

Equation-Free function toolbox for Matlab:

Full Developers Manual

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Abstract

This ‘equation-free toolbox’ empowers the computer-assisted analysis of complex, multiscale systems. Its aim is to enable you to use microscopic simulators to perform system level tasks and analysis. The methodology bypasses the derivation of macroscopic evolution equations by using only short bursts of microscale simulations which are often the best available description of a system ([Kevrekidis & Samaey 2009](#), [Kevrekidis et al. 2004](#), [2003](#), e.g.). This suite of functions should empower users to start implementing such methods—but so far we have only just started.

Contents

1	Introduction	3
2	Projective integration of deterministic ODEs	8
2.1	Introduction	9
2.2	PIRK2(): projective integration of second-order accuracy . . .	10
2.3	egPIMM: Example projective integration of Michaelis–Menton kinetics	16
2.4	PIG(): Projective Integration via a General macroscale integrator	18
2.5	PIRK4(): projective integration of fourth-order accuracy . . .	25
2.6	Example: PI using Runge–Kutta macrosolvers	32
2.7	Example: Projective Integration using General macrosolvers	34
2.8	Explore: Projective Integration using constraint-defined manifold computing	36
2.9	To do/discuss	38
3	Patch scheme for given microscale discrete space system	39
3.1	Introduction	40
3.2	configPatches1(): configures spatial patches in 1D	41
3.3	patchSmooth1(): interface to time integrators	45
3.4	patchEdgeInt1(): sets edge values from interpolation over the macroscale	46
3.5	homogenisationExample: simulate heterogeneous diffusion in 1D	50
3.6	BurgersExample: simulate Burgers’ PDE on patches	54
3.7	ensembleAverageExample: simulate an ensemble of solutions for heterogeneous diffusion in 1D	58
3.8	waterWaveExample: simulate a water wave PDE on patches .	63
3.9	configPatches2(): configures spatial patches in 2D	69
3.10	patchSmooth2(): interface to time integrators	74
3.11	patchEdgeInt2(): 2D patch edge values from 2D interpolation	75

3.12	wave2D: example of a wave on patches in 2D	80
3.13	To do	83
3.14	Miscellaneous tests	83
A	Create, document and test algorithms	88
B	Aspects of developing a ‘toolbox’ for patch dynamics	91
B.1	Macroscale grid	91
B.2	Macroscale field variables	91
B.3	Boundary and coupling conditions	92
B.4	Mesotime communication	92
B.5	Projective integration	92
B.6	Lift to many internal modes	93
B.7	Macroscale closure	93
B.8	Exascale fault tolerance	93
B.9	Link to established packages	94

1 Introduction

This Developers Manual contains complete descriptions of the code in each function in the toolbox, and each example. For concise descriptions of each function, quick start guides, and some basic examples, see the User Manual.

Users Place the folder of this toolbox in a path searched by MATLAB. Then read the section(s) that documents the function of interest.

Quick start Maybe start by adapting one of the included examples. Many of the main functions include, at their start, example code of their use (code which is executed if the function is invoked without any arguments).

- To projectively integrate over time a multiscale, slow-fast, system of ODEs you could use `PIRK2()`: adapt the Michaelis–Menten example at the start of `PIRK2.m` ([Section 2.2.2](#)).
- You may use forward bursts of simulation in order to simulate the slow dynamics backward in time, as in `egPIMM.m` ([Section 2.3](#)).
- To only resolve the slow dynamics in the projective integration, use lifting and restriction functions by adapting the singular perturbation ODE example at the start of `PIG.m` ([Section 2.4.2](#)).
- Consider an evolving system over a large spatial domains when all you have is a microscale code. To efficiently simulate over the large domain, one can simulate in just small patches of the domain, appropriately coupled:
 - in 1D adapt the code at the start of `configPatches1.m` for Burgers’ PDE ([Section 3.2.2](#)), or the staggered patches of 1D water wave equations in `waterWaveExample.m` ([Section 3.8](#));
 - in 2D adapt the code at the start of `configPatches2.m` for non-linear diffusion ([Section 3.9.2](#)), or the regular patches of the 2D wave equation of `wave2D.m` ([Section 3.12](#)).
- The above are for systems that have *smooth* spatial structures on the microscale: when the microscale is ‘rough’ with a known period (so far only in 1D), then adapt the example of `HomogenisationExample.m` ([Section 3.5](#)).

Blackbox scenario Assume that a researcher/practitioner has a detailed and *trustworthy* computational simulation of some problem of interest. Let’s say the simulation is coded in terms of micro-field variable values $\vec{u}_i(t)$ for indices i in some (large) set of integers, and for time t . Perhaps the indices i correspond to ‘spatial’ coordinates $\vec{x}_i(t)$. In lattice problems the positions \vec{x}_i

would be fixed in time (unless employing a moving mesh on the microscale); in particle problems the positions would evolve. The positional coordinates are $\vec{x}_i \in \mathbb{R}^d$ where for spatial problems integer $d = 1, 2, 3$, but it may be more when solving for a distribution of velocities, or pore sizes, or trader's beliefs, etc. The micro-field variables could be in \mathbb{R}^p for any $p = 1, 2, \dots, \infty$. Further, assume that the computational simulation is too expensive over all the desired spatial domain $\mathbb{X} \subset \mathbb{R}^d$. Thus we aim a toolbox to simulate only on macroscale distributed patches.

Contributors The aim of this project is to collectively develop a MATLAB toolbox of equation-free algorithms. Initially the algorithms are basic, and the plan is to subsequently develop more and more capability.

MATLAB appears a good choice for a first version since it is widespread, efficient, supports various parallel modes, and development costs are reasonably low. Further it is built on BLAS and LAPACK so the cache and superscalar CPU are potentially well utilised. We aim to develop functions that work for MATLAB. [Appendix A](#) outlines some details for contributors.

Figure 1.1: The Projective Integration method greatly accelerates simulation/integration of a system exhibiting multiple time scales. The Projective Integration [Chapter 2](#) presents several separate functions, as well as several optional wrapper functions that may be invoked. This chart overviews constructing a Projective Integration simulation, whereas [Figure 1.2](#) roughly guides which top-level Projective Integration functions should be used. [Chapter 2](#) fully details each function.

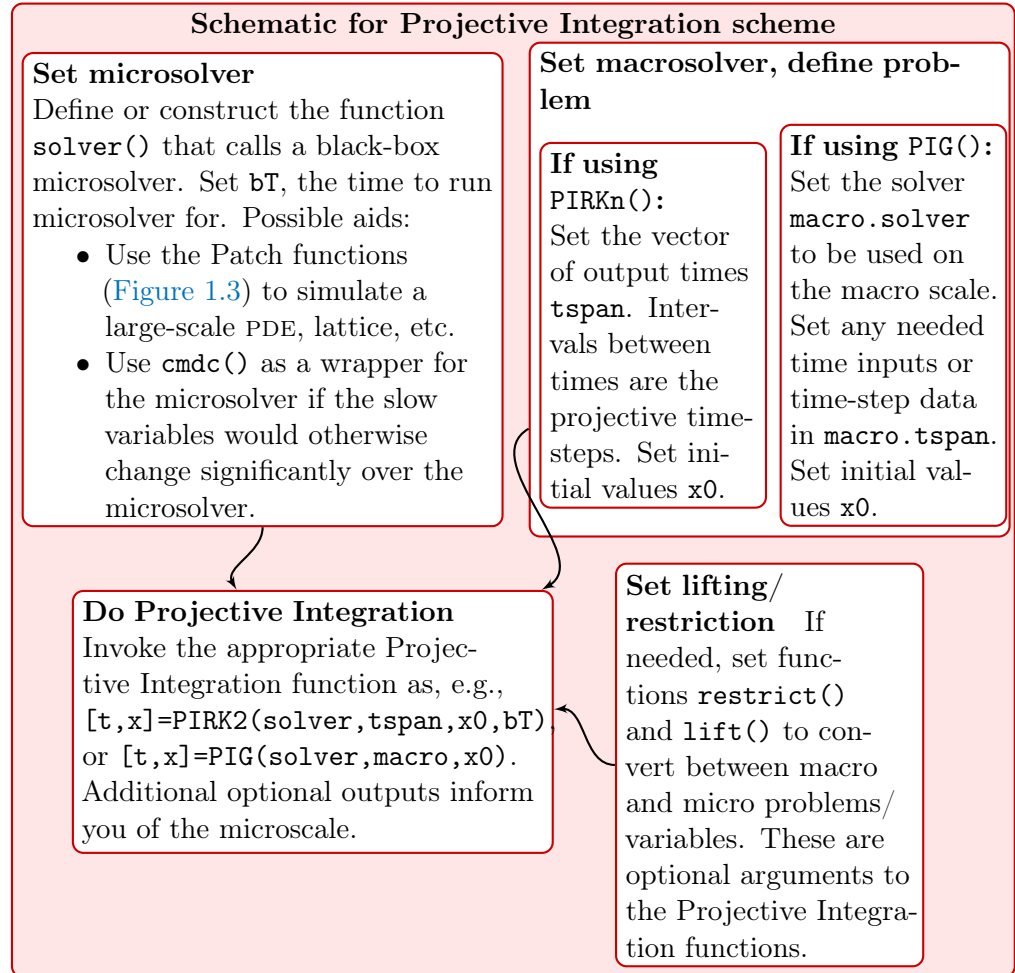


Figure 1.2: The Projective Integration method greatly accelerates simulation/integration of a system exhibiting multiple time scales. In conjunction with Figure 1.1, this chart roughly guides which top-level Projective Integration functions should be used. Chapter 2 fully details each function.

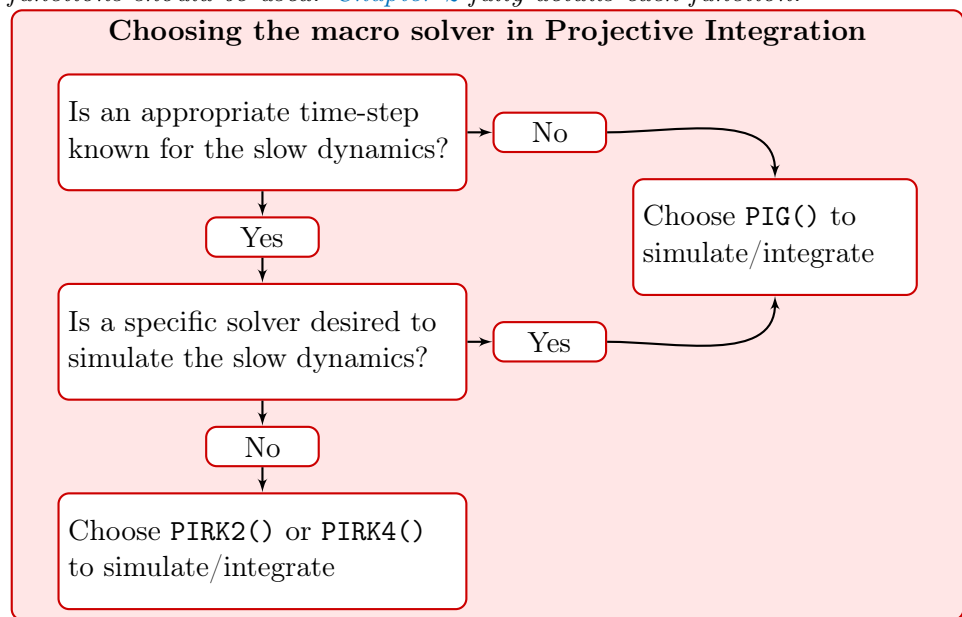
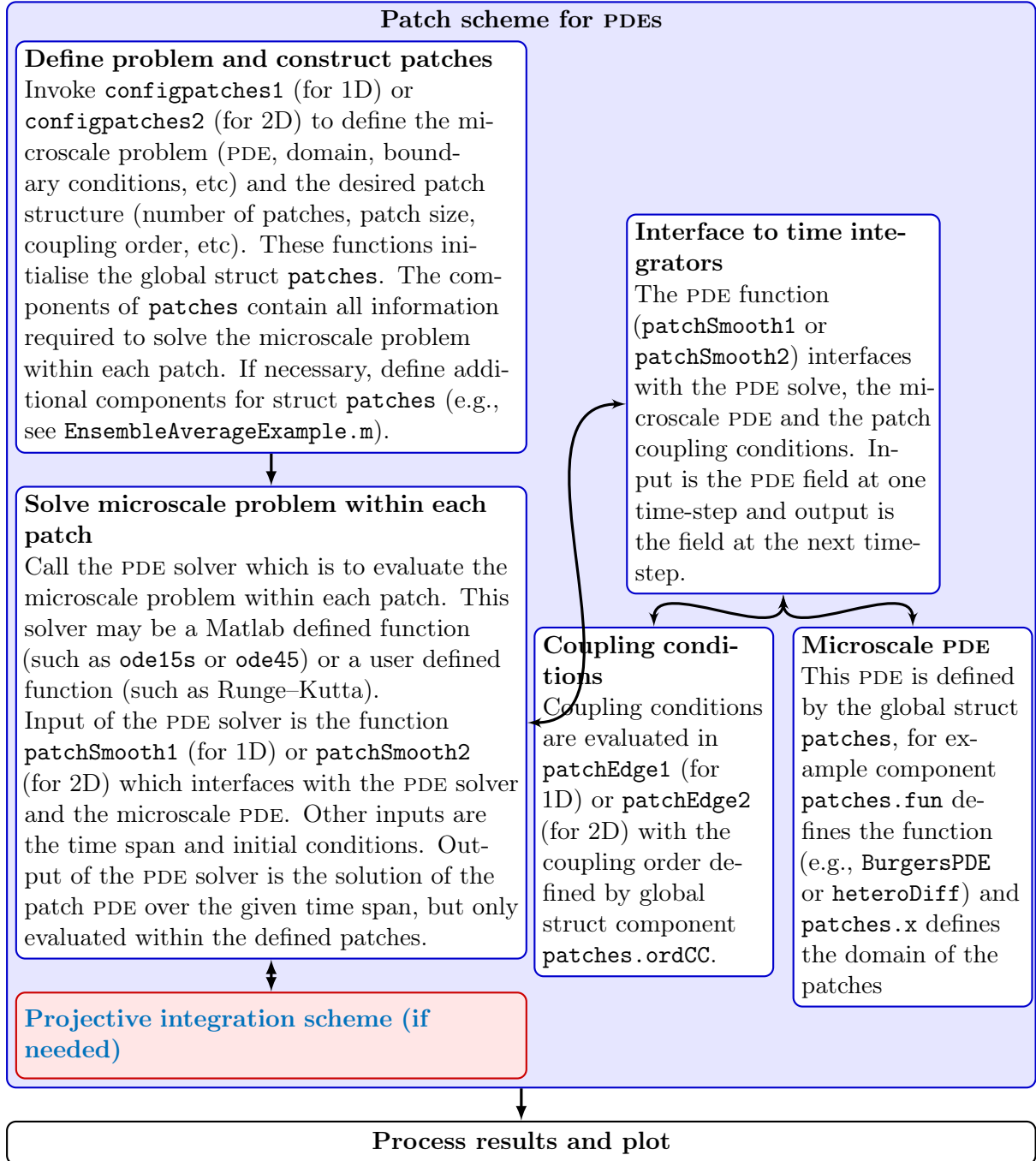


Figure 1.3: The Patch methods, [Chapter 3](#), accelerate simulation/integration of multiscale systems with interesting spatial (or network) structure/patterns. The patch methods use your given microsimulators whether coded from PDEs, lattice systems, or agent/particle microscale simulators. The patch functions require that a user configure the patches, and interface the coupled patches with a time integrator/simulator. This chart overviews the main functions involved and their interrelationships.



2 Projective integration of deterministic ODEs

Chapter contents

2.1	Introduction	9
2.2	PIRK2(): projective integration of second-order accuracy . . .	10
2.2.1	Introduction	10
2.2.2	If no arguments, then execute an example	12
2.2.3	The projective integration code	13
2.2.4	If no output specified, then plot simulation	16
2.3	egPIMM: Example projective integration of Michaelis–Menton kinetics	16
2.3.1	Invoke projective integration	16
2.3.2	Code a burst of Michaelis–Menten enzyme kinetics . . .	18
2.4	PIG(): Projective Integration via a General macroscale integrator	18
2.4.1	Introduction	19
2.4.2	If no arguments, then execute an example	21
2.4.3	The projective integration code	23
2.4.4	If no output specified, then plot simulation.	25
2.5	PIRK4(): projective integration of fourth-order accuracy . . .	25
2.5.1	Introduction	26
2.5.2	The projective integration code	28
2.5.3	If no output specified, then plot simulation	31
2.5.4	cdmc()	31
2.6	Example: PI using Runge–Kutta macrosolvers	32
2.7	Example: Projective Integration using General macrosolvers	34
2.8	Explore: Projective Integration using constraint-defined manifold computing	36
2.9	To do/discuss	38

2.1 Introduction

This section provides some good projective integration functions (Gear & Kevrekidis 2003*b,c*, Givon et al. 2006, Maclean & Gottwald 2015, Sieber et al. 2018, e.g.). The goal is to enable computationally expensive multi-scale dynamic simulations/integrations to efficiently compute over very long time scales.

Quick start Section 2.2.2 shows the most basic use of a projective integration function. Section 2.3 shows how to code more variations of the introductory example of a long time simulation of the Michaelis–Menton multiscale system of differential equations.

Scenario When you are interested in a complex system with many interacting parts or agents, you usually are primarily interested in the self-organised emergent macroscale characteristics. Projective integration empowers us to efficiently simulate such long-time emergent dynamics. We suppose you have coded some accurate, fine scale simulation of the complex system, and call such code a microsolver.

The Projective Integration section of this toolbox consists of several functions. Each function implements over a long-time scale a variant of a standard numerical method to simulate/integrate the emergent dynamics of the complex system. Each function has standardised inputs and outputs.

Main functions

- Projective Integration by second or fourth-order Runge–Kutta, `PIRK2()` and `PIRK4()` respectively. These schemes are suitable for precise simulation of the slow dynamics, provided the time period spanned by an application of the microsolver is not too large.
- Projective Integration with a General solver, `PIG()`. This function enables a Projective Integration implementation of any solver with macroscale time-steps. It does not matter whether the solver is a standard Matlab algorithm, or one supplied by the user. As explored in later examples, `PIG()` should only be used in very stiff systems.
- ‘Constraint-defined manifold computing’, `cdmc()`. This helper function, based on the method introduced in Gear et al. (2005*a*), iteratively applies the microsolver and projects the output backwards in time. The result is to constrain the fast variables close to the slow manifold, without advancing the current time by the duration of an application of the microsolver. This function can be used to reduce errors related to the simulation length of the microsolver in either the `PIRK` or `PIG` functions. In particular, it enables `PIG()` to be used on problems that are not particularly stiff.

The above functions share dependence on a user-specified ‘microsolver’, that accurately simulates some problem of interest.

The following sections describe the `PIRK2()` and `PIG()` functions in detail, providing an example for each. Then `PIRK4()` is very similar to `PIRK2()`. Descriptions for the minor functions follow, and an example of the use of `cdmc()`.

2.2 `PIRK2()`: projective integration of second-order accuracy

Section contents

2.2.1	Introduction	10
2.2.2	If no arguments, then execute an example	12
2.2.3	The projective integration code	13
2.2.4	If no output specified, then plot simulation	16

2.2.1 Introduction

This Projective Integration scheme implements a macroscale scheme that is analogous to the second-order Runge–Kutta Improved Euler integration.

```
21 function [x, tms, xms, rm, svf] = PIRK2(microBurst, tSpan, x0, bT)
```

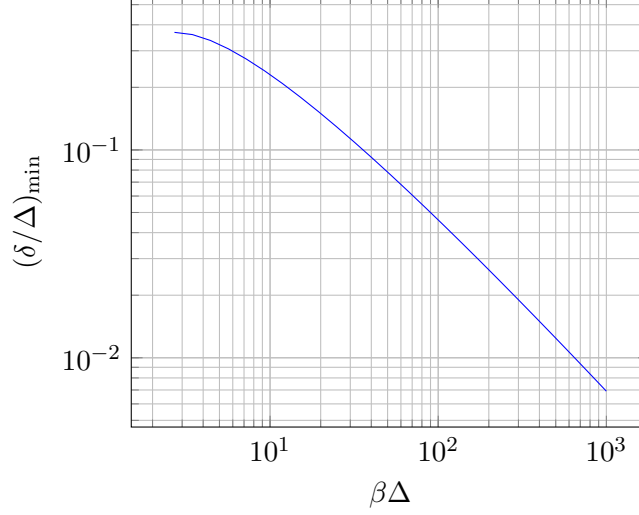
Input If there are no input arguments, then this function applies itself to the Michaelis–Menton example: see the code in [Section 2.2.2](#) as a basic template of how to use.

- `microBurst()`, a user-coded function that computes a short-time burst of the microscale simulation.

```
[tOut, xOut] = microBurst(tStart, xStart, bT)
```

- Inputs: `tStart`, the start time of a burst of simulation; `xStart`, the row n -vector of the starting state; `bT`, optional, the total time to simulate in the burst—if `microBurst()` determines the burst time, then replace `bT` in the argument list by `varargin`.
- Outputs: `tOut`, the column vector of solution times; and `xOut`, an array in which each *row* contains the system state at corresponding times.
- `tSpan` is an ℓ -vector of times at which the user requests output, of which the first element is always the initial time. `PIRK2()` does not use adaptive time-stepping; the macroscale time-steps are (nearly) the steps between elements of `tSpan`.
- `x0` is an n -vector of initial values at the initial time `tSpan(1)`. Elements of `x0` may be `NaN`: they are included in the simulation and output, and often represent boundaries in space fields.

Figure 2.1: Need macroscale step Δ such that $|\alpha\Delta| \lesssim \sqrt{6\varepsilon}$ for given relative error ε and slow rate α , and then $\delta/\Delta \gtrsim \frac{1}{\beta\Delta} \log \beta\Delta$ determines the minimum required burst length δ for given fast rate β .



- **bT**, optional, either missing, or empty (`[]`), or a scalar: if a given scalar, then it is the length of the micro-burst simulations—the minimum amount of time needed for the microscale simulation to relax to the slow manifold; else if missing or `[]`, then `microBurst()` must itself determine the length of a computed burst.

```
69 if nargin<4, bT=[]; end
```

Choose a long enough burst length Suppose: you have some desired relative accuracy ε that you wish to achieve (e.g., $\varepsilon \approx 0.01$ for two digit accuracy); the slow dynamics of your system occurs at rate/frequency of magnitude about α ; and the rate of *decay* of your fast modes are faster than the lower bound β (e.g., if the fast modes decay roughly like e^{-12t} , e^{-34t} , e^{-56t} then $\beta \approx 12$). Then choose

1. a macroscale time-step, $\Delta = \text{diff}(\text{tSpan})$, such that $\alpha\Delta \approx \sqrt{6\varepsilon}$, and
2. a microscale burst length, $\delta = \text{bT} \gtrsim \frac{1}{\beta} \log(\beta\Delta)$ (see [Figure 2.1](#)).

Output If there are no output arguments specified, then a plot is drawn of the computed solution \mathbf{x} versus tSpan .

- \mathbf{x} , an $\ell \times n$ array of the approximate solution vector. Each row is an estimated state at the corresponding time in tSpan . The simplest usage is then `x = PIRK2(microBurst,tSpan,x0,bT)`.

However, microscale details of the underlying Projective Integration computations may be helpful. `PIRK2()` provides two to four optional outputs of the microscale bursts.

- **tms**, optional, is an L dimensional column vector containing microscale times of burst simulations, each burst separated by `NaN`;

- **xms**, optional, is an $L \times n$ array of the corresponding microscale states—this data is an accurate simulation of the state and may help visualise more details of the solution.
- **rm**, optional, a struct containing the ‘remaining’ applications of the microBurst required by the Projective Integration method during the calculation of the macrostep:
 - **rm.t** is a column vector of microscale times; and
 - **rm.x** is the array of corresponding burst states.

The states **rm.x** do not have the same physical interpretation as those in **xms**; the **rm.x** are required in order to estimate the slow vector field during the calculation of the Runge–Kutta increments, and do not in general resemble the true dynamics.

- **svf**, optional, a struct containing the Projective Integration estimates of the slow vector field.
 - **svf.t** is a 2ℓ dimensional column vector containing all times at which the Projective Integration scheme is extrapolated along microBurst data to form a macrostep.
 - **svf.dx** is a $2\ell \times n$ array containing the estimated slow vector field.

