

xSPDE: The Stochastic Toolbox User's Guide v3.2

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1 Introduction

The xSPDE stochastic toolbox is an extensible **S**tochastic **P**artial **D**ifferential **E**quation solver. It can solve any number of simultaneous equations in any number of space dimensions, with averaging and graphical output. These equations include ordinary differential equations, partial differential equations with specified boundaries, stochastic equations and stochastic partial differential equations. This user's guide can be used as the basis for a 3-4 hour self-taught tutorial on solving differential and stochastic equations. Additionally, it includes exercises for the reader. A full-length manual is available online. Research use of this software is encouraged, and should be referenced to the corresponding paper[1]. See: www.github.com/peterddrummond/xspde_matlab.

Stochastic equations are equations with random noise terms [2, 3]. They occur in many fields of biology, chemistry, engineering, economics, physics, meteorology and other disciplines. The general equation solved can be written in differential form as

$$\frac{\partial \mathbf{a}}{\partial t} = \mathbf{A}[\mathbf{a}] + \mathbf{B}[\mathbf{a}] \cdot \mathbf{w}(t, \mathbf{x}) + \mathbf{L}[\nabla, \mathbf{a}] . \quad (1.1)$$

Here, \mathbf{A} is a vector, \mathbf{B} a matrix, $\mathbf{L}[\nabla, \mathbf{a}]$ is a differential operator in arbitrary dimensional real space \mathbf{x} , and \mathbf{w} is a noise vector, delta-correlated in time:

$$\langle w_i(t, \mathbf{x}) w_j(t', \mathbf{x}') \rangle = \delta(t - t') \delta(\mathbf{x} - \mathbf{x}') \delta_{ij} \quad (1.2)$$

Ordinary and stochastic differential equations are described first. Stochastic partial differential equations including space dependence [4, 5] are also numerically soluble with xSPDE. These are described in Chapter 4.

xSPDE is a Matlab based software toolbox. It has functions that can numerically solve both ordinary and partial differential stochastic equations, and graph results of correlations and averages. There are many equations of this type, and the xSPDE toolbox can treat a wide range. The extensible general structure permits drop-in replacements of the functions provided. Different simulations can be carried out sequentially, to simulate the various stages in an experiment or other process. It can even be used with no noise.

The code supports parallelism at both the vector instruction level and at the thread level, using Matlab matrix instructions and the parallel toolbox. It calculates averages of arbitrary functions of any number of complex or real fields, as well as Fourier transforms in time or space. It provides error estimates for both step-size and sampling error.

xSPDE is distributed without any guarantee, under an open-source license. Contributions and bug reports always welcome.

2 xSPDE simulations

The general purpose xSPDE toolbox function is:

- **xspde**, the combined simulation (xSIM) and graphics (xGRAPH) function.
- All required files are available in a toolbox: *xspde.mltbx*, or in a folder: xSPDE3.

2.1 Running xSPDE interactively

The xSPDE code can be run interactively as a script, or as a function in batch mode, either at a local workstation or on a remote cluster. Data can be either plotted immediately, or saved then plotted later. To simulate a stochastic equation interactively, first check that the Matlab xSPDE toolbox is installed.

If you have the toolbox file, *xspde.mltbx*, just open it and click on *install*.

Otherwise the Matlab path must pointing to the xSPDE folder and subfolders. If you have the folders, but not the toolbox, proceed as follows:

- Click on the Matlab HOME tab (top left), then Set Path
- Click on Add with Subfolders
- Find the xSPDE3 folder in the drop-down menu, and select it
- Click on close to save the path.

Next, type *clear* to clear old data, and enter the xSPDE inputs and functions into the command window. For the simplest of examples, do this by cutting and pasting from an electronic file of this manual. You can also use the Matlab built-in editor.

A typical script first defines parameters and function specifications, in a structure, then runs the simulation code with the parameter structure as an input, as follows:

```
r.[label1] = [parameter1]
r.[label2] = ...
r.da = @(a,w,r) [derivative]
xspde(r)
```

Note the following points to remember:

- The notation $r.[label1] = [parameter1]$ defines a parameter or function in the data structure r .

- There are many input possibilities, and they all have default values.
- You don't have to save the data if you just want an immediate plot
- The notation `r.da=@(a,w,r) [derivative] = ..` defines an inline function, da/dt .
- In this example, `a` is the stochastic variable, `w` the random noise, and `r` the internal data structure.
- For example, `r.t` is the current time, and `r.x`, `r.y`, `r.z`, the grid of coordinates.

2.2 Inputs

All xSPDE simulations use a structure *in* for input data. All functions use an internally generated structure *r*, combining both *in* and additional internally generated parameters. Any name will do for either structure, as long as you are consistent.

All xSPDE inputs have default values, which can be changed by typing parameters into the *in* structure. If you only need the first element of a vector or array, just input the value that is required, the remaining elements will be given their default value. All parameters used can be output using the documentation switch, `print = 1`.

2.2.1 Common inputs

The most common xSPDE input parameters are:

Label	Type	Default value	Description
<i>fields</i>	integer vector	1	Stochastic field variables
<i>noises</i>	integer vector	1	Number of noises
<i>dimension</i>	integer	1	Space-time dimensions
<i>name</i>	string	' '	Simulation name
<i>olabels</i>	string	'a'	Observable labels
<i>da</i>	function	0	The stochastic derivative
<i>define</i>	function	0	Defined field definition
<i>initial</i>	function	0	Function to initialize variables
<i>ensembles</i>	integer vector	[1,1,1]	Numbers of stochastic trajectories
<i>observe</i>	function	<i>a</i>	Observable function(s)
<i>linear</i>	function	0	Linear interaction picture function
<i>ranges</i>	real vector	[10,10,..]	Ranges in time and space
<i>steps</i>	integer	[1]	Intermediate steps per time point
<i>points</i>	integer vector	[51,35,..]	Output lattice points in [t,x,y,z]
<i>probability</i>	cell array of vectors	0	Binning ranges for probabilities
<i>images</i>	integer cell array	{0,..}	Movie images in time per 2D observable
<i>transforms</i>	vector cells	{[0 0 0 0],...}	Set to 1 for Fourier transforms in [t,x,y,z]
<i>scatters</i>	integer cell array	{0,..}	Set to s for s scatter plots in the observable
<i>print</i>	integer	0	Set to 1 for parameter outputs

A complete list of user inputs is given in Chapter (6).

2.2.2 Fields

The first concept to understand is the *fields* parameter. All stochastic variables are integrated as matrices. The first input data, *fields*, is the range of the first index, which is the number of variables or field components that are integrated. This has a default value of *fields* = 1. If required, *fields* can have a second value, like [1, 1]. The second index of *fields* gives the number of user defined auxiliary fields, which are sometimes required. These do not have a differential equation, but have a functional definition, and add to the number of field components available in equations or outputs.

Both types of fields are indexed by using the first index of the field variable, like $a(i, j)$. Index numbers i greater than *fields*(1) access the auxiliary fields, to give an extended vector of fields plus auxiliary fields. The second matrix index j accesses all other aspects of the field, including space indices and an ensemble index when several equations are integrated in parallel. These can be unpacked to give a higher-dimensional field array. This is useful if one wishes to address all the space indices independently, but in most cases it is simply not necessary.

2.2.3 Functions

The next concept to understand is the idea of a *function*. All important functionality in xSPDE uses modular functions. Functions can be specified inline, which is short and fast. An example of this is $r.da = @(a, w, r) w$. This defines the function da , which gives the stochastic derivative function. In this example, it simply returns the input value w . This user specified inline function is then known internally by the function handle $r.da$. Alternatively, external function handles can be used to define modular Matlab codes. This is available for complex functions that might have internal logic.

2.2.4 Observables

The final concept to understand is the idea of an *observable*. In a stochastic calculation there are many different averages that might be needed, from the fields themselves to moments, correlations, spectra, or maybe even more complex quantities. These are calculated internally and stored. One can also store the raw trajectories, but this is usually a very costly exercise in terms of memory. The functions that define what is averaged internally are called the *observe* functions. Once data is averaged internally, further functional transforms and plots of the averages are available.

2.2.5 Use the dot

Matlab formulae often require a 'dot' or a 'colon'. To multiply vectors element-wise, like $a_i = b_i c_i$, the notation $a = b .* c$ indicates all elements are multiplied. This is used in xspde to speed up calculations in parallel.

A formula for a stochastic field typically requires you to address the first index - which is the field component - and treat all the other elements in parallel. To do this, use the notation $a(1, :)$. The colon indicates that all the second index elements are being addressed in parallel.

Whenever a formula combines operations over spatial lattices or ensembles, remember to **USE THE DOT**.

2.3 Random walk

The first example is the simplest possible stochastic equation:

$$\dot{a} = w(t), \quad (2.1)$$

This is carried out numerically and graphed using $N = \text{points}(1)$ points. Since the first time point stored is always the initial value, there are $N - 1$ integration steps, each of length $dt = \text{ranges}(1)/(N - 1)$. As a result, the graphs have discrete steps, and more detail is obtained if more time steps are used. The default value is $N = 51$, which is in the *xpreferences* file. This is adjustable by the user. It can also be changed for a simulation, by inputting a new value of *points*.

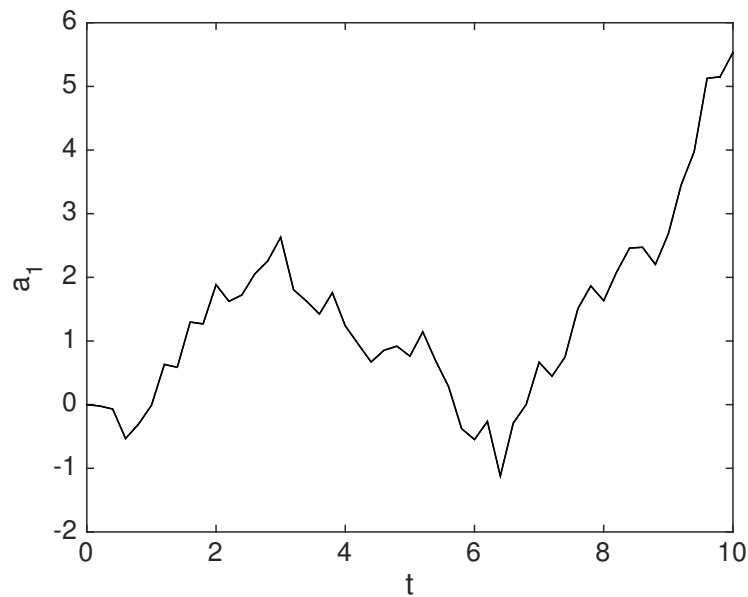


Figure 2.1: The simplest case: a random walk.

Exercises

Unless you type *clear* first, any changes to the input structure are additive; so you will get the sum of all the previous inputs as well as your new input.

- **Run the complete xSPDE script given below in Matlab.**

It is simplest to cut and paste from an electronic file into the Matlab command window. Be careful; pasting can cause subtle changes that may require correction. Some generated characters may be invalid Matlab input characters, and these will need retyping.

You should get the output in Fig (2.1).

```
r.da=@(a,w,r) w; xspde(r);
```

Here *r.da* defines the time derivative function \dot{a} , with *w* being the delta-correlated Gaussian noise that is generated internally.

- **What do you see if you average over 10000 trajectories ?**

```
r.ensembles = 10000;
xspde(r);
```

- **What do you see if you plot the mean square distance? Note that variances should increase linearly with *t*.**

```
r.observe = @(a,r) a.^2;
r.olabels = '<a^2>';
xspde(r);
```

- What if you add a force that takes the particle back to the origin?

$$\dot{a} = -a + w(t), \quad (2.2)$$

```
r.da=@(a,w,r) -a+w;
xspde(r);
```

Mathematical exercise:

- As an offline exercise, try to solve the equivalent diffusion equation for the probability P to find the variance $\langle a^2 \rangle$! The equation is:

$$\frac{\partial P(a)}{\partial t} = \left[\frac{\partial}{\partial a} + \frac{1}{2} \frac{\partial^2}{\partial a^2} \right] P(a)$$

2.4 Laser quantum noise

Next we treat a model for the quantum noise of a single mode laser as it turns on, near threshold:

$$\dot{a} = ga + bw(t) \quad (2.3)$$

where the noise is complex, $w = (w_1 + iw_2)$, so that:

$$\langle w(t)w^*(t') \rangle = 2\delta(t - t'). \quad (2.4)$$

Here the coefficient b describes the quantum noise of the laser, and is inversely proportional to the equilibrium photon number.

