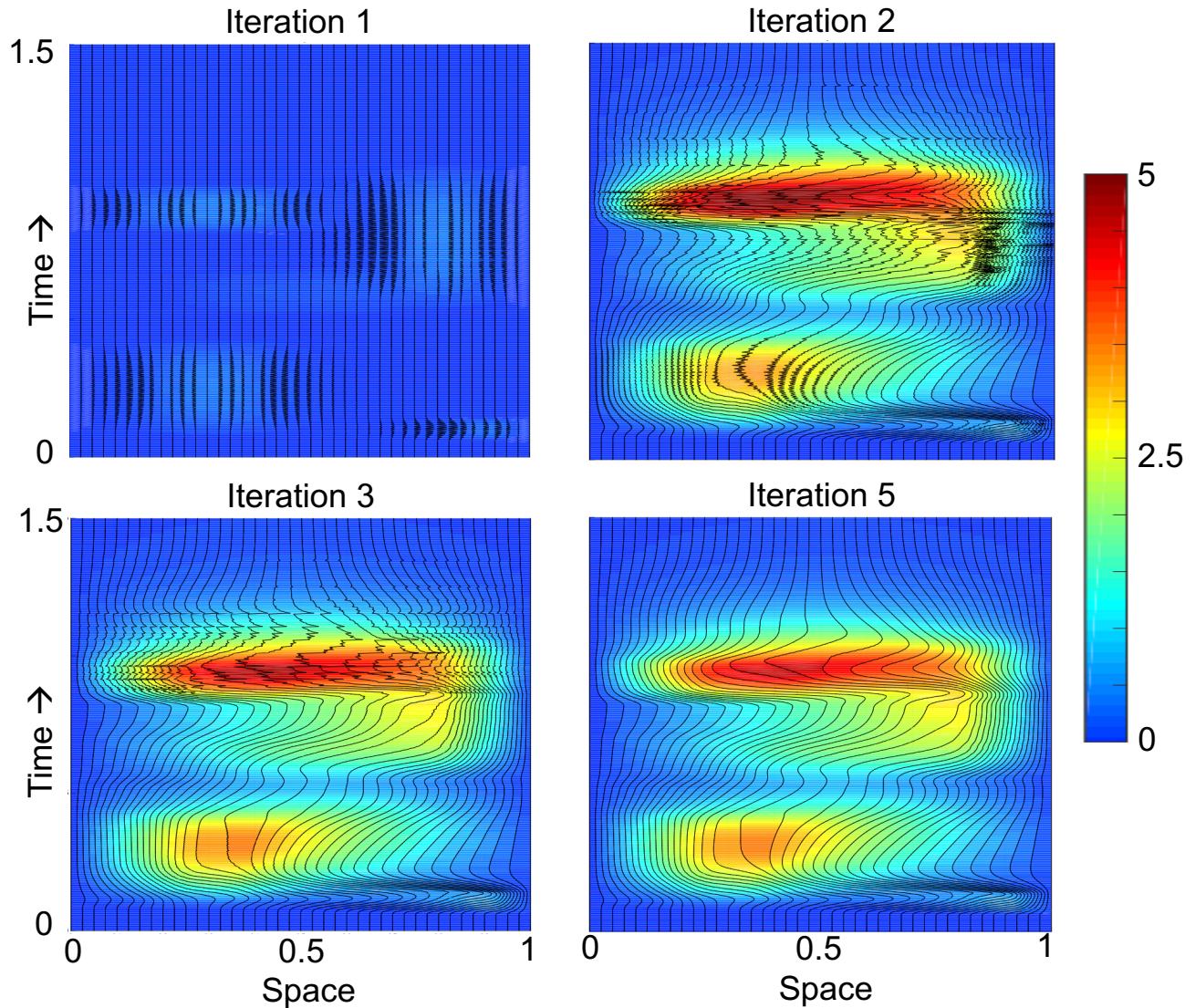
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- **Parallel-in-time solution**
 - Sharp profile is transported over 1100 time steps
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- Future Work: Improve convergence (relaxation, coarse-grid equations)