2.2.2 If no arguments, then execute an example

```
174 if nargin==0
```

Example code for Michaelis–Menton dynamics The Michaelis–Menten enzyme kinetics is expressed as a singularly perturbed system of differential equations for $x(t)$ and $y(t)$ (encoded in function **MMburst** in the next paragraph):

$$\frac{dx}{dt} = -x + (x + \frac{1}{2})y \quad \text{and} \quad \frac{dy}{dt} = \frac{1}{\epsilon} [x - (x + 1)y].$$

With initial conditions $x(0) = 1$ and $y(0) = 0$, the following code computes and plots a solution over time $0 \leq t \leq 6$ for parameter $\epsilon = 0.05$. Since the rate of decay is $\beta \approx 1/\epsilon$ we choose a burst length $\epsilon \log(\Delta/\epsilon)$ as here the macroscale time-step $\Delta = 1$.

```
194 epsilon = 0.05
195 ts = 0:6
196 bT = epsilon*log((ts(2)-ts(1))/epsilon)
197 [x,tms,xms] = PIRK2(@MMburst, ts, [1;0], bT);
198 figure, plot(ts,x,'o:',tms,xms)
199 title('Projective integration of Michaelis--Menten enzyme kinetics')
200 xlabel('time t'), legend('x(t)','y(t)')
```

Upon finishing execution of the example, exit this function.

```
206 return
207 end%if no arguments
```

Example function code for a burst of ODEs Integrate a burst of length `bT` of the ODEs for the Michaelis–Menten enzyme kinetics at parameter ϵ inherited from above. Code ODEs in function `dMMdt` with variables $x = \mathbf{x}(1)$ and $y = \mathbf{x}(2)$. Starting at time `ti`, and state `xi` (row), we here simply use `ode23` to integrate in time.

```

221 function [ts, xs] = MMburst(ti, xi, bT)
222     dMMdt = @(t,x) [ -x(1)+(x(1)+0.5)*x(2)
223                     1/epsilon*( x(1)-(x(1)+1)*x(2) ) ];
224     [ts, xs] = ode23(dMMdt, [ti ti+bT], xi);
225 end

```

2.2.3 The projective integration code

Determine the number of time-steps and preallocate storage for macroscale estimates.

```

242 nT=length(tSpan);
243 x=nan(nT,length(x0));

```

Get the number of expected outputs and set logical indices to flag what data should be saved.

```

251 nArgs=nargout();
252 saveMicro = (nArgs>1);
253 saveFullMicro = (nArgs>3);
254 saveSvf = (nArgs>4);

```

Run a preliminary application of the `microBurst` on the initial conditions to help relax to the slow manifold. This is done in addition to the `microBurst` in the main loop, because the initial conditions are often far from the attracting slow manifold. Require the user to input and output rows of the system state.

```

267 x0 = reshape(x0,1,[]);
268 [relax_t,relax_x0] = microBurst(tSpan(1),x0,bT);

```

Use the end point of the `microBurst` as the initial conditions.

```

276 tSpan(1) = relax_t(end);
277 x(1,:)=relax_x0(end,:);

```

If saving information, then record the first application of the `microBurst`. Allocate cell arrays for times and states for outputs requested by the user, as concatenating cells is much faster than iteratively extending arrays.

```

287 if saveMicro
288     tms = cell(nT,1);
289     xms = cell(nT,1);
290     tms{1} = reshape(relax_t,[],1);
291     xms{1} = relax_x0;
292     if saveFullMicro
293         rm.t = cell(nT,1);
294         rm.x = cell(nT,1);
295         if saveSvf

```

```

296         svf.t = nan(2*nT-2,1);
297         svf.dx = nan(2*nT-2,length(x0));
298     end
299 end
300 end

```

Loop over the macroscale time-steps

```

308 for jT = 2:nT
309     T = tSpan(jT-1);

```

If two applications of the microBurst would cover one entire macroscale time-step, then do so (setting some internal states to NaN); else proceed to projective step.

```

317     if ~isempty(bT) & 2*abs(bT)>=abs(tSpan(jT)-T) & bT*(tSpan(jT)-T)>0
318         [t1,xm1] = microBurst(T, x(jT-1,:), tSpan(jT)-T);
319         x(jT,:) = xm1(end,:);
320         t2=nan; xm2=nan(1,size(xm1,2));
321         dx1=xm2; dx2=xm2;
322     else

```

Run the first application of the microBurst; since this application directly follows from the initial conditions, or from the latest macrostep, this microscale information is physically meaningful as a simulation of the system. Extract the size of the final time-step.

```

333         [t1,xm1] = microBurst(T, x(jT-1,:), bT);
334         del = t1(end)-t1(end-1);

```

Check for round-off error.

```

340         xt=[reshape(t1(end-1:end),[],1) xm1(end-1:end,:)];
341         roundingTol=1e-8;
342         if norm(diff(xt))/norm(xt,'fro') < roundingTol
343             warning(['significant round-off error in 1st projection at T=' num2str(T)
344             end

```

Find the needed time-step to reach time `tSpan(n+1)` and form a first estimate `dx1` of the slow vector field.

```

353         Dt = tSpan(jT)-t1(end);
354         dx1 = (xm1(end,:)-xm1(end-1,:))/del;

```

Project along `dx1` to form an intermediate approximation of `x`; run another application of the microBurst and form a second estimate of the slow vector field (assuming the burst length is the same, or nearly so).

```

364         xint = xm1(end,:) + (Dt-(t1(end)-t1(1)))*dx1;
365         [t2,xm2] = microBurst(T+Dt, xint, bT);
366         del = t2(end)-t2(end-1);
367         dx2 = (xm2(end,:)-xm2(end-1,:))/del;

```

Check for round-off error.


```

373     xt=[reshape(t2(end-1:end),[],1) xm2(end-1:end,:)];
374     if norm(diff(xt))/norm(xt,'fro') < roundingTol
375         warning(['significant round-off error in 2nd projection at T=' num2str(T)
376         end

```

Use the weighted average of the estimates of the slow vector field to take a macrostep.

```

384     x(jT,:) = xm1(end,:) + Dt*(dx1+dx2)/2;

```

Now end the if-statement that tests whether a projective step saves simulation time.

```

392     end

```

If saving trusted microscale data, then populate the cell arrays for the current loop iterate with the time-steps and output of the first application of the microBurst. Separate bursts by NaNs.

```

402     if saveMicro
403         tms{jT} = [reshape(t1,[],1); nan];
404         xms{jT} = [xm1; nan(1,size(xm1,2))];

```

If saving all microscale data, then repeat for the remaining applications of the microBurst.

```

412         if saveFullMicro
413             rm.t{jT} = [reshape(t2,[],1); nan];
414             rm.x{jT} = [xm2; nan(1,size(xm2,2))];

```

If saving Projective Integration estimates of the slow vector field, then populate the corresponding cells with times and estimates.

```

423             if saveSvf
424                 svf.t(2*jT-3:2*jT-2) = [t1(end); t2(end)];
425                 svf.dx(2*jT-3:2*jT-2,:) = [dx1; dx2];
426             end
427         end
428     end

```

Terminate the main loop:

```

434     end

```

Overwrite $x(1,:)$ with the specified initial condition $tSpan(1)$.

```

443     x(1,:) = reshape(x0,1,[]);

```

For additional requested output, concatenate all the cells of time and state data into two arrays.

```

451     if saveMicro
452         tms = cell2mat(tms);
453         xms = cell2mat(xms);
454         if saveFullMicro
455             rm.t = cell2mat(rm.t);
456             rm.x = cell2mat(rm.x);

```

```

457     end
458 end

```

2.2.4 If no output specified, then plot simulation

```

466 if nArgs==0
467     figure, plot(tSpan,x,'o:')
468     title('Projective Simulation with PIRK2')
469 end

    This concludes PIRK2().

476 end

```

2.3 egPIMM: Example projective integration of Michaelis–Menton kinetics

Section contents

2.3.1	Invoke projective integration	16
2.3.2	Code a burst of Michaelis–Menten enzyme kinetics . .	18

The Michaelis–Menten enzyme kinetics is expressed as a singularly perturbed system of differential equations for $x(t)$ and $y(t)$:

$$\frac{dx}{dt} = -x + (x + \tfrac{1}{2})y \quad \text{and} \quad \frac{dy}{dt} = \frac{1}{\epsilon} [x - (x + 1)y].$$

As illustrated in [Figure 2.3](#), the slow variable $x(t)$ evolves on a time scale of one, whereas the fast variable $y(t)$ evolves on a time scale of the small parameter ϵ .

2.3.1 Invoke projective integration

Clear, and set the scale separation parameter ϵ to something small like 0.01. Here use $\epsilon = 0.1$ for clearer graphs.

```

30 clear all, close all
31 global epsilon
32 epsilon = 0.1

```

First, [Section 2.3.2](#) encodes the computation of bursts of the Michaelis–Menten system in a function `MMburst()`. Second, here set macroscale times of computation and interest into vector `ts`. Then, invoke Projective Integration with `PIRK2()` applied to the burst function, say using bursts of simulations of length 2ϵ , and starting from the initial condition for the Michaelis–Menten system of $(x(0), y(0)) = (1, 0)$ (off the slow manifold).

```

47 ts = 0:6
48 xs = PIRK2(@MMburst, ts, [1;0], 2*epsilon)
49 plot(ts,xs,'o:')
50 xlabel('time t'), legend('x(t)', 'y(t)')
51 pause(1)

```

Figure 2.2: Michaelis–Menten enzyme kinetics simulated with the projective integration of `PIRK2()`: macroscale samples.

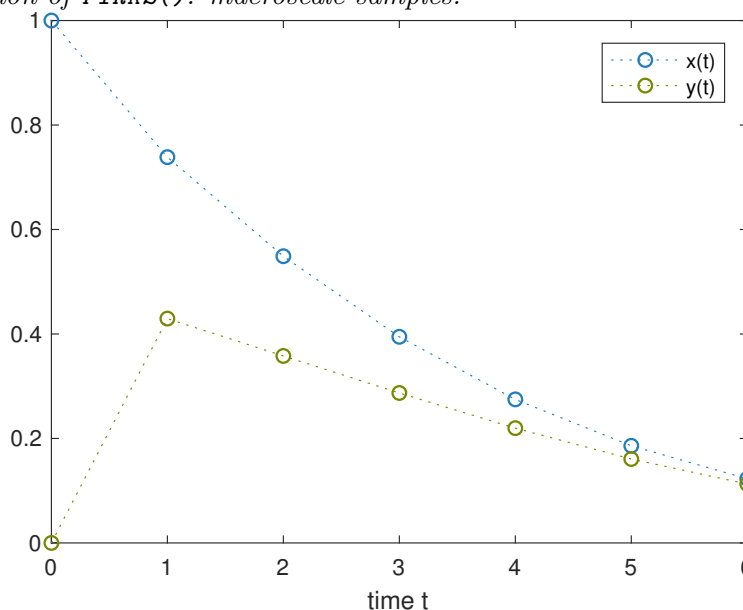


Figure 2.2 plots the macroscale results showing the long time decay of the Michaelis–Menten system on the slow manifold. Sieber et al. (2018) [§4] used this system as an example of their analysis of the convergence of Projective Integration.

Optional: request and plot the microscale bursts Because the initial conditions of the simulation are off the slow manifold, the initial macroscale step appears to ‘jump’ (Figure 2.2). To see the initial transient attraction to the slow manifold we plot some microscale data in Figure 2.3. Two further output variables provide this microscale burst information.

```

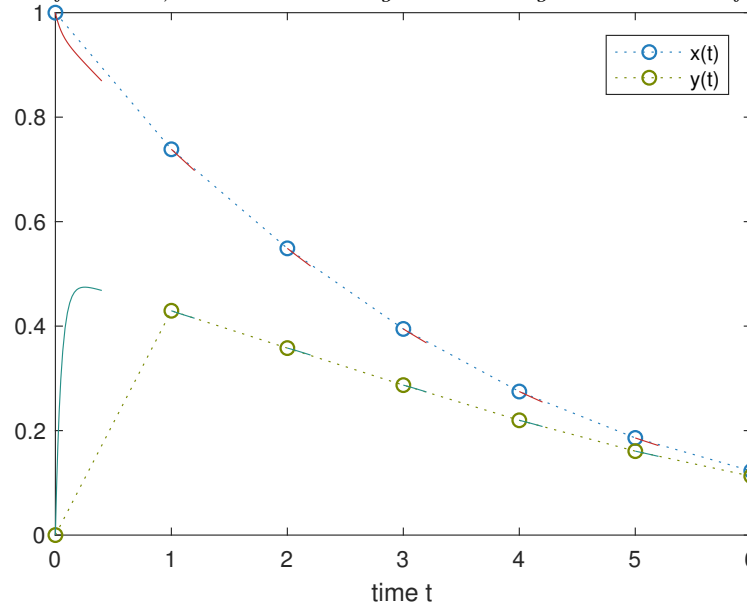
77 [xs,tMicro,xMicro] = PIRK2(@MMburst, ts, [1;0], 2*epsilon);
78 figure, plot(ts,xs,'o:',tMicro,xMicro)
79 xlabel('time t'), legend('x(t)', 'y(t)')
80 pause(1)

```

Figure 2.3 plots the macroscale and microscale results—also showing that the initial burst is by default twice as long. Observe the slow variable $x(t)$ is also affected by the initial transient which indicates that other schemes which ‘freeze’ slow variables are less accurate.

Optional: simulate backward in time Figure 2.4 shows that projective integration even simulates backward in time along the slow manifold using short forward bursts (Gear & Kevrekidis 2003a). Such backward macroscale simulations succeed despite the fast variable $y(t)$, when backward in time, being viciously unstable. However, backward integration appears to need longer bursts, here 3ϵ .

Figure 2.3: Michaelis–Menten enzyme kinetics simulated with the projective integration of `PIRK2()`: the microscale bursts show the initial transients on a time scale of $\epsilon = 0.1$, and then the alignment along the slow manifold.



```

110 ts = 0:-1:-5
111 [xs,tMicro,xMicro] = PIRK2(@MMburst, ts, 0.2*[1;1], 3*epsilon);
112 figure, plot(ts,xs,'o:',tMicro,xMicro)
113 xlabel('time t'), legend('x(t)','y(t)')

```

2.3.2 Code a burst of Michaelis–Menten enzyme kinetics

Say use `ode23()` to integrate a burst of the differential equations for the Michaelis–Menten enzyme kinetics. Code differential equations in function `dMMdt` with variables $x = x(1)$ and $y = x(2)$. For the given start time t_i , and start state x_i , `ode23()` integrates the differential equations for a burst time of bT , and return the simulation data.

```

140 function [ts, xs] = MMburst(ti, xi, bT)
141     global epsilon
142     dMMdt = @(t,x) [ -x(1)+(x(1)+0.5)*x(2)
143                     1/epsilon*( x(1)-(x(1)+1)*x(2) ) ];
144     [ts, xs] = ode23(dMMdt, [ti ti+bT], xi);
145 end

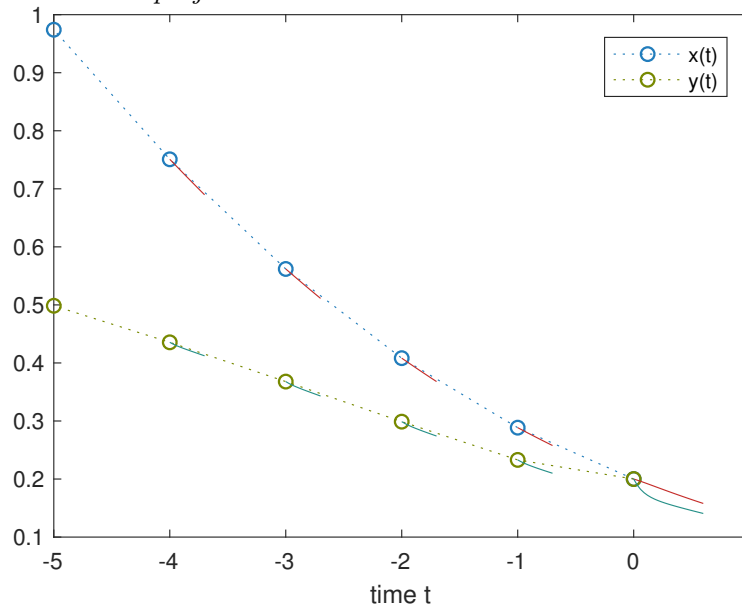
```

2.4 `PIG()`: Projective Integration via a General macroscale integrator

Section contents

2.4.1	Introduction	19
2.4.2	If no arguments, then execute an example	21
2.4.3	The projective integration code	23

Figure 2.4: Michaelis–Menten enzyme kinetics simulated backward with the projective integration of `PIRK2()`: the microscale bursts show the short forward simulations used to project backward in time at $\epsilon = 0.1$.



2.4.4 If no output specified, then plot simulation. 25

2.4.1 Introduction

This is a Projective Integration scheme when the macroscale integrator is any specified coded scheme. The advantage is that one may use MATLAB's inbuilt integration functions, with all their sophisticated error control and adaptive time-stepping, to do the macroscale integration/simulation.

By default, `PIG()` uses 'constraint-defined manifold computing' for the microscale simulations. This algorithm, initiated by [Gear et al. \(2005b\)](#), uses a backwards projection so that the simulation time is unchanged after running the microscale simulator. The implementation is `cdmc()`, described in [Section 2.5.4](#).

```

30 function [T,X,tms,xms,svf] = PIG(macroInt,microBurst,Tspan,x0 ...
31                                ,restrict,lift,cdmcFlag)

```

Inputs:

- `macroInt()`, the numerical method that the user wants to apply on a slow-time macroscale. Either input a standard MATLAB integration function (such as 'ode23' or 'ode45'), or code your own integration function using standard arguments. That is, if you code your own, then it must be

$$[Ts,Xs] = \text{macroInt}(F,Tspan,X0)$$

where

- function $F(T, X)$ notionally evaluates the time derivatives $d\vec{X}/dt$ at any time;
- **Tspan** is either the macro-time interval, or the vector of macroscale times at which macroscale values are to be returned; and
- **X0** are the initial values of \vec{X} at time **Tspan**(1).

Then the i th row of **Xs**, **Xs**($i, :$), is to be the vector $\vec{X}(t)$ at time $t = \text{Ts}(i)$. Remember that in **PIG()** the function $F(T, X)$ is to be estimated by Projective Integration.

- **microBurst()** is a function that produces output from the user-specified code for a burst of microscale simulation. The function must internally specify how long a burst it is to use. Usage

`[tbs,xbs] = microBurst(tb0,xb0)`

Inputs: **tb0** is the start time of a burst; **xb0** is the n -vector microscale state at the start of a burst.

Outputs: **tbs**, the vector of solution times; and **xbs**, the corresponding microscale states.

- **Tspan**, a vector of macroscale times at which the user requests output. The first element is always the initial time. If **macroInt** adaptively selects time steps (e.g., **ode45**), then **Tspan** consists of an initial and final time only.
- **x0**, the n -vector of initial microscale values at the initial time **Tspan**(1).

Optional Inputs: **PIG()** allows for none, two or three additional inputs after **x0**. If you distinguish distinct microscale and macroscale states and your aim is to do Projective Integration on the macroscale only, then lifting and restriction functions must be provided to convert between them. Usage **PIG(...,restrict,lift)**:

- **restrict(x)**, a function that takes an input n -dimensional microscale state \vec{x} and computes the corresponding N -dimensional macroscale state \vec{X} ;
- **lift(X,xApprox)**, a function that converts an input N -dimensional macroscale state \vec{X} to a corresponding n -dimensional microscale state \vec{x} , given that **xApprox** is a recently computed microscale state on the slow manifold.

Either both **restrict()** and **lift()** are to be defined, or neither. If neither are defined, then they are assumed to be identity functions, so that $N=n$ in the following.

If desired, the default constraint-defined manifold computing microsolver may be disabled, via **PIG(...,restrict,lift,cdmcFlag)**

- **cdmcFlag**, any seventh input to **PIG()**, will disable **cdmc()**, e.g., the string `'cdmc off'`.

If the `cdmcFlag` is to be set without using a `restrict()` or `lift()` function, then use empty matrices `[]` for the restrict and lift functions.

Output Between zero and five outputs may be requested. If there are no output arguments specified, then a plot is drawn of the computed solution `X` versus `T`. Most often you would store the first two output results of `PIG()`, via say `[T,X] = PIG(...)`.

- `T`, an L -vector of times at which `macroInt` produced results.
- `X`, an $L \times N$ array of the computed solution: the i th row of `X`, `X(i,:)`, is to be the macro-state vector $\vec{X}(t)$ at time $t = T(i)$.

However, microscale details of the underlying Projective Integration computations may be helpful, and so `PIG()` provides some optional outputs of the microscale bursts, via `[T,X,tms,xms] = PIG(...)`

- `tms`, optional, is an ℓ -dimensional column vector containing microscale times of burst simulations, each burst separated by `NaN`;
- `xms`, optional, is an $\ell \times n$ array of the corresponding microscale states.

In some contexts it may be helpful to see directly how Projective Integration approximates a reduced slow vector field, via `[T,X,tms,xms,svf] = PIG(...)` in which

- `svf`, optional, a struct containing the Projective Integration estimates of the slow vector field.
 - `svf.T` is a \hat{L} -dimensional column vector containing all times at which the microscale simulation data is extrapolated to form an estimate of $d\vec{x}/dt$ in `macroInt()`.
 - `svf.dX` is a $\hat{L} \times N$ array containing the estimated slow vector field.

If `macroInt()` is, for example, the forward Euler method (or the Runge–Kutta method), then $\hat{L} = L$ (or $\hat{L} = 4L$).

2.4.2 If no arguments, then execute an example

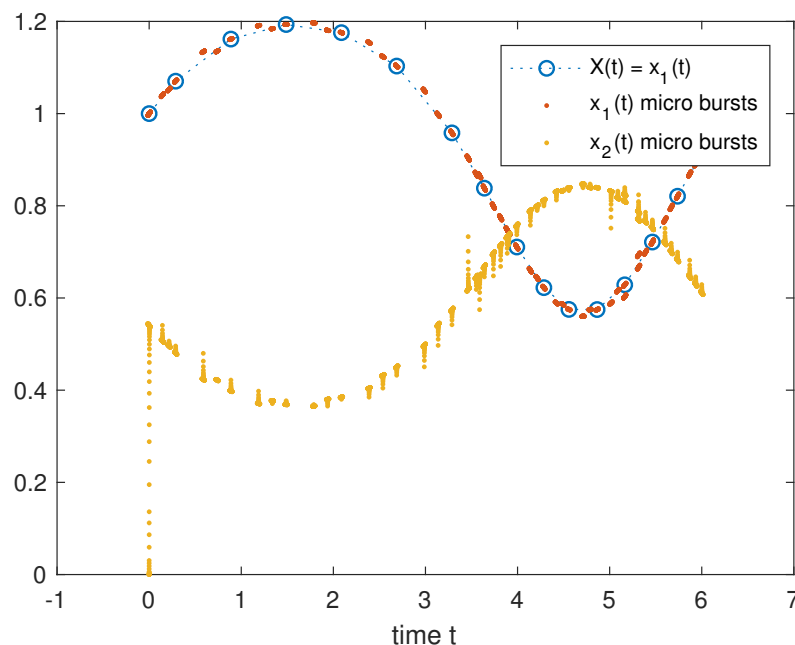
179 `if nargin==0`

As a basic example, consider a microscale system of the singularly perturbed system of differential equations

$$\frac{dx_1}{dt} = \cos(x_1) \sin(x_2) \cos(t) \quad \text{and} \quad \frac{dx_2}{dt} = \frac{1}{\epsilon} [\cos(x_1) - x_2]. \quad (2.1)$$

The macroscale variable is $X(t) = x_1(t)$, and the evolution dX/dt is unclear. With initial condition $X(0) = 1$, the following code computes and plots a solution of the system (2.1) over time $0 \leq t \leq 6$ for parameter $\epsilon = 10^{-3}$ (Figure 2.5). Whenever needed by `microBurst()`, the microscale system (2.1) is initialised ('lifted') using $x_2(t) = x_2^{\text{approx}}$ (yellow dots in Figure 2.5).

Figure 2.5: Projective Integration by PIG of the example system (2.1) in Section 2.4.2. The macroscale solution $X(t)$ is represented by just the blue circles. The microscale bursts are the microscale states $(x_1(t), x_2(t)) = (\text{red}, \text{yellow})$ dots.



First we code the right-hand side function of the microscale system (2.1) of ODEs.

```

213 epsilon = 1e-3;
214 dxdt=@(t,x) [ cos(x(1))*sin(x(2))*cos(t)
215               ( cos(x(1))-x(2) )/epsilon ];

```

Second, we code microscale bursts, here using the standard `ode45()`. We choose a burst length $2\epsilon \log(1/\epsilon)$ as the rate of decay is $\beta \approx 1/\epsilon$ and we do not know the macroscale time-step invoked by `macroInt()`, so blithely assume $\Delta \leq 1$ and then double the usual formula for safety.

```

227 bT = 2*epsilon*log(1/epsilon)
228 microBurst = @(tb0,xb0) ode45(dxdt,[tb0 tb0+bT],xb0);

```

Third, code functions to convert between macroscale and microscale states.

```

235 restrict = @(x) x(1);
236 lift = @(X,xApprox) [X; xApprox(2)];

```

Fourth, invoke PIG to use `ode23()`, say, on the macroscale slow evolution. Integrate the micro-bursts over $0 \leq t \leq 6$ from initial condition $\vec{x}(0) = (1, 0)$. You could set `Tspan=[0 -6]` to integrate backward in macroscale time with forward microscale bursts (Gear & Kevrekidis 2003a).

```

247 Tspan = [0 6];
248 x0 = [1;0];
249 [Ts,Xs,tms,xms] = PIG('ode23',microBurst,Tspan,x0,restrict,lift);

```


Plot output of this projective integration.

```

255 figure, plot(Ts,Xs,'o:',tms,xms,'.')
256 title('Projective integration of singularly perturbed ODE')
257 xlabel('time t')
258 legend('X(t) = x_1(t)', 'x_1(t) micro bursts', 'x_2(t) micro bursts')

```

Upon finishing execution of the example, exit this function.

```

264 return
265 end%if no arguments

```

2.4.3 The projective integration code

If no lifting/restriction functions are provided, then assign them to be the identity functions.

```

282 if nargin < 5 || isempty(restrict)
283     lift=@(X,xApprox) X;
284     restrict=@(x) x;
285 end

```

Get the number of expected outputs and set logical indices to flag what data should be saved.

```

293 nArgs=nargout();
294 saveMicro = (nArgs>2);
295 saveSvf = (nArgs>4);

```

Find the number of time-steps at which output is expected, and the number of variables.

```

303 nT=length(Tspan)-1;
304 nx = length(x0);
305 nX = length(restrict(x0));

```

Reformulate the microsolver to use `cdmc()`, unless flagged otherwise. The result is that the solution from `microBurst` will terminate at the given initial time.

```

314 if nargin<7
315     microBurst = @(t,x) cdmc(microBurst,t,x);
316 else
317     warning(['A ' class(cdmfFlag) ' seventh input to PIG().'\...
318         ' PIG will not use constraint-defined manifold computing.'])
319 end

```

Execute a first application of the `microBurst` on the initial conditions. This is done in addition to the `microBurst` in the main loop, because the initial conditions are often far from the attracting slow manifold.

```

331 [relaxT,x0MicroRelax] = microBurst(Tspan(1),x0);
332 xMicroLast = x0MicroRelax(end,:).';
333 X0Relax = restrict(xMicroLast);

```

Update the initial time.

```
340 Tspan(1) = relaxT(end);
```

Allocate cell arrays for times and states for any of the outputs requested by the user. If saving information, then record the first application of the microBurst. Note that it is unknown a priori how many applications of microBurst will be required; this code may be run more efficiently if the correct number is used in place of $nT+1$ as the dimension of the cell arrays.

```
352 if saveMicro
353     tms=cell(nT+1,1); xms=cell(nT+1,1);
354     n=1;
355     tms{n} = reshape(relaxT,[],1);
356     xms{n} = xOMicroRelax;
357
358     if saveSvf
359         svf.T = cell(nT+1,1);
360         svf.dX = cell(nT+1,1);
361     else
362         svf = [];
363     end
364 else
365     tms = []; xms = []; svf = [];
366 end
```

Define a function of macro simulation The idea of PIG() is to use the output from the microBurst() to approximate an unknown function $F(t, X)$ that computes $d\vec{X}/dt$. This approximation is then used in the system/user-defined ‘coarse solver’ macroInt(). The approximation is computed in

```
378 function [dXdt]=PIFun(t,X)
```

Run a microBurst from the given macroscale initial values.

```
384 x = lift(X,xMicroLast);
385 [tTmp,xMicroTmp] = microBurst(t,reshape(x,[],1));
386 xMicroLast = xMicroTmp(end,:).';
```

Compute the standard Projective Integration approximation of the slow vector field.

```
393 X2 = restrict(xMicroTmp(end,:));
394 X1 = restrict(xMicroTmp(end-1,:));
395 dt = tTmp(end)-tTmp(end-1);
396 dXdt = (X2 - X1).'/dt;
```

Save the microscale data, and the Projective Integration slow vector field, if requested.

```
403 if saveMicro
404     n=n+1;
405     tms{n} = [reshape(tTmp,[],1); nan];
```

```

406         xms{n} = [xMicroTmp; nan(1,nx)];
407         if saveSvf
408             svf.T{n-1} = t;
409             svf.dX{n-1} = dXdt;
410         end
411     end
412 end% PIFun function

```

Invoke the macroscale integration Integrate PIF() with the user-specified simulator macroInt(). For some reason, in MATLAB we need to use a one-line function, PIF, that invokes the above macroscale function, PIFun.

```

424 PIF = @(t,x) PIFun(t,x);
425 [T,X] = feval(macroInt,PIF,Tspan,X0Relax. ');

```

Overwrite X(1,:) and T(1), which the user expect to be X0 and Tspan(1) respectively, with the given initial conditions.

```

434 X(1,:) = restrict(x0);
435 T(1) = Tspan(1);

```

Concatenate all the additional requested outputs into arrays.

```

442 if saveMicro
443     tms = cell2mat(tms);
444     xms = cell2mat(xms);
445     if saveSvf
446         svf.T = cell2mat(svf.T);
447         svf.dX = cell2mat(svf.dX);
448     end
449 end

```

2.4.4 If no output specified, then plot simulation.

```

457 if nArgs==0
458     figure, plot(T,X,'o:')
459     title('Projective Simulation via PIG')
460 end

```

This concludes PIG().

```

468 end

```

2.5 PIRK4(): projective integration of fourth-order accuracy

Section contents

2.5.1	Introduction	26
2.5.2	The projective integration code	28
2.5.3	If no output specified, then plot simulation	31
2.5.4	cdmc()	31

2.5.1 Introduction

This Projective Integration scheme implements a macrosolver analogous to the fourth-order Runge–Kutta method.

```
18 function [x, tms, xms, rm, svf] = PIRK4(microBurst, tSpan, x0, bT)
```

See [Section 2.2](#) as the inputs and outputs are the same as PIRK2().