Exercises

You should type *clear* first when starting new simulations.

- **Solve for the case of $g = 0.1$, $b = 0.01$**

Most lasers have more than 100 photons, and hence less noise!

For this exercise, small error-bars will display on the graph. These are calculated from the difference between using steps of size dt and steps of size $dt/2$. They only appear if greater than a minimum relative size, typically 1% of the graph size, which can be set by the user.

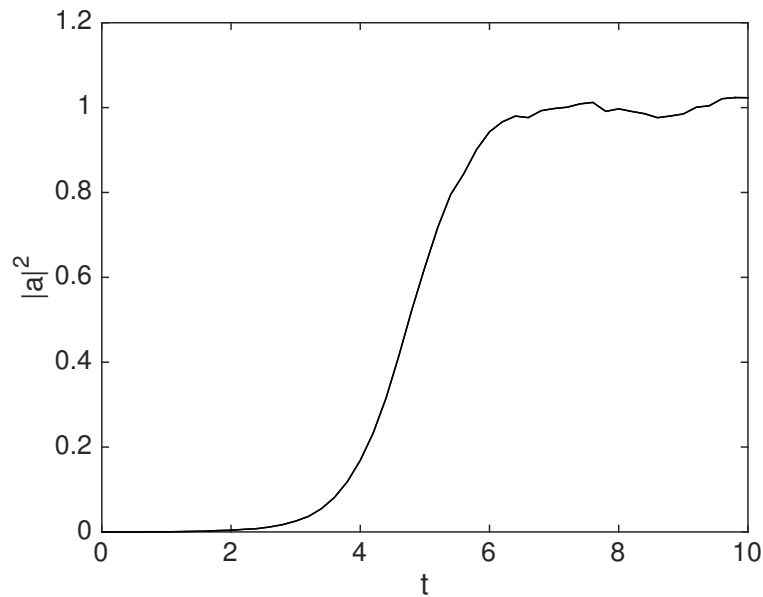


Figure 2.2: Simulation of the stochastic equation describing a laser turning on.

```
clear
r.noises=2;
r.observe = @(a,r) abs(a)^2;
r.olabels = '|a|^2';
r.da=@(a,w,r) a+0.01*(w(1)+i*w(2));
xspde(r);
```

Consider the case where the laser saturates to a steady state:

$$\dot{a} = \left(1 - |a|^2\right) a + bw(t) \quad (2.5)$$

- **Solve for the saturated laser case**

You should get the output graph in Fig (2.2).

```
r.da=@(a,w,r) (1-abs(a)^2)*a+0.01*(w(1)+i*w(2));
xspde(r);
```

2.5 Fourier transforms

Frequency spectra have many uses, especially for understanding the steady-state fluctuations of any physical system. The xSPDE spectral definition is:

$$\tilde{a}(\omega) = \frac{1}{\sqrt{2\pi}} \int e^{-i\omega t} a(t) dt$$

To get an output from a temporally Fourier transformed field, just set $transforms\{n\} = 1$, for the particular observable (n) you need to calculate in transform space. This parameter is a cell array. It can have a different value for every observable and for every dimension in space-time if you have space dimensions as well.

As an example, take the random walk equation in a complex space,

$$\frac{da}{dt} = -a + \zeta(t)$$

where $\zeta(t) = w_1(t) + iw_2(t)$. Use an initial condition of $a = (v_1 + iv_2)/\sqrt{2}$, with $\langle v_i^2 \rangle = 1$, to start near the steady state.

For sufficiently long time-intervals, the solution in frequency space - where $\omega = 2\pi f$ is the angular frequency - is given by:

$$\tilde{a}(\omega) = \frac{\tilde{\zeta}(\omega)}{1 + i\omega}$$

The expectation value of the noise spectrum, in the long time-intervals limit, is therefore:

$$\begin{aligned} \langle |\tilde{a}(\omega)|^2 \rangle &= \frac{1}{2\pi(1 + \omega^2)} \int \int e^{-i\omega(t-t')} \langle \zeta(t)\zeta^*(t') \rangle dt dt' . \\ &= \frac{T_{eff}}{\pi(1 + \omega^2)} \end{aligned}$$

The final result is obtained from the fact that the first integral vanishes unless $t = t'$, since for small steps, $\langle \zeta(t)\zeta^*(t') \rangle \rightarrow 2\delta(t - t')$.

The time integral is carried out numerically as a sum which has $N = points(1)$ time points of interval dt . The definition of dt is therefore $dt = T/(N - 1)$, where $T = ranges(1)$. The ‘effective’ time duration for the Fourier transform time integrals is $T_{eff} = Ndt = T \times N/(N - 1) = T + dt$. This corresponds to a midpoint rule integration, which technically is over the interval $[-dt/2, T + dt/2]$.

Exercises

- **Plot the spectrum over a range of $t = 100$, with 640 points and a random initial equation near the equilibrium value.**

The input parameters are given below. There are parallel operations here, for ensemble averaging, so **USE THE DOT**.

```

clear
r.points = 640;
r.ranges = 100;
r.noises = 2;
r.ensembles = 10000;
r.initial = @(v,r) (v(1,:)+1i*v(2,:))/sqrt(2);
r.da = @(a,w,r) -a + w(1,:)+1i*w(2,:);
r.observe = @(a,r) a.*conj(a);
r.transforms =1;
r.olabels = '|a(\omega)|^2';
xspde(r);

```

Note that `r.transforms =1` tells xSPDE to Fourier transform the field over the time coordinate before averaging, to give a spectrum. The first argument v of the *initial* function is a random field, used optionally to initialize the stochastic variable. You can transform any field in any dimension, for any number of resulting averages. See Chapter 6 for details.

To define as many observables as you like, use a Matlab cell array;

```

r.observe{1} = ..;
r.observe{2} = ..;

```

- **Simulate over a range of $t = 200$. What changes do you see? Why?**
- **Change the equation to the laser noise equations. Why is the spectrum much narrower?**

2.6 Probability density

It is possible to graph probability densities of a real observable. This is achieved through specifying the observable and the binning ranges, by inputting:

$$r.probability\{n\} = o1 : ostep : o2;$$

If present, this returns probability density of variable $o(n)$, through binning into ranges of width *ostep* around centers starting at *o1*, and ending at *o2*. Returns a result of $1/ostep$ in the j -th bin if the trajectory is inside the bin, so that $o(j) - ostep/2 < o < o(j) + ostep/2$, and zero otherwise. This gives a probability graph on output, plotted against time. Note that on graphing, an extra dimension is added for the variable o . Only probabilities of one-dimensional real variables are plotted, for interface simplicity. The probability can be for any function of the stochastic variable if required.

The *xgraph* program provides facilities for extracting slices and windows of the probabilities where required. To plot the probabilities in the first observable for a range of 0:1, add the following input before the *xsim* or *xspde* command:

```

r.probability{1}=0:0.1:1.01;

```

2.7 Ito and Stratonovich

The xSPDE toolbox is mainly designed to treat Stratonovich equations [2], which are the broad-band limit of a finite band-width random noise equation. Another type of stochastic equation is the Ito form, which is a limit where derivatives are evaluated at the start of each step. If you have an Ito equation, you can either use the Euler integration option, and set *r.step = xEuler*, or else apply the Stratonovich correction to the drift, as outlined below.

To emphasize the difference, an Ito equation is often written as a difference equation:

$$d\mathbf{a} = \mathbf{A}^I[\mathbf{a}] + \mathbf{B}[\mathbf{a}] \cdot d\mathbf{w}(t). \quad (2.6)$$

Here $\langle dw_i(\mathbf{x}) dw_j(\mathbf{x}') \rangle = \delta_{ij} dt$.

When \mathbf{B} is constant, the two methods are identical. If \mathbf{B} is not constant, the Ito drift term \mathbf{A}^I is different to the corresponding Stratonovich one. The relationship is:

$$A_i = A_i^I - \frac{1}{2} \sum_{j,m} \frac{\partial B_{ij}}{\partial a_m} B_{mj}. \quad (2.7)$$

2.8 Financial calculus

A well-known Ito stochastic equation is the Black-Scholes equation, used to price financial options. It describes the fluctuations in a stock value:

$$da = \mu a dt + a \sigma dw, \quad (2.8)$$

where $\langle dw^2 \rangle = dt$. Since the noise is multiplicative, the equation is different in Ito and Stratonovich forms of stochastic calculus.

The corresponding Stratonovich equation, as used in xSPDE for the standard default integration routine is:

$$\dot{a} = (\mu - \sigma^2/2) a + a \sigma w(t). \quad (2.9)$$

Exercises

- Solve for a startup with a volatile stock having $\mu = 0.1$, $\sigma = 1$.

An interactive xSPDE script in Matlab is given below with an output graph in Fig (2.3), Note the spiky behaviour, typical of multiplicative noise, and also of the risky stocks in the small capitalization portions of the stock market.

```
clear
r.initial=@(v,r) 1;
r.da=@(a,w,r) -0.4*a+a*w;
xspde(r);
```

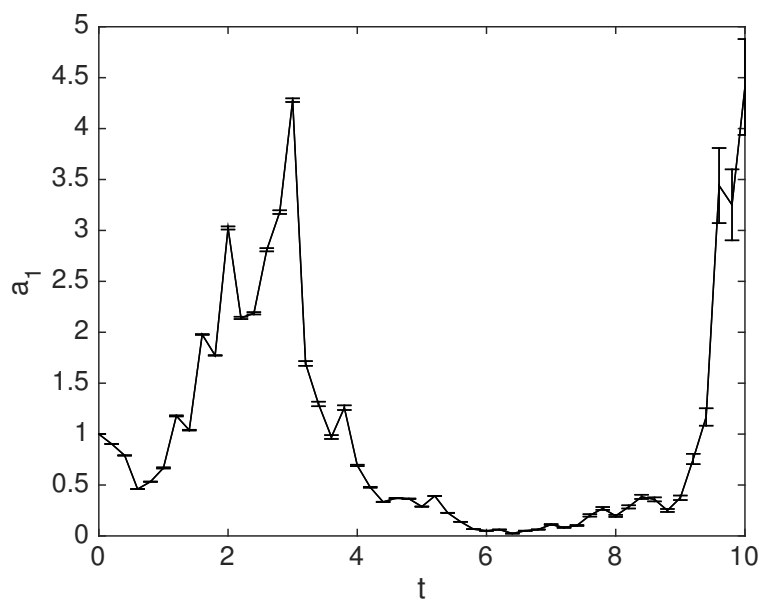


Figure 2.3: Simulation of the Black-Scholes equation describing stock prices.

Note that $r.initial$ describes the initialization function. The first argument of $@(v, r)$ is v , an initial random variable with unit variance. *The error-bars are estimates of step-size error.* Errors can be reduced by using more time-steps.

- **Solve for a more mature stock having $\mu = 0.1$, $\sigma = 0.1$.**

3 xSPDE projects

Projects are larger scale investigations, where files are used to record inputs and data. This requires a new input parameter:

Label	Type	Default value	Description
<i>r.file</i>	string	"	Matlab or HDF5 data file

3.1 Ensembles and errors

Averages over stochastic ensembles are the specialty of xSPDE, which requires specification of the ensemble size. A sophisticated hierarchy of ensemble specifications in three levels is possible, to allow maximum resource utilization, so that:

$$r.ensembles = [ensembles(1), ensembles(2), ensembles(3)].$$

The local ensemble, *ensembles*(1), gives within-thread parallelism, allowing vector instruction use for single-core efficiency. The serial ensemble, *ensembles*(2), gives a number of independent trajectories calculated serially.

The parallel ensemble, *ensembles*(3), gives multi-core parallelism, and requires the Matlab parallel toolbox. This improves speed when there are multiple cores. One should optimally put *ensembles*(3) equal to the available number of CPU cores.