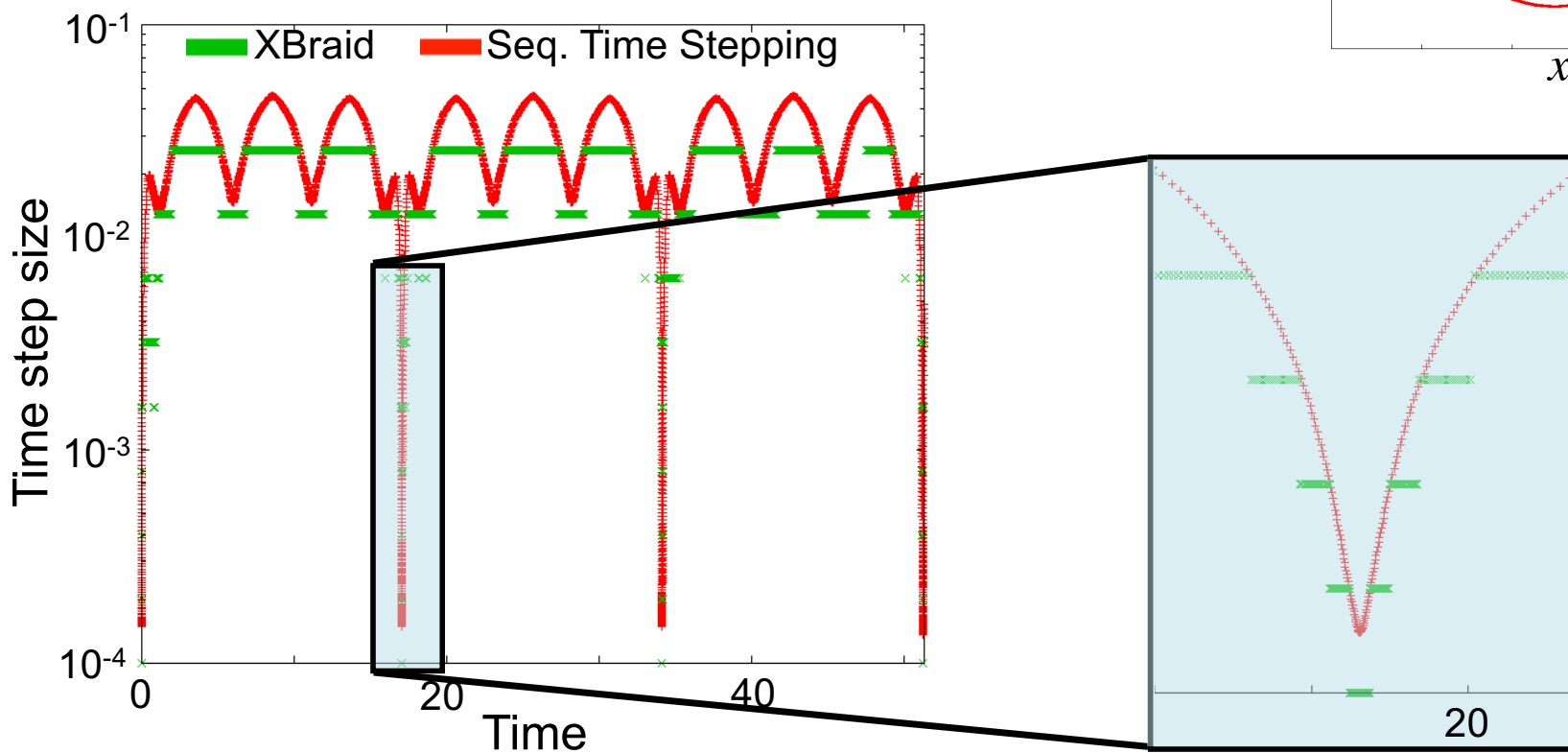
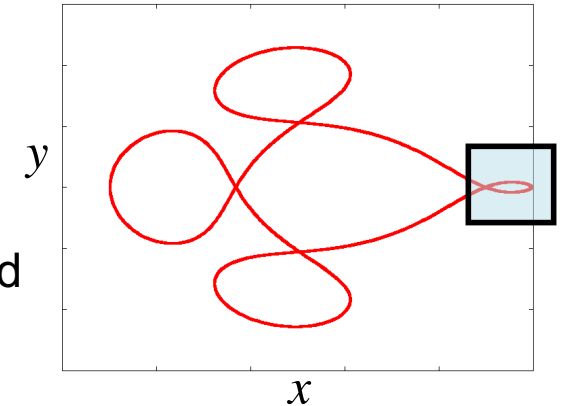
Moving mesh