If no arguments, then execute an example

```
29 if nargin==0
```

Example of Michaelis–Menton backwards in time The Michaelis–Menten enzyme kinetics is expressed as a singularly perturbed system of differential equations for $x(t)$ and $y(t)$ (encoded in function `MMburst`):

$$\frac{dx}{dt} = -x + (x + \frac{1}{2})y \quad \text{and} \quad \frac{dy}{dt} = \frac{1}{\epsilon}[x - (x + 1)y].$$

With initial conditions $x(0) = y(0) = 0.2$, the following code uses forward time bursts in order to integrate backwards in time to $t = -5$. It plots the computed solution over time $-5 \leq t \leq 0$ for parameter $\epsilon = 0.1$. Since the rate of decay is $\beta \approx 1/\epsilon$ we choose a burst length $\epsilon \log(|\Delta|/\epsilon)$ as here the macroscale time-step $\Delta = -1$.

```
50 epsilon = 0.1
51 ts = 0:-1:-5
52 bT = epsilon*log(abs(ts(2)-ts(1))/epsilon)
53 [x,tms,xms,rm,svf] = PIRK4(@MMburst, ts, 0.2*[1;1], bT);
54 figure, plot(ts,x,'o:',tms,xms)
55 xlabel('time t'), legend('x(t)','y(t)')
56 title('Backwards-time projective integration of Michaelis--Menten')
```

Upon finishing execution of the example, exit this function.

```
62 return
63 end%if no arguments
```

Example function code for a burst of ODEs Integrate a burst of length `bT` of the ODEs for the Michaelis–Menten enzyme kinetics at parameter ϵ inherited from above. Code ODEs in function `dMMdt` with variables $x = \mathbf{x}(1)$ and $y = \mathbf{x}(2)$. Starting at time `ti`, and state `xi` (row), we here simply use `ode23` to integrate in time.

```
77 function [ts, xs] = MMburst(ti, xi, bT)
78     dMMdt = @(t,x) [ -x(1)+(x(1)+0.5)*x(2)
79                     1/epsilon*( x(1)-(x(1)+1)*x(2) ) ];
80     [ts, xs] = ode23(dMMdt, [ti ti+bT], xi);
81 end
```

Input

- `microBurst()`, a function that produces output from the user-specified code for microscale simulation.

`[tOut, xOut] = microBurst(tStart, xStart, bT)`

- Inputs: `tStart`, the start time of a burst of simulation; `xStart`, the row n -vector of the starting state; `bT`, optional, the total time to simulate in the burst—if `microBurst()` determines `bT`, then replace `bT` in the argument list by `varargin`.
- Outputs: `tOut`, the column vector of solution times; and `xOut`, an array in which each *row* contains the system state at corresponding times.
- `tSpan` is an ℓ -vector of times at which the user requests output, of which the first element is always the initial time. `PIRK4()` does not use adaptive time-stepping; the macroscale time-steps are (nearly) the steps between elements of `tSpan`.
- `x0` is an n -vector of initial values at the initial time `tSpan(1)`. Elements of `x0` may be `NaN`: they are included in the simulation and output, and often represent boundaries in space fields.
- `bT`, optional, either missing, or empty (`[]`), or a scalar: if a given scalar, then it is the length of the micro-burst simulations—the minimum amount of time needed for the microscale simulation to relax to the slow manifold; else if missing or `[]`, then `microBurst()` must itself determine the length of a computed burst.

130 `if nargin<4, bT=[]; end`

Output If there are no output arguments specified, then a plot is drawn of the computed solution \mathbf{x} versus `tSpan`.

- \mathbf{x} , an $\ell \times n$ array of the approximate solution vector. Each row is an estimated state at the corresponding time in `tSpan`. The simplest usage is then `x = PIRK4(microBurst,tSpan,x0,bT)`.

However, microscale details of the underlying Projective Integration computations may be helpful. `PIRK4()` provides two to four optional outputs of the microscale bursts.

- `tms`, optional, is an L dimensional column vector containing microscale times of burst simulations, each burst separated by `NaN`;
- `xms`, optional, is an $L \times n$ array of the corresponding microscale states—this data is an accurate simulation of the state and may help visualise more details of the solution.
- `rm`, optional, a struct containing the ‘remaining’ applications of the micro-burst required by the Projective Integration method during the calculation of the macrostep:

- `rm.t` is a column vector of microscale times; and
- `rm.x` is the array of corresponding burst states.

The states `rm.x` do not have the same physical interpretation as those in `xms`; the `rm.x` are required in order to estimate the slow vector field during the calculation of the Runge–Kutta increments, and do not in general resemble the true dynamics.

- `svf`, optional, a struct containing the Projective Integration estimates of the slow vector field.
 - `svf.t` is a 4ℓ dimensional column vector containing all times at which the Projective Integration scheme is extrapolated along micro-burst data to form a macrostep.
 - `svf.dx` is a $4\ell \times n$ array containing the estimated slow vector field.

2.5.2 The projective integration code

Determine the number of time-steps and preallocate storage for macroscale estimates.

```
194 nT=length(tSpan);
195 x=nan(nT,length(x0));
```

Get the number of expected outputs and set logical indices to flag what data should be saved.

```
203 nArgs=nargout();
204 saveMicro = (nArgs>1);
205 saveFullMicro = (nArgs>3);
206 saveSvf = (nArgs>4);
```

Run a preliminary application of the micro-burst on the initial conditions to help relax to the slow manifold. This is done in addition to the micro-burst in the main loop, because the initial conditions are often far from the attracting slow manifold. Require the user to input and output rows of the system state.

```
219 x0 = reshape(x0,1,[]);
220 [relax_t,relax_x0] = microBurst(tSpan(1),x0,bT);
```

Use the end point of the micro-burst as the initial conditions.

```
228 tSpan(1) = relax_t(end);
229 x(1,:)=relax_x0(end,:);
```

If saving information, then record the first application of the micro-burst. Allocate cell arrays for times and states for outputs requested by the user, as concatenating cells is much faster than iteratively extending arrays.

```
239 if saveMicro
240     tms = cell(nT,1);
241     xms = cell(nT,1);
242     tms{1} = reshape(relax_t,[],1);
243     xms{1} = relax_x0;
```

```

244     if saveFullMicro
245         rm.t = cell(nT,1);
246         rm.x = cell(nT,1);
247         if saveSvf
248             svf.t = nan(4*nT-4,1);
249             svf.dx = nan(4*nT-4,length(x0));
250         end
251     end
252 end

```

Loop over the macroscale time-steps

```

260 for jT = 2:nT
261     T = tSpan(jT-1);

```

If four applications of the micro-burst would cover the entire macroscale time-step, then do so (setting some internal states to NaN); else proceed to projective step.

```

270     if ~isempty(bT) & 4*abs(bT)>=abs(tSpan(jT)-T) & bT*(tSpan(jT)-T)>0
271         [t1,xm1] = microBurst(T, x(jT-1,:), tSpan(jT)-T);
272         x(jT,:) = xm1(end,:);
273         t2=nan; xm2=nan(1,size(xm1,2));
274         t3=nan; t4=nan; xm3=xm2; xm4 = xm2; dx1=xm2; dx2=xm2;
275     else

```

Run the first application of the micro-burst; since this application directly follows from the initial conditions, or from the latest macrostep, this microscale information is physically meaningful as a simulation of the system. Extract the size of the final time-step.

```

286         [t1,xm1] = microBurst(T, x(jT-1,:), bT);
287         del = t1(end)-t1(end-1);

```

Check for round-off error.

```

293         xt=[reshape(t1(end-1:end),[],1) xm1(end-1:end,:)];
294         roundingTol=1e-8;
295         if norm(diff(xt))/norm(xt,'fro') < roundingTol
296             warning(['significant round-off error in 1st projection at T=' num2str(T)
297         end

```

Find the needed time-step to reach time `tSpan(n+1)` and form a first estimate `dx1` of the slow vector field.

```

306         Dt = tSpan(jT)-t1(end);
307         dx1 = (xm1(end,:)-xm1(end-1,:))/del;

```

Assume burst times are the same length for this macro-step, or effectively so (recall that `bT` may be empty as it may be only coded and known in `microBurst()`).

```

315     abT = t1(end)-t1(1);

```

Project along $\mathbf{dx1}$ to form an intermediate approximation of \mathbf{x} ; run another application of the micro-burst and form a second estimate of the slow vector field.

```

326     xint = xm1(end,:) + (Dt/2-abT)*dx1;
327     [t2,xm2] = microBurst(T+Dt/2, xint, bT);
328     del = t2(end)-t2(end-1);
329     dx2 = (xm2(end,:)-xm2(end-1,:))/del;
330
331     xint = xm1(end,:) + (Dt/2-abT)*dx2;
332     [t3,xm3] = microBurst(T+Dt/2, xint, bT);
333     del = t3(end)-t3(end-1);
334     dx3 = (xm3(end,:)-xm3(end-1,:))/del;
335
336     xint = xm1(end,:) + (Dt-abT)*dx3;
337     [t4,xm4] = microBurst(T+Dt, xint, bT);
338     del = t4(end)-t4(end-1);
339     dx4 = (xm4(end,:)-xm4(end-1,:))/del;

```

Check for round-off error.

```

345     xt=[reshape(t2(end-1:end),[],1) xm2(end-1:end,:)];
346     if norm(diff(xt))/norm(xt,'fro') < roundingTol
347         warning(['significant round-off error in 2nd projection at T=' num2str(T)
348         end

```

Use the weighted average of the estimates of the slow vector field to take a macrostep.

```

356     x(jT,:) = xm1(end,:) + Dt*(dx1 + 2*dx2 + 2*dx3 + dx4)/6;

```

Now end the if-statement that tests whether a projective step saves simulation time.

```

364     end

```

If saving trusted microscale data, then populate the cell arrays for the current loop iterate with the time-steps and output of the first application of the micro-burst. Separate bursts by NaNs.

```

374     if saveMicro
375         tms{jT} = [reshape(t1,[],1); nan];
376         xms{jT} = [xm1; nan(1,size(xm1,2))];

```

If saving all microscale data, then repeat for the remaining applications of the micro-burst.

```

384     if saveFullMicro
385         rm.t{jT} = [reshape(t2,[],1); nan;...
386                   reshape(t3,[],1); nan;...
387                   reshape(t4,[],1); nan];
388         rm.x{jT} = [xm2; nan(1,size(xm2,2));...
389                   xm3; nan(1,size(xm2,2));...
390                   xm4; nan(1,size(xm2,2))];

```


If saving Projective Integration estimates of the slow vector field, then populate the corresponding cells with times and estimates.

```

399         if saveSvf
400             svf.t(4*jT-7:4*jT-4) = [t1(end); t2(end); t3(end); t4(end)];
401             svf.dx(4*jT-7:4*jT-4,:) = [dx1; dx2; dx3; dx4];
402         end
403     end
404 end

```

Terminate the main loop:

```

410 end

```

Overwrite $x(1,:)$ with the specified initial condition $tSpan(1)$.

```

419 x(1,:) = reshape(x0,1,[]);

```

For additional requested output, concatenate all the cells of time and state data into two arrays.

```

427 if saveMicro
428     tms = cell2mat(tms);
429     xms = cell2mat(xms);
430     if saveFullMicro
431         rm.t = cell2mat(rm.t);
432         rm.x = cell2mat(rm.x);
433     end
434 end

```

2.5.3 If no output specified, then plot simulation

```

442 if nArgs==0
443     figure, plot(tSpan,x,'o:')
444     title('Projective Simulation with PIRK4')
445 end

```

This concludes `PIRK4()`.

```

452 end

```

2.5.4 `cdmc()`

`cdmc()` iteratively applies the micro-burst and then projects backwards in time to the initial conditions. The cumulative effect is to relax the variables to the attracting slow manifold, while keeping the final time for the output the same as the input time.

```

13 function [ts, xs] = cdmc(microBurst,t0,x0)

```

Input

- `microBurst()`, a black-box micro-burst function suitable for Projective Integration. See any of `PIRK2()`, `PIRK4()`, or `PIG()` for a description of `microBurst()`.
- `t0`, an initial time
- `x0`, an initial state

Output

- `ts`, a vector of times. `tout(end)` will equal `t`.
- `xs`, an array of state estimates produced by `microBurst()`.

This function is a wrapper for the micro-burst. For instance if the problem of interest is a dynamical system that is not too stiff, and which can be simulated by the `microBurst sol(t,x,T)`, one would define

```
cSol = @(t,x) cdmc(sol,t,x) |
```

and thereafter use `csol()` in place of `sol()` as the `microBurst` for any Projective Integration scheme. The original `microBurst sol()` could create large errors if used in a Projective Integration scheme, but the output of `cdmc()` should not.

Begin with a standard application of the micro-burst.

```
41 [t1,x1] = feval(microBurst,t0,x0);
42 bT = t1(end)-t1(1);
```

Project backwards to before the initial time, then simulate just one burst forward to obtain a simulation burst that ends at the original `t0`.

```
50 dxdt = (x1(end,:) - x1(end-1,:))/(t1(end,:) - t1(end-1,:));
51 x0 = x1(end,:)-2*bT*dxdt;
52 t0 = t1(1)-bT;
53 [t2,x2] = feval(microBurst,t0,x0.');
```

Return both sets of output, though only `(t2,x2)` will be used in PI.

```
59 ts = [t1; t2];
60 xs = [x1; x2];
```

2.6 Example: PI using Runge–Kutta macrosolvers

This script is a demonstration of the `PIRK()` schemes, that use a Runge–Kutta macrosolver, applied to a simple linear system with some slow and fast directions.

Clear workspace and set a seed.

```
16 clear
17 rng(1)
18 global dxdt
```

The majority of this example involves setting up details for the microsolver. We use a simple function `gen_linear_system()` that outputs a function $f(t, x) = A\vec{x} + \vec{b}$, where matrix A has some eigenvalues with large negative real part, corresponding to fast variables and some eigenvalues with real part close to zero, corresponding to slow variables. The function `gen_linear_system()` requires that we specify bounds on the real part of the strongly stable eigenvalues,

```
33 fastband = [-5e2; -1e2];
```

and bounds on the real part of the weakly stable/unstable eigenvalues,

```
40 slowband = [-0.002; 0.002];
```

We now generate a random linear system with seven fast and three slow variables.

```
47 dxdt = gen_linear_system(7,3,fastband,slowband);
```

Set the macroscale times at which we request output from the PI scheme and the initial conditions.

```
57 tSpan = 0: 1 : 20;
```

```
58 x0 = linspace(-10,10,10)';
```

We implement the PI scheme, saving the coarse states in `x`, the ‘trusted’ applications of the microsolver in `tms` and `xms`, and the additional applications of the microsolver in `rm` (the second, third and fourth outputs are optional).

```
71 [x, tms, xms, rm] = PIRK4(@linearBurst, tSpan, x0);
```

To verify, we also compute the trajectories using a standard integrator.

```
78 [tt,ode45x] = ode45(dxdt,tSpan([1,end]),x0);
```

Figure 2.6 plots the output.

```
94 clf()
```

```
95 hold on
```

```
96 PI_sol=plot(tSpan,x,'bo');
```

```
97 std_sol=plot(tt,ode45x,'r');
```

```
98 plot(tms,xms,'k.', rm.t,rm.x,'g.');
```

```
99 legend([PI_sol(1),std_sol(1)],'PI Solution',...
```

```
100         'Standard Solution','Location','NorthWest')
```

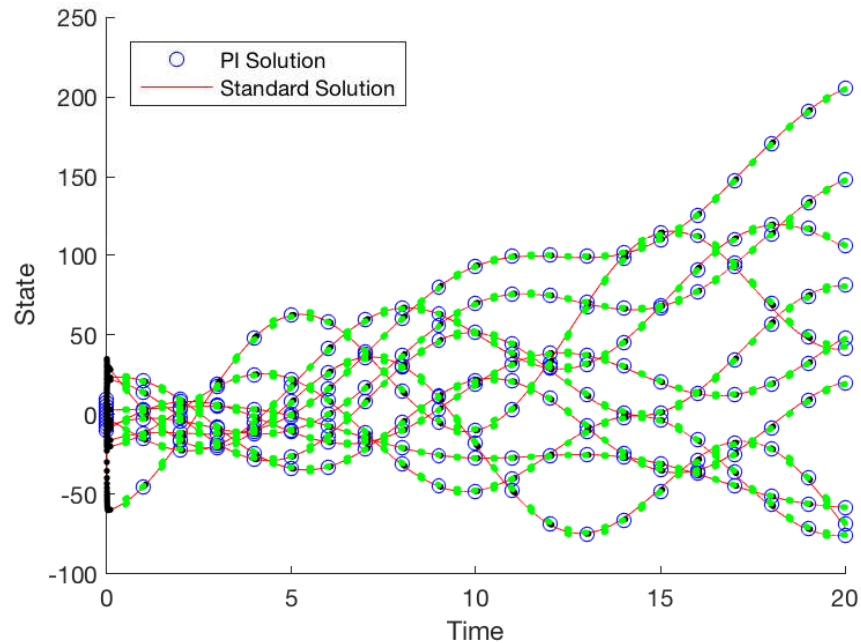
```
101 xlabel('Time'), ylabel('State')
```

Save plot to a file.

```
107 set(gcf,'PaperPosition',[0 0 14 10]), print('-depsc2','PIRK')
```

Code the micro-burst function using simple Euler steps. As a rule of thumb, the time-steps Δt should satisfy $\Delta t \leq 1/|\text{fastband}(1)|$ and the time to simulate with each application of the microsolver, ΔT , should be larger than or equal to $1/|\text{fastband}(2)|$. We set the integration scheme to be used in the microsolver. Since the time-steps are so small, we just use the forward Euler scheme

Figure 2.6: Demonstration of `PIRK4()`. From initial conditions, the system rapidly transitions to an attracting invariant manifold. The PI solution accurately tracks the evolution of the variables over time while requiring only a fraction of the computations of the standard solver.



```

121 function [ts, xs] = linearBurst(ti, xi, varargin)
122 global dxdt
123 dt = 0.001;
124 ts = ti+(0:dt:0.05)';
125 nts = length(ts);
126 xs = NaN(nts,length(xi));
127 xs(1,:)=xi;
128 for k=2:nts
129     xi = xi + dt*dxdt(ts(k),xi.').';
130     xs(k,:)=xi;
131 end
132 end

```

2.7 Example: Projective Integration using General macrosolvers

In this example the Projective Integration-General scheme is applied to a singularly perturbed ordinary differential equation. The aim is to use a standard non-stiff numerical integrator, such as `ode45()`, on the slow, long-time macroscale. For this stiff system, `PIG()` is an order of magnitude faster than ordinary use of `ode45`.

```

18 clear all, close all

```

Set time scale separation and model.

```

25  epsilon = 1e-4;
26  dxdt=@(t,x) [ cos(x(1))*sin(x(2))*cos(t)
27               (cos(x(1))-x(2))/epsilon ];

Set the 'black-box' microsolver to be an integration using a standard solver,
and set the standard time of simulation for the microsolver.

36  bT = epsilon*log(1/epsilon);
37  microBurst = @(tb0, xb0) ode45(dxdt,[tb0 tb0+bT],xb0);

Set initial conditions, and the time to be covered by the macrosolver.

45  x0 = [1 1.4];
46  tSpan=[0 15];

Now time and integrate the above system over tspan using PIG() and, for
comparison, a brute force implementation of ode45(). Report the time taken
by each method (in seconds).

55  tic
56  [ts,xs,tms,xms] = PIG('ode45',microBurst,tSpan,x0);
57  secsPIGusingODE45asMacro = toc
58  tic
59  [t45,x45] = ode45(dxdt,tSpan,x0);
60  secsODE45alone = toc

Plot the output on two figures, showing the truth and macrosteps on both,
and all applications of the microsolver on the first figure.

70  figure
71  h = plot(ts,xs,'o', t45,x45,'-', tms,xms,'.');
72  legend(h(1:2:5),'PI Solution','ode45 Solution','PI microsolver')
73  xlabel('Time'), ylabel('State')
74
75  figure
76  h = plot(ts,xs,'o', t45,x45,'-');
77  legend(h([1 3]),'PI Solution','ode45 Solution')
78  xlabel('Time'), ylabel('State')
79  %set(gcf,'PaperPosition',[0 0 14 10]), print('-depsc2','PIGExample')

```

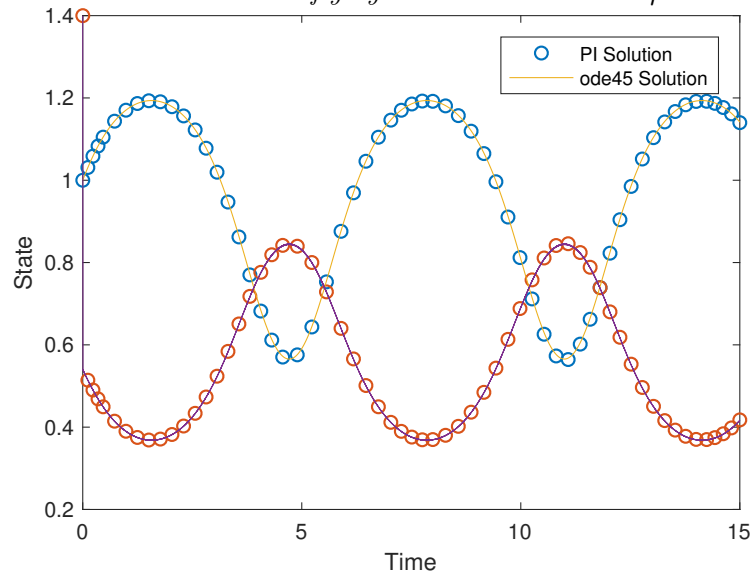
Figure 2.7 plots the output.

- The problem may be made more, or less, stiff by changing the time-scale separation parameter $\epsilon = \text{epsilon}$. The compute time of `PIG()` is almost independent of ϵ , whereas that of `ode45()` is proportional to $1/\epsilon$.

But if the problem is insufficiently stiff (larger ϵ), then `PIG()` produces nonsense. This nonsense is overcome by `cdmc()` (Section 2.8).

- The mildly stiff problem in Section 2.6 may be efficiently solved by a standard solver (e.g., `ode45()`). The stiff but low dimensional problem in this example can be solved efficiently by a standard stiff solver (e.g., `ode15s()`). The real advantage of the Projective Integration schemes is

Figure 2.7: Accurate simulation of a stiff nonautonomous system by `PIG()`. The microsolver is called on-the-fly by the macrosolver `ode45`.



in high dimensional stiff problems, that cannot be efficiently solved by most standard methods.

2.8 Explore: Projective Integration using constraint-defined manifold computing

In this example the Projective Integration-General scheme is applied to a singularly perturbed ordinary differential equation in which the time scale separation is not large. The results demonstrate the value of the default `cdmc()` wrapper for the microsolver.

```
16 clear all, close all
```

Set a weak time scale separation, and model.

```
23 epsilon = 0.01;
24 dxdt=@(t,x) [ cos(x(1))*sin(x(2))*cos(t)
25               (cos(x(1))-x(2))/epsilon ];
```

Set the microsolver to be an integration using a standard solver, and set the standard time of simulation for the microsolver.

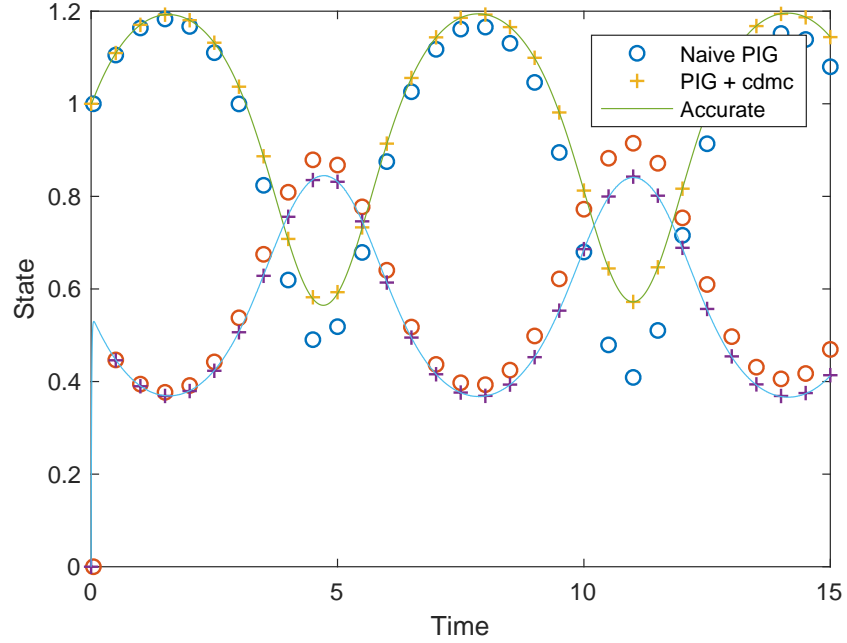
```
34 bT = epsilon*log(1/epsilon);
35 microBurst = @(tb0,xb0) ode45(dxdt,[tb0 tb0+bT],xb0);
```

Set initial conditions, and the time to be covered by the macrosolver.

```
43 x0 = [1 0];
44 tSpan=0:0.5:15;
```

Simulate using `PIG()`, first without the default treatment of `cdmc` for the microsolver and second with. Generate a trusted solution using standard numerical methods.