The *total* number of stochastic trajectories or samples is

$$ensembles(1) \times ensembles(2) \times ensembles(3).$$

Either *ensembles*(2) or *ensembles*(3) are required if sampling error-bars are to be calculated, owing to the sub-ensemble averaging method used in xSPDE to calculate sampling errors accurately.

3.2 xSIM and xGRAPH

An XPDE session can either run simulations interactively, described in Chapter 2, or else using a function file called a project file. This allows xSPDE to run in a batch mode, as needed for longer projects which involve large ensembles. In either case, the Matlab path must include the xSPDE folder. For generating graphs automatically, the script input or project function should end with the combined function **xspde**.

With batch jobs it is often useful to divide xSPDE into its simulation function, xSIM, and its graphics function, xGRAPH, to allow graphs to be made at a later time from

the simulation. In this case the function *xsim* runs the simulation, and *xgraph* makes the graphs. The two-stage option is better for running batch jobs, which you can graph at a later time.

To create a data file, you must enter the filename when running the simulation, using the *r.file = filename* input. A typical xSPDE project function of this type is as follows:

```
function [e,ec] = project.m
    r[label1] = [parameter1];
    r[label2] = ...;
    r.file = '[my file].mat';
    [e,r] = xsim(r);
    ec = xgraph(r.file);
end
```

After preparing a project file using the editor, click on the Run arrow above the editor window.

A batch job workflow is as follows:

- Create the metadata *r*, and include a file name, *r.file*.
- Change the Matlab directory path to home using *cd ~*.
- Run the simulation with **[e,r]=xsim(r)**, or just **xsim(r)**;
- Note: *xsim* changes the file-name if it already exists, and returns the new name for plotting!
- Run **xgraph(r.file)**, or **xgraph(r.file,r)** if you want to edit the inputs in the structure *r*, and the data will be graphed.
- You can use either Matlab (.mat) or standard HDF5 (.h5) file-types.

3.3 Kubo project

To get started on more complex programs, we next simulate the Kubo oscillator, which is an oscillator with a random frequency:

$$\dot{a} = iaw \tag{3.1}$$

Exercises

- **Simulate the Kubo oscillator using a file, *Kubo.m*, with two ensembles to allow sampling error estimates. The error vector *error* gives the time-step error plus the sampling error.**

```

function [error] = Kubo()
r.name = 'Kubo oscillator';
r.ensembles = [400,16];
r.initial = @(v,r) 1+0*v;
r.da = @(a,w,r) i*a.*w;
r.olabels = {'<a_1>'};
r.file = 'kubo.mat';
[error,r]=xsim(r);
xgraph(r.file);
end

```

The initial value of 1 is modified by adding $0 * v$. This is necessary, so that it returns an array whose first dimension equals the ensemble size.

This function is designed to generate a data file, `kubo.mat`. If you run this more than once without deleting the earlier file, you will get a warning and a new file-name, `kubo1.mat`, which is stored in the in structure. Under these circumstance, `xGRAPH` will graph the data in the most recent file saved, with the new file-name. This occurs because `xSPDE` will **not** overwrite old files, to protect your previous data.

You can also include modified graphics parameters as a second input when running `xGRAPH`, just in case the first graphs you generate need changes.

3.4 Challenge problem #1

This is a harder example, involving a full nonlinear quantum phase-space simulation. The method can also be used to investigate quantum non-equilibrium phase transitions, tunneling in open systems, quantum entanglement, Einstein-Podolsky-Rosen paradoxes, Bell violations, and many other problems treated in the literature[2, 3].

The following new ideas are introduced:

1. **ranges is the space-time integration range.**
2. **steps gives integration steps per time point, for increased accuracy.**

A simple case is the nonlinear driven quantum ‘time crystal’ or subharmonic generator - for example, an optomechanical, superconducting or nonlinear optical medium in a driven cavity.

This has a Stratonovich equation, which is the form treated by the `xspde` software:

$$\begin{aligned}\frac{da_1}{d\tau} &= -(c + \frac{1}{2})a_1 + a_2\lambda [1 - a_1^2] + \sqrt{1 - a_1^2}\eta_1(\tau) \\ \frac{da_2}{d\tau} &= -(c + \frac{1}{2})a_2 + a_1\lambda [1 - a_2^2] + \sqrt{1 - a_2^2}\eta_2(\tau)\end{aligned}$$

There is a bistable region, which leads to a discrete time symmetry breaking. The solution in the steady-state is

$$P = (1 - a_1^2)^c (1 - a_2^2)^c e^{2\lambda a_1 a_2}$$

The integration manifold is the region of real a_1, a_2 , such that $a_1^2 \leq 1, a_2^2 \leq 1$. There are two physically possible metastable values of the amplitudes. The physically observed quantity is the amplitude and number:

$$\begin{aligned}\langle \hat{a} \rangle &= \frac{1}{2} \langle a_1 + a_2 \rangle \\ \langle \hat{n} \rangle &= \langle a_1 a_2 \rangle\end{aligned}$$

Typical parameters that show bistable behaviour are $c = 1, \lambda = 4$.

Exercises

- **Simulate the nonlinear oscillator by creating a file, *Nonlinear.m***
- **Can you observe quantum tunneling in the bistable regime?**
- **Do you see transient Schrodinger ‘cat states’ with a negative n value?**

Since tunneling is random, for real experimental comparisons, one would have to measure correlation functions and spectra. However, a tunneling event in a simulation indicates that it is likely in experiment too. These calculations require long time scales, **in.ranges**, to observe tunneling, and a large number of time steps per plotted time point, **in.steps**, to maintain accuracy in the quantum simulations.

4 Stochastic PDEs

A stochastic partial differential equation or *SPDE* for a complex vector field is defined in both time t and space dimensions \mathbf{x} . The total number of *fields* or internal indices is unlimited. The total space-time *dimension*, D in xSPDE is also unlimited. It includes time ($D = 1$) and space ($D > 1$). Large numbers of space-time dimensions are memory-intensive and slow, with a currently practical limit of about $D = 6$ unless you have a very large memory and fast computer!

The general equation solved can be written in differential form as

$$\frac{\partial \mathbf{a}}{\partial t} = \mathbf{A}[\mathbf{a}] + \mathbf{B}[\mathbf{a}] \cdot \mathbf{w}(t) + \mathbf{L}[\nabla, \mathbf{a}] . \quad (4.1)$$

Here \mathbf{a} is a real or complex vector field, with '*fields*' components, and $\mathbf{w} = [\mathbf{w}^x, \mathbf{w}^k]$ is a random field, with '*noises*' components. This is a vector such that: *noises*(1)' is the number of x-correlated noises, \mathbf{w}^x , and *noises*(2)' is the number of k-correlated noises, \mathbf{w}^k . These correlations are user-adjustable, explained below.

Initial conditions are set at the initial time of $t = o_1$ with a user-defined function so that:

$$a(o_1) = \text{initial}(v, r)$$

The *initial* function includes '*randoms*' initial random fields $v = [v^x, v^k]$. Their correlations are also defined either in x-space or in k-space. To allow this, the input parameter '*randoms*' is a vector such that: '*randoms*(1)' is the number of x-correlated random fields, v^x , and '*randoms*(2)' is the number of k-correlated random fields, v^k . All random fields in the *initial* function, even if defined with momentum correlations, are transformed to position space before use. If there is no scaling of the momentum noise, v^x and v^k have the same correlations.

Transverse *boundaries* are of three types: Neumann (specified derivative), periodic, or Dirichlet (specified field). These are indicated using *boundaries* = -1, 0, 1, which can be specified for each space dimension, field index and boundary, as explained below in greater detail.

The spatial derivatives or other functions linking different spatial points can be specified either in the functions A , B , or else in the *linear* function, if the derivative terms are linear functions of the fields. Use of the *linear* function allows an interaction picture algorithm, with increased efficiency. Currently, the *linear* function is only available in conjunction with a Fourier transform propagator.

4.1 Coordinates and indices

The spatial interval used is from a preset input *origin* \mathbf{o} , and has predetermined input *ranges* \mathbf{r} . If the *origin* or the *ranges* are not input, they have default values. The default origin of the first (time) coordinate starts at $t = 0$. Spatial coordinates have default origins that give symmetric intervals, with the origin at $-\mathbf{r}/2$. Dimensions are indexed with the first (time) dimension as $i = 1$. As an example, for the x-dimension, the problem is solved on an interval of $x = [o_2, o_2 + r_2]$, with a default of $x = [-r_2/2, r_2/2]$.

In order to discretize the problem for solution the n_i lattice points are fitted into the spatial interval so that $dx_i = r_i/(n_i - 1)$, ie:

$$x_i = o_i + (i - 1)dx_i. \quad (4.2)$$

Stochastic data To store stochastic data like field variables that are used in the stochastic equations, the following convention is always used. The logical array index ordering is $fields(i, \mathbf{j}, e)$, where:

- The first index is a field index (i).
- The d indices $j_1, \dots, j_d \equiv \mathbf{j}$ are for time and space, or momentum space if Fourier transformed.
- The last index is an ensembles index (e), used to index over field trajectories.
- For storing space coordinates like $r.x$, the first two indices are $i, j_1 = 1$, and are not used.
- Transient arrays in calculations use a reduced space-time index $\mathbf{j}' = [1, j_2, \dots]$, with time index equal to one.
- Arrays in calculations are packed to have indices $a(i, J)$, where $J = (\mathbf{j}', e)$ is a flattened index.

The dimensionality of the fields being integrated is logically stored in the array shape $r.d.fields = (i, \mathbf{j}', e)$, which is packed for calculations into a matrix labelled \mathbf{a} . The use of a two-dimensional index in internal storage simplifies many functions, and this two-dimensional shape is stored in the internal matrix shape $r.d.a = (i, J)$. Where Fourier transforms are used internally for linear propagation, the momentum arrays start at $k = 0$ in the first index.

A **packed** matrix, a , can always be unpacked into its full index array form using:

$$fields = reshape(a, r.d.fields)$$

An **unpacked** array, $fields$, can always be packed into its compressed array form using:

$$a = reshape(fields, r.d.a)$$

Averaged data The array index ordering for averages generated by the *observe* functions is still logically (i, j, e) , although there are differences, as follows:

- Averaged data for storing or graphing is stored in unpacked form, including time for stored data, so that all indices can be readily accessed for graphing.
- Data obtained from functions of fields retains the first index to indicate different output lines on a graph. This also allows a vector average to be calculated.
- Averages that are defined in momentum space have the momentum origin at $k = 0$ is shifted to become the central index for graphing.
- Data that is ensemble averaged has no ensemble index, so the last index is used for storing error-bars instead.
- Where integrals or spatial averages are used, the entire array is filled to keep them homogeneous, although this will be changed in future.
- Where data is binned to get distributions, an extra 'space' dimension is added to j to store the binning index in each space-time location.

4.2 Boundary conditions

Boundary conditions must be given for all partial differential equations. The default boundary conditions are periodic. The implicit setting of this is that periodicity is enforced such that $a(o_i - dx_i/2) = a(o_i + r_i + dx_i/2)$, which is the usual discrete Fourier transform requirement.

Otherwise, the differential equation boundaries are specified at $a(o_i)$, $a(o_i + r_i)$, using the cell-array input $boundaries\{d\}(i, j)$, which is defined per space dimension ($d = 2, 3..$), field index ($i = 1, 2..$) and boundary $j = (1, 2)$. Here $d > 1$ is the transverse dimension, not including time, which only has an initial condition. The available boundary types are:

Neumann: For specified *derivative* boundaries, $boundaries\{d\}(i, j) = -1$

Periodic: For *periodic* boundaries, $boundaries\{d\}(i, j) = 0$

Dirichlet: For specified *field* boundaries, $boundaries\{d\}(i, j) = 1$

These are specified in a cell array: $boundaries\{d\}(i, 1)$ sets the lower boundary type, while $boundaries\{d\}(i, 2)$ gives the upper boundary type. Each space dimension, variable and boundary is set independently. Note that in xspde, the equations are always initial value problems in time, so the time dimension boundary specification for $d = 1$ is not included. The default option is periodic boundaries if *boundaries* is not specified. The boundary values returned by the default boundary function, *xboundfun*, are all zero.