- 1D space moving mesh proof-of-concept
- Mesh points move towards regions with a rapidly changing solution
- Fast convergence and scalable iteration counts
- More complicated moving mesh problems coming...



Temporal adaptivity proof-of-concept

- Classic ODE modeling satellite orbit around earth and moon (2 variables in space, x and y)
- One region of orbit requires very fine time steps
 - Carry out 4 periods of orbit, refining step size as needed



Nearly 50 years of research exists, but has only scratched the surface

- Earliest work goes back to 1964 by Nievergelt
 - Led to multiple shooting methods, Keller (1968)
- Space-time multigrid methods for parabolic problems
 - Hackbusch (1984); Horton (1992); Horton and Vandewalle (1995)
 - The latter is one of the first optimal & fully parallelizable methods to date
- Parareal was introduced by Lions, Maday, and Turincini in 2001
 - Probably the most widely studied method
 - Gander and Vandewalle (2007) show that parareal is two-level FAS multigrid
- Discretization specific work includes
 - Minion, Williams (2008, 2010) – PFASST, spectral deferred correction / parareal
 - DeSterck, Manteuffel, McCormick, Olson (2004, 2006) – FOSLS
- Research on these methods is ramping up!
 - Ruprecht, Krause, Speck, Emmett, Langer, ... this is not an exhaustive list



Summary and conclusions

- Sequential time integration bottleneck is real
 - Parallel in time is needed for future architectures
 - This is a major paradigm shift
- XBraid applies multigrid reduction to the time dimension
 - Multigrid is ideal for exascale (optimal, resilient, ...)
 - Result is a flexible and non-intrusive approach
- The more intrusive XBraid is allowed to be, the more efficient the algorithm is.
- There is much future work to be done!
 - More problem types, more complicated discretizations, performance improvements, adaptive meshing, ...



Thank You! Any Questions?

A good read, *Parallel Time Integration with Multigrid*, SIAM J. Sci. Comp.

Open Source XBraid Code

- <http://llnl.gov/casc/xbraid>
- Supports C, C++, F90



Our Team



Veselin
Dobrev



Rob
Falgout



Matthieu
Lecouvez



Tzanio
Kolev



Jacob
Schroder



Scott
MacLachlan



Hans
De Sterck



Stephanie
Friedhoff

Collaborators

CU Boulder ([Manteuffel](#), [McCormick](#), [O'Neill](#)), U Wuppertal ([Friedhoff](#)), U Stuttgart ([Roehrle](#), [Hessenthaler](#)), Kaiserslautern ([Guenther](#)), LLNL ([Woodward](#), [Top](#))

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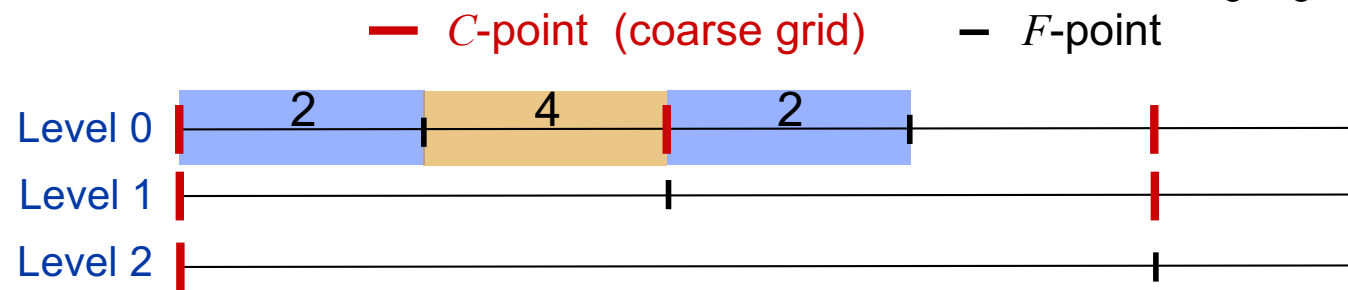
Appendix: Advanced XBraid features