Figure 2.8: Accurate simulation of a weakly stiff non-autonomous system by `PIG()` using `cdmc()`, and an inaccurate solution using a naive application of `PIG()`.



```

55 [nt,nx] = PIG('ode45',microBurst,tSpan,x0,[],[],'no cdmc');
56 [ct,cx] = PIG('ode45',microBurst,tSpan,x0);
57 [t45,x45] = ode45(dxdt,tSpan([1 end]),x0);

```

Figure 2.8 plots the output.

```

73 figure
74 h = plot(nt,nx,'rx', ct,cx,'bo', t45,x45,'-');
75 legend(h(1:2:5),'Naive PIG','default: PIG + cdmc','Accurate')
76 xlabel('Time'), ylabel('State')
77 %set(gcf,'PaperPosition',[0 0 14 10]), print('-depsc2','PIGExplore')

```

The source of the error in the standard `PIG()` scheme is the burst length `bT`, that is significant on the slow time scale. Set `bT` to `20*epsilon` or `50*epsilon`¹ to worsen the error in both schemes. This example reflects a general principle, that most Projective Integration schemes will incur a global error term which is proportional to the simulation time of the microsolver and independent of the order of the microsolver. The `PIRK()` schemes have been written to minimise, if not eliminate entirely, this error, but by design `PIG()` works with any user-defined macrosolver and cannot reduce this error. The function `cdmc()` reduces this error term by attempting to mimic the microsolver without advancing time.

¹ This example is quite extreme: at `bT=50*epsilon`, it would be computationally much cheaper to simulate the entire length of `tSpan` using the microsolver alone.

2.9 To do/discuss

- Can we implement for Octave? We would like to use nested functions for some examples, because the function code then inherits parameter(s) from the main function. However, in Octave we cannot then use handles to these nested functions due to the error “handles to nested functions are not yet supported”—which apparently is not going to be fixed anytime soon (as at March 2019).
- could implement Projective Integration by ‘arbitrary’ Runge–Kutta scheme; that is, by having the user input a particular Butcher table—surely only specialists would be interested.
- can ‘reverse’ the order of projection and microsolver applications with a little fiddling. Then output at each user-requested coarse time is the end point of an application of the microsolver—better predictions for fast variables.
- Can maybe implement microsolvers that terminate a burst when the fast dynamics have settled using, for example, the ‘Events’ function handle in ode23.
- Need projective integration of systems with fast oscillations, perhaps by DMD.
- Need projective integration for stochastic systems.

3 Patch scheme for given microscale discrete space system

Chapter contents

3.1	Introduction	40
3.2	<code>configPatches1()</code> : configures spatial patches in 1D	41
3.2.1	Introduction	41
3.2.2	If no arguments, then execute an example	42
3.2.3	The code to make patches and interpolation	43
3.3	<code>patchSmooth1()</code> : interface to time integrators	45
3.3.1	Introduction	45
3.4	<code>patchEdgeInt1()</code> : sets edge values from interpolation over the macroscale	46
3.4.1	Introduction	46
3.5	<code>homogenisationExample</code> : simulate heterogeneous diffusion in 1D	50
3.5.1	Script to simulate via stiff or projective integration	51
3.5.2	<code>heteroDiff()</code> : heterogeneous diffusion	53
3.5.3	<code>heteroBurst()</code> : a burst of heterogeneous diffusion	54
3.6	<code>BurgersExample</code> : simulate Burgers' PDE on patches	54
3.6.1	Script code to simulate a microscale space-time map	54
3.6.2	<code>burgersMap()</code> : discretise the PDE microscale	57
3.6.3	<code>burgerBurst()</code> : code a burst of the patch map	57
3.7	<code>ensembleAverageExample</code> : simulate an ensemble of solutions for heterogeneous diffusion in 1D	58
3.7.1	Script to simulate via stiff or projective integration	59
3.7.2	<code>heteroDiff()</code> : heterogeneous diffusion	63
3.7.3	<code>heteroBurst()</code> : a burst of heterogeneous diffusion	63
3.8	<code>waterWaveExample</code> : simulate a water wave PDE on patches	63
3.8.1	Script code to simulate wave systems	65
3.8.2	<code>idealWavePDE()</code> : ideal wave PDE	67
3.8.3	<code>waterWavePDE()</code> : water wave PDE	68

3.9	<code>configPatches2()</code> : configures spatial patches in 2D	69
3.9.1	Introduction	69
3.9.2	If no arguments, then execute an example	70
3.9.3	The code to make patches	72
3.10	<code>patchSmooth2()</code> : interface to time integrators	74
3.10.1	Introduction	74
3.11	<code>patchEdgeInt2()</code> : 2D patch edge values from 2D interpolation	75
3.12	<code>wave2D</code> : example of a wave on patches in 2D	80
3.12.1	Check on the linear stability of the wave PDE	80
3.12.2	Execute a simulation	81
3.12.3	Example of simple wave PDE inside patches	82
3.13	To do	83
3.14	Miscellaneous tests	83
3.14.1	<code>patchEdgeInt1test</code> : test the spectral interpolation	83
3.14.2	<code>patchEdgeInt2test</code> : tests 2D spectral interpolation	86

3.1 Introduction

The patch scheme applies to spatio-temporal systems where the spatial domain is larger than what can be computed in reasonable time in a given complicated microscale code. In the scheme we compute the microscale details only on small patches of the space-time domain, and produce correct macroscale predictions by craftily coupling the patches across unsimulated space (Hyman 2005, Samaey et al. 2005, 2006, Roberts & Kevrekidis 2007, Liu et al. 2015, e.g.). The resulting macroscale predictions were generally proved to be consistent with the microscale dynamics, to some specified order of accuracy, in a series of papers: 1D-space dissipative systems (Roberts & Kevrekidis 2007, Bunder et al. 2017); 2D-space dissipative systems (Roberts et al. 2014); and 1D-space wave-like systems (Cao & Roberts 2016b).

The microscale spatial structure is to be on a lattice such as obtained from finite difference approximation of a PDE. Usually continuous in time.

Quick start See Sections 3.2.2 and 3.9.2 which respectively list example basic code that uses the provided functions to simulate the 1D Burgers' PDE, and a 2D nonlinear 'diffusion' PDE.

3.2 configPatches1(): configures spatial patches in 1D

Section contents

3.2.1	Introduction	41
3.2.2	If no arguments, then execute an example	42
3.2.3	The code to make patches and interpolation	43

3.2.1 Introduction

Makes the struct `patches` for use by the patch/gap-tooth time derivative function `patchSmooth1()`. [Section 3.2.2](#) lists an example of its use.

```

17 function configPatches1(fun,Xlim,BCs,nPatch,ordCC,ratio,nSubP ...
18                               ,nEdge)
19 global patches
```

Input If invoked with no input arguments, then executes an example of simulating Burgers' PDE—see [Section 3.2.2](#) for the example code.

- `fun` is the name of the user function, `fun(t,u,x)`, that computes time derivatives (or time-steps) of quantities on the patches.
- `Xlim` give the macro-space domain of the computation: patches are equi-spaced over the interior of the interval `[Xlim(1),Xlim(2)]`.
- `BCs` somehow will define the macroscale boundary conditions. Currently, `BCs` is ignored and the system is assumed macro-periodic in the domain.
- `nPatch` is the number of equi-spaced spaced patches.
- `ordCC` is the 'order' of interpolation across empty space of the macroscale mid-patch values to the edge of the patches for inter-patch coupling: currently must be ≥ -1 .
- `ratio` (real) is the ratio of the half-width of a patch to the spacing of the patch mid-points: so `ratio` = $\frac{1}{2}$ means the patches abut; and `ratio` = 1 is overlapping patches as in holistic discretisation.
- `nSubP` is the number of equi-spaced microscale lattice points in each patch. Must be odd so that there is a central lattice point.
- `nEdge`, optional, for each patch, the number of edge values set by interpolation at the edge regions of each patch. May be omitted. The default is one (suitable for microscale lattices with only nearest neighbour interactions).

Output The *global* struct `patches` is created and set with the following components.

- `.fun` is the name of the user's function `fun(u,t,x)` that computes the time derivatives (or steps) on the patchy lattice.

- `.ordCC` is the specified order of inter-patch coupling.
- `.alt` is true for interpolation using only odd neighbouring patches as for staggered grids, and false for the usual case of all neighbour coupling.
- `.Cwtsr` and `.Cwtsl` are the `ordCC`-vector of weights for the inter-patch interpolation onto the right and left edges (respectively) with patch:macroscale ratio as specified.
- `.x` is `nSubP × nPatch` array of the regular spatial locations x_{ij} of the microscale grid points in every patch.
- `.nEdge` is, for each patch, the number of edge values set by interpolation at the edge regions of each patch.

3.2.2 If no arguments, then execute an example

```
100 if nargin==0
```

The code here shows one way to get started: a user's script may have the following three steps (left-right arrows denote function recursion).

1. `configPatches1`
2. `ode15s` integrator \leftrightarrow `patchSmooth1` \leftrightarrow user's `burgersPDE`
3. process results

Establish global patch data struct to interface with a function coding Burgers' PDE: to be solved on 2π -periodic domain, with eight patches, spectral interpolation couples the patches, each patch of half-size ratio 0.2, and with seven microscale points forming each patch.

```
119 configPatches1(@BurgersPDE,[0 2*pi], nan, 8, 0, 0.2, 7);
```

Set an initial condition, with some randomness, and simulate in time using a standard stiff integrator and the interface function `patchsmooth1()` (Section 3.3).

```
128 u0=0.3*(1+sin(patches.x))+0.1*randn(size(patches.x));
129 [ts,ucts]=ode15s(@patchSmooth1,[0 0.5],u0(:));
```

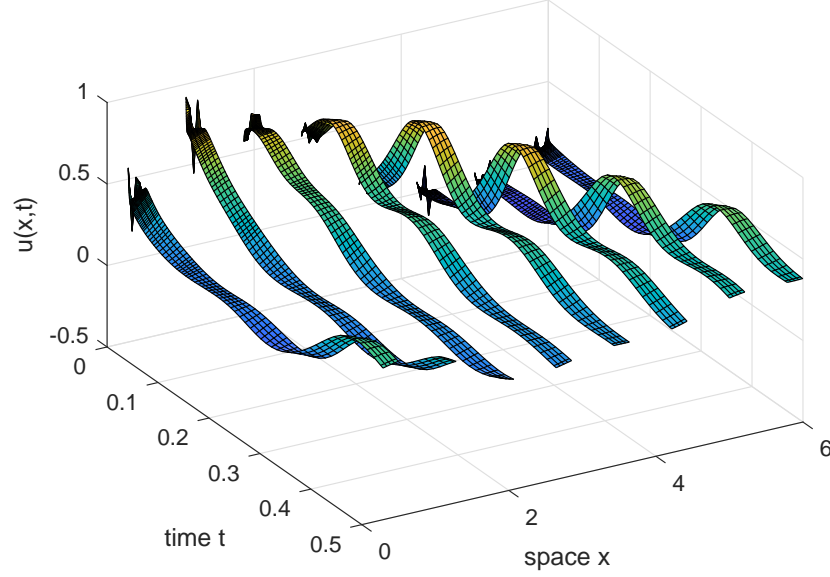
Plot the simulation using only the microscale values interior to the patches: set x -edges to `nan` to leave the gaps. Figure 3.1 illustrates an example simulation in time generated by the patch scheme applied to Burgers' PDE.

```
139 figure(1),clf
140 patches.x([1 end],:)=nan;
141 surf(ts,patches.x(:),ucts'), view(60,40)
142 title('Example of Burgers PDE on patches in space')
143 xlabel('time t'), ylabel('space x'), zlabel('u(x,t)')
```

Upon finishing execution of the example, exit this function.

```
154 return
155 end%if no arguments
```

Figure 3.1: field $u(x,t)$ of the patch scheme applied to Burgers' PDE.
Example of Burgers PDE on patches in space



Example of Burgers PDE inside patches As a microscale discretisation of Burgers' PDE $u_t = u_{xx} - 30uu_x$, here code $\dot{u}_{ij} = \frac{1}{\delta x^2}(u_{i+1,j} - 2u_{i,j} + u_{i-1,j}) - 30u_{ij}\frac{1}{2\delta x}(u_{i+1,j} - u_{i-1,j})$.

```

165 function ut=BurgersPDE(t,u,x)
166     dx=diff(x(1:2)); % microscale spacing
167     i=2:size(u,1)-1; % interior points in patches
168     ut=nan(size(u)); % preallocate storage
169     ut(i,:)=diff(u,2)/dx^2 ...
170         -30*u(i,:).*(u(i+1,:)-u(i-1,:))/(2*dx);
171 end

```

3.2.3 The code to make patches and interpolation

Set one edge-value to compute by interpolation if not specified by the user.
 Store in the struct.

```

192 if nargin<8, nEdge=1; end
193 if nEdge>1, error('multi-edge-value interp not yet implemented'), end
194 if 2*nEdge+1>nSubP, error('too many edge values requested'), end
195 patches.nEdge=nEdge;

```

First, store the pointer to the time derivative function in the struct.

```

202 patches.fun=fun;

```

Second, store the order of interpolation that is to provide the values for the inter-patch coupling conditions. Spectral coupling is ordCC of 0 and -1.

```

210 if (ordCC<-1) | ~(floor(ordCC)==ordCC)
211     error('ordCC out of allowed range integer>-2')
212 end

```

For odd `ordCC` do interpolation based upon odd neighbouring patches as is useful for staggered grids.

```
219 patches.alt=mod(ordCC,2);
220 ordCC=ordCC+patches.alt;
221 patches.ordCC=ordCC;
```

Check for staggered grid and periodic case.

```
227 if patches.alt & (mod(nPatch,2)==1)
228     error('Require an even number of patches for staggered grid')
229 end
```

Might as well precompute the weightings for the interpolation of field values for coupling. (Could sometime extend to coupling via derivative values.)

```
237 patches.Cwtsr=zeros(ordCC,1);
238 if patches.alt % eqn (7) in \cite{Cao2014a}
239     patches.Cwtsr(1:2:ordCC)=[1 ...
240         cumprod((ratio^2-(1:2:(ordCC-2)).^2)/4)./ ...
241         factorial(2*(1:(ordCC/2-1)))];
242     patches.Cwtsr(2:2:ordCC)=[ratio/2 ...
243         cumprod((ratio^2-(1:2:(ordCC-2)).^2)/4)./ ...
244         factorial(2*(1:(ordCC/2-1))+1)*ratio/2];
245 else %
246     patches.Cwtsr(1:2:ordCC)=(cumprod(ratio^2- ...
247         (((1:(ordCC/2))-1)).^2)./factorial(2*(1:(ordCC/2))-1)/ratio);
248     patches.Cwtsr(2:2:ordCC)=(cumprod(ratio^2- ...
249         (((1:(ordCC/2))-1)).^2)./factorial(2*(1:(ordCC/2))));
250 end
251 patches.Cwtsl=(-1).^((1:ordCC)'-patches.alt).*patches.Cwtsr;
```

Third, set the centre of the patches in a the macroscale grid of patches assuming periodic macroscale domain.

```
258 X=linspace(Xlim(1),Xlim(2),nPatch+1);
259 X=X(1:nPatch)+diff(X)/2;
260 DX=X(2)-X(1);
```

Construct the microscale in each patch, assuming Dirichlet patch edges, and a half-patch length of $\text{ratio} \cdot \text{DX}$.

```
268 if mod(nSubP,2)==0, error('configPatches1: nSubP must be odd'), end
269 i0=(nSubP+1)/2;
270 dx=ratio*DX/(i0-1);
271 patches.x=bsxfun(@plus,dx*(-i0+1:i0-1)',X); % micro-grid
272 end% function
```

Fin.

3.3 patchSmooth1(): interface to time integrators

Section contents

3.3.1 Introduction	45
------------------------------	----

3.3.1 Introduction

To simulate in time with spatial patches we often need to interface a user's time derivative function with time integration routines such as `ode15s` or `PIRK2`. This function provides an interface. It assumes that the sub-patch structure is *smooth* so that the patch centre-values are sensible macroscale variables, and patch edge values are determined by macroscale interpolation of the patch-centre values. Communicate patch-design variables to this function using the previously established global struct `patches` (Section 3.2).

```

25 function dudt=patchSmooth1(t,u)
26 global patches

```

Input

- `u` is a vector of length `nSubP · nPatch · nVars` where there are `nVars` field values at each of the points in the `nSubP × nPatch` grid.
- `t` is the current time to be passed to the user's time derivative function.
- `patches` a struct set by `configPatches1()` with the following information used here.
 - `.fun` is the name of the user's function `fun(t,u,x)` that computes the time derivatives on the patchy lattice. The array `u` has size `nSubP × nPatch × nVars`. Time derivatives must be computed into the same sized array, although herein the patch edge values are overwritten by zeros.
 - `.x` is `nSubP × nPatch` array of the spatial locations x_{ij} of the microscale grid points in every patch. Currently it *must* be an equi-spaced lattice on both macro- and microscale.

Output

- `dudt` is `nSubP · nPatch · nVars` vector of time derivatives, but with patch edge values set to zero.

Reshape the fields `u` as a 2/3D-array, and sets the edge values from macroscale interpolation of centre-patch values. Section 3.4 describes `patchEdgeInt1()`.

```

76 u=patchEdgeInt1(u);

```

Ask the user function for the time derivatives computed in the array, overwrite its edge values with the dummy value of zero, then return to a to the user/integrator as column vector.

```

86 dudt=patches.fun(t,u,patches.x);
87 dudt([1 end],:,:)=0;
88 dudt=reshape(dudt,[],1);

```

Fin.

3.4 patchEdgeInt1(): sets edge values from interpolation over the macroscale

Section contents

3.4.1 Introduction 46

3.4.1 Introduction

Couples 1D patches across 1D space by computing their edge values from macroscale interpolation of either the mid-patch value or the patch-core average. This function is primarily used by `patchSmooth1()` but is also useful for user graphics. A spatially discrete system could be integrated in time via the patch or gap-tooth scheme (Roberts & Kevrekidis 2007). Assumes that the core averages are in some sense *smooth* so that these averages are sensible macroscale variables. Then patch edge values are determined by macroscale interpolation of the core averages (Bunder et al. 2017). Communicate patch-design variables via the global struct `patches`.

```

27 function u=patchEdgeInt1(u)
28 global patches

```

Input

- `u` is a vector of length $nSubP \cdot nPatch \cdot nVars$ where there are $nVars$ field values at each of the points in the $nSubP \times nPatch$ grid.
- `patches` a struct set by `configPatches1()` which includes the following.
 - `.x` is $nSubP \times nPatch$ array of the spatial locations x_{ij} of the microscale grid points in every patch. Currently it *must* be an equi-spaced lattice on both macro- and microscale.
 - `.ordCC` is order of interpolation integer ≥ -1 .
 - `.alt` in $\{0,1\}$ is one for staggered grid (alternating) interpolation.
 - `.Cwtsr` and `.Cwtsl` define the coupling.

Output

- `u` is $nSubP \times nPatch \times nVars$ 2/3D array of the fields with edge values set by interpolation of patch core averages.

Determine the sizes of things. Any error arising in the reshape indicates `u` has the wrong size.


```

66 [nSubP,nPatch] = size(patches.x);
67 nVars = round(numel(u)/numel(patches.x));
68 if numel(u)~=nSubP*nPatch*nVars
69     nSubP=nSubP, nPatch=nPatch, nVars=nVars, sizeu=size(u)
70 end
71 u = reshape(u,nSubP,nPatch,nVars);

```

Compute lattice sizes from inside the patches as the edge values may be NaNs, etc.

```

78 dx = patches.x(3,1)-patches.x(2,1);
79 DX = patches.x(2,2)-patches.x(2,1);

```

If the user has not defined the patch core, then we assume it to be a single point in the middle of the patch. For `patches.nCore` $\neq 1$ the half width ratio is reduced, as described by [Bunder et al. \(2017\)](#).

```

88 if ~isfield(patches,'nCore')
89     patches.nCore = 1;
90 end
91 r = dx*(nSubP-1)/2/DX*(nSubP - patches.nCore)/(nSubP - 1);

```

For the moment assume the physical domain is macroscale periodic so that the coupling formulas are simplest. Should eventually cater for periodic, odd-mid-gap, even-mid-gap, even-mid-patch, Dirichlet, Neumann etc. These index vectors point to patches and their two immediate neighbours.

```

102 j = 1:nPatch; jp = mod(j,nPatch)+1; jm = mod(j-2,nPatch)+1;

```

Calculate centre of each patch and the surrounding core (`nSubP` and `nCore` are both odd).

```

109 i0 = round((nSubP+1)/2);
110 c = round((patches.nCore-1)/2);

```

Lagrange interpolation gives patch-edge values Consequently, compute centred differences of the patch core averages for the macro-interpolation of all fields. Assumes the domain is macro-periodic.

```

120 if patches.ordCC>0 % then non-spectral interpolation
121     if patches.EnsAve
122         uCore = sum(mean(u((i0-c):(i0+c),j,:),3),1)';
123         dmu = zeros(patches.ordCC,nPatch);
124     else
125         uCore = reshape(sum(u((i0-c):(i0+c),j,:),1),nPatch,nVars);
126         dmu = zeros(patches.ordCC,nPatch,nVars);
127     end;
128     if patches.alt % use only odd numbered neighbours
129         dmu(1,.,.) = (uCore(jp,.)+uCore(jm,.))/2; % \mu
130         dmu(2,.,.) = (uCore(jp,.)-uCore(jm,.)); % \delta
131         jp = jp(jp); jm = jm(jm); % increase shifts to \pm 2
132     else % standard
133         dmu(1,j,.) = (uCore(jp,.)-uCore(jm,.))/2; % \mu\delta

```

```

134     dmu(2,j,:) = (uCore(jp,:)-2*uCore(j,:)+uCore(jm,:))/2; % \delta^2
135     end% if odd/even

```

Recursively take δ^2 of these to form higher order centred differences (could unroll a little to cater for two in parallel).

```

143     for k = 3:patches.ordCC
144         dmu(k,,:,) = dmu(k-2,jp,:)-2*dmu(k-2,j,:)+dmu(k-2,jm,:);
145     end

```

Interpolate macro-values to be Dirichlet edge values for each patch ([Roberts & Kevrekidis 2007](#), [Bunder et al. 2017](#)), using weights computed in `configPatches1()`. Here interpolate to specified order.

```

154     if patches.EnsAve
155         u(nSubP,j,:) = repmat(uCore(j)'*(1-patches.alt) ...
156             +sum(bsxfun(@times,patches.Cwtsr,dmu)), [1,1,nVars]) ...
157             -sum(u((nSubP-patches.nCore+1):(nSubP-1),:,:),1);
158         u(1,j,:) = repmat(uCore(j)'*(1-patches.alt) ...
159             +sum(bsxfun(@times,patches.Cwtsl,dmu)), [1,1,nVars]) ...
160             -sum(u(2:patches.nCore,:,:),1);
161     else
162         u(nSubP,j,:) = uCore(j,:)*(1-patches.alt) ...
163             + reshape(-sum(u((nSubP-patches.nCore+1):(nSubP-1),j,:),1) ...
164                 +sum(bsxfun(@times,patches.Cwtsr,dmu)), nPatch,nVars);
165         u(1,j,:) = uCore(j,:)*(1-patches.alt) ...
166             + reshape(-sum(u(2:patches.nCore,j,:),1) ...
167                 +sum(bsxfun(@times,patches.Cwtsl,dmu)), nPatch,nVars);
168     end;

```

Case of spectral interpolation Assumes the domain is macro-periodic. As the macroscale fields are N -periodic, the macroscale Fourier transform writes the centre-patch values as $U_j = \sum_k C_k e^{ik2\pi j/N}$. Then the edge-patch values $U_{j\pm r} = \sum_k C_k e^{ik2\pi/N(j\pm r)} = \sum_k C'_k e^{ik2\pi j/N}$ where $C'_k = C_k e^{ikr2\pi/N}$. For `nPatch` patches we resolve ‘wavenumbers’ $|k| < \text{nPatch}/2$, so set row vector $\mathbf{k}s = k2\pi/N$ for ‘wavenumbers’ $k = (0, 1, \dots, k_{\max}, -k_{\max}, \dots, -1)$ for odd N , and $k = (0, 1, \dots, k_{\max}, \pm(k_{\max} + 1), -k_{\max}, \dots, -1)$ for even N .

```

186     else% spectral interpolation

```

Deal with staggered grid by doubling the number of fields and halving the number of patches (`configPatches1()` tests that there are an even number of patches). Then the patch-ratio is effectively halved. The patch edges are near the middle of the gaps and swapped.

```

196     if patches.alt % transform by doubling the number of fields
197         v = nan(size(u)); % currently to restore the shape of u
198         u = cat(3,u(:,1:2:nPatch,:),u(:,2:2:nPatch,:));
199         altShift = reshape(0.5*[ones(nVars,1);-ones(nVars,1)],1,1,[]);
200         iV = [nVars+1:2*nVars 1:nVars]; % scatter interp to alternate field
201         r = r/2; % ratio effectively halved
202         nPatch = nPatch/2; % halve the number of patches

```

```

203     nVars = nVars*2;    % double the number of fields
204     else % the values for standard spectral
205         altShift = 0;
206         iV = 1:nVars;
207     end

```

Now set wavenumbers.

```

213     kMax = floor((nPatch-1)/2);
214     ks = 2*pi/nPatch*(mod((0:nPatch-1)+kMax,nPatch)-kMax);

```

Test for reality of the field values, and define a function accordingly.

```

221     if imag(u(i0,:,:))==0, uclean=@(u) real(u);
222     else                    uclean=@(u) u;
223     end

```

Compute the Fourier transform of the patch centre-values for all the fields. If there are an even number of points, then zero the zig-zag mode in the FT and add it in later as cosine.

```

232     Ck = fft(u(i0,:,:));
233     if mod(nPatch,2)==0
234         Czz = Ck(1,nPatch/2+1,:)/nPatch;
235         Ck(1,nPatch/2+1,:) = 0;
236     end

```

The inverse Fourier transform gives the edge values via a shift a fraction r to the next macroscale grid point. Enforce reality when appropriate.

```

244     u(nSubP,:,iV) = uclean(ifft(bsxfun(@times,Ck ...
245         ,exp(1i*bsxfun(@times,ks,altShift+r)))));
246     u( 1,:,iV) = uclean(ifft(bsxfun(@times,Ck ...
247         ,exp(1i*bsxfun(@times,ks,altShift-r)))));

```

For an even number of patches, add in the cosine mode.

```

253     if mod(nPatch,2)==0
254         cosr = cos(pi*(altShift+r+(0:nPatch-1)));
255         u(nSubP,:,iV) = u(nSubP,:,iV)+uclean(bsxfun(@times,Czz,cosr));
256         cosr = cos(pi*(altShift-r+(0:nPatch-1)));
257         u( 1,:,iV) = u( 1,:,iV)+uclean(bsxfun(@times,Czz,cosr));
258     end

```

Restore staggered grid when appropriate. Is there a better way to do this??