4.2.1 Setting the boundary values

For non-vanishing, specified boundary conditions, the boundary function $boundfun(a, d, r)$ is called. This returns the boundary values used for the fields or derivatives in dimension $d > 1$ as an array of dimension $b(i, J, j, K)$.

Here i is the field index, J is a flattened dimension equal to the product of field array dimensions for $d_1 < d$, and K is a flattened dimension equal to the product of field array dimensions for $d_2 > d$, including the ensemble dimension. Crucially, j is the index of the dimension whose boundary values are specified, so only two values are needed: $j = 1, 2$ for the upper and lower boundary values, which could either be field values or their derivatives.

Boundary values can be a function of both the fields (a) and internal variables like the current time ($r.t$). These may have predefined or stochastic initial values at $t = 0$ which are calculated only once. In such cases the boundary values must first be initialized, so the routine $boundfun(a, d, r)$ is internally initialized with time $t < origin(1)$, and with random Gaussian values in the input field a . These are delta-correlated in space, i.e., with the same definition as the “randoms”. The xSDPE program stores the returned values b for the boundaries in an internal cell array, $boundval\{d\}$, for later use if required.

The default initial boundary value is zero, set by the default boundary function $xboundfun(a, d, r)$. Subsequent boundary values returned by $xboundfun(a, d, r)$, are set equal to the initial boundary values stored in $boundval\{d\}$.

4.3 Fourier transforms

It is often useful to Fourier transform a field before it is averaged. This is controlled independently for each *observe* function and space-time dimension by *transforms*. The xSPDE Fourier transforms or spectrum definitions in space-time are given by the symmetric Fourier transform definition:

$$\begin{aligned}\tilde{a}(\omega, \mathbf{k}) &= \mathcal{F}(a(t, \mathbf{x})) \\ &= \frac{1}{[2\pi]^{d/2}} \int e^{i(\omega t - \mathbf{k} \cdot \mathbf{x})} a(t, \mathbf{x}) dt d\mathbf{x}\end{aligned}\quad (4.3)$$

The inverse Fourier transform is the function:

$$\begin{aligned}a(t, \mathbf{x}) &= \mathcal{F}^{-1}(\tilde{a}) \\ &= \frac{1}{[2\pi]^{d/2}} \int e^{i(\mathbf{k} \cdot \mathbf{x} - \omega t)} \tilde{a}(\omega, \mathbf{k}) d\omega d\mathbf{k}\end{aligned}$$

This is translated into a sum over the lattice points using a discrete Fourier transform at the lattice points x_i , so that:

$$\tilde{a}(\omega_i, \mathbf{k}_i) = \frac{dt d\mathbf{x}}{[2\pi]^{d/2}} \sum_{j_1 \dots j_d} \exp[i(\omega_{i_1} t_{j_1} - \mathbf{k}_i \cdot \mathbf{x}_j)] a(t_{j_1}, \mathbf{x}_j) \quad (4.4)$$

The momenta k_i have an interval of

$$dk_i = \frac{2\pi}{n_i dx_i} \quad (4.5)$$

with k_i values given for even n by:

$$k_i = \left(1 - \frac{n_i}{2}\right) dk_i, \dots \frac{n_i}{2} dk_i \quad (4.6)$$

and for odd n by:

$$k_i = \frac{1 - n_i}{2} dk_i, \dots \frac{n_i - 1}{2} dk_i \quad (4.7)$$

Once Fourier transformed, the *observe* function can be used to take any further functions or combinations of Fourier transformed fields prior to averaging. Important points to keep in mind are as follows:

- Fourier transforms are specified for the k -th *observe* function independently of all other functions, by specifying $transforms\{k\} = [\ell_1, \dots, \ell_d]$,
- Here $\ell_j = 0, 1$ is a logical switch, set to $\ell_j = 1$ if the j -th dimension requires a Fourier transform, and $\ell_j = 0$ if there is no Fourier transform.
- The internal fields $r.k\{1\}, \dots, r.k\{d\}$ are available for use in making functions of momentum for use with observations.
- In propagation calculations, the momentum lattice values start with $k = 0, \dots$, following standard Matlab and FFT conventions.
- For storing and graphing, momentum lattice values are reordered to start with $k = -k_{max}, \dots$, following standard graphics and mathematical conventions.

A stochastic partial differential equation or *SPDE* for a complex vector field is defined in both time t and space dimensions \mathbf{x} . The total *dimension* d in xSPDE is unlimited, and includes time and space. Large d is memory-intensive and slow! The general equation solved can be written in differential form as

$$\frac{\partial \mathbf{a}}{\partial t} = \mathbf{A}[\mathbf{a}] + \mathbf{B}[\mathbf{a}] \cdot \mathbf{w}(t) + \mathbf{L}[\nabla, \mathbf{a}]. \quad (4.8)$$

The linear function L is input as a separate function in xSPDE, to allow for greatest efficiency and use of an interaction picture. Note that it depends on momentum space coordinates, and this involves Fourier transforms.

4.3.1 Specifying the equations

Details of how to input the equations are as follows:

- The initial conditions are specified by the user from the *initial*(v, r) function, where $\mathbf{v} = [\mathbf{v}^x, \mathbf{v}^k]$ is an array of random initial values, delta-correlated in space or momentum space. For space correlations, set *randoms*(1), for momentum correlations set *randoms*(2). Here $f^{(r)}(\mathbf{v}(\mathbf{k}))$ is an arbitrary momentum filter, for correlated noise. Set this using the *rfilter*(v, r) function, specified by the user.
- $\mathbf{A}[\mathbf{a}]$ and $\mathbf{B}[\mathbf{a}]$ are vector and matrix functions of \mathbf{a} , set by the user using the *da*(a, w, r) function.
- Here, $\mathbf{w} = [\mathbf{w}^x, \mathbf{w}^k]$ are real noise fields, delta-correlated in time, and also in space or momentum, as explained below.
- The *linear* function L is input as a function in xSPDE, to allow for greatest efficiency and use of an interaction picture. Here $\mathbf{L}[\nabla, \mathbf{a}]$ is a matrix of *linear* terms including derivatives, diagonal in the array indices. Note that it depends on momentum space coordinates, and generally involves Fourier transforms. If $d = 1$, there are no space dimensions or derivatives. The term \mathbf{L} may be omitted if not required.

4.3.2 Initial random fields and noise transforms

Fourier transforms are available for use both on initial random values and on noise fields during time-evolution. This is controlled by the second element of *randoms* and *noises* respectively.

When *randoms*(1) > 0, an initial random field \mathbf{v}^x is generated with delta-correlations in x -space. When *randoms*(2) > 0, an initial random field $\tilde{\mathbf{v}}^k$ is generated with delta-correlations in k -space. This is filtered with a user-specified filter function to give $\tilde{\mathbf{v}}^{kf}$, then inverse Fourier transformed to give \mathbf{v}^k . Both random fields are passed to the *initial* function as an extended vector $[\mathbf{v}^x, \mathbf{v}^k]$, for field *initialization* in space.

There is a user specified filter function available, to modify random fields $\tilde{\mathbf{v}}^k$, that are delta-correlated in momentum space using a filter function, '*rfilter*' so that $\tilde{v}_i^{kf}(\mathbf{k}) = f_i^{(r)}(\mathbf{v}^k(\mathbf{k}))$, before being used. The corresponding correlations are:

$$\begin{aligned}
\langle v_i^x(\mathbf{x}) v_j^x(\mathbf{x}') \rangle &= \delta(\mathbf{x} - \mathbf{x}') \delta_{ij} \sim \frac{1}{\Delta V} \delta_{\mathbf{x}, \mathbf{x}'} \delta_{ij} \\
\langle \tilde{v}_i^k(\mathbf{k}) \tilde{v}_j^k(\mathbf{k}') \rangle &= \delta(\mathbf{k} - \mathbf{k}') \delta_{ij} \sim \frac{1}{\Delta K} \delta_{\mathbf{k}, \mathbf{k}'} \delta_{ij} \\
\langle \tilde{v}_i^{kf}(\mathbf{k}) \tilde{v}_j^{kf}(\mathbf{k}') \rangle &= \langle f_i^{(r)}(\tilde{\mathbf{v}}^k(\mathbf{k})) f_j^{(r)}(\tilde{\mathbf{v}}^k(\mathbf{k}')) \rangle.
\end{aligned} \tag{4.9}$$

Note that on a lattice, we replace the Dirac continuous delta-function by a discrete Kronecker delta function scaled by an inverse volume element either in space (ΔV) or momentum (ΔK). The xSPDE Fourier transforms are given by a symmetric Fourier transform definition, so that if we inverse Fourier-transform the k -randoms, without filtering, then:

$$v^k(\mathbf{x}) = \frac{1}{[2\pi]^{(d-1)/2}} \int e^{i\mathbf{k} \cdot \mathbf{x}} \tilde{\mathbf{v}}^k(\mathbf{k}) d\mathbf{k} \tag{4.10}$$

These have random initial values that are real and delta-correlated in space, so that:

$$\langle v^x(\mathbf{x}) v^x(\mathbf{x}') \rangle = \delta(\mathbf{x} - \mathbf{x}'). \quad (4.11)$$

The corresponding noises in position space are correlated according to:

$$\begin{aligned} \langle v^k(\mathbf{x}) (v^k(\mathbf{x}'))^* \rangle &= \frac{1}{[2\pi]^{(d-1)}} \int e^{i(\mathbf{k} \cdot \mathbf{x} - \mathbf{k}' \cdot \mathbf{x}')} \langle \tilde{v}^k(\mathbf{k}) \tilde{v}^k(\mathbf{k}') \rangle d\mathbf{k} d\mathbf{k}' \\ &= \frac{1}{[2\pi]^{(d-1)}} \int e^{i(\mathbf{x} - \mathbf{x}') \cdot \mathbf{k}} d\mathbf{k} \\ &= \delta(\mathbf{x} - \mathbf{x}'). \end{aligned} \quad (4.12)$$

Similarly, if we don't conjugate the k-noise, then:

$$\langle v^k(\mathbf{x}) v^k(\mathbf{x}') \rangle = \delta(\mathbf{x} + \mathbf{x}'). \quad (4.13)$$

However, if we define $\tilde{v}^c(\mathbf{k}) = [\tilde{v}_1^k(\mathbf{k}) + i\tilde{v}_2^k(\mathbf{k})] / \sqrt{2}$, then we obtain complex noise is only delta correlated when conjugated.

$$\begin{aligned} \langle v^c(\mathbf{x}) (v^c(\mathbf{x}'))^* \rangle &= \delta(\mathbf{x} - \mathbf{x}') \\ \langle v^c(\mathbf{x}) v^c(\mathbf{x}') \rangle &= 0. \end{aligned} \quad (4.14)$$

This is obtainable with the x-space noise as well, but the utility of the k-space noise is that it can be filtered to have nonlocal correlations in space if required.

During propagation in time, $\mathbf{w} = [\mathbf{w}^x, \mathbf{w}^k]$ are real noise fields that are delta-correlated in space-time. They are calculated in an analogous way, except with an additional factor of $1/\sqrt{dt}$ because they are delta correlated in time as well. There is a user specified scaling function available, to take random noises w^k in momentum space that are then scaled using a filter function, 'nfilter' so that $w_i^{kf}(\mathbf{k}) = f_i^{(n)}(\mathbf{w}^k(\mathbf{k}))$, before being used:

$$\begin{aligned} \langle w_i^x(t, \mathbf{x}) w_j^x(t, \mathbf{x}') \rangle &= \delta(\mathbf{x} - \mathbf{x}') \delta(t - t') \delta_{ij} \\ \langle \tilde{w}_i^k(t, \mathbf{k}) \tilde{w}_j^k(t, \mathbf{k}') \rangle &= \delta(\mathbf{k} - \mathbf{k}') \delta(t - t') \delta_{ij} \\ \langle \tilde{w}_i^{kf}(t, \mathbf{k}) \tilde{w}_j^{kf}(t', \mathbf{k}') \rangle &= \langle f_i^{(n)}(\tilde{\mathbf{w}}^k(t, \mathbf{k})) f_j^{(n)}(\tilde{\mathbf{w}}^k(t', \mathbf{k}')) \rangle. \end{aligned} \quad (4.15)$$

4.4 Stochastic Ginzburg-Landau

Including two space dimensions, or space-time dimensions of $d = 3$, an example of a SPDE is the stochastic Ginzburg-Landau equation. This describes symmetry breaking. The system develops a spontaneous phase which varies spatially as well. The model is used to describe lasers, magnetism, superconductivity, superfluidity and particle physics:

$$\dot{a} = (1 - |a|^2) a + bw(t) + c\nabla^2 a \quad (4.16)$$

where

$$\langle w(x)w^*(x') \rangle = 2\delta(t-t')\delta(x-x'). \quad (4.17)$$

The following new ideas are introduced for this problem:

1. dimension is the space-time dimension.
2. The 'dot' notation used for parallel operations over lattices.
3. linear is the linear operator - a laplacian in these cases.
4. images produces movie-style images at discrete time slices.
5. Dx indicates a derivative operation, $\partial/\partial x$.
6. $-5 < x < 5$ is the default xSPDE coordinate range in space.