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- *Shell-vectors and BDF-k*
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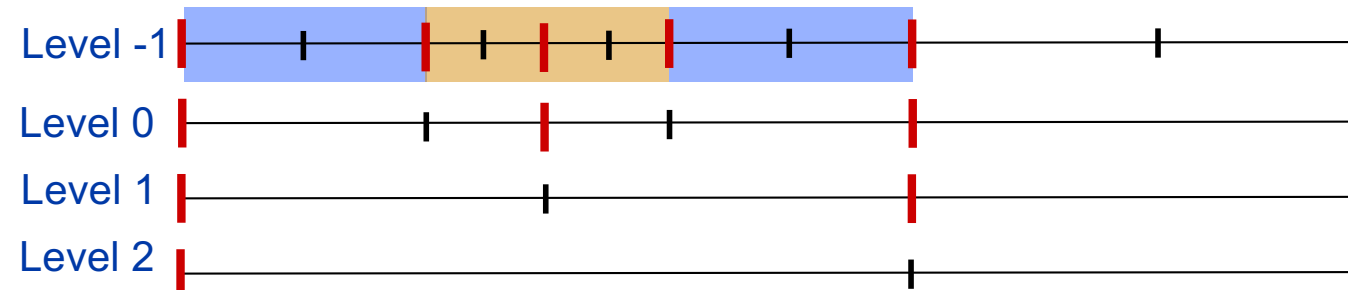


Advanced feature: FMG allows for adaptivity in time and space

- User returns refinement factor in Step ()
- Example time grid hierarchy



- User requests refinement factors on the finest grid which generates a new grid and hierarchy



Advanced feature: adaptivity in time

- File: examples/ex-01-refinement.c Solves: $u_t = \lambda u$
 - Do adaptive temporal refinement with a local truncation error estimate
 - Call `braid_StepStatusSetRFactor(status, k)` to refine a interval `k` times
 - Called from inside of `Step()`

```
$ make ex-01-refinement
$ ./ex-01-refinement -nt 32 -mi 4 -refine 2 -max_rfactor 4

Braid: || r_0 || not available, wall time = ...
Braid: || r_1 || = 1.282307e-01, conv factor = 1.00e+00, wall time =...
Braid: Temporal refinement occurred, 128 time steps

Braid: || r_1 || = 1.361139e-02, conv factor = 1.00e+00, wall time =...
Braid: Temporal refinement occurred, 512 time steps
```



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Braid: Temporal refinement occurred, 4096 time steps
```



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Braid: || r_2 || = 7.711806e-06, conv factor = 1.40e-01, wall time =...
Braid: || r_3 || = 6.265552e-07, conv factor = 8.12e-02, wall time =...
```