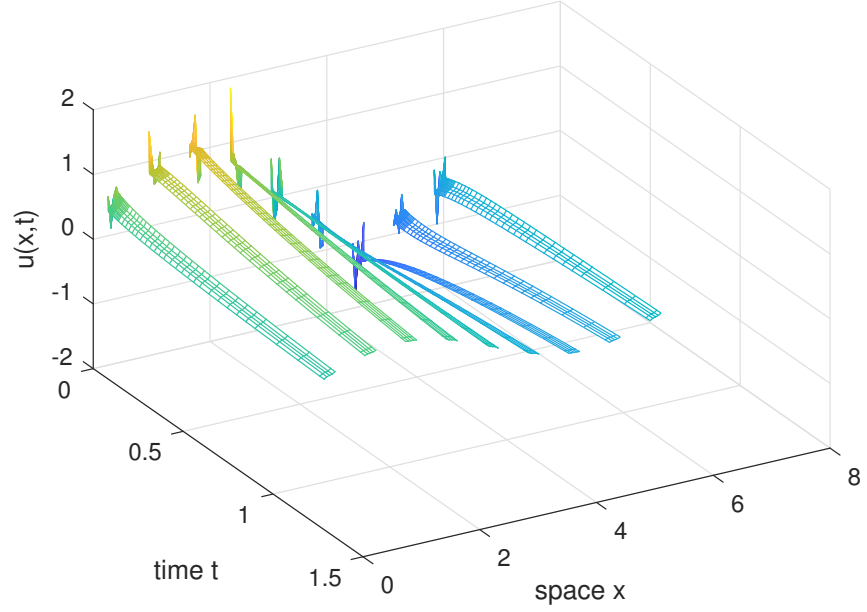
```

265     if patches.alt
266         nVars = nVars/2;  nPatch = 2*nPatch;
267         v(:,1:2:nPatch,:) = u(:, :, 1:nVars);
268         v(:,2:2:nPatch,:) = u(:, :, nVars+1:2*nVars);
269         u = v;
270     end
271     end% if spectral

```

Fin, returning the 2/3D array of field values.

Figure 3.2: the diffusing field $u(x, t)$ in the patch (gap-tooth) scheme applied to microscale heterogeneous diffusion (Section 3.5).



3.5 homogenisationExample: simulate heterogeneous diffusion in 1D on patches

Section contents

3.5.1	Script to simulate via stiff or projective integration . .	51
3.5.2	heteroDiff(): heterogeneous diffusion	53
3.5.3	heteroBurst(): a burst of heterogeneous diffusion . .	54

Figure 3.2 shows an example simulation in time generated by the patch scheme function applied to heterogeneous diffusion. That such simulations of heterogeneous diffusion makes valid predictions was established by Bunder et al. (2017) who proved that the scheme is accurate when the number of points in a patch is one more than a multiple of the microscale periodicity.

The first part of the script implements the following gap-tooth scheme (left-right arrows denote function recursion).

1. configPatches1
2. ode15s ↔ patchSmooth1 ↔ heteroDiff
3. process results

Consider a lattice of values $u_i(t)$, with lattice spacing dx , and governed by the heterogeneous diffusion

$$\dot{u}_i = [c_{i-1/2}(u_{i-1} - u_i) + c_{i+1/2}(u_{i+1} - u_i)]/dx^2. \quad (3.1)$$

In this 1D space, the macroscale, homogenised, effective diffusion should be the harmonic mean of these coefficients.

3.5.1 Script to simulate via stiff or projective integration

Set the desired microscale periodicity, and microscale diffusion coefficients (with subscripts shifted by a half).

```
50 clear all
51 mPeriod = 3
52 cDiff = exp(randn(mPeriod,1))
53 cHomo = 1/mean(1./cDiff)
```

Establish global data struct `patches` for heterogeneous diffusion solved on 2π -periodic domain, with nine patches, each patch of half-size 0.2. A user can add information to `patches` in order to communicate to the time derivative function. Quadratic (fourth-order) interpolation `ordCC = 4` provides values for the inter-patch coupling conditions. The odd integer `patches.nCore = 3` defines the size of the patch core (this must be larger than zero and less than `nSubP`), where a core of size zero indicates that the value in the centre of the patch gives the macroscale. The introduction of a finite width core requires a redefinition of the half-patch ratio, as described by [Bunder et al. \(2017\)](#). We evaluate the patch coupling by interpolating the core.

```
73 global patches
74 nPatch = 9
75 ratio = 0.2
76 nSubP = 2*mPeriod+1
77 Len = 2*pi;
78 ordCC = 4;
79 configPatches1(@heteroDiff,[0 Len],nan,nPatch ...
80               ,ordCC,ratio,nSubP);
```

Add to the global data struct `patches` for use by the time derivative function (for example): here include the diffusivity coefficients, repeated to fill up a patch

```
87 patches.c= repmat(cDiff,(nSubP-1)/mPeriod,1);
```

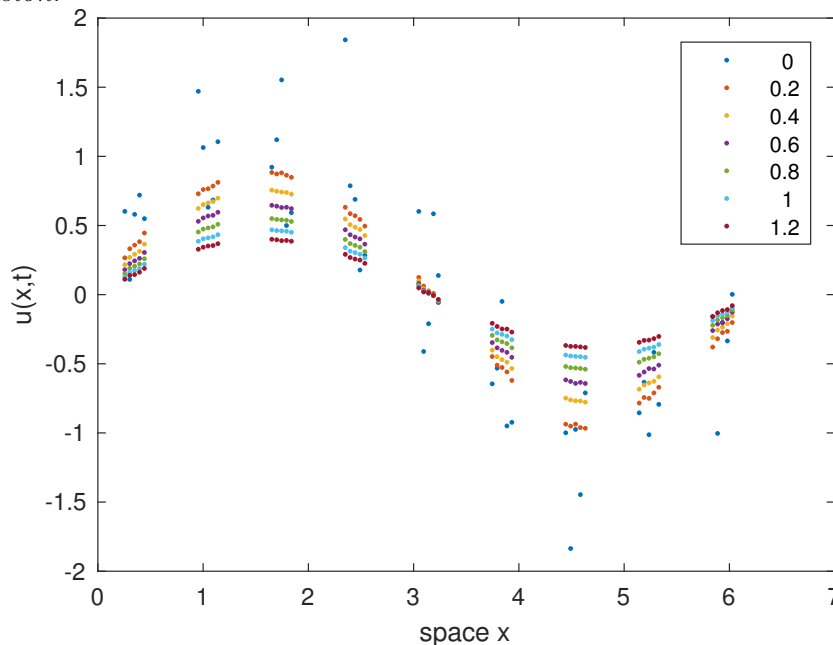
For comparison: conventional integration in time Set an initial condition, and here integrate forward in time using a standard method for stiff systems—because of the simplicity of linear problems this method works quite efficiently here. Integrate the interface `patchSmooth1` ([Section 3.3](#)) to the microscale differential equations.

```
100 u0 = sin(patches.x)+0.4*randn(nSubP,nPatch);
101 [ts,ucts] = ode15s(@patchSmooth1, [0 2/cHomo], u0(:));
102 ucts = reshape(ucts,length(ts),length(patches.x(:)),[]);
```

Plot the simulation in [Figure 3.2](#).

```
109 figure(1),clf
110 xs = patches.x; xs([1 end],:) = nan;
111 mesh(ts,xs(:),ucts'), view(60,40)
112 xlabel('time t'), ylabel('space x'), zlabel('u(x,t)')
```

Figure 3.3: field $u(x, t)$ shows basic projective integration of patches of heterogeneous diffusion: different colours correspond to the times in the legend. This field solution displays some fine scale heterogeneity due to the heterogeneous diffusion.



```

113 set(gcf,'PaperPosition',[0 0 14 10]);% cm
114 print('-depsc2','homogenisationCtsU')

```

Use projective integration in time Now take `patchSmooth1`, the interface to the time derivatives, and wrap around it the projective integration `PIRK2` (Section 2.2), of bursts of simulation from `heteroBurst` (Section 3.7.3), as illustrated by Figure 3.3.

This second part of the script implements the following design, where the micro-integrator could be, for example, `ode45` or `rk2int`.

1. `configPatches1` (done in first part)
2. `PIRK2` \leftrightarrow `heteroBurst` \leftrightarrow micro-integrator \leftrightarrow `patchSmooth1` \leftrightarrow `heteroDiff`
3. process results

Mark that edge of patches are not to be used in the projective extrapolation by setting initial values to NaN.

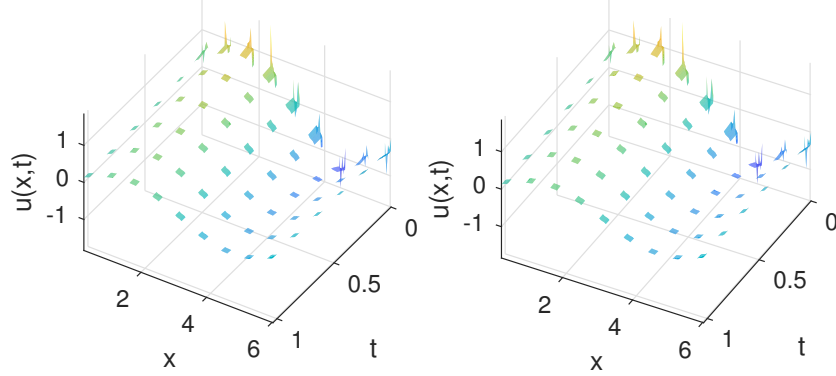
```

146 u0([1 end], :) = nan;

```

Set the desired macro- and microscale time-steps over the time domain: the macroscale step is in proportion to the effective mean diffusion time on the macroscale; the burst time is proportional to the intra-patch effective diffusion time; and lastly, the microscale time-step is proportional to the diffusion time between adjacent points in the microscale lattice.

Figure 3.4: stereo pair of the field $u(x,t)$ during each of the microscale bursts used in the projective integration.



```

158 ts = linspace(0,2/cHomo,7)
159 bT = 3*( ratio*Len/nPatch )^2/cHomo
160 addpath(' ../ProjInt', ' ../SandpitPlay/RKint')
161 [us,tss,uss] = PIRK2(@heteroBurst, ts, u0(:), bT);

```

Plot the macroscale predictions to draw [Figure 3.3](#).

```

168 figure(2),clf
169 plot(xs(:),us','.')
170 ylabel('u(x,t)'), xlabel('space x')
171 legend(num2str(ts',3))
172 set(gcf,'PaperPosition',[0 0 14 10]);% cm
173 print('-depsc2','homogenisationU')

```

Also plot a surface detailing the microscale bursts as shown in [Figure 3.4](#).

```

186 figure(3),clf
187 for k = 1:2, subplot(1,2,k)
188     surf(tss,xs(:),uss', 'EdgeColor','none')
189     ylabel('x'), xlabel('t'), zlabel('u(x,t)')
190     axis tight, view(126-4*k,45)
191 end
192 set(gcf,'PaperPosition',[0 0 14 6]);% cm
193 print('-depsc2','homogenisationMicro')

```

End of the script.

3.5.2 heteroDiff(): heterogeneous diffusion

This function codes the lattice heterogeneous diffusion inside the patches. For 2D input arrays u and x (via edge-value interpolation of `patchSmooth1`, [Section 3.3](#)), computes the time derivative (3.2) at each point in the interior of a patch, output in ut . The column vector (or possibly array) of diffusion coefficients c_i have previously been stored in struct `patches`.

```

212 function ut = heteroDiff(t,u,x)
213     global patches

```

```

214     dx = diff(x(2:3)); % space step
215     i = 2:size(u,1)-1; % interior points in a patch
216     ut = nan(size(u)); % preallocate output array
217     ut(i, :, :) = diff(patches.c.*diff(u))/dx^2;
218 end% function

```

3.5.3 heteroBurst(): a burst of heterogeneous diffusion

This code integrates in time the derivatives computed by `heteroDiff` from within the patch coupling of `patchSmooth1`. Try `ode23`, although `ode45` may give smoother results.

```

232 function [ts, ucts] = heteroBurst(ti, ui, bT)
233     [ts, ucts] = ode23(@patchSmooth1, [ti ti+bT], ui(:));
234 end

```

Fin.

3.6 BurgersExample: simulate Burgers' PDE on patches

Section contents

3.6.1	Script code to simulate a microscale space-time map	54
	Alternatively use projective integration	56
3.6.2	<code>burgersMap()</code> : discretise the PDE microscale	57
3.6.3	<code>burgerBurst()</code> : code a burst of the patch map	57

Figure 3.1 shows an example simulation in time generated by the patch scheme function applied to Burgers' PDE. This code similarly applies the Equation-Free functions to a microscale space-time map (Figure 3.5), a map that happens to be derived as a microscale space-time discretisation of Burgers' PDE. Then this example applies projective integration to simulate further in time.

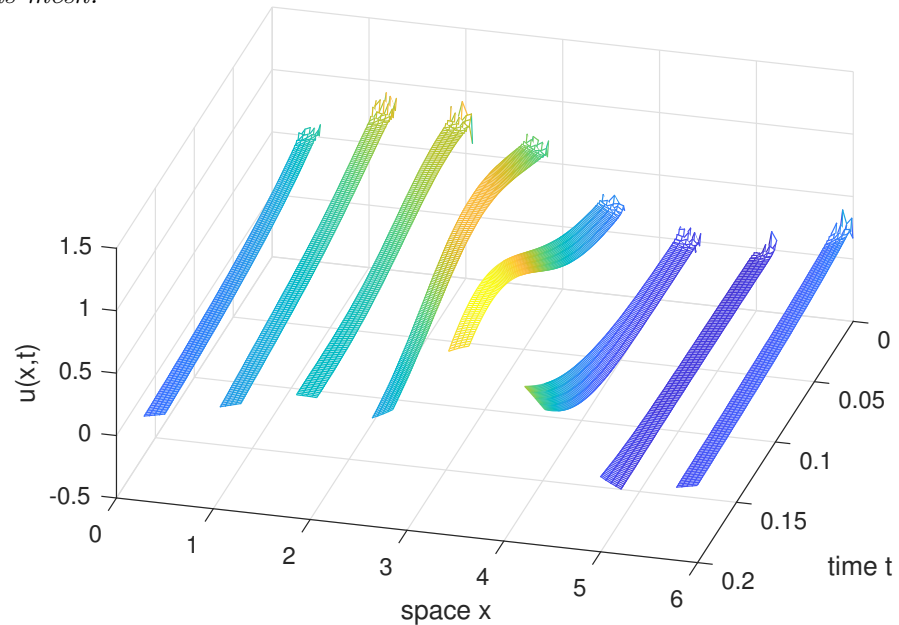
The first part of the script implements the following gap-tooth scheme (left-right arrows denote function recursion).

1. `configPatches1`
2. `burgerBurst` \leftrightarrow `patchSmooth1` \leftrightarrow `burgersMap`
3. process results

3.6.1 Script code to simulate a microscale space-time map

Establish global data struct for the Burgers' map (Section 3.6.2) solved on 2π -periodic domain, with eight patches, each patch of half-size ratio 0.2, with seven points within each patch, and say fourth-order interpolation provides edge-values that couple the patches.

Figure 3.5: a short time simulation of the Burgers' map (Section 3.6.2) on patches in space. It requires many very small time-steps only just visible in this mesh.



```

48 clear all
49 global patches
50 nPatch = 8
51 ratio = 0.2
52 nSubP = 7
53 interpOrd = 4
54 Len = 2*pi
55 configPatches1(@burgersMap,[0 Len],nan,nPatch,interpOrd,ratio,nSubP);

```

Set an initial condition, and simulate a burst of the microscale space-time map over a time 0.2 using the function `burgerBurst()` (Section 3.6.3).

```

63 u0 = 0.4*(1+sin(patches.x))+0.1*randn(size(patches.x));
64 [ts,us] = burgerBurst(0,u0,0.2);

```

Plot the simulation. Use only the microscale values interior to the patches by setting the edges to `nan` in order to leave gaps.

```

72 figure(1),clf
73 xs = patches.x; xs([1 end],:) = nan;
74 mesh(ts,xs(:,),us')
75 xlabel('time t'), ylabel('space x'), zlabel('u(x,t)')
76 view(105,45)

```

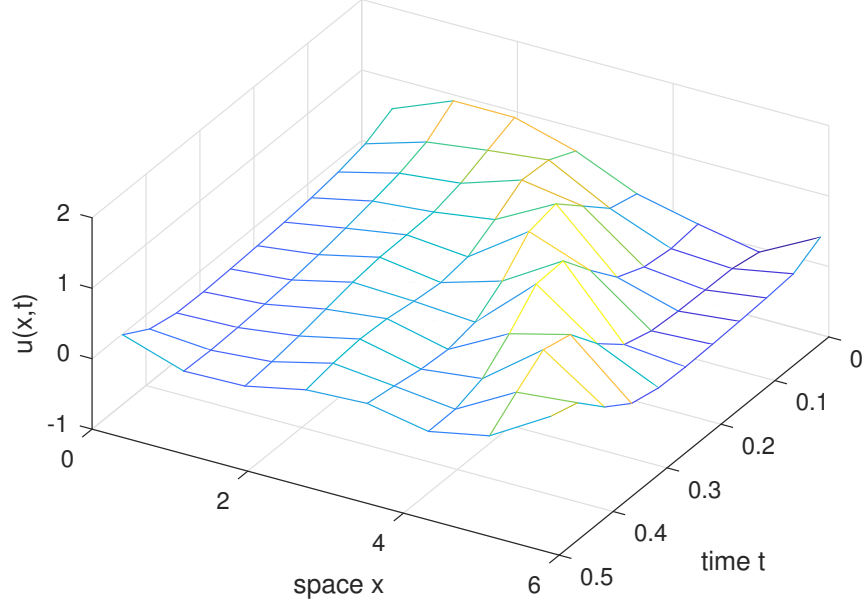
Save the plot to file to form Figure 3.5.

```

82 set(gcf,'paperposition',[0 0 14 10])
83 print('-depsc2','BurgersMapU')

```

Figure 3.6: macroscale space-time field $u(x,t)$ in a basic projective integration of the patch scheme applied to the microscale Burgers' map.



Alternatively use projective integration

Around the microscale burst `burgerBurst()`, wrap the projective integration function `PIRK2()` of Section 2.2. Figure 3.6 shows the macroscale prediction of the patch centre values on macroscale time-steps.

This second part of the script implements the following design.

1. `configPatches1` (done in first part)
2. `PIRK2` \leftrightarrow `burgerBurst` \leftrightarrow `patchSmooth1` \leftrightarrow `burgersMap`
3. process results

Mark that edge-values of patches are not to be used in the projective extrapolation by setting initial values to NaN.

```
115 u0([1 end], :) = nan;
```

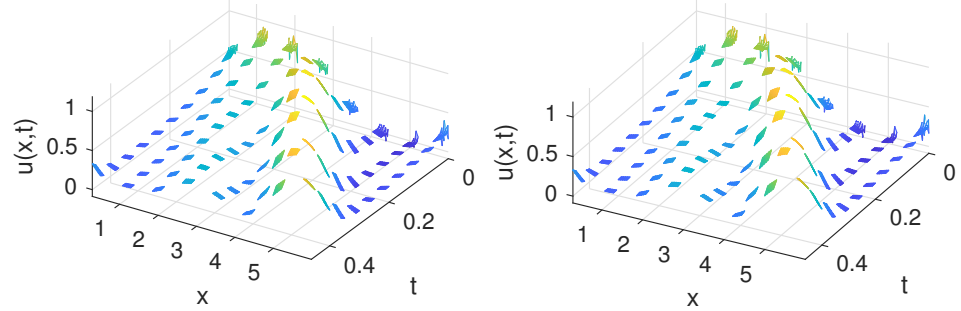
Set the desired macroscale time-steps, and microscale burst length over the time domain. Then projectively integrate in time using `PIRK2()` which is (roughly) second-order accurate in the macroscale time-step.

```
124 ts = linspace(0,0.5,11);
125 bT = 3*(ratio*Len/nPatch/(nSubP/2-1))^2
126 addpath(' ../ProjInt')
127 [us,tss,uss] = PIRK2(@burgerBurst,ts,u0(:),bT);
```

Plot and save the macroscale predictions of the mid-patch values to give the macroscale mesh-surface of Figure 3.6 that shows a progressing wave solution.

```
136 figure(2),clf
137 midP = (nSubP+1)/2;
138 mesh(ts,xs(midP,:),us(:,midP:nSubP:end)')
```

Figure 3.7: the field $u(x,t)$ during each of the microscale bursts used in the projective integration. View this stereo pair cross-eyed.



```

139 xlabel('time t'), ylabel('space x'), zlabel('u(x,t)')
140 view(120,50)
141 set(gcf,'paperposition',[0 0 14 10])
142 print('-depsc2','BurgersU')

```

Then plot and save the microscale mesh of the microscale bursts shown in Figure 3.7 (a stereo pair). The details of the fine microscale mesh are almost invisible.

```

157 figure(3), clf
158 for k = 1:2, subplot(2,2,k)
159     mesh(tss,xs(:),uss')
160     ylabel('x'), xlabel('t'), zlabel('u(x,t)')
161     axis tight, view(126-4*k,50)
162 end
163 set(gcf,'paperposition',[0 0 17 12])
164 print('-depsc2','BurgersMicro')

```

3.6.2 burgersMap(): discretise the PDE microscale

This function codes the microscale Euler integration map of the lattice differential equations inside the patches. Only the patch-interior values mapped (patchSmooth1() overrides the edge-values anyway).

```

181 function u = burgersMap(t,u,x)
182     dx = diff(x(2:3));
183     dt = dx^2/2;
184     i = 2:size(u,1)-1;
185     u(i,:) = u(i,:) + dt*( diff(u,2)/dx^2 ...
186         -20*u(i,:).*(u(i+1,:)-u(i-1,:))/(2*dx) );
187 end

```

3.6.3 burgerBurst(): code a burst of the patch map

```

197 function [ts, us] = burgerBurst(ti, ui, bT)

```

First find and set the number of microscale time-steps.

```

203     global patches
204     dt = diff(patches.x(2:3))^2/2;
205     ndt = ceil(bT/dt -0.2);
206     ts = ti+(0:ndt)*dt;

```

Use `patchSmooth1()` (Section 3.3) to apply the microscale map over all time-steps in the burst. The `patchSmooth1()` interface provides the interpolated edge-values of each patch. Store the results in rows to be consistent with ODE and projective integrators.

```

216     us = nan(ndt+1,numel(ui));
217     us(1,:) = reshape(ui,1,[]);
218     for j = 1:ndt
219         ui = patchSmooth1(ts(j),ui);
220         us(j+1,:) = reshape(ui,1,[]);
221     end

```

Linearly interpolate (extrapolate) to get the field values at the precise final time of the burst. Then return.

```

228     ts(ndt+1) = ti+bT;
229     us(ndt+1,:) = us(ndt,:) ...
230         + diff(ts(ndt:ndt+1))/dt*diff(us(ndt:ndt+1,:));
231 end

```

Fin.

3.7 ensembleAverageExample: simulate an ensemble of solutions for heterogeneous diffusion in 1D on patches

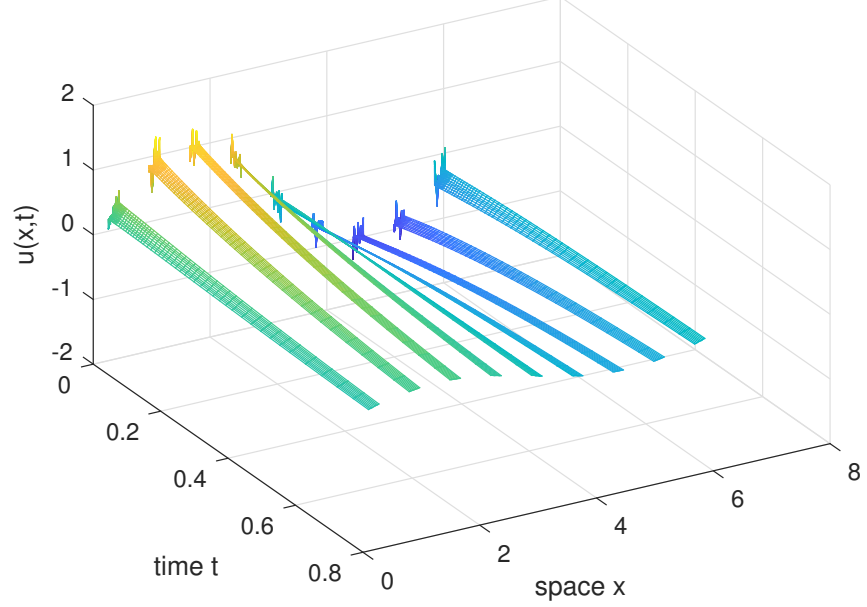
Section contents

3.7.1	Script to simulate via stiff or projective integration . . .	59
3.7.2	<code>heteroDiff()</code> : heterogeneous diffusion	63
3.7.3	<code>heteroBurst()</code> : a burst of heterogeneous diffusion . . .	63

This example is an extension of the homogenisation example of Section 3.5 for heterogeneous diffusion. In cases where the periodicity of the heterogeneous diffusion is known, then Section 3.5 provides a efficient patch dynamics simulation. However, if the diffusion is not completely known or is stochastic, then we cannot choose ideal patch and core sizes as described by Bunder et al. (2017) and applied in Section 3.5. In this case, Bunder et al. (2017) recommend constructing an ensemble of diffusivity configurations and then computing an ensemble of field solutions, finally averaging over the ensemble of fields to obtain the ensemble averaged field solution.

For a first comparison, we present a very similar example to that presented in Section 3.5, but whereas Section 3.5 simulates using only one diffusivity configuration, here we simulate over an ensemble. For example, Figure 3.8 is similar to Figure 3.2, but the former is an ensemble average of an ensemble

Figure 3.8: the diffusing field $u(x,t)$ in the patch (gap-tooth) scheme applied to microscale heterogeneous diffusion with an ensemble average. The ensemble average smooths out the heterogeneous diffusion.



of eight different simulations with different diffusivity configurations and the latter is simulated from just one diffusivity configuration. The main difference between these two is that the average over the ensemble removes any heterogeneity in the solution.

Much of this script is similar to that of [Section 3.5](#), but with some additions to manage the ensemble. The first part of the script implements the following gap-tooth scheme (left-right arrows denote function recursion).