Exercises

- Solve the stochastic G-L equation for $b = 0.001$ and $c = 0.01i$.
- Change to a real diffusion so that $c = 0.1$.

In the first case, you should get the output graphed in Fig (4.1) .

```
clear;
r.name = 'Extended laser gain equation';
r.noises=2;
r.dimension=3;
r.steps=10;
r.linear = @(r) i*0.01*(r.Dx.^2+r.Dy.^2);
r.observe = @(a,~) abs(a).^2;
r.images =6;
r.olabels = '|a|^2';
r.da=@(a,w,~) (1-abs(a(1,:)).^2).*a+0.001*(w(1,:)+i*w(2,:));
xspde(r)
```

Here the notation $z(1,:)$ is like the dot: the colon means that the operation is repeated over all values of the second index, which is the spatial lattice index.

4.5 Soliton

The famous nonlinear Schrödinger equation (NLSE) is:

$$\frac{da}{dt} = \frac{i}{2} [\nabla^2 a - a] + ia|a|^2$$

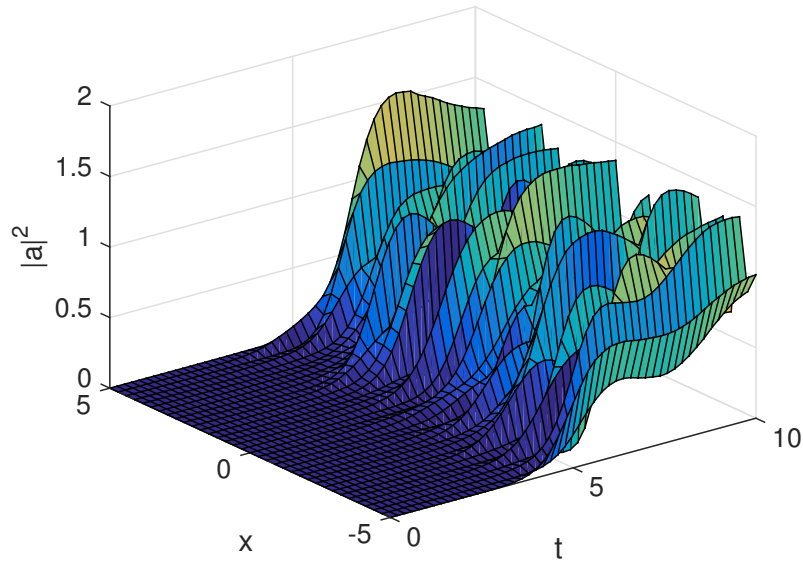


Figure 4.1: Simulation of the stochastic equation describing symmetry breaking in two dimensions. Spatial fluctuations are caused by the different phase-domains that interfere. The graph obtained here is projected onto the $y = 0$ plane.

Together with the initial condition that $a(0, x) = \text{sech}(x)$, this has a soliton, an exact solution that doesn't change in time:

$$a(t, x) = \text{sech}(x)$$

The Fourier transform at $k = 0$ is simply:

$$\tilde{a}(t, 0) = \frac{1}{\sqrt{2\pi}} \int \text{sech}(x) dx = \sqrt{\frac{\pi}{2}}$$

Exercises

- Solve the NLSE for a soliton

```
function [e] = Soliton()
r.name = 'NLS soliton';
r.dimension = 2;
r.initial = @(v,r) sech(r.x);
r.da = @(a,~,r) i*a.*(conj(a).*a);
r.linear = @(r) 0.5*i*(r.Dx.^2-1.0);
r.olabels = {'a_1(x)'};
r.compare{1}= @(t,~) 1;
e = xspde(r);
end
```

- Add an additive complex noise of $0.01(w_1 + iw_2)$ to the differential equation, then replot with an average over 1000 samples.

4.6 Gaussian diffraction

Free diffraction of a Gaussian wave-function in three dimensions, is given by

$$\frac{da}{dt} = \frac{i}{2} \nabla^2 a$$

The xSPDE spectral definition in space-time is:

$$\tilde{a}(\omega, \mathbf{k}) = \frac{1}{[2\pi]^{d/2}} \int e^{i(\mathbf{k} \cdot \mathbf{x} - \omega t)} a(t, \mathbf{x}) dt d\mathbf{x}$$

Together with the initial condition that $a(0, x) = \exp(-|\mathbf{x}|^2/2)$, this has an exact solution for the diffracted intensity in either ordinary space or momentum space is therefore:

$$\begin{aligned} |a(t, \mathbf{x})|^2 &= \frac{1}{(1+t^2)^{3/2}} \exp(-|\mathbf{x}|^2 / (1+t^2)) \\ |\tilde{a}(t, \mathbf{k})|^2 &= \exp(-|\mathbf{k}|^2) \end{aligned}$$

To get an output from a spatially Fourier transformed field in three-dimensions, but without a Fourier transform in time, just set *transforms* = [0, 1, 1, 1], for the particular observable you need to calculate in transform space.

Exercises

- Solve the diffraction equation without noise
- Note that the example below stores data in a standardized HDF5 file.

```
function [e] = Gaussian()
r.dimension = 4;
r.initial = @(v,r) exp(-0.5*(r.x.^2+r.y.^2+r.z.^2));
r.linear = @(r) 1i*0.05*(r.Dx.^2+r.Dy.^2+r.Dz.^2);
r.observe = @(a,r) a.*conj(a);
r.olabels = '|a(x)|^2';
r.file = 'Gaussian.h5';
r.images = 4;
[e,r] = xsim(r);
e = xgraph(r.file);
end
```

- Add an additive complex noise of $0.01(w_1 + iw_2)$ to the Gaussian differential equation, then replot with an average over 10 samples.

Note that for this, you'll need to define $r.da = @(a,w,r)0.01 * (w(1,:) + i * w(2,:))$

4.7 Planar noise

The next example is growth of thermal noise of a two-component complex field in a plane, given by the equation

$$\frac{d\mathbf{a}}{dt} = \frac{i}{2}\nabla^2\mathbf{a} + \mathbf{w}(t, \mathbf{x})$$

where ζ is a delta-correlated complex noise vector field:

$$w_j(t, \mathbf{x}) = [w_j^{re}(t, \mathbf{x}) + i\zeta_j^{im}(t, \mathbf{x})] / \sqrt{2},$$

with the initial condition that the initial noise is delta-correlated in position space

$$a(0, \mathbf{x}) = \zeta^{(in)}(\mathbf{x})$$

where:

$$\zeta^{(in)}(\mathbf{x}) = [\zeta^{re(in)}(\mathbf{x}) + i\zeta^{im(in)}(\mathbf{x})] / \sqrt{2}$$

This has an exact solution for the noise intensity in either ordinary space or momentum space:

$$\begin{aligned} \langle |a_j(t, \mathbf{x})|^2 \rangle &= (1+t)/dV \\ \langle |\tilde{a}_j(t, \mathbf{k})|^2 \rangle &= (1+t)/dV_k \\ \langle \tilde{a}_1(t, \mathbf{k}) \tilde{a}_2^*(t, \mathbf{k}) \rangle &= 0 \end{aligned}$$

Here, the noise is delta-correlated, and dV , dV_k are the cartesian space and momentum space lattice cell volumes respectively. Suppose that $n = n_x n_y$ is the total number of spatial points, and there are $n_{x(y)}$ points in the $x(y)$ -direction, so then:

$$\begin{aligned} dV &= dx dy \\ dV_k &= dk_x dk_y = \frac{(2\pi)^2}{ndV}. \end{aligned}$$

In the simulations, two planar noise fields are propagated, one using delta-correlated noise, the other with noise transformed to momentum space to allow filtering. This allows use of finite correlation lengths when needed, by including a frequency filter function that is used to multiply the noise in Fourier-space. The Fourier-space noise variance is the square of the filter function.

The first noise index, $r.noises(1)$, indicates how many noise fields are generated, while $r.noises(2)$ indicates how many of these are spatially correlated, via Fourier transform, filter and inverse Fourier transform. These appear to the user as additional noises, so the total is $r.noises(1) + r.noises(2)$. The filtered noises have a finite correlation length. They are correlated with the first $r.noises(1)$ x-space noises they are generated from, as this can be useful.

Exercises

- Solve the planar noise growth equation

```
function [e] = Planar()
r.name = 'Planar noise growth';
r.dimension = 3;
r.fields = 2;
r.ranges = [1,5,5];
r.steps = 2;
r.noises = [2,2];
r.ensembles = [10,4,4];
r.initial = @Initial;
r.da = @Da;
r.linear = @Linear;
r.observe = @(a,r) a(1,:).*conj(a(1,:));
r.olabels= '<|a_1(x)|^2>';
r.compare = @(t,r) [1+t]/r.dv;
r.images = 4;
e = xspde(r);
end

function a0 = Initial(v,r)
a0(1,:) = (v(1,:)+1i*v(2,:))/sqrt(2);
a0(2,:) = (v(3,:)+1i*v(4,:))/sqrt(2);
end

function da = Da(a,z,r)
da(1,:) = (z(1,:)+1i*z(2,:))/sqrt(2);
da(2,:) = (z(3,:)+1i*z(4,:))/sqrt(2);
end

function L = Linear(r)
lap = r.Dx.^2+r.Dy.^2;
L(1,:) = 1i*0.5*lap(:);
L(2,:) = 1i*0.5*lap(:);
end
```

- Add a decay rate of $-a$ to the differential equation, then replot
- Add growth and nonlinear saturation terms

4.8 GPE

The next example is a Gross-Pitaevskii equation in two dimensions,

$$\frac{da}{dt} = \frac{i}{2} \nabla^2 a - ia(V(r) + |a|^2) + \epsilon \eta$$

where η is a correlated complex noise vector field:

$$\eta(t, \mathbf{x}) = w_1(t, \mathbf{x}) + iw_2(t, \mathbf{x}),$$

with the initial condition that the initial random field and the noise are both filtered in momentum space

$$a(0, \mathbf{x}) = a_0(\mathbf{x}) + \epsilon \zeta^{(in)}(\mathbf{x})$$

where:

$$\zeta^{(in)}(\mathbf{x}) = v_1(\mathbf{x}) + iv_2(\mathbf{x})$$

We add a Gaussian filter in momentum space for both the initial random field and noise so that, if $\tilde{w}(\mathbf{k})$ is a delta-correlated noise in momentum space:

$$\begin{aligned} w(\mathbf{k}) &= \tilde{w}(\mathbf{k}) \exp\left(-|\mathbf{k}|^2\right) \\ v(\mathbf{k}) &= \tilde{v}(\mathbf{k}) \exp\left(-|\mathbf{k}|^2\right) \end{aligned}$$

This allows use of finite correlation lengths when needed, by including a frequency filter function that is used to multiply the noise in Fourier-space. The Fourier-space noise variance is the square of the filter function.

The first noise index, $r.noises(1)$, indicates how many noise fields are generated that are delta-correlated in x , while $r.noises(2)$ indicates how many of these are spatially correlated, via Fourier transform, filter and inverse Fourier transform. These appear to the user as additional noises, so the total is $r.noises(1) + r.noises(2)$. The filtered noises have a finite correlation length.