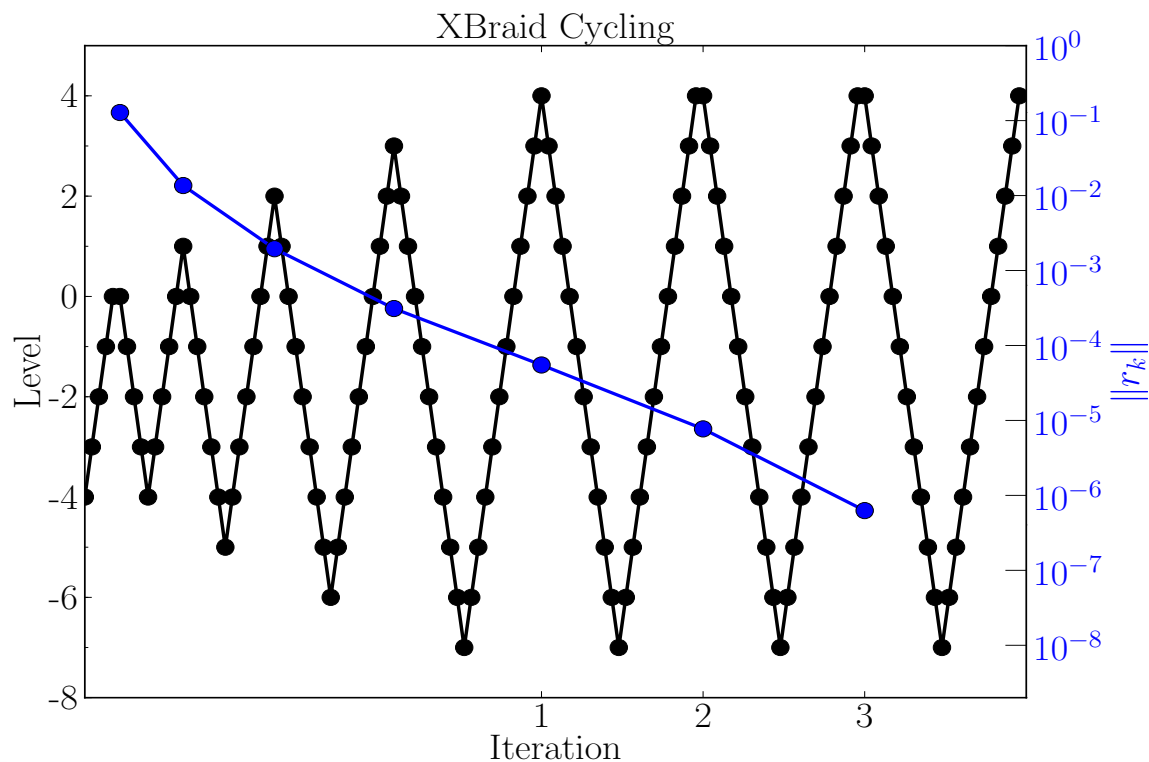


Advanced feature: adaptivity in time

- File: examples/ex-01-refinement.c Solves: $u_t = \lambda u$

- Now, visualize the cycling
- Observe how the new levels (and time-points) are added
- This causes an uneven reduction in the residual

```
$ python ../user_utils/cycleplot.py
```



Refinement here is with a V-cycle. But can also be done with FMG cycles.

Advanced feature: residual function

- File: examples/ex-01-expanded.c Solves: $u_t = \lambda u$

Observe how turning on the residual function changes convergence

```
$ ./ex-01-expanded -ntime 128 -res
...
iterations          = 7

$ ./ex-01-expanded -ntime 128
...
iterations          = 6
```

- File: examples/ex-03.c Solves: $u_t = -u_{xx} - u_{yy}$

```
$ make ex-03
$ ./ex-03 -nt 128 -nx 9 9 -mi 3 -res
  Braid: || r_1 || = 5.231464e-01, conv factor = 1.00e+00, wall time = ...
  Braid: || r_2 || = 6.067546e-02, conv factor = 1.16e-01, wall time = ...

$ ./ex-03 -nt 128 -nx 9 9 -mi 3
  Braid: || r_1 || = 5.002967e-01, conv factor = 1.00e+00, wall time = ...
  Braid: || r_2 || = 2.701758e-02, conv factor = 5.40e-02, wall time = ...
```

Understanding the residual feature

- XBraid computes the FAS residual in a block-row fashion for the space-time system

$$A_i(\mathbf{u}_i, \mathbf{u}_{i-1}) = f_i$$

- Consider, for example, the common additive form of a user residual:

User option to define $\longrightarrow A_i(\mathbf{u}_i, \mathbf{u}_{i-1}) = -\Phi(\mathbf{u}_{i-1}) + \Psi(\mathbf{u}_i)$

FAS residual computed internally $\longrightarrow r_i = f_i + \Phi(\mathbf{u}_{i-1}) - \Psi(\mathbf{u}_i)$

- Default setting:** Step () = $\Phi(\mathbf{u}_i)$ and $\Psi = I$

- XBraid can compute the rest of the residual on its own

- Residual setting:** user defines a new function $\text{Residual}(\mathbf{u}_i, \mathbf{u}_{i-1}) = A_i(\mathbf{u}_i, \mathbf{u}_{i-1})$

- This function defines the equation to be solved, implying that Step () must be compatible.
- Step () must now compute $u_i = \Psi^{-1}(f_i + \Phi(u_{i-1}))$
- Notice how Step () must now account for f_i , that is, fstop in Step () is no longer NULL!