1. configPatches1
2. ode15s \leftrightarrow patchSmooth1 \leftrightarrow heteroDiff
3. process results

Consider a lattice of values $u_i(t)$, with lattice spacing dx , and governed by the heterogeneous diffusion

$$\dot{u}_i = [c_{i-1/2}(u_{i-1} - u_i) + c_{i+1/2}(u_{i+1} - u_i)]/dx^2. \quad (3.2)$$

In this 1D space, the macroscale, homogenised, effective diffusion should be the harmonic mean of these coefficients. But we do not have full knowledge of these coefficients.

3.7.1 Script to simulate via stiff or projective integration

Say we only know four diffusivities in our diffusion problem, as defined here (which are the same as those given in [Section 3.5](#)).

```

75 clear all
76 mPeriod = 4
77 rng('default'); rng(1);

```

```

78 cDiff = exp(4*rand(mPeriod,1))
79 cHomo = 1/mean(1./cDiff)

```

The chosen parameters are the same as [Section 3.5](#), but here we also introduce the Boolean `patches.EnsAve` which determines whether or not we construct an ensemble average of diffusivity configurations. Setting `patches.EnsAve=0` simulates the same problem as in [Section 3.5](#).

```

91 global patches
92 nPatch = 9
93 ratio = 0.2
94 nSubP = 11
95 Len = 2*pi;
96 ordCC = 4;
97 patches.nCore = 3;
98 patches.ratio = ratio*(nSubP - patches.nCore)/(nSubP - 1);
99 configPatches1(@heteroDiff,[0 Len],nan,nPatch ...
100               ,ordCC,patches.ratio,nSubP);
101 patches.EnsAve = 1;

```

In the case of ensemble averaging, `nVars` is the size of the ensemble (for the case of no ensemble averaging `nVars` is the number of different field variables, which in this example is `nVars = 1`) and we use the ensemble described by [Bunder et al. \(2017\)](#) which includes all reflected and translated configurations of `patches.cDiff`. We must increase the size of the diffusivity matrix to $(nSubP-1) \times nPatch \times nVars$.

```

116 patches.cDiff = cDiff((mod(round(patches.x(1:(end-1)),:) ...
117   /(patches.x(2)-patches.x(1))-0.5),mPeriod)+1));
118 if patches.EnsAve
119     nVars = mPeriod+(mPeriod>2)*mPeriod;
120     patches.cDiff = repmat(patches.cDiff,[1,1,nVars]);
121     for sx = 2:mPeriod
122         patches.cDiff(:,:,sx) = circshift( ...
123             patches.cDiff(:,:,sx-1),[sx-1,0]);
124     end;
125     if nVars>2
126         patches.cDiff(:,:,mPeriod+1:end) = flipud( ...
127             patches.cDiff(:,:,1:mPeriod));
128     end;
129 end

```

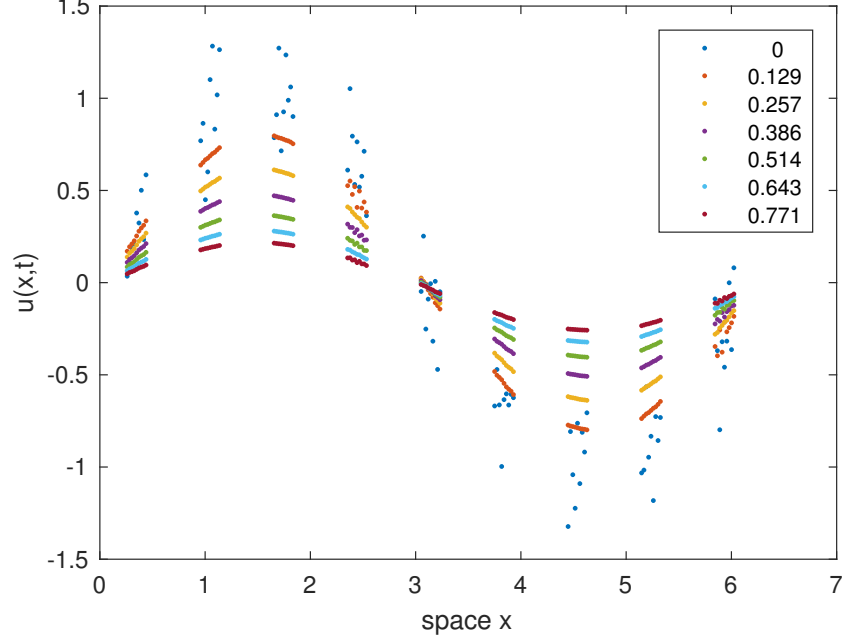
Conventional integration in time Set an initial condition, and here integrate forward in time using a standard method for stiff systems. Integrate the interface `patchSmooth1` ([Section 3.3](#)) to the microscale differential equations.

```

140 u0 = sin(patches.x)+0.2*randn(nSubP,nPatch);
141 if patches.EnsAve
142     u0 = repmat(u0,[1,1,nVars]);
143 end

```

Figure 3.9: field $u(x,t)$ shows basic projective integration of patches of heterogeneous diffusion with an ensemble average: different colours correspond to the times in the legend. Once transients have decayed, this field solution is smooth due to the ensemble average.



```

144 [ts,ucts] = ode15s(@patchSmooth1, [0 2/cHomo], u0(:));
145 ucts = reshape(ucts,length(ts),length(patches.x(:)),[]);

```

Plot the ensemble averaged simulation in [Figure 3.8](#).

```

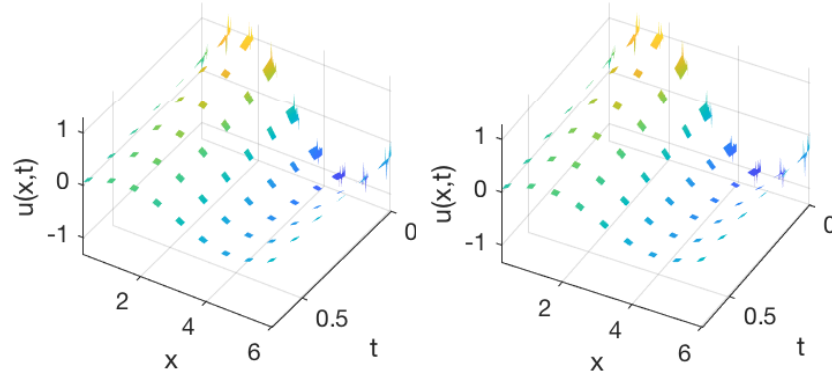
153 if patches.EnsAve % calculate the ensemble average
154     uctsAve = mean(ucts,3);
155 else
156     uctsAve = ucts;
157 end
158 figure(1),clf
159 xs = patches.x; xs([1 end],:) = nan;
160 mesh(ts,xs(:),uctsAve'), view(60,40)
161 xlabel('time t'), ylabel('space x'), zlabel('u(x,t)')
162 set(gcf,'PaperPosition',[0 0 14 10]);% cm
163 print('-depsc2','ensAveExCtsU')

```

Use projective integration in time Now take `patchSmooth1`, the interface to the time derivatives, and wrap around it the projective integration `PIRK2` ([Section 2.2](#)), of bursts of simulation from `heteroBurst` ([Section 3.7.3](#)), as illustrated by [Figure 3.9](#). The rest of this code follows that of [Section 3.5](#), but as we now evaluate an ensemble of field solutions, our final step is always an ensemble average.

Mark that edge of patches are not to be used in the projective extrapolation by setting initial values to NaN.

Figure 3.10: stereo pair of ensemble averaged fields $u(x,t)$ during each of the microscale bursts used in the projective integration.



```
190 u0([1 end],:) = nan;
```

Set the desired macro- and microscale time-steps over the time domain.

```
197 ts = linspace(0,2/cHomo,7)
198 bT = 3*( ratio*Len/nPatch )^2/cHomo
199 addpath(' ../ProjInt', ' ../SandpitPlay/RKint')
200 [us,tss,uss] = PIRK2(@heteroBurst, ts, u0(:), bT);
```

Plot an average of the ensemble of macroscale predictions to draw [Figure 3.9](#).

```
207 usAve = mean(reshape(us,size(us,1),length(xs(:)),nVars),3);
208 ussAve = mean(reshape(uss,length(tss),length(xs(:)),nVars),3);
209 figure(2),clf
210 plot(xs(:),usAve','.')
211 ylabel('u(x,t)'), xlabel('space x')
212 legend(num2str(ts',3))
213 set(gcf,'PaperPosition',[0 0 14 10]);% cm
214 print('-depsc2','ensAveExU')
```

Also plot a surface detailing the ensemble average microscale bursts as shown [Figure 3.10](#).

```
229 figure(3),clf
230 for k = 1:2, subplot(1,2,k)
231     surf(tss,xs(:),ussAve', 'EdgeColor','none')
232     ylabel('x'), xlabel('t'), zlabel('u(x,t)')
233     axis tight, view(126-4*k,45)
234 end
235 set(gcf,'PaperPosition',[0 0 14 6]);% cm
236 print('-depsc2','ensAveExMicro')
```

End of the script.

3.7.2 heteroDiff(): heterogeneous diffusion

This function codes the PDE of the heterogeneous diffusion inside the patches, coding it as ODEs on a microscale lattice. For 2D input arrays u and x (via edge-value interpolation of `patchSmooth1`, [Section 3.3](#)), computes the time derivative (3.2) at each point in the interior of a patch, output in ut . The column vector (or possibly array) of diffusion coefficients c_i have previously been stored in struct `patches`.

```

255 function ut = heteroDiff(t,u,x)
256     global patches
257     dx = diff(x(2:3)); % space step
258     i = 2:size(u,1)-1; % interior points in a patch
259     ut = nan(size(u)); % preallocate output array
260     ut(i, :, :) = diff(patches.cDiff.*diff(u))/dx^2;
261 end% function

```

3.7.3 heteroBurst(): a burst of heterogeneous diffusion

This code integrates in time the derivatives computed by `heteroDiff` from within the patch coupling of `patchSmooth1`. Try four possibilities:

- `ode23` generates ‘noise’ that is unsightly at best and may be ruinous;
- `ode45` is similar to `ode23`, but with reduced noise;
- `ode15s` does not cater for the NaNs in some components of u ;
- `rk2int` simple specified step integrator, but may require inefficiently small time-steps.

```

284 function [ts, ucts] = heteroBurst(ti, ui, bT)
285     switch '45'
286     case '23', [ts,ucts] = ode23(@patchSmooth1,[ti ti+bT],ui(:));
287     case '45', [ts,ucts] = ode45(@patchSmooth1,[ti ti+bT],ui(:));
288     case '15s', [ts,ucts] = ode15s(@patchSmooth1,[ti ti+bT],ui(:));
289     case 'rk2', ts = linspace(ti,ti+bT,200)';
290                 ucts = rk2int(@patchSmooth1,ts,ui(:));
291     end
292 end

```

Fin.

3.8 waterWaveExample: simulate a water wave PDE on patches

Section contents

3.8.1	Script code to simulate wave systems	65
3.8.2	idealWavePDE(): ideal wave PDE	67
3.8.3	waterWavePDE(): water wave PDE	68

Figure 3.11: water depth $h(x, t)$ (above) and velocity field $u(x, t)$ (below) of the gap-tooth scheme applied to the ideal wave PDE (3.3), linearised. The microscale random component to the initial condition has long lasting effects on the simulation—but the macroscale wave still propagates.

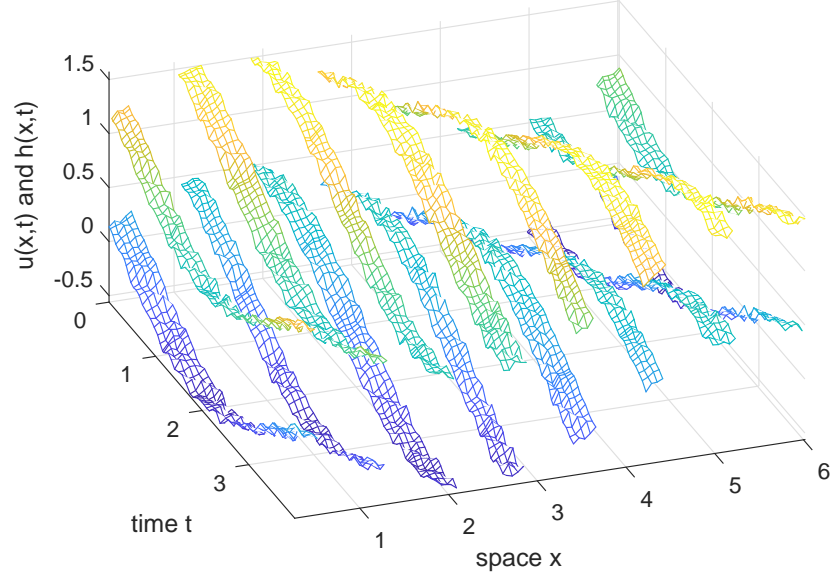


Figure 3.11 shows an example simulation in time generated by the patch scheme function applied to an ideal wave PDE (Cao & Roberts 2013). The inter-patch coupling is realised by spectral interpolation of the mid-patch values to the patch edges.

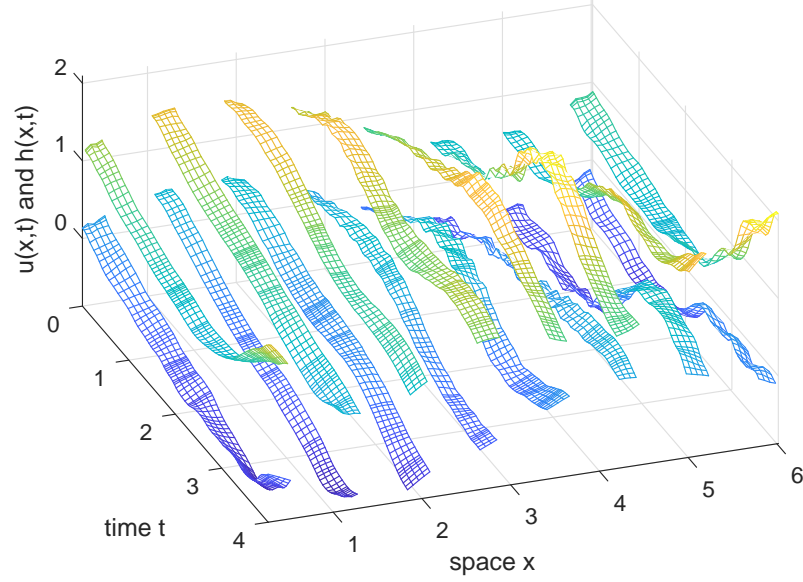
This approach, based upon the differential equations coded in Section 3.8.2, may be adapted by a user to a wide variety of 1D wave and wave-like systems. For example, the differential equations of Section 3.8.3 describes the nonlinear microscale simulator of the nonlinear shallow water wave PDE derived from the Smagorinski model of turbulent flow (Cao & Roberts 2012, 2016a).

Often, wave-like systems are written in terms of two conjugate variables, for example, position and momentum density, electric and magnetic fields, and water depth $h(x, t)$ and mean longitudinal velocity $u(x, t)$ as herein. The approach developed in this section applies to any wave-like system in the form

$$\frac{\partial h}{\partial t} = -c_1 \frac{\partial u}{\partial x} + f_1[h, u] \quad \text{and} \quad \frac{\partial u}{\partial t} = -c_2 \frac{\partial h}{\partial x} + f_2[h, u], \quad (3.3)$$

where the brackets indicate that the nonlinear functions f_ℓ may involve various spatial derivatives of the fields $h(x, t)$ and $u(x, t)$. For example, Section 3.8.3 encodes a nonlinear Smagorinski model of turbulent shallow water (Cao & Roberts 2012, 2016a, e.g.) along an inclined flat bed: let x measure position along the bed and in terms of fluid depth $h(x, t)$ and depth-averaged

Figure 3.12: water depth $h(x, t)$ (above) and velocity field $u(x, t)$ (below) of the gap-tooth scheme applied to the Smagorinski shallow water wave PDEs (3.4). The microscale random initial component decays where the water speed is non-zero due to ‘turbulent’ dissipation.



longitudinal velocity $u(x, t)$ the model PDEs are

$$\frac{\partial h}{\partial t} = -\frac{\partial(hu)}{\partial x}, \quad (3.4a)$$

$$\frac{\partial u}{\partial t} = 0.985 \left(\tan \theta - \frac{\partial h}{\partial x} \right) - 0.003 \frac{u|u|}{h} - 1.045u \frac{\partial u}{\partial x} + 0.26h|u| \frac{\partial^2 u}{\partial x^2}, \quad (3.4b)$$

where $\tan \theta$ is the slope of the bed. Equation (3.4a) represents conservation of the fluid. The momentum PDE (3.4b) represents the effects of turbulent bed drag $u|u|/h$, self-advection $u\partial u/\partial x$, nonlinear turbulent dispersion $h|u|\partial^2 u/\partial x^2$, and gravitational hydrostatic forcing $(\tan \theta - \partial h/\partial x)$. Figure 3.12 shows one simulation of this system—for the same initial condition as Figure 3.11.

For such wave systems, let’s implement a staggered microscale grid and staggered macroscale patches as introduced by Cao & Roberts (2016b) in their Figures 3 and 4, respectively.

3.8.1 Script code to simulate wave systems

This script implements the following gap-tooth scheme (left-right arrows denote function recursion).

1. configPatches1, and add micro-information
2. ode15s \leftrightarrow patchSmooth1 \leftrightarrow idealWavePDE
3. process results
4. ode15s \leftrightarrow patchSmooth1 \leftrightarrow waterWavePDE
5. process results

Establish the global data struct `patches` for the PDEs (3.3) (linearised) solved on 2π -periodic domain, with eight patches, each patch of half-size ratio 0.2, with eleven points within each patch, and spectral interpolation (-1) to provide edge-values of the inter-patch coupling conditions.

```

115 clear all
116 global patches
117 nPatch = 8
118 ratio = 0.2
119 nSubP = 11 %of the form 4*n-1
120 Len = 2*pi;
121 configPatches1(@idealWavePDE,[0 Len],nan,nPatch,-1,ratio,nSubP);

```

Identify which microscale grid points are h or u values on the staggered micro-grid. Also store the information in the struct `patches` for use by the time derivative function.

```

131 uPts = mod( bsxfun(@plus,(1:nSubP)',(1:nPatch)) ,2);
132 hPts = find(1-uPts);
133 uPts = find(uPts);
134 patches.hPts = hPts; patches.uPts = uPts;

```

Set an initial condition of a progressive wave, and check evaluation of the time derivative. The capital letter U denotes an array of values merged from both u and h fields on the staggered grids (possibly with some optional microscale wave noise).

```

145 U0 = nan(nSubP,nPatch);
146 U0(hPts) = 1+0.5*sin(patches.x(hPts));
147 U0(uPts) = 0+0.5*sin(patches.x(uPts));
148 U0 = U0+0.02*randn(nSubP,nPatch);

```

Conventional integration in time Integrate in time using standard MATLAB stiff integrators. Here do the two cases of the ideal wave and the water wave equations in the one loop.

```

158 for k = 1:2

```

When using `ode15s` we subsample the results because sub-grid scale waves do not dissipate and so the integrator takes very small time-steps for all time.

```

166 [ts,Ucts] = ode15s(@patchSmooth1,[0 4],U0(:));
167 ts = ts(1:5:end);
168 Ucts = Ucts(1:5:end,:);

```

Plot the simulation.

```

174 figure(k),clf
175 xs = patches.x; xs([1 end],:) = nan;
176 mesh(ts,xs(hPts),Ucts(:,hPts)'),hold on
177 mesh(ts,xs(uPts),Ucts(:,uPts)'),hold off
178 xlabel('time t'), ylabel('space x'), zlabel('u(x,t) and h(x,t)')
179 axis tight, view(70,45)

```

Save the plot to file.

```

185     set(gcf,'paperposition',[0 0 14 10])
186     if k==1, print('-depsc2','ps1WaveCtsUH')
187     else print('-depsc2','ps1WaterWaveCtsUH')
188     end

```

For the second time through the loop, change to the Smagorinski turbulence model (3.4) of shallow water flow, keeping other parameters and the initial condition the same.

```

198     patches.fun = @waterWavePDE;
199 end

```

Use projective integration As yet a simple implementation appears to fail, so it needs more exploration and thought. End of the main script.

3.8.2 idealWavePDE(): ideal wave PDE

This function codes the staggered lattice equation inside the patches for the ideal wave PDE system $h_t = -u_x$ and $u_t = -h_x$. Here code for a staggered microscale grid, index i , of staggered macroscale patches, index j : the array

$$U_{ij} = \begin{cases} u_{ij} & i+j \text{ even,} \\ h_{ij} & i+j \text{ odd.} \end{cases}$$

The output U_t contains the merged time derivatives of the two staggered fields. So set the micro-grid spacing and reserve space for time derivatives.

```

297 function Ut = idealWavePDE(t,U,x)
298     global patches
299     dx = diff(x(2:3));
300     Ut = nan(size(U)); ht = Ut;

```

Compute the PDE derivatives at interior points of the patches.

```

306     i = 2:size(U,1)-1;

```

Here ‘wastefully’ compute time derivatives for both PDEs at all grid points—for ‘simplicity’—and then merges the staggered results. Since $\dot{h}_{ij} \approx -(u_{i+1,j} - u_{i-1,j})/(2 \cdot dx) = -(U_{i+1,j} - U_{i-1,j})/(2 \cdot dx)$ as adding/subtracting one from the index of a h -value is the location of the neighbouring u -value on the staggered micro-grid.

```

318     ht(i,:) = -(U(i+1,:)-U(i-1,:))/(2*dx);

```

Since $\dot{u}_{ij} \approx -(h_{i+1,j} - h_{i-1,j})/(2 \cdot dx) = -(U_{i+1,j} - U_{i-1,j})/(2 \cdot dx)$ as adding/subtracting one from the index of a u -value is the location of the neighbouring h -value on the staggered micro-grid.

```

328     Ut(i,:) = -(U(i+1,:)-U(i-1,:))/(2*dx);

```

Then overwrite the unwanted \dot{u}_{ij} with the corresponding wanted \dot{h}_{ij} .

```

335     Ut(patches.hPts) = ht(patches.hPts);
336 end

```

3.8.3 waterWavePDE(): water wave PDE

This function codes the staggered lattice equation inside the patches for the nonlinear wave-like PDE system (3.4). Also, regularise the absolute value appearing the the PDEs via the one-line function `rabs()`.

```

351 function Ut = waterWavePDE(t,U,x)
352     global patches
353     rabs = @(u) sqrt(1e-4 + u.^2);

```

As before, set the micro-grid spacing, reserve space for time derivatives, and index the patch-interior points of the micro-grid.

```

361     dx = diff(x(2:3));
362     Ut = nan(size(U));  ht = Ut;
363     i = 2:size(U,1)-1;

```

Need to estimate h at all the u -points, so into V use averages, and linear extrapolation to patch-edges.

```

371     ii = i(2:end-1);
372     V = Ut;
373     V(ii,:) = (U(ii+1,:)+U(ii-1,:))/2;
374     V(1:2,:) = 2*U(2:3,:)-V(3:4,:);
375     V(end-1:end,:) = 2*U(end-2:end-1,:)-V(end-3:end-2,:);

```

Then estimate $\partial(hu)/\partial x$ from u and the interpolated h at the neighbouring micro-grid points.

```

382     ht(i,:) = -(U(i+1,:).*V(i+1,:)-U(i-1,:).*V(i+1,:))/(2*dx);

```

Correspondingly estimate the terms in the momentum PDE: u -values in U_i and $V_{i\pm 1}$; and h -values in V_i and $U_{i\pm 1}$.

```

390     Ut(i,:) = -0.985*(U(i+1,:)-U(i-1,:))/(2*dx) ...
391               -0.003*U(i,:).*rabs(U(i,:)/V(i,:)) ...
392               -1.045*U(i,:).*(V(i+1,:)-V(i-1,:))/(2*dx) ...
393               +0.26*rabs(V(i,:).*U(i,:)).*(V(i+1,:)-2*U(i,:)+V(i-1,:))/dx^2/2;

```

where the mysterious division by two in the second derivative is due to using the averaged values of u in the estimate:

$$\begin{aligned}
 u_{xx} &\approx \frac{1}{4\delta^2}(u_{i-2} - 2u_i + u_{i+2}) \\
 &= \frac{1}{4\delta^2}(u_{i-2} + u_i - 4u_i + u_i + u_{i+2}) \\
 &= \frac{1}{2\delta^2} \left(\frac{u_{i-2} + u_i}{2} - 2u_i + \frac{u_i + u_{i+2}}{2} \right) \\
 &= \frac{1}{2\delta^2} (\bar{u}_{i-1} - 2u_i + \bar{u}_{i+1}).
 \end{aligned}$$

Then overwrite the unwanted \dot{u}_{ij} with the corresponding wanted \dot{h}_{ij} .

```

409     Ut(patches.hPts) = ht(patches.hPts);
410 end

```

Fin.

3.9 configPatches2(): configures spatial patches in 2D

Section contents

3.9.1	Introduction	69
3.9.2	If no arguments, then execute an example	70
3.9.3	The code to make patches	72

3.9.1 Introduction

Makes the struct `patches` for use by the patch/gap-tooth time derivative function `patchSmooth2()`. [Section 3.9.2](#) lists an example of its use.

```

19 function configPatches2(fun,Xlim,BCs,nPatch,ordCC,ratio,nSubP,nEdge)
20 global patches

```

Input If invoked with no input arguments, then executes an example of simulating a nonlinear diffusion PDE relevant to the lubrication flow of a thin layer of fluid—see [Section 3.9.2](#) for the example code.

- `fun` is the name of the user function, `fun(t,u,x,y)`, that computes time derivatives (or time-steps) of quantities on the patches.
- `Xlim` array/vector giving the macro-space domain of the computation: patches are distributed equi-spaced over the interior of the rectangle $[Xlim(1), Xlim(2)] \times [Xlim(3), Xlim(4)]$: if `Xlim` is of length two, then use the same interval in both directions.
- `BCs` somehow will define the macroscale boundary conditions. Currently, `BCs` is ignored and the system is assumed macro-periodic in the domain.
- `nPatch` determines the number of equi-spaced spaced patches: if scalar, then use the same number of patches in both directions, otherwise `nPatch(1:2)` give the number in each direction.
- `ordCC` is the ‘order’ of interpolation across empty space of the macroscale mid-patch values to the edge of the patches for inter-patch coupling: currently must be in $\{0\}$.
- `ratio` (real) is the ratio of the half-width of a patch to the spacing of the patch mid-points: so `ratio` = $\frac{1}{2}$ means the patches abut; and `ratio` = 1 would be overlapping patches as in holistic discretisation: if scalar, then use the same ratio in both directions, otherwise `ratio(1:2)` give the ratio in each direction.
- `nSubP` is the number of equi-spaced microscale lattice points in each patch: if scalar, then use the same number in both directions, otherwise `nSubP(1:2)` gives the number in each direction. Must be odd so that there is a central lattice point.

- **nEdge** is, for each patch, the number of edge values set by interpolation at the edge regions of each patch. May be omitted. The default is one (suitable for microscale lattices with only nearest neighbours interactions).