Exercises

- Solve the GPE equation

```

function [e] = GPE()
% e = GPE() tests xSPDE for a vortex in two space dimensions
% Tests a three-dimensional noisy GPE in a harmonic well by
% (1) Modifying points and ranges in 3 dimensions
% (2) Using momentum filters on noises generated
% (3) Setting the noise and random transform filter in k-space
% (4) Changing the movie image type to give a contour plot
% (5) Using auxiliary functions defined inline
% (6) Graphing data from results saved in an HDF5 file
% xSPDE functions are licensed by Peter D. Drummond, (2015) - see
License
cd ~
r.name = 'GPE';
r.dimension = 3;
r.points = [101,64,64];
r.ranges = [1,20,20];
r.noises = [0,2];
r.rfilter = @(w,r) w.*[exp(- r.kx.^2-r.ky.^2);exp(- r.kx.^2-r.ky.^2)];
r.nfilter = @(v,r) v.*[exp(- r.kx.^2-r.ky.^2);exp(- r.kx.^2-r.ky.^2)];
b = @(xi) .1*(xi(1,:)+1i*xi(2,:));
r.initial = @(w,r) (r.x+1i*r.y)./(1+10*(r.x.^2 +r.y.^2))+b(w);
V = @(r) 0.01*(r.x.^2 +r.y.^2)-0.001*1i*(r.x.^2 +r.y.^2).^2;
r.da = @(a,xi,r) -1i*a.*(V(r)+conj(a).*a)+b(xi);
r.linear = @(r) 0.5*1i*(r.Dx.^2+r.Dy.^2);
r.observe{1} = @(a,r) a.*conj(a);
r.images = {2};
r.imagetype = {2};
r.olabels = {'|a|^2'};
r.file = 'GPE.mat';
%r.print =2;
[e,r] = xsim(r); %%sim program
xgraph(r.file,r); %%graph program
end %%end of function

```

4.9 Characteristic equation

The next example is the characteristic equation for a traveling wave at constant velocity. It is included to illustrate what happens at periodic boundaries, when Fourier-transform methods are used for propagation. There are a number of methods known to prevent this effect, including addition of absorbers - often called apodization - at the boundaries. The equation is:

$$\frac{da}{dt} + \frac{da}{dx} = 0$$

Together with the initial condition that $a(0, x) = \text{sech}(2x + 5)$, this has an exact solution that propagates at a constant velocity:

$$a(t, x) = \text{sech}(2(x - t) + 5)$$

The time evolution at $x = 0$ is simply:

$$a(t, 0) = \text{sech}(2(t - 5/2))$$

Exercises

- Solve the characteristic equation given above, noting the effects of periodic boundaries.

```
function [e] = Characteristic()
r.name = 'Characteristic'
r.dimension = 2;
r.initial = @(v,r) sech(2.*(r.x+2.5));
r.da = @(a,z,r) 0*a;
r.linear = @(r) -r.Dx;
r.olabels = {'a_1(x)'};
r.compare = @(t,r) sech(2.*(t-2.5));
e = xspde(r);
end
```

- Recalculate with the opposite velocity, and a new exact solution.

4.10 Challenge problem #2

Counter-intuitively, a random potential *prevents* normal wave-packet spreading in quantum-mechanics. This is Anderson localization: a famous property of quantum mechanics in a random potential. A typical experimental method is to confine an ultra-cold Bose-Einstein condensate (BEC) in a trap, then release the BEC in a random external potential produced by a laser. The expansion rate of the BEC is reduced by the Anderson localization due to the random potential. Physically, the observable quantity is the particle density $n = |\psi|^2$.

This can be treated either using a Schrodinger equation with a random potential, at low density, or using the Gross-Pitaevskii (GP) equation to include atom-atom interactions at the mean field level. In this example of a problem where strong localization occurs, the general equations are:

$$\frac{\partial \psi}{\partial t} = \frac{1}{i\hbar} \left[-\frac{\hbar^2}{2m} \nabla^2 + V(\mathbf{r}) + g|\psi|^2 \right] \psi$$

In calculations, it is best to use a dimensionless form by rescaling coordinates and fields. A simple way to simulate this with xSPDE is to treat ψ as a scaled field $a(1)$, and to assume the random potential field $V(\mathbf{r})$ as caused by interactions with second random field $|a(2)|^2$. This has the advantage that it is similar to the actual experiment, and allows one to treat time-dependent potentials as well, if desired.

With the rescaling, this simplifies to:

$$\frac{\partial a_1}{\partial \tau} = i \left[\frac{\partial^2}{\partial \zeta^2} - |a_2|^2 - |a_1|^2 \right] a_1$$

A convenient initial condition is to use:

$$\begin{aligned} a_1 &= a_0 \exp(-\zeta^2) \\ \langle a_2(\zeta) a_2(\zeta') \rangle &= v \delta(\zeta - \zeta') \end{aligned}$$

Exercise

- **Solve Schrodinger's equation without a random potential, to observe expansion.**
- **Include a random potential v , to observe localization.**
- **Experiment with nonlinear terms and higher dimensions.**

Note that the GP equation is a mean field approximation; this is still not a full solution of the many-body problem! Also, the experiments are somewhat more complicated than this, and actually observe the momentum distribution.

5 Logic and data

The simulation program logic is straightforward. It is a very compact function called **xspde**. This calls **xSIM**, for the simulation, then **xGRAPH** for the graphics. Most of the work is done by other specialized functions. Input parameters come from an **input** array, output is saved either in a **data** array, or else in a specified file. During the simulation, global averages and error-bars are calculated for both time-step and sampling errors. When completed, timing and error results are printed.

5.1 Simulations: xSIM

5.1.1 Input and data structures

To explain xSPDE in full detail,

- Simulation inputs are stored in the **input** cell array.
- This describes a *sequence* of simulations, so that **input**=**{r1,r2,...}**.
- Each structure **r1,r2,...** describes a simulation, whose output is the input of the next.
- The main simulation function is called using **xsim(input)**.
- Averages are recorded sequentially in the **data** cell array.
- Raw trajectory data is stored in the **raw** cell array if required.

The sequence **input** has a number of individual simulation objects, with parameters that specify the simulation functions and give the equations and observables. If there is only one simulation, just one data structure is needed, without requiring a cell array. In addition, xSPDE generates graphs with its own graphics program.

5.1.2 User definable simulation functions

The most commonly used functions are indicated with an asterisk. Others have automatic defaults that are not usually changed, except in special cases.

***initial** is used to initialize each integration in time. This is a user-defined function, which can involve random numbers if there is an initial probability distribution. This creates a stochastic field on the lattice, called **a**. The default is **xinitial**, which sets fields to zero. The returned first dimension should be **fields(1)**.

- *observe** is the initial observation function, whose output is averaged over the ensembles, called from **xpath**. The default, **xobserve**, returns the real part of the amplitudes.
- *function** is an optional function used to take arbitrary transforms of the initial observe outputs, after averaging over the *first* ensemble. The default returns the averages unchanged.
- *linear** is the linear response, including transverse derivatives in space. The default, **xlinear**, sets this to zero.
- *da** is called by **step** to calculate derivatives of the stochastic fields at every step in the process, including the stochastic terms. The default, **xstep**, sets this to zero. The returned first dimension should be `fields(1)`. All stochastic fields are accessible as `a(n,:)`, where $n \leq \text{fields}(1)$.
- *define** is called by **step** to calculate any user-defined auxiliary fields. The default, **xdefine**, sets these to zero. The returned first dimension should be `fields(2)`. If auxiliary fields are used they are accessible as `a(n,:)`, where $n > \text{fields}(1)$.
- transfer** is a function used to initialize sequential simulations, where previous or output stochastic field values are automatically included as input to the next stage in the intergration sequence. The default, **xtransfer**, uses the output of the previous simulation for the input to the next stage. This can be changed by the user.
- step** is the algorithm that computes each step in time. This also generates the random numbers at each time-step. Options include **xEuler**, **xRK2**, **xRK4**, for Euler and Runge-Kutta integration methods. This can be user-modified by initializing the handle `step` to add a new integration function. The default is **xRK4**, which is widely applicable. Another good option is **xMP**, which uses a midpoint or central-difference technique in the interaction picture.
- rfilter** returns a momentum filter for random initial fields generated in momentum space. The default, **xrfilter**, sets this to unity.
- nfilter** returns a momentum filter for random noises generated in momentum space. The default, **xnfilter**, sets this to unity.
- grid** calculates the spatial integration grid. The default, **xgrid**, returns a rectangular grid in ordinary and momentum space.
- prop** is the spatial propagation function. The default, **xprop**, takes a Fourier transform of `a`, multiplies by `propfactor` to propagate in time, then takes an inverse Fourier transform.
- randomgen** generates the initial random fields. The default, **xrandomgen**, returns Gaussian real fields that are delta-correlated in space or momentum space.

noisegen generates the initial random fields. The default, **xnoisegen**, returns Gaussian real noises that are delta-correlated in time and also in space or momentum space.

propfactor returns the interaction picture propagation factors. The default, **xpropfactor**, uses data from the **linear** function to calculate this.

5.2 Arrays and errors

Knowing the details of array indexing inside xSPDE isn't usually necessary. Yet it becomes important if you want to write your own functions to extend xSPDE, interface xSPDE with other functions, or read and write xSPDE data files with external programs. It also helps to understand how the program works.

5.2.1 Simulation data in xSIM

In xSIM, the space-time dimension d is unlimited, although xGRAPH can only plot up to three selected choices of axes. All data, including grid coordinates, fields and average results is stored in real or complex numerical arrays of implicit or explicit rank $2 + d$. During the stochastic simulation, these arrays are flattened to matrices, to shorten the Matlab code in the user functions. This preserves the same implicit index ordering, and these flattened matrices can be expanded into full arrays.

The array index ordering in all xSPDE functions always (i, \mathbf{j}, e) , where:

- The first index is a field index (i). In cases like coordinates where this isn't needed, the first index is $i = 1$.
- The next d indices $j_1, \dots, j_d \equiv \mathbf{j}$ are for time and space, where \mathbf{j} indicates a d -dimensional index
- To conserve memory, transient arrays during calculations use $\mathbf{j}' = [1, j_2, \dots]$, which indicates a d dimensional index with time index set to one.
- The last index is an ensembles or checks index (e). For fields, this is used to store field trajectories.

When the fields, noises or coordinates are integrated by the xSIM integration functions, the field and data arrays are always flattened to a matrix. This is for reasons of convenience in defining mathematical operations. The first index is the field or line index, and the combined second index covers all the rest: $(i, \mathbf{j}, e) \rightarrow (i, J)$. This is done internally, but not in externally stored data, nor in xGRAPH, which operates on the full data arrays.

Stored data uses heterogenous cell arrays to package numerical arrays with additional high level indices. The first is always the sequence index, s . Raw data also requires indices for checks and high level ensembles. Inside a sequence, data cell arrays have an additional graph index n . This distinguishes the different averages generated for output graphs and data.

In summary, the xSPDE internal arrays are as follows:

- **Field** arrays $a(i, \mathbf{j}', e)$ - these have a field index, a low-level ensemble index $e = 1, \dots, \text{ensembles}(1)$, one time-point, and a full spatial lattice.
- **Random** and **noise** arrays $w(i, \mathbf{j}', e)$ - these are like field arrays. They are initial random fields or noise fields for the stochastic equations. The first index can have a different range to the field index.
- **Coordinate** arrays $x(1, \mathbf{j}', 1)$ - these arrays are the values of coordinates at grid-points, with labels x, y, z . Numeric labels $x\{l\}$ are used with dimensions $d > 4$, depending on the axis number, $l = 2, \dots, d$. The same sizes are used for:
 - momentum coordinates kx, ky, kz (alternatively labelled $k\{2\}, k\{3\}, \dots$) -
 - spectral derivative arrays Dx, Dy, Dz (alternatively labelled $D\{2\}, D\{3\}, \dots$)
- **Raw** arrays $r\{s, c, h\}(i, \mathbf{j}, e)$ These are cell array of fields with all sequences, error-checking, trajectories and time points stored. They are used optionally, as they take up large amounts of memory. These are saved in cell arrays with indices s for the sequence, c for the time-step error-check and h for high level ensemble index.
 - $s = 1, \dots, S$ for the sequence number,
 - $c = 1, 2$ for the error-checking time-step used, first coarse then fine,
 - $h = 1, \dots, \text{ensemble}(2) * \text{ensemble}(3)$ for a high level combined parallel and serial ensemble index.
- **Data** arrays $d\{g\}(\ell, \mathbf{j}, c)$ - these are first generated in xSIM by the observe functions, internally transformed if required, then passed to xGRAPH to store generated average data at all time points, with a *check* index $c = 1, 2, 3$. Check indices are $c = 1$ for the estimated average, $c = 2$ for the estimated time-step error, and $c = 3$ for the estimated statistical error or standard deviation. The cell index g is the *graph* index, and the first array index ℓ is a user-generated graphics *line* index. This may or may not correspond to a field index: it is a user option.