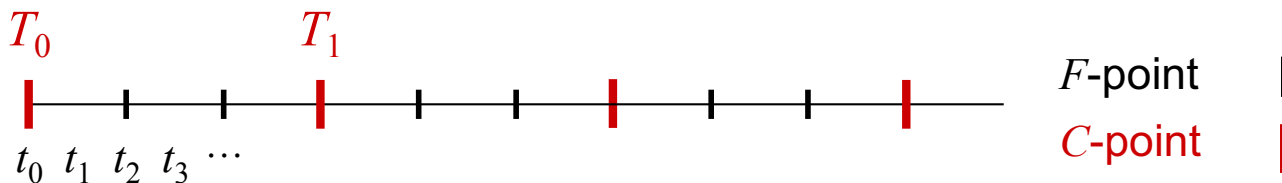
- Computational savings:** consider the heat equation and backward Euler

- Default:** Step () implements Φ , a full implicit solve for an accurate residual

- Residual:** Step () implements a very weak inexact solve (only used for relaxation)
Residual () uses $\Phi = I$ and Ψ is just a sparse matrix (very cheap!)

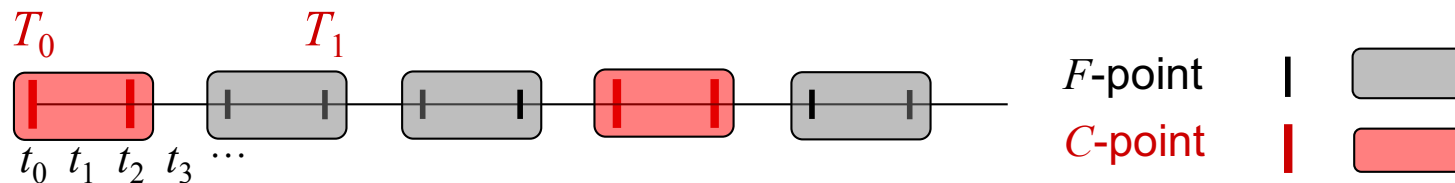
Advanced feature: shell-vectors & BDF-k

- File: examples/ex-01-expanded-bdf2.c Solves: $u_t = \lambda u$
- XBraid is designed for one-step methods. This is the standard way to partition the time-line.



Advanced feature: shell-vectors & BDF-k

- File: examples/ex-01-expanded-bdf2.c Solves: $u_t = \lambda u$
- XBraid is designed for one-step methods. The new way to partition so that BDF- k looks “one-step” is to group k time-steps together (here, $k = 2$).



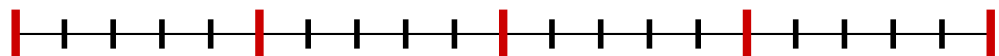
- Creates non-uniform time-step sizes on coarse grids
- The shell-vector feature allows for the storage of meta-data at every time point, including F-points that are otherwise not stored.
 - This meta-data allows for tracking the irregular time-grid spacing
- Other BDF- k strategies, like reducing order on coarse-grids, are possible
- To use the shell option, you must define new shell functions for allocating, copying, and freeing vector shells

Advanced feature: extra storage

- File: examples/ex-03.c

Solves: $u_t = -u_{xx} - u_{yy}$

- Set a storage value k (default is -1)
 - For $level \geq k \geq 0$, store all points; for $level < k$, store only C-points
 - $k = 0$ storage at all points on all levels
 - $k = -1$ special value, storage only at C-points on all levels



- F-point (fine grid only)
- C-point (coarse & fine grid)