Output The *global* struct **patches** is created and set with the following components.

- **.fun** is the name of the user's function **fun(u,t,x,y)** that computes the time derivatives (or steps) on the patchy lattice.
- **.ordCC** is the specified order of inter-patch coupling.
- **.alt** is true for interpolation using only odd neighbouring patches as for staggered grids, and false for the usual case of all neighbour coupling—not yet implemented.
- **.Cwtsr** and **.Cwtsl** are the **ordCC**-vector of weights for the inter-patch interpolation onto the right and left edges (respectively) with patch:macroscopic ratio as specified.
- **.x** is **nSubP(1) × nPatch(1)** array of the regular spatial locations x_{ij} of the microscale grid points in every patch.
- **.y** is **nSubP(2) × nPatch(2)** array of the regular spatial locations y_{ij} of the microscale grid points in every patch.
- **.nEdge** is, for each patch, the number of edge values set by interpolation at the edge regions of each patch.

3.9.2 If no arguments, then execute an example

```
123 if nargin==0
```

The code here shows one way to get started: a user's script may have the following three steps (arrows indicate function recursion).

1. **configPatches2**
2. **ode15s** integrator \leftrightarrow **patchSmooth2** \leftrightarrow user's **nonDiffPDE**
3. process results

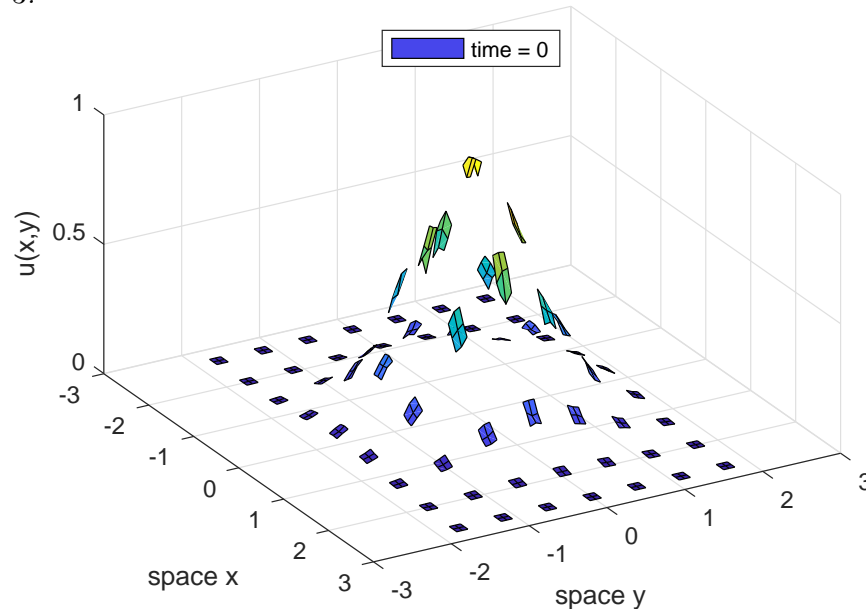
Establish global patch data struct to interface with a function coding a nonlinear 'diffusion' PDE: to be solved on 6×4 -periodic domain, with 9×7 patches, spectral interpolation (0) couples the patches, each patch of half-size ratio 0.25, and with 5×5 points within each patch. [Roberts et al. \(2014\)](#) established that this scheme is consistent with the PDE (as the patch spacing decreases).

```
145 nSubP = 5;
146 configPatches2(@nonDiffPDE,[-3 3 -2 2], nan, [9 7], 0, 0.25, nSubP);
```

Set a Gaussian initial condition using auto-replication of the spatial grid.

```
153 x = reshape(patches.x,nSubP,1,[],1);
154 y = reshape(patches.y,1,nSubP,1,[],1);
```


Figure 3.13: initial field $u(x, y, t)$ at time $t = 0$ of the patch scheme applied to a nonlinear ‘diffusion’ PDE: Figure 3.14 plots the computed field at time $t = 3$.



```

155 u0 = exp(-x.^2-y.^2);
156 u0 = u0.*(0.9+0.1*rand(size(u0)));

```

Initiate a plot of the simulation using only the microscale values interior to the patches: set x and y -edges to `nan` to leave the gaps.

```

164 figure(1), clf
165 x = patches.x; y = patches.y;
166 x([1 end], :) = nan; y([1 end], :) = nan;

```

Start by showing the initial conditions of Figure 3.13 while the simulation computes.

```

173 u = reshape(permute(u0,[1 3 2 4]), [numel(x) numel(y)]);
174 hsurf = surf(x(:),y(:),u');
175 axis([-3 3 -3 3 -0.001 1]), view(60,40)
176 legend('time = 0','Location','north')
177 xlabel('space x'), ylabel('space y'), zlabel('u(x,y)')
178 drawnow

```

Save the initial condition to file for Figure 3.13.

```

184 set(gcf,'PaperPosition',[0 0 14 10])
185 print('-depsc2','configPatches2ic')

```

Integrate in time using standard functions.

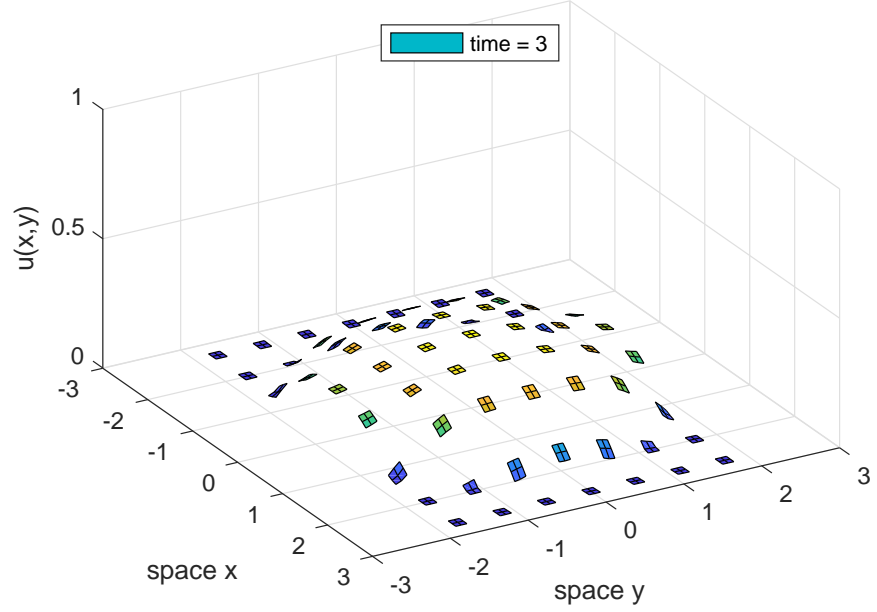
```

199 disp('Wait while we simulate h_t=(h^3)_xx+(h^3)_yy')
200 [ts,ucts] = ode15s(@patchSmooth2,[0 3],u0(:));

```

Animate the computed simulation to end with Figure 3.14.

Figure 3.14: field $u(x,y,t)$ at time $t = 3$ of the patch scheme applied to a nonlinear ‘diffusion’ PDE with initial condition in Figure 3.13.



```

207 for i = 1:length(ts)
208     u = patchEdgeInt2(ucts(i,:));
209     u = reshape(permute(u,[1 3 2 4]), [numel(x) numel(y)]);
210     hsurf.ZData = u';
211     legend(['time = ' num2str(ts(i),2)])
212     pause(0.1)
213 end
214 print('-depsc2','configPatches2t3')

```

Upon finishing execution of the example, exit this function.

```

229 return
230 end%if no arguments

```

Example of nonlinear diffusion PDE inside patches As a microscale discretisation of $u_t = \nabla^2(u^3)$, code $\dot{u}_{ijkl} = \frac{1}{\delta x^2}(u_{i+1,j,k,l}^3 - 2u_{i,j,k,l}^3 + u_{i-1,j,k,l}^3) + \frac{1}{\delta y^2}(u_{i,j+1,k,l}^3 - 2u_{i,j,k,l}^3 + u_{i,j-1,k,l}^3)$.

```

241 function ut = nonDiffPDE(t,u,x,y)
242     dx = diff(x(1:2)); dy = diff(y(1:2)); % microscale spacing
243     i = 2:size(u,1)-1; j = 2:size(u,2)-1; % interior points in patches
244     ut = nan(size(u)); % preallocate storage
245     ut(i,j, :, :) = diff(u(:,j, :, :).^3,2,1)/dx^2 ...
246                     +diff(u(i, :, :, :).^3,2,2)/dy^2;
247 end

```

3.9.3 The code to make patches

Initially duplicate parameters as needed.

```

263 if numel(Xlim)==2, Xlim = repmat(Xlim,1,2); end
264 if numel(nPatch)==1, nPatch = repmat(nPatch,1,2); end
265 if numel(ratio)==1, ratio = repmat(ratio,1,2); end
266 if numel(nSubP)==1, nSubP = repmat(nSubP,1,2); end

Set one edge-value to compute by interpolation if not specified by the user.
Store in the struct.

274 if nargin<8, nEdge = 1; end
275 if nEdge>1, error('multi-edge-value interp not yet implemented'), end
276 if 2*nEdge+1>nSubP, error('too many edge values requested'), end
277 patches.nEdge = nEdge;

First, store the pointer to the time derivative function in the struct.

286 patches.fun = fun;

Second, store the order of interpolation that is to provide the values for the
inter-patch coupling conditions. Spectral coupling is ordCC of 0 or -1.

295 if ~ismember(ordCC,[0])
296     error('ordCC out of allowed range [0]')
297 end

For odd ordCC do interpolation based upon odd neighbouring patches as is
useful for staggered grids.

304 patches.alt = mod(ordCC,2);
305 ordCC = ordCC+patches.alt;
306 patches.ordCC = ordCC;

Might as well precompute the weightings for the interpolation of field values
for coupling. (Could sometime extend to coupling via derivative values.)

322 ratio = ratio(:)'; % force to be row vector
323 if patches.alt % eqn (7) in \cite{Cao2014a}
324     patches.Cwtsr = [1
325         ratio/2
326         (-1+ratio.^2)/8
327         (-1+ratio.^2).*ratio/48
328         (9-10*ratio.^2+ratio.^4)/384
329         (9-10*ratio.^2+ratio.^4).*ratio/3840
330         (-225+259*ratio.^2-35*ratio.^4+ratio.^6)/46080
331         (-225+259*ratio.^2-35*ratio.^4+ratio.^6).*ratio/645120 ];
332 else %
333     patches.Cwtsr = [ratio
334         ratio.^2/2
335         (-1+ratio.^2).*ratio/6
336         (-1+ratio.^2).*ratio.^2/24
337         (4-5*ratio.^2+ratio.^4).*ratio/120
338         (4-5*ratio.^2+ratio.^4).*ratio.^2/720
339         (-36+49*ratio.^2-14*ratio.^4+ratio.^6).*ratio/5040
340         (-36+49*ratio.^2-14*ratio.^4+ratio.^6).*ratio.^2/40320 ];
341 end

```

```

342 patches.Cwtsr = patches.Cwtsr(1:ordCC,:);
343 % should avoid this next implicit auto-replication
344 patches.Cwtsl = (-1).^((1:ordCC)'-patches.alt).*patches.Cwtsr;

    Third, set the centre of the patches in a the macroscale grid of patches
    assuming periodic macroscale domain.

353 X = linspace(Xlim(1),Xlim(2),nPatch(1)+1);
354 X = X(1:nPatch(1))+diff(X)/2;
355 DX = X(2)-X(1);
356 Y = linspace(Xlim(3),Xlim(4),nPatch(2)+1);
357 Y = Y(1:nPatch(2))+diff(Y)/2;
358 DY = Y(2)-Y(1);

    Construct the microscale in each patch, assuming Dirichlet patch edges, and
    a half-patch length of ratio(1) · DX and ratio(2) · DY.

366 nSubP = nSubP(:)'; % force to be row vector
367 if mod(nSubP,2)==[0 0], error('configPatches2: nSubP must be odd'), end
368 i0 = (nSubP(1)+1)/2;
369 dx = ratio(1)*DX/(i0-1);
370 patches.x = bsxfun(@plus,dx*(-i0+1:i0-1)',X); % micro-grid
371 i0 = (nSubP(2)+1)/2;
372 dy = ratio(2)*DY/(i0-1);
373 patches.y = bsxfun(@plus,dy*(-i0+1:i0-1)',Y); % micro-grid
374 end% function

    Fin.

```

3.10 patchSmooth2(): interface to time integrators

Section contents

3.10.1 Introduction 74

3.10.1 Introduction

To simulate in time with spatial patches we often need to interface a users time derivative function with time integration routines such as `ode15s` or `PIRK2`. This function provides an interface. It assumes that the sub-patch structure is *smooth* so that the patch centre-values are sensible macroscale variables, and patch edge values are determined by macroscale interpolation of the patch-centre values. Communicate patch-design variables to this function via the previously established global struct `patches`.

```

25 function dudt = patchSmooth2(t,u)
26 global patches

```

Input

- **u** is a vector of length $\text{prod}(\text{nSubP}) \cdot \text{prod}(\text{nPatch}) \cdot \text{nVars}$ where there are **nVars** field values at each of the points in the $\text{nSubP}(1) \times \text{nSubP}(2) \times \text{nPatch}(1) \times \text{nPatch}(2)$ grid.
- **t** is the current time to be passed to the user's time derivative function.
- **patches** a struct set by `configPatches2()` with the following information used here.
 - **.fun** is the name of the user's function `fun(t,u,x,y)` that computes the time derivatives on the patchy lattice. The array **u** has size $\text{nSubP}(1) \times \text{nSubP}(2) \times \text{nPatch}(1) \times \text{nPatch}(2) \times \text{nVars}$. Time derivatives must be computed into the same sized array, but herein the patch edge values are overwritten by zeros.
 - **.x** is $\text{nSubP}(1) \times \text{nPatch}(1)$ array of the spatial locations x_{ij} of the microscale grid points in every patch. Currently it *must* be an equi-spaced lattice on both macro- and microscale.
 - **.y** is similarly $\text{nSubP}(2) \times \text{nPatch}(2)$ array of the spatial locations y_{ij} of the microscale grid points in every patch. Currently it *must* be an equi-spaced lattice on both macro- and microscale.

Output

- **dudt** is $\text{prod}(\text{nSubP}) \cdot \text{prod}(\text{nPatch}) \cdot \text{nVars}$ vector of time derivatives, but with patch edge values set to zero.

Reshape the fields **u** as a 4/5D-array, and sets the edge values from macroscale interpolation of centre-patch values. [Section 3.11](#) describes `patchEdgeInt2()`.

```
84 u = patchEdgeInt2(u);
```

Ask the user function for the time derivatives computed in the array, overwrite its edge values with the dummy value of zero, then return to a to the user/integrator as column vector.

```
94 dudt = patches.fun(t,u,patches.x,patches.y);
95 dudt([1 end],:,:,:) = 0;
96 dudt(:, [1 end],:,:) = 0;
97 dudt = reshape(dudt, [], 1);
```

Fin.

3.11 patchEdgeInt2(): sets 2D patch edge values from 2D macroscale interpolation

Section contents

Couples 2D patches across 2D space by computing their edge values via macroscale interpolation. Assumes that the sub-patch structure is *smooth* so that the patch centre-values are sensible macroscale variables, and patch

edge values are determined by macroscale interpolation of the patch-centre values. Communicate patch-design variables via the global struct `patches`.

```
20 function u = patchEdgeInt2(u)
21 global patches
```

Input

- `u` is a vector of length $nx \cdot ny \cdot Nx \cdot Ny \cdot nVars$ where there are `nVars` field values at each of the points in the $nx \times ny \times Nx \times Ny$ grid on the $Nx \times Ny$ array of patches.
- `patches` a struct set by `configPatches2()` which includes the following information.
 - `.x` is $nx \times Nx$ array of the spatial locations x_{ij} of the microscale grid points in every patch. Currently it *must* be an equi-spaced lattice on both macro- and microscale.
 - `.y` is similarly $ny \times Ny$ array of the spatial locations y_{ij} of the microscale grid points in every patch. Currently it *must* be an equi-spaced lattice on both macro- and microscale.
 - `.ordCC` is order of interpolation, currently only $\{0\}$.
 - `.Cwtsr` and `.Cwtsl`—not yet used

Output

- `u` is $nx \times ny \times Nx \times Ny \times nVars$ array of the fields with edge values set by interpolation.

Determine the sizes of things. Any error arising in the reshape indicates `u` has the wrong size.

```
75 [ny,Ny] = size(patches.y);
76 [nx,Nx] = size(patches.x);
77 nVars = round(numel(u)/numel(patches.x)/numel(patches.y));
78 if numel(u) ~= nx*ny*Nx*Ny*nVars
79     nSubP=[nx ny], nPatch=[Nx Ny], nVars=nVars, sizeu=size(u)
80 end
81 u = reshape(u,[nx ny Nx Ny nVars]);
```

With Dirichlet patches, the half-length of a patch is $h = dx(n_\mu - 1)/2$ (or -2 for specified flux), and the ratio needed for interpolation is then $r = h/\Delta X$. Compute lattice sizes from inside the patches as the edge values may be NaNs, etc.

```
91 dx = patches.x(3,1)-patches.x(2,1);
92 DX = patches.x(2,2)-patches.x(2,1);
93 rx = dx*(nx-1)/2/DX;
94 dy = patches.y(3,1)-patches.y(2,1);
95 DY = patches.y(2,2)-patches.y(2,1);
96 ry = dy*(ny-1)/2/DY;
```

For the moment assume the physical domain is macroscale periodic so that the coupling formulas are simplest. Should eventually cater for periodic, odd-mid-gap, even-mid-gap, even-mid-patch, Dirichlet, Neumann, Robin?? These index vectors point to patches and their two immediate neighbours.

```
107 %i=1:Nx; ip=mod(i,Nx)+1; im=mod(j-2,Nx)+1;
108 %j=1:Ny; jp=mod(j,Ny)+1; jm=mod(j-2,Ny)+1;
```

The centre of each patch (as nx and ny are odd) is at

```
115 i0 = round((nx+1)/2);
116 j0 = round((ny+1)/2);
```

Lagrange interpolation gives patch-edge values So compute centred differences of the mid-patch values for the macro-interpolation, of all fields. Assumes the domain is macro-periodic.

```
126 if patches.ordCC>0 % then non-spectral interpolation
127 error('non-spectral interpolation not yet implemented')
128 dmu=nan(patches.ordCC,nPatch,nVars);
129 % if patches.alt % use only odd numbered neighbours
130 % dmu(1,:,:)=(u(i0,jp,:)+u(i0,jm,:))/2; % \mu
131 % dmu(2,:,:) = u(i0,jp,:)-u(i0,jm,:); % \delta
132 % jp=jp(jp); jm=jm(jm); % increase shifts to \pm 2
133 % else % standard
134 % dmu(1,:,:)=(u(i0,jp,:)-u(i0,jm,:))/2; % \mu\delta
135 % dmu(2,:,:)=(u(i0,jp,:)-2*u(i0,j,:)+u(i0,jm,:)); % \delta^2
136 % end% if odd/even
```

Recursively take δ^2 of these to form higher order centred differences (could unroll a little to cater for two in parallel).

```
144 for k=3:patches.ordCC
145     dmu(k,:,:) = dmu(k-2,jp,:)-2*dmu(k-2,j,:)+dmu(k-2,jm,:);
146 end
```

Interpolate macro-values to be Dirichlet edge values for each patch ([Roberts & Kevrekidis 2007](#)), using weights computed in `configPatches2()`. Here interpolate to specified order.

```
154 u(nSubP,j,:)=u(i0,j,:)*(1-patches.alt) ...
155     +sum(bsxfun(@times,patches.Cwtsr,dmu));
156 u( 1,j,:)=u(i0,j,:)*(1-patches.alt) ...
157     +sum(bsxfun(@times,patches.Cwtsl,dmu));
```

Case of spectral interpolation Assumes the domain is macro-periodic. We interpolate in terms of the patch index j , say, not directly in space. As the macroscale fields are N -periodic in the patch index j , the macroscale Fourier transform writes the centre-patch values as $U_j = \sum_k C_k e^{ik2\pi j/N}$. Then the edge-patch values $U_{j\pm r} = \sum_k C_k e^{ik2\pi/N(j\pm r)} = \sum_k C'_k e^{ik2\pi j/N}$ where $C'_k = C_k e^{ikr2\pi/N}$. For N patches we resolve ‘wavenumbers’ $|k| < N/2$, so set

row vector $\mathbf{k}_s = k2\pi/N$ for ‘wavenumbers’ $k = (0, 1, \dots, k_{\max}, -k_{\max}, \dots, -1)$ for odd N , and $k = (0, 1, \dots, k_{\max}, \pm(k_{\max} + 1) - k_{\max}, \dots, -1)$ for even N .

179 else% spectral interpolation

Deal with staggered grid by doubling the number of fields and halving the number of patches (configPatches2 tests there are an even number of patches). Then the patch-ratio is effectively halved. The patch edges are near the middle of the gaps and swapped.

```
189 % if patches.alt % transform by doubling the number of fields
190 % error('staggered grid not yet implemented')
191 % v=nan(size(u)); % currently to restore the shape of u
192 % u=cat(3,u(:,1:2:nPatch,:),u(:,2:2:nPatch,:));
193 % altShift=reshape(0.5*[ones(nVars,1);-ones(nVars,1)],1,1,[]);
194 % iV=[nVars+1:2*nVars 1:nVars]; % scatter interp to alternate field
195 % r=r/2; % ratio effectively halved
196 % nPatch=nPatch/2; % halve the number of patches
197 % nVars=nVars*2; % double the number of fields
198 % else % the values for standard spectral
199 % altShift = 0;
200 % iV = 1:nVars;
201 % end
```

Now set wavenumbers in the two directions. In the case of even N these compute the +-case for the highest wavenumber zig-zag mode, $k = (0, 1, \dots, k_{\max}, +(k_{\max} + 1) - k_{\max}, \dots, -1)$.

```
210 kMax = floor((Nx-1)/2);
211 krx = rx*2*pi/Nx*(mod((0:Nx-1)+kMax,Nx)-kMax);
212 kMay = floor((Ny-1)/2);
213 kry = ry*2*pi/Ny*(mod((0:Ny-1)+kMay,Ny)-kMay);
```

Test for reality of the field values, and define a function accordingly.

```
220 if imag(u(i0,j0,:,:))==0, uclean = @(u) real(u);
221 else uclean = @(u) u; end
```

Compute the Fourier transform of the patch centre-values for all the fields. If there are an even number of points, then zero the zig-zag mode in the FT and add it in later as cosine.

```
230 Ck = fft2(squeeze(u(i0,j0,:,:)));
```

The inverse Fourier transform gives the edge values via a shift a fraction $\mathbf{r}_x/\mathbf{r}_y$ to the next macroscale grid point. Initially preallocate storage for all the IFFTs that we need to cater for the zig-zag modes when there are an even number of patches in the directions.

```
241 nFTx = 2-mod(Nx,2);
242 nFTy = 2-mod(Ny,2);
243 unj = nan(1,ny,Nx,Ny,nVars,nFTx*nFTy);
244 u1j = nan(1,ny,Nx,Ny,nVars,nFTx*nFTy);
```



```

245 uin = nan(nx,1,Nx,Ny,nVars,nFTx*nFTy);
246 ui1 = nan(nx,1,Nx,Ny,nVars,nFTx*nFTy);

```

Loop over the required IFFTs.

```

252 iFT = 0;
253 for iFTx = 1:nFTx
254   for iFTy = 1:nFTy
255     iFT = iFT+1;

```

First interpolate onto x -limits of the patches. (It may be more efficient to product exponentials of vectors, instead of exponential of array—only for $N > 100$. Can this be vectorised further??)

```

264 for jj = 1:ny
265   ks = (jj-j0)*2/(ny-1)*kry; % fraction of kry along the edge
266   unj(1,jj,::,iV,iFT) = ifft2( bsxfun(@times,Ck ...
267     ,exp(1i*bsxfun(@plus,altShift+krx',ks))));
268   u1j(1,jj,::,iV,iFT) = ifft2( bsxfun(@times,Ck ...
269     ,exp(1i*bsxfun(@plus,altShift-krx',ks))));
270 end

```

Second interpolate onto y -limits of the patches.

```

276 for i = 1:nx
277   ks = (i-i0)*2/(nx-1)*krx; % fraction of krx along the edge
278   uin(i,1,::,iV,iFT) = ifft2( bsxfun(@times,Ck ...
279     ,exp(1i*bsxfun(@plus,ks',altShift+kry))));
280   ui1(i,1,::,iV,iFT) = ifft2( bsxfun(@times,Ck ...
281     ,exp(1i*bsxfun(@plus,ks',altShift-kry))));
282 end

```

When either direction have even number of patches then swap the zig-zag wavenumber to the conjugate.

```

289 if nFTy==2, kry(Ny/2+1) = -kry(Ny/2+1); end
290 end% iFTy-loop
291 if nFTx==2, krx(Nx/2+1) = -krx(Nx/2+1); end
292 end% iFTx-loop

```

Put edge-values into the u -array, using `mean()` to treat a zig-zag mode as cosine. Enforce reality when appropriate via `uclean()`.

```

300 u(end,::,::,iV) = uclean( mean(unj,6) );
301 u( 1 ,::,::,iV) = uclean( mean(u1j,6) );
302 u(:,end,::,iV) = uclean( mean(uin,6) );
303 u(:, 1 ,::,iV) = uclean( mean(ui1,6) );

```

Restore staggered grid when appropriate. Is there a better way to do this??

```

310 %if patches.alt
311 % nVars=nVars/2; nPatch=2*nPatch;
312 % v(:,1:2:nPatch,:)=u(:,::,1:nVars);
313 % v(:,2:2:nPatch,:)=u(:,::,nVars+1:2*nVars);
314 % u=v;

```

```

315 %end
316 end% if spectral
317 end% function patchEdgeInt2

```

Fin, returning the 4/5D array of field values with interpolated edges.

3.12 wave2D: example of a wave on patches in 2D

Section contents

3.12.1 Check on the linear stability of the wave PDE	80
3.12.2 Execute a simulation	81
3.12.3 Example of simple wave PDE inside patches	82

For $u(x, y, t)$, test and simulate the simple wave PDE in 2D space:

$$\frac{\partial^2 u}{\partial t^2} = \nabla^2 u.$$

This script shows one way to get started: a user's script may have the following three steps (left-right arrows denote function recursion).