Although the indices always occur in the same order, the fields, noises and output data can have different first dimensions. They are specified using different input parameters.

5.2.2 Graphics data in xGRAPH

After averaging, the data no longer needs an ensemble index. However, averaged data uses an index in the same location, to specify the error bars. Similarly, the field index is replaced by a line index, which is used as the first index of averaged data to generate a different line on the graph. These are then combined into a sequence, each with multiple graphs, and therefore need more indices.

- **Graphics** data arrays $gd\{s\}\{g\}(\ell, \mathbf{j}, c)$ - these store the complete sequence of *data* that is plotted, and may even have further functional transformations when processed in xGRAPH, if required. Graphics cell data requires high-level cell

indices $\{s\}$ for the *sequence* index, and $\{g\}$ for the *graph* index, which normally corresponds to the index $\{g\}$ of the observe function used to generate averages, unless the index was changed internally by an xSIM function. One can have several data arrays to make a number of distinct output graphs labelled g . Each graph may have multiple averages, giving different lines ℓ for comparisons.

The output line index o in a graphics or data array describes different lines on a graph. Similarly, the last ensemble index e in a field array has a different range to c , the check index, in a data array.

5.2.3 Data indexing summary

The following detailed information is not required to use xSPDE, but it helps to give insight to how data is stored, and how to customize and extend the code. There are several different types of arrays used. These are as follows:

Label	Indices	Description
a	(i, \mathbf{j}', e)	Stochastic fields
w	(i, \mathbf{j}', e)	Random and noise fields
x	$(1, \mathbf{j}', e)$	Space coordinates
r	$\{s, c, h\}(i, \mathbf{j}, e)$	Raw data
o	$\{g\}(\ell, \mathbf{j}, c)$	Observed average data
g	$\{s\}\{g\}(\ell, \mathbf{j}, c)$	Graph data

Here:

- i is the field index,
- \mathbf{j} is the full time-space index, \mathbf{j}' is the space index with a time index set to 1,
- e is the low-level ensemble or error index,
- s is the sequence index,
- c is the error-checking index,
- h is the high-level ensemble index,
- g is the graph index,
- ℓ is the graphics line index.

5.2.4 Step-size errors

Errors caused by the finite time-domain step-size, as well as those caused by the finite ensemble, are checked automatically. Those caused by the spatial lattice are not checked in the xSIM code. They must be checked by manually, by comparing results with different transverse lattice ranges and step-size.

Errors due to a finite step-size are estimated by comparing two simulations with different step-sizes and the same random sequence, to make sure the final results are accurate. The final 2D output graphs will have error-bars if *checks* = 1 was specified, which is also the default parameter setting. Error-bars below a minimum relative size compared to the vertical range of the plot, specified by the graphics variable *minbar*, are not plotted.

There is a clear strategy if the errors are too large. Either increase the *points*, which gives more plotted points and lower errors, or increase the *steps*, which reduces the step size without changing the graphical resolution. The default algorithm and extrapolation order can also be changed, please read the full xSPDE manual when doing this. Error bars on the graphs can be removed by setting *checks* = 0 or increasing *minbar* in final graphs.

5.2.5 Sampling errors

Sampling error estimation in xSIM uses sub-ensemble averaging. This generally leads to more reliable sampling error estimates, and makes efficient use of the vector instruction sets that are used by Matlab. Ensembles are specified in three levels. The first, *ensemble(1)*, is called the number of samples for brevity. All computed quantities returned by the **observe** functions are first averaged over the samples, which are calculated efficiently using a parallel vector of trajectories. By the central limit theorem, these low-level sample averages are distributed as a normal distribution at large sample number.

Next, the sample averages are averaged **again** over the two higher level ensembles, if specified. This time, the variance is accumulated. The variance of these distributions is used to estimate a standard deviation in the mean, since each computed quantity is now a normally distributed result. This method is applied to all the *graphs* observables. The two lines generated represent $\bar{o} \pm \sigma$, where *o* is the observe function output, and σ is the standard deviation in the mean.

The highest level ensemble, *ensemble(3)*, is used for parallel simulations. This requires the Matlab parallel toolbox. Either type of high-level ensemble, or both together, can be used to calculate sampling errors.

Note that one standard deviation is not a strong bound; errors are expected to exceed this value in 32% of observed measurements. Another point to remember is that stochastic errors are often correlated, so that a group of points may all have similar errors due to statistical sampling.

If *ensembles(2)* > 1 or *ensembles(3)* > 1, which allows xSPDE to calculate sampling errors, it will plot upper and lower limits of one standard deviation. If the sampling errors are too large, try increasing *ensembles(1)*, which increases the trajectories in a single thread. An alternative is to increase *ensembles(2)*. This is slower, but is only limited by the compute time, or else to increase *ensembles(3)*, which gives higher level parallelization. Each is limited in different ways; the first by memory, and the second by time, the third by the number of available cores. Sampling error control helps ensures accuracy.

5.3 Graphics: xGRAPH

The main functions involved for graphics are:

xGRAPH is called by xSPDE when the ensemble loops finished. The results are graphed and output if required.

5.3.1 xGRAPH inputs

The input to *xGRAPH* can either come from a file, or it can be from data generated directly with *xSIM*. The data is a cell array, where each member of the cell array is a numerical graphics array, defining one independent set of averaged data. the observed data averages, stored in a cell array of size $d\{n\}(o, j, c)$.

5.3.2 User definable graphics functions

It is possible to simply run **xgraph** as is, without much intervention. However, there are many customization options, including two user defined functions which have useful applications. These are as follows:

***gfunction(d,r)** is a cell array of functions whose output is plotted. There is one graphics function for each multidimensional dataset that is plotted. The default value is just the average stored in the data array generated by the simulation observe function.

An arbitrary number of functions of these observables can also be plotted, including vector observables. The input to graphics functions is the observed data averages or functions of averages, stored in a cell array of size $d\{n\}(o, j, c)$.

***compare(r)** is a cell array of functions used to obtain comparison results. These are calculated from the user-specified **compare** handle, giving a dashed line on the generated time-domain graphs, and an error summary which is printed.

The results from the two graphics functions plotted using the xGRAPH routines.

5.3.3 xGRAPH outputs

This routine is prewritten to cover a range of useful graphs, but can be modified to suit the user.

The code will cascade down from higher to lower dimensions, generating different types of user-defined graphs. Each type of graph is generated once for each specified graphics function. The graphics axes that are used for plotting, and the points plotted, are defined using the optional **axes** input parameters, where $axes\{n\}$ indicates the n-th specified graphics function or set of generated graphical data.

If there are no **axes** inputs, or the inputs are zero - for example, $axes\{1\} = \{0, 0, 0\}$ - only the lowest dimensions are plotted, up to 3. If the **axes** inputs project out a

single point in a given dimension, - for example, $axes\{1\} = \{0, 31, -1, 0\}$, these axes are suppressed in the plots. This reduces the effective dimension of the data - in this case to two dimensions.

If the *steps* option is used on input - for example $steps = [2, 3, 4]$, this adds additional integration points between the plotted points, to improve accuracy. These extra points are treated differently in space compared to time. The extra points in time are simply not stored or plotted, to conserve memory. Extra points in space are stored, in case greater detail is needed, but the default values of the *axes* inputs are such that the intermediate points are skipped on plotting. This can be changed by using *axes* inputs for more detail.

Examples:

- $axes\{1\} = \{0\}$ - For function 1, plot all the time points; higher dimensions get defaults.
- $axes\{2\} = \{-1, 0\}$ - For function 2, plot the maximum time (the default), and all x-points.
- $axes\{3\} = \{1 : 4 : 51, 32, 64\}$ - For function 3, plot every 4-th time point at x point 32, y point 64
- $axes\{4\} = \{0, 2 : 4 : 48, 0\}$ - For function 4, plot every time point, every 4-th x point, and all y-points.

Note that -1 indicates a default ‘typical’ point, which is the last point on the time axis, and the midpoint on the other axes. The last example could also be achieved in a simpler way, except it would be applied to all default axes, by using an input of:

- $steps = [1, 4, 1]$; $points = [100, 12, 48]$ - plot all the t points; every 4th x-point, and all the y points.

The *pdimension* input can also be used to reduce dimensionality, as this sets the maximum effective *plotted dimension*. For example, $pdimension\{1\} = 1$ means that only plots vs time are output for the first function plotted. Note that in the following, t, x, y, z may be replaced by corresponding higher dimensions if there are *axes* that are suppressed. Slices can be taken at any desired point, not just the midpoint. Using the standard notation of, for example, $axes\{1\} = \{6 : 3 : 81\}$, can be used to modify the starting, interval, and finishing points for complete control on the plot points.

The graphics results depend on the resulting **effective** dimension, which is equal to the actual space-time dimension unless there is an *axes* suppression, described above. Since the plot has to include a data axis, the plot itself will have an extra data axis, as well as the space-time lattice *dimension* axes.

We can plot only three axes directly using standard graphics tools. The strategy to deal with the higher dimensionality is as follows:

dimension=4.. For higher lattice dimensions, only a slice through the chosen point, or the default midpoint, is available, reducing the lattice dimension to 3.

dimension=3 For three lattice dimensions, if *images* > 1, a movie of distinct 3D graphic images of observable *vs x,y* are plotted as *images* slices - usually in time. Otherwise, only a slice through the chosen point, or the default midpoint, is available for the highest dimension, reducing the lattice dimension to 2.

dimension=2 For two lattice dimensions, one 3D image of observable *vs x,t* is plotted. A movie of distinct 2D graphic plots is also possible. Otherwise, only a slice through $x = 0$ is available, reducing the lattice dimension to 1.

dimension=1 For one lattice dimension, a 2D plot of observable *vs t* is plotted, with data at each lattice point in time. Exact results, error bars and sampling error bounds are included if available.

In addition to time-dependent graphs, the **xGRAPH** function can generate *images* (3D) and *transverse* (2D) plots at specified points in time, up to a maximum given by the number of time points specified. The number of these can be individually specified for each graphics output. The images available are specified as *imagetype*= 1,...4, giving 3D perspective plots (the default), grey-scale colour plots, contour plots, and pseudo-color plots respectively.

Error bars, sampling errors and multiple lines are only graphed for 2D plots. Error-bars are not plotted when they are below a user-specified size, with a default of 1% of the maximum range, to improve graphics quality. Higher dimensional graphs do not include this data, for visibility reasons, but they are still recorded in the data files.

5.4 Architecture

5.4.1 xSPDE data objects

The xSPDE data objects include parameters and methods, but with a very open type of object-oriented architecture to allow extensibility. All xSPDE functions are modular and easily replaceable. In many cases this is as easy as just defining a new function handle to replace the default value. To define your own integration function, include in the xSPDE input the line:

```
r.step=@Mystep;
```

Next, include anywhere on your Matlab path the function definition, for example:

```
function a = Mystep(a,w,r)
% a = Mystep(a,w,r) propagates a step my way.
..
a = ...;
end
```

5.4.2 xSPDE hints

- When using xSPDE, it is a good idea to run the batch test script, Batchtest.m.
- Batchtest.m tests your parallel toolbox installation. If you have no license for this, omit the third ensemble setting.
- To create a project file, it is often easiest to start with an existing project function with a similar equation: see Examples folder.
- Graphics parameters can also be included in the structure **in**, to modify graphs.
- Comparison functions can be included if you want to compare with analytic results.
- A full list of functionality is listed next.