- The extra storage critically gives improved initial guesses to implicit solvers
- The extra storage changes the problem being solved
 - The operator Φ changes as the initial guess changes
- Look at the residual histories with

```
$ make ex-03
$ ./ex-03 -nx 17 17 -nt 128 -storage -1
$ ./ex-03 -nx 17 17 -nt 128 -storage 0
$ ./ex-03 -nx 17 17 -nt 128 -storage 1
```

$$\begin{pmatrix} I & & & \\ -\Phi & I & & \\ & \ddots & \ddots & \\ & & -\Phi & I \end{pmatrix} \begin{pmatrix} \mathbf{u}_0 \\ \mathbf{u}_1 \\ \vdots \\ \mathbf{u}_N \end{pmatrix} = \begin{pmatrix} \mathbf{g}_0 \\ \mathbf{g}_1 \\ \vdots \\ \mathbf{g}_N \end{pmatrix}$$

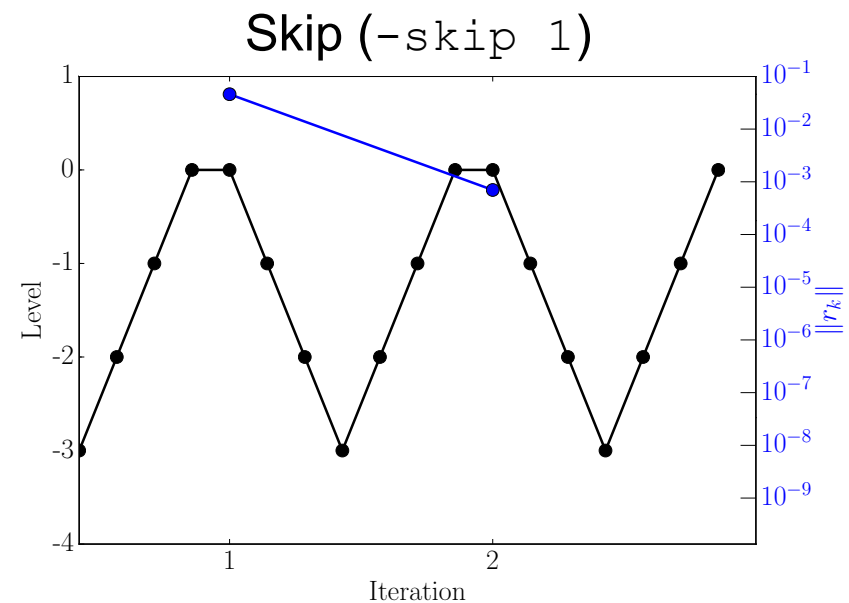
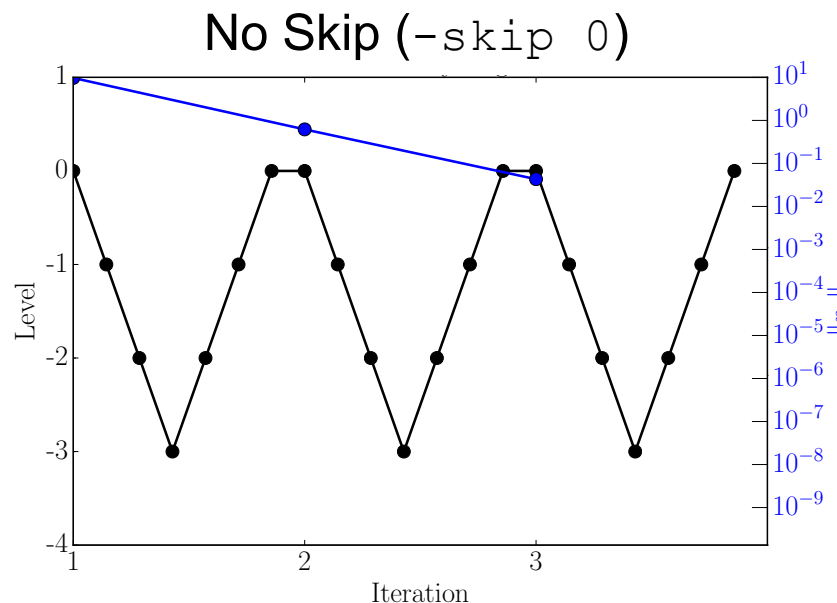
Advanced feature: skip option