1. configPatches2
2. ode15s integrator \leftrightarrow patchSmooth2 \leftrightarrow wavePDE
3. process results

Establish global patch data struct to interface with a function coding the wave PDE: to be solved on 2π -periodic domain, with 9×9 patches, spectral interpolation (0) couples the patches, each patch of half-size ratio 0.25, and with 5×5 points within each patch.

```

34 clear all, close all
35 global patches
36 nSubP = 5;
37 nPatch = 9;
38 configPatches2(@wavePDE, [-pi pi], nan, nPatch, 0, 0.25, nSubP);

```

3.12.1 Check on the linear stability of the wave PDE

Set a zero equilibrium as basis. Then find the indices of patch-interior points as the only ones to vary in order to construct the Jacobian.

```

50 disp('Check linear stability of the wave scheme')
51 uv0 = zeros(nSubP, nSubP, nPatch, nPatch, 2);
52 uv0([1 end], :, :, :, :) = nan;
53 uv0(:, [1 end], :, :, :) = nan;
54 i = find(~isnan(uv0));

```

Now construct the Jacobian. Since linear wave PDE, use large perturbations.

```

61 small = 1;
62 jac = nan(length(i));
63 sizeJacobian = size(jac)
64 for j = 1:length(i)
65     uv = uv0(:);
66     uv(i(j)) = uv(i(j))+small;
67     tmp = patchSmooth2(0,uv)/small;
68     jac(:,j) = tmp(i);
69 end

```

Now explore the eigenvalues a little: find the ten with the biggest real-part; if small enough, then the method may be good.

```

77 evals = eig(jac);
78 nEvals = length(evals)
79 [~,k] = sort(-abs(real(evals)));
80 evalsWithBiggestRealPart = evals(k(1:10))
81 if abs(real(evals(k(1))))>1e-4
82     warning('eigenvalue failure: real-part > 1e-4')
83     return, end

```

Check eigenvalues close to true waves of the PDE (not yet the micro-discretised equations).

```

90 kwave = 0:(nPatch-1)/2;
91 freq = sort(reshape(sqrt(kwave'.^2+kwave.^2),1,[]));
92 freq = freq(diff([-1 freq])>1e-9);
93 freqerr = [freq; min(abs(imag(evals)-freq))]

```

3.12.2 Execute a simulation

Set a Gaussian initial condition using auto-replication of the spatial grid: here u_0 and v_0 are in the form required for computation: $n_x \times n_y \times N_x \times N_y$.

```

108 x = reshape(patches.x,nSubP,1,[],1);
109 y = reshape(patches.y,1,nSubP,1,[]);
110 u0 = exp(-x.^2-y.^2);
111 v0 = zeros(size(u0));

```

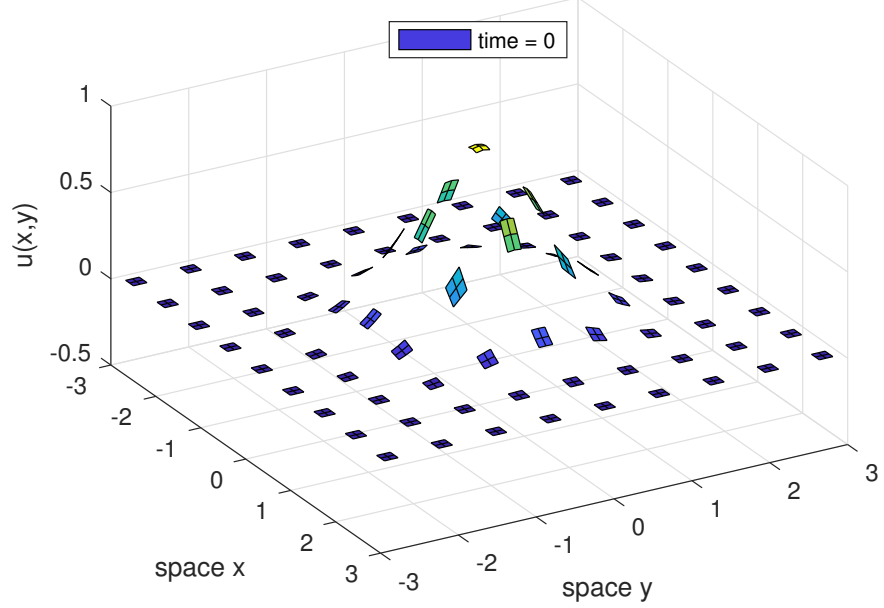
Initiate a plot of the simulation using only the microscale values interior to the patches: set x and y -edges to **nan** to leave the gaps. Start by showing the initial conditions of [Figure 3.13](#) while the simulation computes. To mesh/surf plot we need to ‘transpose’ to size $n_x \times N_x \times n_y \times N_y$, then reshape to size $n_x \cdot N_x \times n_y \cdot N_y$.

```

123 x = patches.x; y = patches.y;
124 x([1 end],:) = nan; y([1 end],:) = nan;
125 u = reshape(permute(u0,[1 3 2 4]), [numel(x) numel(y)]);
126 usurf = surf(x(:),y(:),u');
127 axis([-3 3 -3 3 -0.5 1]), view(60,40)
128 xlabel('space x'), ylabel('space y'), zlabel('u(x,y)')
129 legend('time = 0','Location','north')

```

Figure 3.15: initial field $u(x, y, t)$ at time $t = 0$ of the patch scheme applied to the simple wave PDE: Figure 3.16 plots the computed field at time $t = 2$.



```

130 drawnow
131 set(gcf,'paperposition',[0 0 14 10])
132 print('-depsc','wave2Dic')

Integrate in time using standard functions.

145 disp('Wait while we simulate u_t=v, v_t=u_xx+u_yy')
146 [ts, uvs] = ode15s(@patchSmooth2, [0 2], [u0(:); v0(:)]);

```

Animate the computed simulation to end with Figure 3.16. Because of the very small time-steps, subsample to plot at most 200 times.

```

154 di = ceil(length(ts)/200);
155 for i = [1:di:length(ts)-1 length(ts)]
156     uv = patchEdgeInt2(uvs(i,:));
157     uv = reshape(permute(uv, [1 3 2 4 5]), [numel(x) numel(y) 2]);
158     usurf.ZData = uv(:, :, 1)';
159     legend(['time = ' num2str(ts(i), 2)])
160     pause(0.1)
161 end
162 print('-depsc', ['wave2Dt' num2str(ts(end))])

```

3.12.3 Example of simple wave PDE inside patches

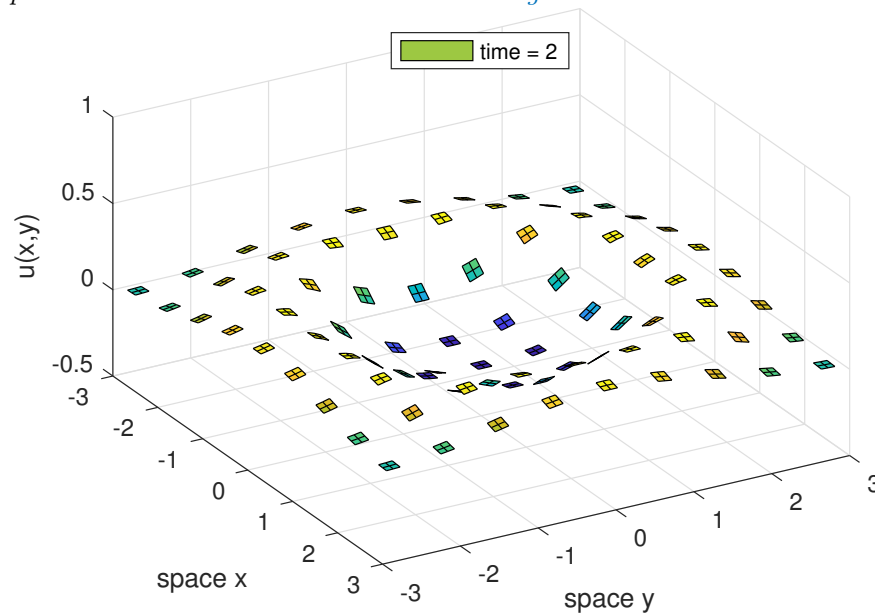
As a microscale discretisation of $u_{tt} = \nabla^2(u)$, so code $\dot{u}_{ijkl} = v_{ijkl}$ and $\dot{v}_{ijkl} = \frac{1}{\delta x^2}(u_{i+1,j,k,l} - 2u_{i,j,k,l} + u_{i-1,j,k,l}) + \frac{1}{\delta y^2}(u_{i,j+1,k,l} - 2u_{i,j,k,l} + u_{i,j-1,k,l})$.

```

183 function uvt = wavePDE(t, uv, x, y)
184     if ceil(t+1e-7)-t < 2e-2, simTime = t, end %track progress
185     dx = diff(x(1:2)); dy = diff(y(1:2)); % microscale spacing
186     i = 2:size(uv,1)-1; j = 2:size(uv,2)-1; % interior patch-points

```

Figure 3.16: field $u(x, y, t)$ at time $t = 2$ of the patch scheme applied to the simple wave PDE with initial condition in Figure 3.15.



```

187     uvt = nan(size(uv)); % preallocate storage
188     uvt(i,j,:,:1) = uv(i,j,:,:2);
189     uvt(i,j,:,:2) = diff(uv(:,j,:,:1),2,1)/dx^2 ...
190                     +diff(uv(i,:,:,:),2,2)/dy^2;
191     end

```

3.13 To do

- Testing needs to be quantitative.
- more than two space dimensions??
- Heterogeneous microscale via averaging regions—but I suspect should be separated from simple homogenisation
- Parallel processing versions.
- ??
- Adapt to maps in micro-time? Surely easy, just an example.

3.14 Miscellaneous tests

3.14.1 patchEdgeInt1test: test the spectral interpolation

A script to test the spectral interpolation of function `patchEdgeInt1()` Establish global data struct for the range of various cases.

```

13     clear all
14     global patches

```

```

15  nSubP=3
16  i0=(nSubP+1)/2; % centre-patch index

```

Test standard spectral interpolation Test over various numbers of patches, random domain lengths and random ratios.

```

24  for nPatch=5:10
25  nPatch=nPatch
26  Len=10*rand
27  ratio=0.5*rand
28  configPatches1(@sin,[0,Len],nan,nPatch,0,ratio,nSubP);
29  kMax=floor((nPatch-1)/2);

```

Test single field Set a profile, and evaluate the interpolation.

```

37  for k=-kMax:kMax
38      u0=exp(1i*k*patches.x*2*pi/Len);
39      ui=patchEdgeInt1(u0(:));
40      normError=norm(ui-u0);
41      if abs(normError)>5e-14
42          normError=normError
43          error(['failed single var interpolation k=' num2str(k)])
44      end
45  end

```

Test multiple fields Set a profile, and evaluate the interpolation. For the case of the highest wavenumber, squash the error when the centre-patch values are all zero.

```

54  for k=1:nPatch/2
55      u0=sin(k*patches.x*2*pi/Len);
56      v0=cos(k*patches.x*2*pi/Len);
57      uvi=patchEdgeInt1([u0(:);v0(:)]);
58      normuError=norm(uvi(:,1)-u0)*norm(u0(i0,:));
59      normvError=norm(uvi(:,2)-v0)*norm(v0(i0,:));
60      if abs(normuError)+abs(normvError)>2e-13
61          normuError=normuError, normvError=normvError
62          error(['failed double field interpolation k=' num2str(k)])
63      end
64  end

```

End the for-loop over various geometries.

```

71  end

```

Now test spectral interpolation on staggered grid Must have even number of patches for a staggered grid.

```

79  for nPatch=6:2:20
80  nPatch=nPatch
81  ratio=0.5*rand

```

```

82  nSubP=3; % of form 4*N-1
83  Len=10*rand
84  configPatches1(@simpleWavepde,[0,Len],nan,nPatch,-1,ratio,nSubP);
85  kMax=floor((nPatch/2-1)/2)

```

Identify which microscale grid points are h or u values.

```

91  uPts=mod( bsxfun(@plus,(1:nSubP)',(1:nPatch)) ,2);
92  hPts=find(1-uPts);
93  uPts=find(uPts);

```

Set a profile for various wavenumbers. The capital letter U denotes an array of values merged from both u and h fields on the staggered grids.

```

100 fprintf('Single field-pair test.\n')
101 for k=-kMax:kMax
102     U0=nan(nSubP,nPatch);
103     U0(hPts)=rand*exp(+1i*k*patches.x(hPts)*2*pi/Len);
104     U0(uPts)=rand*exp(-1i*k*patches.x(uPts)*2*pi/Len);
105     Ui=patchEdgeInt1(U0(:));
106     normError=norm(Ui-U0);
107     if abs(normError)>5e-14
108         normError=normError
109         error(['failed single sys interpolation k=' num2str(k)])
110     end
111 end

```

Test multiple fields Set a profile, and evaluate the interpolation. For the case of the highest wavenumber zig-zag, squash the error when the alternate centre-patch values are all zero. First shift the x -coordinates so that the zig-zag mode is centred on a patch.

```

121 fprintf('Two field-pairs test.\n')
122 x0=patches.x((nSubP+1)/2,1);
123 patches.x=patches.x-x0;
124 for k=1:nPatch/4
125     U0=nan(nSubP,nPatch); V0=U0;
126     U0(hPts)=rand*sin(k*patches.x(hPts)*2*pi/Len);
127     U0(uPts)=rand*sin(k*patches.x(uPts)*2*pi/Len);
128     V0(hPts)=rand*cos(k*patches.x(hPts)*2*pi/Len);
129     V0(uPts)=rand*cos(k*patches.x(uPts)*2*pi/Len);
130     UVi=patchEdgeInt1([U0(:);V0(:)]);
131     normuError=norm(UVi(:,1:2:nPatch,1)-U0(:,1:2:nPatch))*norm(U0(i0,2:2:nPatch)
132         +norm(UVi(:,2:2:nPatch,1)-U0(:,2:2:nPatch))*norm(U0(i0,1:2:nPatch));
133     normvError=norm(UVi(:,1:2:nPatch,2)-V0(:,1:2:nPatch))*norm(V0(i0,2:2:nPatch)
134         +norm(UVi(:,2:2:nPatch,2)-V0(:,2:2:nPatch))*norm(V0(i0,1:2:nPatch));
135     if abs(normuError)+abs(normvError)>2e-13
136         normuError=normuError, normvError=normvError
137         error(['failed double field interpolation k=' num2str(k)])
138     end
139 end

```

```

End for-loop over patches
146 end

Finish If no error messages, then all OK.
157 fprintf('\nIf you read this, then all tests were passed\n')

```

3.14.2 patchEdgeInt2test: tests 2D spectral interpolation

Try 99 realisations of random tests.

```

11 clear all, close all
12 global patches
13 for realisation=1:99

Choose and configure random sized domains, random sub-patch resolution,
random size-ratios, random number of periodic-patches.

19 Lx=1+3*rand, Ly=1+3*rand
20 nSubP=1+2*randi(3,1,2)
21 ratios=rand(1,2)/2
22 nPatch=2+randi(4,1,2)
23 configPatches2(@sin,[0 Lx 0 Ly],nan,nPatch,0,ratios,nSubP)

Choose a random number of fields, then generate trigonometric shape with
random wavenumber and random phase shift.

29 nV=randi(3)
30 [nx,Nx]=size(patches.x);
31 [ny,Ny]=size(patches.y);
32 u0s=nan(nx,ny,Nx,Ny,nV);
33 for iV=1:nV
34     kx=randi([0 ceil((nPatch(1)-1)/2)])
35     ky=randi([0 ceil((nPatch(2)-1)/2)])
36     phix=pi*rand*(2*kx~nPatch(1))
37     phiy=pi*rand*(2*ky~nPatch(2))
38     % generate 2D array via auto-replication
39     u0=sin(2*pi*kx*patches.x(:)/Lx+phix) ...
40     .*sin(2*pi*ky*patches.y(:)/Ly+phiy);
41     % reshape into 4D array
42     u0=reshape(u0,[nx Nx ny Ny]);
43     u0=permute(u0,[1 3 2 4]);
44     % store into 5D array
45     u0s(:,:,,:,iV)=u0;
46 end

Copy and NaN the edges, then interpolate

52 u=u0s; u([1 end],:,:,:,)=nan; u(:,[1 end],:,:,:,)=nan;
53 u=patchEdgeInt2(u(:));

```

If there is an error in the interpolation then abort the script for checking:
record parameter values and inform.

```
59  err=u-u0s;  
60  normerr=norm(err(:))  
61  if normerr>1e-12, error('2D interpolation failed'), end  
62  end
```

Appendix A Create, document and test algorithms

For developers to create and document the various functions, we use an idea due to Neil D. Lawrence of the University of Sheffield.

- Each class of toolbox functions is located in separate folders in the repository, say `Dir`.
- Create a LaTeX file `Dir/funs.tex`: establish as one LaTeX chapter that `\input{Dir/*.m}`s the files of the functions in the class, example scripts of use, and possibly test scripts, [Table A.1](#).
- Each such `Dir/funs.tex` file is to be included from the main LaTeX file `Doc/docBody.tex` so that people can most easily work on one chapter at a time:
 - put `\include{funs}` into `Doc/docBody.tex`;
 - to include, create a ‘link’ file `Doc/funs.tex` whose only active content is the command `\input{../Dir/funs.tex}`;
 - in `Doc/docBody.tex` modify the `\graphicspath` command to include `{../Dir/Figs}`.
- Each toolbox function is documented as a separate section, within its chapter, with tests and examples as separate sections.
- Each function-section and test-section is to be created as a MATLAB `Dir/*.m` file, say `Dir/fun1.m`, so that users simply invoke the function in MATLAB as usual by `fun1(...)`.

Some editors may need to be told that `fun1.m` is a LaTeX file. For example, TexShop on the Mac requires one to execute in a Terminal

```
defaults write TexShop OtherTeXExtensions -array-add ".m"
```

- [Table A.2](#) gives the template for the `Dir/*.m` function-sections. The format for a example/test-section is similar.
- Any figures from examples should be generated and then saved for later inclusion with the following (which finally works properly for MATLAB 2017+)

```
set(gcf,'PaperPosition',[0 0 14 10]);% cm
print('-depsc2','filename')
```

If it is a suitable replacement for an existing graphic, then move it into the `Dir/Figs` folder. Include such a graphic into the LaTeX document with (do *not* postfix with `.eps` or `.pdf`)

```
\includegraphics[scale=0.9]{filename}
```

Table A.1: example `Dir/*.tex` file to typeset in the master document a function-section, say `fun.m`, and maybe the test/example-sections.

```

1 % input *.m files for ... Author, date
2 %!TEX root = ../Doc/eqnFreeDevMan.tex
3 \chapter{...}
4 \label{sec:...}
5 \localtableofcontents
6 \section{Introduction}
7 introduction...
8 \input{../Dir/fun.m} % prefix associated files with 'fun'
9 \input{../Dir/funExample.m}
10 ...
11 \begin{devMan}
12 \section{To do}
13 ...
14 \section{Miscellaneous tests}
15 \input{../Dir/funTest.m}
16 ...
17 \end{devMan}

```

- In figures and other graphics, do *not* resize/scale fixed width constructs: instead use `\linewidth` to configure large-scale layout, `em` for small-widths, and `ex` for small-heights.
- For every function, generally include at the start of the function a simple example of its use. The example is only to be executed when the function is invoked with no input arguments (`if nargin==0`).

When appropriate, if a function is invoked with no output arguments (`if nargout==0`), then draw some reasonable graph of the results.

- In all MATLAB code, prefer camel case for variable names (not under-scores).
- In the documentation (e.g., [Higham 1998](#), Ch. 4): write actively, not passively (e.g., avoid “-tion” words); avoid wishy-washy “can”; use the present tense; cross-reference precisely; and so on.

Table A.2: template for a function-section Dir/*.m file.

```

1  % Short explanation for users typing "help fun"
2  % Author, date
3  %!TEX root = ../Doc/eqnFreeDevMan.tex
4  %{
5  \section{\texttt{...}: ...}
6  \label{sec:...}
7  \localtableofcontents
8  \subsection{Introduction}
9  Overview LaTeX explanation.
10 \begin{matlab}
11 %}
12 function ...
13 %{
14 \end{matlab}
15 \paragraph{Input} ...
16 \paragraph{Output} ...
17 \begin{devMan}
18 Repeated as desired:
19 LaTeX between end-matlab and begin-matlab
20 \begin{matlab}
21 %}
22 Matlab code between %} and %{
23 %{
24 \end{matlab}
25 Concluding LaTeX before following final lines.
26 \end{devMan}
27 %}

```

Appendix B Aspects of developing a ‘toolbox’ for patch dynamics

Chapter contents

B.1	Macroscale grid	91
B.2	Macroscale field variables	91
B.3	Boundary and coupling conditions	92
B.4	Mesotime communication	92
B.5	Projective integration	92
B.6	Lift to many internal modes	93
B.7	Macroscale closure	93
B.8	Exascale fault tolerance	93
B.9	Link to established packages	94

This appendix documents sketchy further thoughts on aspects of the development.

B.1 Macroscale grid

The patches are to be distributed on a macroscale grid: the j th patch ‘centred’ at position $\vec{X}_j \in \mathbb{X}$. In principle the patches could move, but let’s keep them fixed in the first version. The simplest macroscale grid will be rectangular (`meshgrid`), but we plan to allow a deformed grid to secondly cater for boundary fitting to quite general domain shapes \mathbb{X} . And plan to later allow for more general interconnect networks for more topologies in application.

B.2 Macroscale field variables

The researcher/practitioner has to know an appropriate set of macroscale field variables $\vec{U}(t) \in \mathbb{R}^{d_{\vec{U}}}$ for each patch. For example, first they might be a simple average over a core of a patch of all of the micro-field variables; second, they might be a subset of the average micro-field variables; and third in general the macro-variables might be a nonlinear function of the micro-field variables (such as temperature is the average speed squared). The core might be just one point, or a sizeable fraction of the patch.

The mapping from microscale variable to macroscale variables is often termed the restriction.

In practice, users may not choose an appropriate set of macro-variables, so will eventually need to code some diagnostic to indicate a failure of the assumed closure.

B.3 Boundary and coupling conditions

The physical domain boundary conditions are distinct from the conditions coupling the patches together. Start with physical boundary conditions of periodicity in the macroscale.

Second, assume the physical boundary conditions are that the macro-variables are known at macroscale grid points around the boundary. Then the issue is to adjust the interpolation to cater for the boundary presence and shape. The coupling conditions for the patches should cater for the range of Robin-like boundary conditions, from Dirichlet to Neumann. Two possibilities arise: direct imposition of the coupling action ([Roberts & Kevrekidis 2007](#)), or control by the action.

Third, assume that some of the patches have some edges coincident with the boundary of the macroscale domain \mathbb{X} , and it is on these edges that macroscale physical boundary conditions are applied. Then the interpolation from the core of these edge patches is the same as the second case of prescribed boundary macro-variables. An issue is that each boundary patch should be big enough to cater for any spatial boundary layers transitioning from the applied boundary condition to the interior slow evolution.

Alternatively, we might have the physical boundary condition constrain the interpolation between patches.

Often microscale simulations are easiest to write when ‘periodic’ in microscale space. To cater for this we should also allow a control at perhaps the quartiles of a micro-periodic simulator.

B.4 Mesotime communication

Since communication limits large scale parallelism, a first step in reducing communication will be to implement only updating the coupling conditions when necessary. Error analysis indicates that updating on times longer the microscale times and shorter than the macroscale times can be effective ([Bunder et al. 2016](#)). Implementations can communicate one or more derivatives in time, as well as macroscale variables.

At this stage we can effectively parallelise over patches: first by simply using Matlab’s `parfor`. Probably not using a GPU as we probably want to leave GPUs for the black-box to utilise within each patch.

B.5 Projective integration

To take macroscale time-steps, invoke several possible projective integration schemes: simple Euler projection, Heun-like method, etc ([Samaey et al. 2010](#)). Need to decide how long a microscale burst needs to be.

Should not need an implicit scheme as the fast dynamics are meant to be only in the micro variables, and the slow dynamics only in the macroscale variables. However, it could be that the macroscale variables have fast oscillations and it is only the amplitude of the oscillations that are slow. Perhaps need to detect and then fix or advise.

A further stage is to implement a projective integration scheme for stochastic macroscale variables: this is important because the averaging over a core of microscale roughness will almost invariably have at least some stochastic legacy effect. [Calderon \(2007\)](#) did some useful research on stochastic projective intergration.

B.6 Lift to many internal modes

In most problems the number of macroscale variables at any given position in space, $d_{\vec{J}}$, is less than the number of microscale variables at a position, $d_{\vec{u}}$; often much less ([Kevrekidis & Samaey 2009](#), e.g.). In this case, every time we start a patch simulation we need to provide $d_{\vec{u}} - d_{\vec{J}}$ data at each position in the patch: this is lifting. The first methodology is to first guess, then run repeated short bursts with reinitialisation, until the simulation reaches a slow manifold. Then run the real simulation.

If the time taken to reach a local quasi-equilibrium is too long, then it is likely that the macroscale closure is bad and the macroscale variables need to be extended.

A second step is to cater for cases where the slow manifold is stochastic or is surrounded by fast waves: when it is hard to detect the slow manifold, or the slow manifold is not attractive.

B.7 Macroscale closure

In some circumstances a researcher/practitioner will not code the appropriately set of macroscale variables for a complete closure of the macroscale. For example, in thin film fluid dynamics at low Reynolds number the only macroscale variable is the fluid depth; however, at higher Reynolds number, circa ten, the inertia of the fluid becomes important and the macroscale variables must additionally include a measure of the mean lateral velocity/momentum ([Roberts & Li 2006](#), e.g.).

At some stage we need to detect any flaw in the closure, and perhaps suggest additional appropriate macroscale variables, or at least their characteristics. Indeed, a poor closure and a stochastic slow manifold are really two faces of the same problem: the problem is that the chosen macroscale variables do not have a unique evolution in terms of themselves. A good resolution of the issue will account for both faces.

B.8 Exascale fault tolerance

Matlab is probably not an appropriate vehicle to deal with real exascale faults. However, we should cater by coding procedures for fault tolerance

and testing them at least synthetically. Eventually provide hooks to a user routine to be invoked under various potential scenarios. The nature of fault tolerant algorithms will vary depending upon the scenario, even assuming that each patch burst is executed on one CPU (or closely coupled CPUs): if there are much more CPUs than patches, then maybe simply duplicate all patch simulations; if much less CPUs than patches, then an asynchronous scheduling of patch bursts should effectively cater for recomputation of failed bursts; if comparable CPUs to patches, then more subtle action is needed.

Once mesotime communication and projective integration is provided, a recomputation approach to intermittent hardware faults should be effective because we then have the tools to restart a burst from available macroscale data. Should also explore proceeding with a lower order interpolation that misses the faulty burst—because an isolated lower order interpolation probably will not affect the global order of error (it does not in approximating some boundary conditions ([Gustafsson 1975](#), [Svard & Nordstrom 2006](#)))

B.9 Link to established packages

Several molecular/particle/agent based codes are well developed and used by a wide community of researchers. Plan to develop hooks to use some such codes as the microscale simulators on patches. First, plan to connect to LAMMPS ([Plimpton et al. 2016](#)). Second, will evaluate performance, issues, and then consider what other established packages are most promising.

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