6 Metadata

The input data, which we label *input*, is a sequence of data structures that we label *in*. All of the input data is later passed to the *xgraph* function. The input data can be numbers, vectors, strings, functions and cell arrays, which just lists of data enclosed in curly brackets, `{}`. All xSPDE metadata has preferred values, so only changes from the preferences need to be input. The resulting combined input including preferred values, is stored internally as a sequence of structures in a cell array to describe the simulation sequence.

Simulation metadata, including all preferred default values that were used in a particular simulation, is also stored in the xSPDE output files. This is done in both the Matlab *.mat* and the HDF5 *.h5* output files, so the entire simulation can be easily reconstructed or changed. Note that in the current version of xSPDE, one must use a Matlab formatted *.mat* file to store raw data which includes all stochastic trajectories. Either the Matlab or the HDF5 files can be used to store both input and averaged output data.

Some conventions are used to simplify inputs, as follows:

- All input data has default values
- Vector inputs of numbers are enclosed in square brackets, `[...]`.
- Where multiple inputs of strings, functions or vectors are needed they should be enclosed in curly brackets, `{...}`, to create a cell array.
- Vector or cell array inputs with only one member don't require brackets.
- Incomplete or partial vector or cell array inputs are filled in with the last applicable default value.
- New function definitions can be handles pointing elsewhere.

6.1 Simulation data

The simulation data of the *in* structure is passed to the *xsim* program, to define the simulation details. User application constants can be included, but must not be reserved names. However, names starting with 'c' or any capital letter like 'A...' - except the reserved 'D' for derivatives - are always available to users. Use of globals is **not** recommended. It is incompatible with the Matlab parallel toolbox. Graphics data can be included, but it is simply stored for the graphics program to use.

6.1.1 Simulation parameters

All simulation data has default values as distributed, which are also user-modifiable through the *xpreferences* function. Any defaults can be checked by typing *print = 2*, then running *xSIM(r)* or *xspde(r)*. All parameters must be input as part of a structure passed to xSPDE. If multiple structures are passed, they are executed in sequence, with the output of the first simulation passed to the second, then the third, and so on.

Label	Default value	Description
<i>version</i>	'xSIM3.2'	Current version number
<i>name</i>	"	Simulation name
<i>dimension</i>	1	Space-time dimension
<i>fields</i>	[1, 0]	Number of [integrated, defined] fields
<i>ranges</i>	[10, ..]	Range of coordinates in [t,x,y,z,..]
<i>origin</i>	[0]	Origin of coordinates in [t,x,y,z,..]
<i>points</i>	51	Output lattice points in [t,x,y,z,..]
<i>noises</i>	[1 0]	Number of noise fields in [x,k]
<i>randoms</i>	[1 0]	Initial random fields in [x,k]
<i>ensembles</i>	[1 1 1]	Ensembles used for averaging
<i>steps</i>	1	Integration steps per output point
<i>iterations</i>	4	Maximum iterations
<i>order</i>	0	Extrapolation order
<i>checks</i>	1	Check time-step errors?
<i>seed</i>	0	Seed for random number generator
<i>file</i>	"	File-name: 'f.mat' = Matlab, 'f.h5' = HDF5
<i>boundaries</i>	[0, 0, 0; 0, 0, 0]	Boundary: '-1,0,1'=Neum,periodic,Dirich: per boundary.
<i>probability</i>	0	Binning ranges for probabilities: see text
<i>ipsteps</i>	2	IP transforms per time-step
<i>numberaxis</i>	0	If 1, forces use of numerical axis labels
<i>print</i>	0	Print: 1 for medium, 2 for high output
<i>c</i>	-	User specified static parameters
<i>olabels</i>	{ 'a_1', .. }	Observable labels
<i>averages</i>	1	Maximum number of observables
<i>transforms</i>	{ [0 0 0 0], .. }	Fourier transforms in [t,x,y,z,..] per observable
<i>raw</i>	0	Raw data switch: 1 for raw output
<i>scatters</i>	{ 0, .. }	Specify to obtain scatter plots, not averages
<i>octave</i>	0	Force octave syntax: 1 for octave

6.1.2 Common functions

The most common functions, output field dimensions and calling arguments, with user-definable functions marked using an asterisk, are:

Label	Arguments	Purpose
<i>*da</i>	(a, z, r)	Stochastic derivative
<i>*initial</i>	(v, r)	Function to initialize fields
<i>*linear</i>	(r)	Linear derivative function
<i>*rfilter</i>	(v, r)	Random filter function in k-space
<i>*nfilter</i>	(w, r)	Noise filter function in k-space
<i>*transfer</i>	$(v, r, a0, r0)$	Transfer in sequence (returns a,r)
<i>*step</i>	(a, z, dt, r)	Calculation of a time-step
<i>*grid</i>	(r)	Grid calculator for lattice
<i>*prop</i>	(a, r)	Interaction picture propagator
<i>*propfactor</i>	(nc, r)	Propagator array calculation
<i>*observe</i>	(a, r)	Observable function cell array
<i>*function</i>	$@(av,)av1(:, :, :).^2, ..$	Functions stored of averages
<i>*define</i>	(a, z, r)	Defines an auxiliary field
<i>randomgen</i>	(r)	Initial random generator
<i>noisegen</i>	(r)	Noise generator
<i>rfilter</i>	(v, r)	Initial random kspace filter
<i>nfilter</i>	$w, (r)$	Noise kspace filter
<i>xint</i>	$(o, [dx,]r)$	Integrates over a spatial lattice
<i>xbin</i>	$(o, [dx,]r)$	Bins results onto x-axis
<i>xint</i>	$(a, [dk,]r)$	Integrates over a reciprocal lattice
<i>xave</i>	$(a, [av,]r)$	Averages over a spatial lattice
<i>xd</i>	$(a, [D,]r)$	Spectral spatial derivative
<i>xd1</i>	$(a, [dir,]r)$	1st derivative, finite diff.
<i>xd2</i>	$(a, [dir,]r)$	2nd derivative, finite diff.

The arguments in square brackets are optional. For more details of use of integration and differentiation functions, see the user manual. In the case of the function *xint*, it is possible to integrate either with respect to dx or dk , in either ordinary space or momentum space, by changing the the second argument passed to *xint* as required. However, the fields that are passed to *xint* must be transformed first. This happens automatically when the observe function is used with transforms selected.

6.2 Internal data

Internally, xSPDE data is stored in a cell array, *latt*, of the data structures passed to functions. This includes the data given above from the input structures. In addition, it includes the computed parameters given below, which includes various internal array dimensions. The stochastic variable arrays like a, w, v are internally flattened to become two-dimensional matrices where possible, to reduce dimensionality and simplify parallel operations. Momenta are stored in two grids. The propagation momentum grid is used while propagating, the graphics grid is used when averages are calculated and graphed.

For more than four total dimensions, the spatial grid, momentum grid and deriva-

tive grid notation of $t, x, y, z, \omega, kx, ky, kz$ and Dx, Dy, Dz is changed to use numerical labels that correspond to the dimension numbers, i.e., $x\{1\}, \dots, x\{d\}, k\{1\}, \dots, k\{d\}, D\{2\}, \dots, D\{d\}$. These are simpler to handle with large numbers of space dimensions. This numeric labeling preference can be used even for lower dimensionality by setting the input switch *numberaxis* = 1. The graph labels will use the same notation also, unless specified otherwise.

Label	Type	Typical value	Description
t	real	5	Current value of time, t
w	real	0.6	Frequency passed to in.observe : ω
x, y, z	array	-	Space grid of x, y, z
kx, ky, kz	array	-	Graphics momentum grid of k_x, k_y, k_z
Dx, Dy, Dz	array	-	Derivative grid of D_x, D_y, D_z
$r\{1\}, \dots, r\{d\}$	array	-	Space grid of r_1, \dots, r_d
$k\{1\}, \dots, k\{d\}$	array	-	Graphics momentum grid of k_1, \dots, k_d
$D\{2\}, \dots, D\{d\}$	array	-	Derivative grid of $D_2, \dots, D_d = ik_1, \dots, ik_d$
dx	vector	0.2	Steps in $[t, x, y, z]$
dk	vector	0.6160	Steps in $[\omega, k_x, k_y, k_z]$
dt	double	0.2000	Output time-step
dtr	double	0.1000	Reduced, internal time-step
v	real	1	Spatial lattice volume
kv	real	1	Momentum lattice volume
dv	real	1	Spatial cell volume
dkv	real	1	Momentum cell volume
xc	vector cells	{[0,... 10]}	Space-time coordinate axes in $[t, x, y, z]$
kc	vector cells	{[0,..15,-15...]}	Computational axes in $[\omega, k_x, k_y, k_z]$
$s.dx$	double	1	Initial stochastic normalization
$s.dxt$	double	3.1623	Propagating stochastic normalization
$s.dk$	double	1	Initial k stochastic normalization
$s.dkt$	double	3.1623	Propagating k stochastic normalization
$n\text{space}$	integer	35	Number of spatial lattice points
$n\text{lattice}$	integer	3500	Total lattice: ensembles(1) x n.space
$n\text{copies}$	integer	20	ensembles(2) x ensembles(3)
$randoms$	integer	2	Number of initial random fields
$noises$	integer	2	Number of noise fields
$d.int$	vector	[1 1]	Dimensions for lattice integration
$d.a$	vector	[1 1]	Dimensions for a field (flattened)
$d.r$	vector	[1 1]	Dimensions for coordinates (flattened)
$d.aplus$	vector	[1 1 1]	Dimensions for integrated plus defined fields
$d.k$	vector	[0 1 1]	Dimensions for noise transforms
$d.obs$	vector	[1 1 1 1]	Dimensions for observations
$d.data$	vector	[1 3 51 1]	Dimensions for average data (flattened)
$d.raw$	vector	[1 1 51 1]	Dimensions for raw data (flattened)

6.3 Graphics data

The simulation data of the *in* structure is also passed to the *xgraph* program, along with optional graphics parameters for each observable, to create an extended graphics data structure.

These are all cell arrays when there is more than one type of graph or observable, so that the graphics parameters can be individually set for each output that is plotted, using the cell index $\{n\}$ in a curly bracket. The axis labels, axis points and axis transformations of each graph are also cell arrays, as there can be any number of axes: so these are all nested cell arrays. If only a single input is specified, the cell index is automatically supplied. Graphics defaults are also user-modifiable by editing the *xgpreferences* function.

The plotted result can be an arbitrary function of the generated average data, by using the optional input *gfunctions*. If this is omitted, the generated average data from *xsim* is plotted.

Label	Default value	Description
<i>gversion</i>	'xGRAPH3.1'	Current version number
<i>in.olabels</i> { <i>n</i> }	'a'	Cell array of output data labels
<i>xlabel</i> s	{ 't' 'x' 'y' 'z' ... }	Generic axis labels
<i>klabel</i> s	{ '\omega' 'k_x' 'k_y' 'k_z' ... }	Transformed axis labels
<i>minbar</i> { <i>n</i> }	0.01	Minimum relative error-bar
<i>*gfunction</i> { <i>n</i> }	@(d,~) d{ <i>n</i> }	Functions plotted of averages
<i>font</i> { <i>n</i> }	18	Font size for graph labels
<i>headers</i> { <i>n</i> }	"	Graph headers
<i>images</i> { <i>n</i> }	0	Number of movie images
<i>imagetype</i> { <i>n</i> }	0	Type of movie images
<i>transverse</i> { <i>n</i> }	0	Number of transverse plots
<i>pdimension</i> { <i>n</i> }	3	Maximum plot dimensions
<i>*compare</i> { <i>n</i> }	-	Comparison functions
<i>*xfunctions</i> { <i>n</i> }	-	Axis transformations
<i>axes</i> { <i>n</i> }	{0,..}	Points plotted for each axis
<i>glabel</i> s{ <i>n</i> }	{ 't' 'x' 'y' 'z' }	Graph-specific axis labels
<i>graphs</i>	-	Maximum number of graphs

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