- File: examples/ex-03.c

Solves: $u_t = -u_{xx} - u_{yy}$

- Skip allows XBraid to skip (typically useless) relaxations on the 1st down cycle
 - By default, skip is turned on**
- Compare the residual histories for

```
$ ./ex-03 -nx 17 17 -nt 128 -mi 4 -skip 0  
$ ./ex-03 -nx 17 17 -nt 128 -mi 4 -skip 1
```



Advanced feature: parallel-run

- File: examples/ex-03.c

Solves: $u_t = -u_{xx} - u_{yy}$

Run in parallel!

```
$ ./mpirun -np 8 ex-03 -pgrid 2 2 2 -nt 256 -nx 17 17
Braid: || r_0 || not available, wall time = ...
Braid: || r_1 || = 6.166798e-01, conv factor = 1.00e+00, wall time = ...
Braid: || r_2 || = 2.319985e-02, conv factor = 3.76e-02, wall time = ...
Braid: || r_3 || = 6.972052e-04, conv factor = 3.01e-02, wall time = ...
Braid: || r_4 || = 1.135286e-05, conv factor = 1.63e-02, wall time = ...
...
```



Advanced feature: spatial coarsening

- File: examples/ex-02.c

Solves: $u_t = -u_{xx}$

Here, we use simple bilinear interpolation (and its transpose) for spatial coarsening

```
$ ./ex-02 -ntime 64 -nspatial 17 -ml 3 -sc
Braid: || r_0 || = 2.935397e+00, conv factor = 1.00e+00, wall time = ...
Braid: || r_1 || = 1.483600e-01, conv factor = 5.05e-02, wall time = ...
Braid: || r_2 || = 3.884625e-03, conv factor = 2.62e-02, wall time = ...
Braid: || r_3 || = 1.315185e-04, conv factor = 3.39e-02, wall time = ...
...
level      dx          dt        dt/dx^2
-----
0   |  1.96e-01    9.82e-02    2.55e+00
1   |  3.93e-01    1.96e-01    1.27e+00
2   |  7.85e-01    3.93e-01    6.37e-01
```

Spatial coarsening is active research and can (sometimes) negatively impact convergence.

```
$ ./ex-02 -ntime 64 -nspatial 17 -ml 3
Braid: || r_0 || = 2.935397e+00, conv factor = 1.00e+00, wall time = ...
Braid: || r_1 || = 1.666814e-01, conv factor = 5.68e-02, wall time = ...
Braid: || r_2 || = 8.328760e-03, conv factor = 5.00e-02, wall time = ...
Braid: || r_3 || = 2.844685e-04, conv factor = 3.42e-02, wall time = ...
...
level      dx          dt        dt/dx^2
-----
0   |  1.96e-01    9.82e-02    2.55e+00
1   |  1.96e-01    1.96e-01    5.09e+00
2   |  1.96e-01    3.93e-01    1.02e+01
```

Advanced feature: coarsening factor

- File: examples/ex-02.c Solves: $u_t = -u_{xx}$

- Changing the coarsening factor does not change convergence (much)
- This powerful fact applies to parabolic problems in general
 - Allows for a great deal of performance tuning
 - Requires that FCF-relaxation or F-cycles be used

```
$ ./ex-02 -ntime 1024 -nspatial 128 -cf 16 -ml 10
```

```
...
```

```
iterations                  = 7
```

```
$ ./ex-02 -ntime 1024 -nspatial 128 -cf 2 -ml 10
```

```
...
```

```
iterations                  = 8
```

Fortran90 interface

- File: examples/ex-01-expanded-f.f90 Solves: $u_t = \lambda u$

Uses Fortran90 modules to define the App and Vector Types

```
module braid_types

  type my_vector
    double precision val
  end type my_vector
  ...

```

User-defined wrapper functions are the same, only written in Fortran90

```
subroutine braid_Sum_F90(app, alpha, x, beta, y)
  ! Braid types
  use braid_types
  implicit none
  type(my_vector) :: x, y
  type(my_app) :: app

  double precision alpha, beta
  y%val = alpha*(x%val) + beta*(y%val)
end subroutine braid_Sum_F90
```



